

# Multiomics analysis provides insights into flavonoids accumulation and biosynthesis in different planting years and localities of *Gongronemopsis tenacissima* (Dai-Bai-Jie) (#109469)

1

Second revision

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# Multiomics analysis provides insights into flavonoids accumulation and biosynthesis in different planting years and localities of *Gongronemopsis tenacissima* ( Dai-Bai-Jie )

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The dried root of *Gongronemopsis tenacissima* ( Roxb. ) S.Reuss, Liede & Meve is a traditional medicine utilized by the Dai ethnic group, commonly known as Dai-Bai-Jie , primarily for detoxification purposes in traditional medicine . Due to the extensive utilization, the wild resources are becoming increasingly scarce. The plants have been domesticated in China. However, the accumulation patterns of secondary metabolites, particularly flavonoids - the primary detoxifying components - along with their biosynthesis, remain unclear.

The differences in flavonoid accumulation and transcriptional regulatory mechanisms underlying the differential accumulation of flavonoids in Dai-Bai-Jie, cultivated for one, two, and three years at high altitudes , as well as three years at low altitudes , were investigated using transcriptome and widely targeted metabolome methods. A total of 1,495 1495 metabolites were identified through

Ultra Performance Liquid Chromatography coupled with Tandem Mass Spectrometry ( UPLC-MS/MS ) from Dai-Bai-Jie, and 943 differential accumulation metabolites were detected among the four groups. All the flavonoids were grouped into six clusters by k-means cluster analysis . The total metabolite content in two, and three years was relatively abundant, and flavonoid levels were generally higher in two, and three years. It is recommended that harvesting at two years of age be considered the optimal strategy . All flavonoids were organized into six clusters through k-means cluster analysis.

A regulatory relationship was observed between genes such as phenylalanine ammonia-lyase, CYP73A, 4-Coumarate: Coenzyme A Ligase, and lavonol synthase and the flavonoid components in Dai-Bai-Jie. However, significant differences in the Shannon, Chao1, or Abundance Coverage Estimator ( ACE ) indices of rhizosphere microorganisms were detected across different planting years and localities were not detected. This study elucidates the accumulation mechanisms of flavonoids and the scientificity of harvesting

years for Dai-Bai-Jie . The results provide a crucial scientific foundation for guiding the large-scale introduction and cultivation of Dai-Bai-Jie as a supplement or alternative to the use of wild resources.

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4 Jie)

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19

20 **Abstract****ABSTRACT**

21 The dried root of *Gongronemopsis tenacissima* (Roxb.) S.Reuss, Liede & Meve is a traditional  
22 medicine utilized by the Dai ethnic group, commonly known as Dai-Bai-Jie, primarily for  
23 detoxification purposes in traditional medicine. Due to the extensive utilization, the wild  
24 resources are becoming increasingly scarce. The plants have been domesticated in China.  
25 However, the accumulation patterns of secondary metabolites, particularly flavonoids-the  
26 primary detoxifying components-along with their biosynthesis, remain unclear.

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44 accumulation mechanisms of flavonoids and the scientificity of harvesting years for Dai-Bai-Jie.  
45 The results provide a crucial scientific foundation for guiding the large-scale introduction and  
46 cultivation of Dai-Bai-Jie -as a supplement or alternative to -the use of wild resources.  
47

48 Keywords: *Gongronemopsis tenacissima*; metabolome; transcriptome; rhizosphere microbes;  
49 flavonoids  
50

## 51 **Introduction****INTRODUCTION**

52 *Gongronemopsis Marsdenia tenacissima* (Roxb.) Moon S.Reuss, Liede & Meve- is a traditional  
53 medicine utilized by the Dai ethnic group, commonly known as Dai-Bai-Jie-, and -holds  
54 significant value in the ethnomedical traditions of Southeast Asia. In the Dai language, it is  
55 referred to as "Ya Jie Xian Da," symbolizing its ability to purge the body of numerous toxins.  
56 This medicinal herb has long been utilized in Dai-inhabited regions such as Xishuangbanna,  
57 Dehong, Ximeng, Menglian, Xinping, Yuanjiang, Mojiang, and Puer in China, as well as  
58 neighbouring countries like Laos and Myanmar (Li *et al.*, 1995). The root of *G. tenacissima* is  
59 employed in folkloric medicine named Dai-Bai-Jie for detoxification purposes. -It is recognized  
60 for its efficacy in counteracting toxicities resulting from various sources, including food,  
61 - animals, and environmental factors such as heat, water, and fire burns. Additionally, it is -used  
62 to relieve throat discomfort and swelling caused by excessive heat toxicity. With a rich  
63 historical background in traditional medicine, *G. tenacissima* (Dai-Bai-Jie )has been  
64 incorporated -into contemporary hospital preparations at institutions like the Xishuangbanna Dai  
65 Hospital. These formulations -include Bai-jie Capsules, Ya-jie Gahan, and Banna Coolant.  
66 Modern pharmacological research has showned that "Dai-Bai-Jie" ~~possessees~~possesses  
67 inhibitory effects on cancer cells, protects against liver damage caused by certain drugs,  
68 demonstrates anti-HIV activity, possesses antioxidant properties, and exhibits antibacterial  
69 activities (Gao *et al.*, 2014; Li *et al.*, 2021).  
70

71 Currently, various bioactive compounds have been isolated from *G. tenacissima*,Dai-Bai-Jie,  
72 including organic acids, polyoxyprogesterone glycosides, volatile oils, and pyrrole alkaloids  
73 (Liao *et al.*, 2016; Pang *et al.*, 2018; Song *et al.*, 2018; Song *et al.*, 2021). These discoveries  
74 not only deepen our understanding of the medicinal properties of this herb but also open up  
75 potential avenues for therapeutic applications in modern medicine.

76 For a long time-, Dai-Bai-Jie- was incorrectly identified as the dried root of *Dregea sinensis*  
77 Hemsl., which belongs to the genus *Dregea* of the Asclepiadaceae family (Lin *et al.*, 2003).  
78 However, a pivotal study conducted in 2014 revealed that Dai-Bai-Jie is, in fact, the dried root  
79 of *M. tenacissima*, a species within -the genus *Marsdenia* (Li *et al.*, 2014; Li -et al., 2023).  
80 This identification was based on comprehensive molecular and morphological analyses utilizing

80 DNA fragments such as *psbD-trnT*, *trnLF*, and ITS, along with observations of leaf morphology  
81 and floral characteristics. It is important to note that the “tong-guan-teng” mentioned in the  
82 Chinese Pharmacopoeia, known for its broad-spectrum anticancer properties, is associated with  
83 *M. cavaleriei* (Chen et al., 2022; Li et al., 2014). Current scientific investigations have  
84 revealed significant differences in the chemical composition and therapeutic effects of these two  
85 species. Specifically, Dai-Bai-Jie is primarily indicated for antidotal properties and management  
86 of gastrointestinal disease, while anticancer activity is chiefly attributed to *M. cavaleriei*. In  
87 2022, *M. tenacissima* was reclassified into the genus *Gongronemopsis* and is now referred to as  
88 *Gongronemopsis tenacissima* (Roxb.) (Liede-Schumann et al., 2022).  
89 Flavonoids are secondary metabolites found widely in plants, possessing a variety of functions  
90 including antioxidant, anti-inflammatory, antitumor, antiviral, antibacterial, anti-vascular  
91 sclerosis, and anti-liver fibrosis activities (referred to as *M. cavaleriei* at the time, Fang et al.,  
92 2023; Wang et al., 2020; Zhang et al., 2023). Recent studies suggested that their protective  
93 effect on intestinal mucosal barrier function may play a role in detoxification mechanisms (Yang  
94 et al., 2020). According to Dai medical theory, the occurrence of disease is closely linked to  
95 imbalances among the four cosmic elements within the body, which can be triggered by the  
96 presence of toxins (Zhang et al., 2023). Such imbalances may stem from disturbances in  
97 antioxidant defences and from disparities between pro- and anti-inflammatory factors.  
98 Notably, recent studies have demonstrated a correlation between the levels of total flavonoids  
99 and total polyphenols in Dai-Bai-Jie with its antioxidant and anti-inflammatory activities (Zhang  
100 et al., 2023). Therefore, flavonoids may represent the most significant active component for  
101 detoxification properties of *G. tenacissima*.  
102 Due to the extensive utilization of *Dai-Bai-Jie* and *G. tenacissima*, wild resources are becoming  
103 increasingly scarce. Fortunately, significant advancements have been made in the artificial  
104 cultivation technology for *G. tenacissima*, leading to small-scale cultivation in Xishuangbanna,  
105 Yunnan. Under natural conditions, the harvesting period for the roots of *G. tenacissima* is typically  
106 determined by empirical knowledge and generally occurs after at least two years  
107 of growth. Similarly, under cultivation conditions, the harvest period is usually 2-3 years,  
108 primarily considering the biomass of the roots.  
109 Despite these advancements, the accumulation patterns of flavonoids in *Dai-Bai-Jie* and *G. tenacissima*  
110 under varying cultivation conditions remain unclear. To address this knowledge gap,  
111 this study investigated the flavonoids accumulation patterns and influencing factors of *Dai-Bai-*  
112 *Jie* and *G. tenacissima* from the multi-omics perspective, which may lead to a better understanding  
113 of the metabolic accumulation mechanism of *Dai-Bai-Jie* and *G. tenacissima* and facilitate the  
114 scientific determination of optimal harvesting years for this medicinal plant.

## 115 116 **MATERIALS AND METHODS****Materials & Methods**

117

### 118 **Plant materials and sampling**

119 The roots of one-year-old (CR1), two-year-old (CR2), and three-year-old (CR3) cultivated *G.*  
120 *tenacissima* (Dai-Bai-Jie) were collected from Menghun County, Xishuangbanna Dai  
121 Autonomous Prefecture, Yunnan, China (E 100.38°, N 21.82°; 1179 m) in November 2022  
122 (Fig. 1) . Additionally, the roots of three-year-old Dai-Bai-Jie (CR4) cultivated in South  
123 Medicine Garden (E100.79°, N22.00°; 533.57m) also located in Xishuangbanna Dai  
124 Autonomous Prefecture, Yunnan Province, China, were gathered. Each plant was divided into  
125 two sections: one for transcriptome sequencing and the other for metabolome analysis, with three  
126 biological replicates per sample. Furthermore, the rhizosphere soil (CM1, CM2, CM3, CM4)  
127 corresponding to each plant (CR1, CR2, CR3, CR4) was collected and utilized for 16S rRNA  
128 and ITS analysis.

### 129 **Metabolome analysis**

130 After the freeze-dried samples were crushed (30 Hz, 1.5 minutes), the extraction solution (70%  
131 methanol water pre-cooled to -20°C) was added, and the mixture was vortexed for 30 seconds.  
132 Subsequently, the samples were vortexed six times (once every 30 minutes) and centrifuged at  
133 12,000 rpm for 3 minutes. The supernatant was then filtered through a microporous filter  
134 membrane with a pore size of 0.22 µm and stored in an injection vial for Ultra Performance  
135 Liquid Chromatography (UPLC-MS/MS) analysis.

136 Ultra High Performance Liquid Chromatography (ExionLC™ AD) was employed for sample  
137 collection and analysis, utilizing an Agilent SB-C18 column (1.8 µm, 2.1 mm × 100 mm). The  
138 mobile phase A consisted of 0.1% formic acid in water, while the mobile phase B was  
139 acetonitrile containing 0.1% formic acid. The column temperature was maintained at 40°C, and  
140 the automatic sampler temperature was set to 4°C. The flow rate was adjusted to 0.35 mL/min,  
141 and the injection volume was 2 µL. Applied Biosystems 6500 QTRAP was used for analysis.  
142 The typical ion source parameters were as follows: electrospray ionization (ESI) temperature of  
143 500°C; ion spray voltage (IS) of 5500 V in positive ion mode and -4500 V in negative ion mode;  
144 ion source gas I (GSI), gas II (GSII), and curtain gas (CUR) were set to 50, 60, and 25 psi,  
145 respectively. The collision-induced dissociation parameters were set to high. SCIEX Analyst  
146 workstation software (version 1.6.3) was used for Multiple Reaction Monitoring (MRM) data  
147 collection and processing.

148 Using MS-Converter, MS raw data files were converted into TXT format for further analysis. An  
149 internal R program, along with a specialized database, was employed for peak detection and  
150 annotation. Prior to analysis, the raw data underwent preprocessing to filter out low-quality ion  
151 signals.

152 After obtaining the organized data, SIMCA (version 16.0.2) software was used for Principal  
153 Component Analysis (PCA) and Orthogonal Partial Least Squares Discriminant Analysis  
154 (OPLS-DA), which were used to explore the metabolic patterns and identify differential  
155 metabolites **Metabolites** (DAMs) with p-values < 0.05 and VIP (variable importance in  
156 projection) >1.

### 157 **RNA-seq processing and data analysis**

158 Total RNA was extracted and purified from the above samples. The extracted RNA was tested  
159 for purity, concentration, and integrity. After the samples were qualified, the mRNA was isolated  
160 and purified by Oligo (dt) for the construction of the cDNA library. Illumina Novaseq 6000  
161 sequencing was performed after the library was qualified. Fastp software (Chen et al., 2018)  
162 was used for quality control on the raw data.

163 After obtaining Clean Reads, Trinity assembly software is used to splice the Clean Reads to  
164 obtain reference sequences for subsequent analysis, trinity assembly software was used to stitch  
165 the clean reads to obtain reference sequences for subsequent analysis.

166 The RSeQC software (*Wang et al.*, 2012) was used to evaluate the quality of transcriptome data,  
167 and to analyze the sequencing data after passing the quality evaluation. Fragments per Kilobase  
168 Million (FPKM) (*Trapnell et al.* 2010) was used to estimate gene expression level. The  
169 transcriptome assembly was assessed in terms of their completeness and the percentage of  
170 complete, fragmented, and missing fragments by using the BUSCO 5.3.2 (<https://busco.ezlab.org>,  
171 *Simão et al.*, 2015). DESeq2 (*Love et al.*, 2014; *Varet et al.*, 2016) was used for differential  
172 expression analysis between samples. The corrected p-value and FDR (False Discovery Rate) were  
173 used as the key indicators for the screening of differentially expressed genes (DEGs). Weighted  
174 Geneco-expression Network Analysis (WGCNA) was used to find the gene modules that are co-  
175 expressed and constructed the hierarchical clustering tree. The statistical power of this  
176 experimental design, calculated in RNASeqPower is 0.70.

177 The whole transcript data set can be found in the National Center for Biotechnology Information  
178 (NCBI) database (BioProject ID: PRJNA996325 ).

179

#### 180 **RT-qPCR validation**

181 We selected five genes associated with flavonoids synthesis for RT-qPCR according to FPKM  
182 value (*Forkmann et al.*, 2001; *Zou et al.*, 2016 ) . GAPDH was used as a reference gene and all  
183 genes used in this study are listed in Table 1. cDNA was synthesized using MonScript™ RTIII  
184 All-in-One Mix with ds DNase (Monad, China). According to the instructions of QuantiNova  
185 SYBR Green PCR Kit (Qiagen, China), RT-qPCR was performed. The total volume of the system  
186 was 10  $\mu$ L, including 5 $\mu$ L 2x SYBR Green PCR Master Mix, 0.7  $\mu$ L upstream Primer with 0.7  
187  $\mu$ M, 0.7  $\mu$ L down-stream primer with 0.7 $\mu$ M, 1 $\mu$ L cDNA with $\leq$ 100ng/reaction, 2.55  $\mu$ L RNase-  
188 free water, 0.05  $\mu$ L QN ROX Reference Dye.

189 Microbial DNA extraction, 16S rRNA and ITS gene sequencing

190 Genomic DNA was extracted using CTAB (Nobleryder, China). 30  $\mu$ L PCR amplification  
191 system was as follows: Phusion® High-Fidelity PCR Master and high fidelity polymerase Mix  
192 (New England Biolabs) 15 $\mu$ L, Primer 1  $\mu$ L, DNA5-10 ng, ddH<sub>2</sub>O. 16S V4 Regional primer  
193 (GTGCCAGCMGCCGCGGGGTAA and 806R GGACTACHVGGGTWTCTAAT) was used  
194 for to identify bacterial diversity. ITS5-1737F 5'-GGAAGTAAAAGTCGTAACAAGG-3'  
195 and ITS2-2043R 5'-GCTGCGTTCTTCATCGATGC-3' was used for identified fungal  
196 diversity. Reaction procedure was set at 98 °C for 1 min, followed by 40 cycles at 98 °C for 10  
197 s, 0 °C for 38 s, and 72 °C for 30 s, 72 °C extension for 5 minutes finally. PCR products was  
198 sequenced on the NovaSeq6000 platform (Maiwei Biotechnology Company).

## 199 **Results**

### 200 **3.1 RNA-seq analysis and DEGs identification**

201 We performed high-throughput transcriptome sequencing on the CR1, CR2, CR3, and CR4 of  
202 Dai-Bai-Jie, with three biological replicates per sample. In total, we obtained 78.27 GB of clean  
203 data. The clean Data of all sample was not less than 6 GB of clean data. The percentages of  
204 bases with a Q30 quality score was greater than 90% for all samples. After assembling and  
205 splicing, 85,346 unigenes were obtained. A BUSCO analysis was performed to evaluate the  
206 completeness, recovering 253 out of 255 conserved eukaryotic genes (99.2%) (Fig. 2A) .

207 Using the criteria of  $|\log_2\text{Fold Change}| \geq 1$  and  $\text{FDR} < 0.05$ , we screened for DEGs. The results  
208 revealed that 15,255, 8,170, 10,529, and 8,225 DEGs were identified in the comparisons of CR1  
209 vs. CR2, CR1 vs. CR3, CR2 vs. CR3, and CR3 vs. CR4, respectively. Among these, 654  
210 common DEGs were shared across CR1, CR2, CR3, and CR4. Specifically, There were 6,043  
211 unique DEGs identified in the comparison of CR1 vs. CR2, 1,243 unique DEGs in CR1 vs. CR3,  
212 2,720 unique DEGs in CR2 vs. CR3, and 2,957 unique DEGs in CR3 vs. CR4 (Fig. 2B).  
213 The DEGs in the four groups were analyzed using the Kyoto Encyclopedia of Genes and  
214 Genomes(KEGG) metabolic pathway. The results showed that the DEGs of CR1 vs. CR2, CR1  
215 vs. CR3, CR2 vs. CR3, and CR3 vs. CR4 were annotated to 144, 140, 143, and 140 KEGG  
216 metabolic and biosynthetic pathways, respectively. Notably, the "Metabolic pathways" category  
217 emerged as the most frequently annotated, encompassing 2492, 1428, 1669, and 1432 genes in  
218 each comparison, respectively. Closely following was the "biosynthesis of secondary  
219 metabolites" category, which annotated 1375, 800, 930, and 808 genes, respectively. The "Plant-  
220 pathogen interaction" pathway was annotated to 514, 271, 364, and 401 genes (Fig. 3).  
221 WGCNA displayed that DEGs are divided into 27 co-expression modules of CR1, CR2, CR3,  
222 and CR4. Among them, the turquoise module has the highest number of genes with 11313,  
223 followed by the blue module with 5550 genes, and the least is the white module, which has 101  
224 genes (Fig. 2C).

### 225 **3.2 RT-qPCR validation**

226 The RT-qPCR results for the five targeted genes indicated that four of them (excluding cluster-  
227 60047.2) displayed a general consistency with the relative transcript abundance observed in the  
228 transcriptome analysis. This concordance validates the reliability of the RNA-seq data (Fig. 4).

### 229 **3.3 Metabolomic profiling**

230 A total of 1495 metabolites were identified from Dai-Bai-Jie using UPLC-MS/MS . These  
231 included 378 amino acids and their derivatives (25.28%) , 265 phenolic acids (17.73%), 168  
232 lipids (11.24%), 114 flavonoids (7.63%), 103 organic acids (6.89%), 92 alkaloids (6.15%), 80  
233 nucleotides and their derivatives (5.35%), 55 lignans and coumarins (3.68%), and 42 terpenoids  
234 (2.81%), 23 steroid (1.54%) and 75 metabolites belonging to other categories (11.71%)  
235 (Fig. 5A). Notably, the flavonoid category was further subdivided into 9 chalcones, 17  
236 dihydroflavonoids, 8 dihydroflavonols, 36 flavonoids, 40 flavonols, and 4 flavanols.

237 PCA was employed to illuminate the overall metabolite differences among the different groups.  
238 The results showed that principal component 1( PC1,38.39%) and principal component (PC2,  
239 23.73%) accounted for 62.12% of the variance in the metabolic profile, indicating significant  
240 differences across four groups. The three samples within each group demonstrated high  
241 aggregation and good repeatability (Fig. 5B).

242 A total of 943 Differential metabolites (DAMs) were detected using  $\text{FC} \geq 2$  or  $\leq 0.5$  and  $\text{VIP} > 1$  as  
243 screening conditions, including 255 amino acids and their derivatives, 174 phenolic acids, 45  
244 nucleotides and their derivatives, 79 flavonoids, 42 lignans and coumarins, 64 alkaloids, 30  
245 terpenoids, 44 organic acids, 20 steroids and 83 lipids. Among them, there were one common  
246 DAMs shared of CR1, CR2, CR3, and CR4. Specifically, there were five unique DAMs in the  
247 comparison of CR1 vs. CR2, 273 unique DAMs in CR1 vs. CR3, 172 unique DAMs in CR2 vs.  
248 CR3, and 46 unique DAMs in CR3 vs. CR4 (Fig. 5D).

249 In the comparison of CR1 vs CR2, a total of 627 DAMs were detected, of which 183 were  
250 down-regulated and 444 were up-regulated. Compared to CR1, the metabolite that significantly  
251 decreased in CR2 was gofruside, whereas the metabolite that significantly increased was 4-O-

252 (2"-O-acetyl-6"-P-coumaroyl- $\beta$ -D-glucopyranosyl)-P-coumaric acid (Fig. 6A) . The metabolite  
253 protocatechuic acid 4-O-(2"-O-Vanillyl) glucoside significantly decreased in CR3 compared to  
254 CR1, while eugenol significantly increased (Fig. 6B). 449 DAMs were detected in CR2 vs CR3,  
255 with 377 down-regulated and 72 up-regulated. The metabolite 6,7-dimethoxy-2-[2-(4'-hydroxy-  
256 3'-methoxyphenyl)ethyl]chromone was significantly reduced in CR3 relative to CR2, while  
257 sinapine was significantly increased (Fig. 6C).—Lastly, a total of 259 DAMs were found in  
258 the comparison between CR3 vs CR4, with 117 down-regulated and 142 up-regulated. The  
259 metabolite that showed a significant decrease in CR4 was rutin, while exhibited a significant  
260 increase when compared to CR3 (Fig. 6D). Cluster analysis was performed on the DAMs across  
261 the four groups. The differences among the four samples groups were pronounced; specifically,  
262 the phenolic acids were commonly more abundant in CR2, and flavonoids were commonly  
263 higher in the CR1 and CR2 compared to in the other groups. Additionally, the levels of amino  
264 acids and their derivatives were higher at CR3, while the contents of terpenes, nucleotides and  
265 their derivatives were higher in CR4 (Fig. 5C).

266 To gain a deeper understanding of the accumulation patterns of metabolites in Dai-Bai-Jie across  
267 different planting ages and altitudes, we employed k-means cluster analysis to categorize all the  
268 metabolites. The analysis revealed that the metabolites clustered into six distinct groups (Fig.  
269 6E). Notably, classes 1 and 6 exhibited the highest concentration of metabolites in CR2, with  
270 class 6 containing the largest number of metabolites among all six classes. Classes 2 and 4, on  
271 the other hand, demonstrated the highest abundance of metabolites in CR3. Class 3 was  
272 characterized by the highest amount of metabolites in CR4, while class 5 displayed the highest  
273 concentration of metabolites in CR1. This categorization provides valuable insights into the  
274 specific patterns of metabolite accumulation within each growth year and altitude, enabling us to  
275 further investigate their potential biological significance.

### 276 **3.4 Comparative metabolomic analysis aiming to flavonoids and flavonoid biosynthesis- 277 related genes among the different plantation age and locality.**

278 A total of 114 flavonoids were detected from Dai-Bai-Jie, including 34.21% flavonols, 31.58%  
279 flavonoids, 14.91% dihydroflavonoids, 7.02% dihydroflavonols, 7.89% chalcone, 3.50%  
280 flavanols, 0.88% flavonols, of which 79 flavonoids were differentially accumulated. Based on K-  
281 means analysis, nine flavonoids, including 3',5-Dihydroxy-4',6,7-trimethoxyflavanone, acacitin-  
282 7-O-galactide, robinin-7-O-galactoside, phelamurin, huangbaioside, eriodictyol-7-O-glucoside,  
283 exhibited a relatively high accumulation in class 2 for CR2. 15 flavonoids including 3',4',7-  
284 trihydroxyflavone, cirsimarin, hesperetin-7-O-glucoside, quercetin, exhibited a relatively high  
285 accumulation in class 6 for CR2. Six flavonoids including kaempferol-7-O-glucuronid,  
286 hesperetin-7-O-(6"-malonyl) glucoside, quercetin-3-O-(6"-O-galloyl) galactoside, myricetin-3-  
287 O-rhamnoside (Myricitrin), diosmetin-7-O-glucuronide, syringetin-7-O-glucoside, exhibited a  
288 relatively high accumulation in class 2 for CR3 . Ten flavonoids including Rutin, hesperetin-5-  
289 O-glucoside,isorhamnetin-3-O-rhamnoside, quercetin-3-O-robinobioside, exhibited a  
290 relatively high accumulation in class 4 for CR3. Five flavonoids including 3-Hydroxy-4',5,7-  
291 trimethoxyflavanone, aromadendrin-7-O-glucoside, eriodictyol-8-C-glucoside,  
292 dihydromyricetin-3-O-glucoside, taxifolin-3'-O-glucoside, exhibited a relatively high  
293 accumulation in class 3 for CR4. 34 flavonoids including rhamnazin, quercetin-3,4'-dimethyl

294 Ether, limocitrin-7-O-glucoside, kumatakenin, exhibited relatively high accumulation in class 5  
295 for CR1.

296 To gain a deeper understanding of the molecular mechanisms underlying the differential  
297 accumulation of flavonoids across various planting year and planting environments, we  
298 conducted a comprehensive analysis of the expression patterns of genes involved in flavonoid  
299 metabolism. KEGG analysis revealed that the 15 flavonoids exhibiting differential accumulation  
300 were mapped to multiple biosynthetic pathways, including the flavonoid biosynthesis pathway  
301 (KO00941), flavonol biosynthesis pathway (KO00944), as well as the broader metabolic  
302 pathway (KO01100) and secondary metabolite biosynthesis pathway (KO01110) (Fig. 7A) .  
303 Correlation analysis was conducted between DAMs mapped to the KEGG pathway and the  
304 corresponding DEGs on the pathway, and the correlation  $> 0.8$  or  $<-0.8$  and the P-value  $<0.05$  as  
305 the screening conditions. The analysis revealed complex regulatory relationship among  
306 phenylalanine ammonia-lyase (PAL Cluster-63886.0, Cluster-63886.1), 4-Coumarate: Coenzyme  
307 A Ligase (4CL, Cluster-58688.4, Cluster-62808.3), lavonol synthase (FLS, Cluster-46899.18,  
308 Cluster-46899.5, Cluster-50957.2, Cluster-57391.0, C12RT1(Cluster-45854.0) and the  
309 metabolites hyperin, Ionicerin, vicenin-2, nicotiflorin, querceti, luteolin-7-O-(6"-malonyl)  
310 glucoside, and hesperetin-7-O-glucoside (Fig. 7B).

### 3.5 Taxonomic features of the rhizosphere microbes of Dai-Bai-Jie

311 Plants recruit specific root-associated microbes that enable them to deliver photosynthates and  
312 root exudates to their root microbiome, thereby stimulating plant growth and productivity  
313 (*Lareen et al.*, 2016). Studies have indicated that the composition of microbial communities at  
314 roots, the so-called root microbiome, can have significant impacts both on plant development  
315 and their stress tolerance (*Mendes et al.*, 2011; *Panke-Buisse et al.*, 2015).

316 The coverage index between the bacterial and fungal sample groups exceeded 0.965, indicating  
317 that the sequencing was representative and accurately reflected the bacterial and fungal diversity  
318 of the samples. The four groups of rhizosphere soil bacteria involved a total of 40 phyla, 71  
319 classes, 154 orders, 300 families, and 695 genera, and fungi comprised 13 phyla, 61 classes, 168  
320 orders, 406 families, and 875 genera. The dominant bacterial phyla in the rhizosphere soils  
321 included Crenarchaeota, Acidobacteriota, Chloroflexi, Firmicutes, Proteobacteria were  
322 the dominant bacteria in the rhizosphere soils, whereas the predominant fungal phyla were  
323 Ascomycota, Basidiomycota, Mortierellomycota, Glomeromycota, Chytridiomycota,  
324 and Rozellomycota.

325 We investigated the richness indices (alpha diversity, ACE, Chao1) and the Shannon diversity  
326 index of the microbial community, as well as the number of operational taxonomic units (OTUs)  
327 across all samples. There were no significant differences in the Shannon, Chao1, and ACE indices  
328 of rhizosphere microorganisms among the four groups (Table 2).

329 A total of 1952 bacterial operational taxonomic units (OTUs) and 5230 fungi were detected in  
330 the rhizosphere microbiome. The co-possessed bacteria in the four rhizosphere soils are 2986  
331 OTUs, 721 are unique to CM1, 406 are unique to CM2, 497 are unique to CM3, and 620 are  
332 unique to CM4 (Fig. 8A). The co-possessed fungi in the four rhizosphere soils are 5677 OTUs,  
333 383 are unique to CM1, 223 are unique to CM2, 263 are unique to CM3, and 406 are unique to  
334 CM4 (Fig. 8B).

335 Community composition analysis revealed that the compositions were similar among all the  
336 twelve rhizosphere soils samples at the phylum level. Excluding CM3.3, the abundance of  
337 Acidobacteriota in CM2 and CM3 was significantly higher than in CM1 and CM4. (Fig. 8C, D).

339 However, the community compositions presented some differences ~~S~~ among all ~~-~~twelve  
340 rhizosphere soils at the genus level (Fig. 8E, F).

## 341 **DISCUSSION**

342 The growth duration is the most critical factor affecting the quality of medicinal plants. Until  
343 now, the harvesting period of Dai-Bai-Jie has primarily centered on biomass accumulation, with  
344 the accumulation of bioactive components remaining unknown. Despite numerous research  
345 reports have examined the metabolites and anti-tumor properties of *G. tenacissima*, majority of  
346 these studies have not specifically targeted Dai-Bai-Jie, ~~S~~ largely due to inaccuracies in plant  
347 identification (Li *et al.*, 2014; Li *et al.*, 2023). Up to now, little is known about the chemical  
348 composition and active ingredients of Dai-Bai-Jie (Liao *et al.*, 2016; Pang *et al.*, 2018; Zhang *et*  
349 *al.*, 2016; Li *et al.*, 2017). This highlights the necessity for further scientific investigation to  
350 comprehensively understand the growth patterns and accumulation of bioactive components in  
351 Dai-Bai-Jie.

352 In this study, a comprehensive metabolic profiling of Dai-Bai-Jie was conducted using UPLC-  
353 MS/MS widely-targeted metabolomics analysis. A total of 1495 metabolites were successfully  
354 identified, signifying the rich metabolite content of Dai-Bai-Jie. These metabolites are likely to  
355 form the pharmacological material basis for the medicinal properties of Dai-Bai-Jie.

356 Additionally, 943 DAMs were detected from four group samples obtained from distinct locations  
357 and three different planting age, which suggests quality variations among them.

358 Flavonoids and total polyphenols were major contributors for detoxification of Dai-Bai-Jie  
359 (Zhang *et al.*, 2023). We detected a diverse array of secondary metabolites, including  
360 flavonoids, phenolic acids, alkaloids, and terpenoids, which may potentially contribute to its  
361 antioxidant and anti-inflammatory activities.

362 When comparing the accumulation of metabolites across different planting ages, it was observed  
363 that the total metabolite content in CR2 and CR3 was relatively abundant. Additionally,  
364 flavonoid levels were generally higher in CR1 and CR2. To achieve an optimal balance between  
365 biomass, economic benefits, and the biological activity of Dai-Bai-Jie, it is recommended that  
366 two-year harvesting serves as the optimal strategy.

367 Despite originating from the same planting age, samples CR3 and CR4 exhibited inconsistent  
368 trends in metabolite accumulation, revealing a total of 259 DAMs. This variation can be  
369 attributed to diverse environmental factors, including altitude, temperature, and soil conditions.

370 Although the number of DAMs identified was ~~few~~lower compared to those observed between  
371 different years, it nonetheless underscores the significant impact of the environment on the  
372 accumulation of secondary metabolites in Dai-Bai-Jie. Furthermore, it suggests that cultivation at  
373 lower altitudes may result in a ~~-~~diminished abundance of secondary metabolites. This could be  
374 due to the influence of lower temperatures at higher altitudes, which may induce the expression  
375 of resistance genes, thereby promoting the accumulation of secondary metabolites.

376 Consequently, in the future large-scale introduction and cultivation of Dai-Bai-Jie, high-altitude  
377 conditions should be carefully considered.

378 Based on our ~~W~~idely targeted metabolome analysis, flavonoids are identified as the predominant secondary metabolites in Dai-Bai-Jie. Notably, the  
379 flavonoid content is significantly greater in plants cultivated for ~~-~~two and three years compared to  
380 those cultivated for one year. This finding is generally consistent with the flavonoid accumulation patterns observed in most medicinal plants (Kuang *et al.*, 2020; Yuan *et al.*, 2022).

384 Numerous flavonoids isolated from Dai-Bai-Jie have exhibited significant biological activities.  
385 Specifically, hesperetin-7-O-glucoside has been demonstrated to effectively modulate the gut  
386 microbiota composition and bile acid metabolism in murine models (Wu *et al.*, 2022). The  
387 antioxidative, antihypertensive, antidiabetic, anti-inflammatory, and cardioprotective activities of  
388 rutin were reported, while rutin pretreatment before administration of ethanol can afford  
389 significant protection against mucosal hyperemia, necrosis, edema, and mucosal or submucosal  
390 hemorrhage (Akash *et al.*, 2024; Chua, 2013; Nicola *et al.*, 2024). Quercetin is known to  
391 possess both mast cell stabilizing and gastrointestinal cytoprotective activity (Anand David *et*  
392 *al.*, 2016; Catalina *et al.*, 2016).  
393 The flavonoid content in Dai-Bai-Jie varies significantly with its plantation age, which may be the  
394 result of DEGs patterns of genes involved in flavonoid biosynthesis. To date  
395 , flavonoid biosynthetic pathway has been extensively studied, with the genes encoding enzymes  
396 involved in this pathway and their respective functions having been verified in numerous plants.  
397 Flavonoids, flavonols, and lignin are synthesized through various branching pathways originating  
398 from the phenylpropane biosynthetic pathway (Froemel *et al.*, 1985). We screened nine DEGs  
399 related to flavonoid biosynthesis from Dai-Bai-Jie, *PAL*, *4CL*, *FLS*, and *C12RT1* includinged. *PAL*  
400 catalyzes the first step in the phenylpropanoid pathway and plays an important role in  
401 the biosynthesis of phenylpropanoid and flavonoid compounds (Levy *et al.*, 2018). *4CL* is the  
402 last enzyme in the general biosynthetic pathway of phenylpropane compounds, which catalyzes  
403 cinnamic acid and its hydroxyl or methoxy derivatives to generate corresponding coenzyme A  
404 esters (Cao *et al.*, 2023; Lavhale *et al.*, 2018). These intermediate products then enter the  
405 biosynthetic pathway of phenylpropane derivatives (Tian *et al.*, 2017). *FLS* is a key enzyme  
406 specific to the flavonol pathway, which converts dihydroflavonol into the corresponding flavonol  
407 by introducing a double bond between C-2 and C-3 of the C-ring (Forkmann *et al.*, 1986; Shi *et*  
408 *al.*, 2021).  
409 Correlation analysis conducted on flavonoid DAMs mapped to the KEGG pathway revealed that  
410 the expression patterns of the genes *PAL*, *4CL*, and *FLS* exhibited a consistent trend with the  
411 accumulation of nicotiflorin and lonicerin. Similarly, hesperetin-7-O-glucoside displayed a  
412 comparable trend with *C12RT1*. These DEGs may serve as key regulators of the distinct  
413 accumulation patterns of flavonoid metabolites in Dai-Bai-Jie.  
414 The RT-qPCR results showed that the expression trend of the key enzyme genes in the biosynthetic  
415 pathway of flavonoids in Dai-Bai-Jie werewas consistent with the results of transcriptome  
416 sequencing, thereby confirming the reliability of the transcriptome data.  
417 In general, the age of plantation has been shown to induce changes in soil nutrient content and  
418 pH, subsequently affecting the composition and diversity of soil bacterial and fungal communities.  
419 For instance, Na *et al.* (2016) reported that fungal diversity decreased with the cultivation going on  
420 from 5 a to 10 a of *Lycium barbarum* L. whereas bacterial diversity remained relatively unchanged.  
421 Conversely, Li *et al.* (2020) observed a significant increase in bacterial diversity and a decrease in  
422 fungal diversity in lily soil with increasing planting years. However, in our study on Dai-Bai-Jie,  
423 we did not detect any significant differences in the Shannon, Chao1, or ACE indices of rhizosphere  
424 microorganisms across different plantation ages and localities. This inconsistency suggests that  
425 the underlying mechanisms governing microbial community dynamics in the rhizospheres of Dai-  
426 Bai-Jie may differ from those observed in other plant species, possibly due to the relatively short  
427 introduction period of Dai-Bai-Jie.  
428 The absence of significant changes in microbial diversity warrants further investigation,  
429 particularly from the perspectives of soil nutrients, pH, and moisture content.

430 In summary, this study comprehensively characterized the disparities in flavonoid metabolite  
431 profiles and abundances across varying cultivation environments and plantation age through  
432 integrated transcriptome and metabolome analyses. Key genes intricately associated with the  
433 differential accumulation of flavonoids were identified. The results laid a foundation for further  
434 regulation of the effective components and support the formulation of scientifically harvesting  
435 practices for Dai-Bai-Jie.

## 436 **Conclusions**CONCLUSIONS

437 In summary, this study thoroughly characterised the disparities in -metabolites and flavonoid  
438 metabolite profiles and abundances across varying cultivation environments and plantation ages  
439 through integrated transcriptome and metabolome analyses. A total of 1,495 metabolites were  
440 identified using UPLC-MS/MS from Dai-Bai-Jie across three different planting durations (one  
441 year, two years, and three years) at two distinct localities. Among these, 943 DAMs were detected.  
442 A total of 114 flavonoids were identified, of which 79 exhibited differential accumulation. The  
443 total metabolite content in CR2 and CR3 was relatively abundant, and flavonoid levels were  
444 generally higher in CR2 and CR3. Therefore, it is recommended that harvesting at two years of  
445 age be considered the optimal strategy. Key genes intricately associated with the differential  
446 accumulation of flavonoids were identified. We found a complex regulatory relationship among  
447 phenylalanine ammonia-lyase (PAL; Cluster-63886.0, Cluster-63886.1), 4-Coumarate: Coenzyme  
448 A Ligase (4CL; Cluster-58688.4, Cluster-62808.3), flavonol synthase (FLS; Cluster-46899.18,  
449 Cluster-46899.5, Cluster-50957.2, Cluster-57391.0, C12RT1; Cluster-45854.0), and the  
450 metabolites hyperin, lonicerin, vicenin-2, nicotiflorin, quercetin, luteolin-7-O-(6"-malonyl)  
451 glucoside, and hesperetin-7-O-glucoside. Different planting ages and localities did not result in  
452 significant differences in the Shannon, Chao1, or ACE indices of the rhizosphere microorganisms  
453 associated with Dai-Bai-Jie. The results establish a foundation for further regulation of  
454 pharmacological components and provide support for the development of scientific harvesting  
455 practices for Dai-Bai-Jie.

## 456 **Acknowledgements**ACKNOWLEDGEMENTS

457 We acknowledge South Medicine Garden of Xishuangbanna Dai Autonomous Prefecture for  
458 providing the material collection site.

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460

## 461 **Funding Statement**

462 This study was financially supported by the the Xishuangbanna Prefecture Science and  
463 Technology Plan Project (No. 202401001) .

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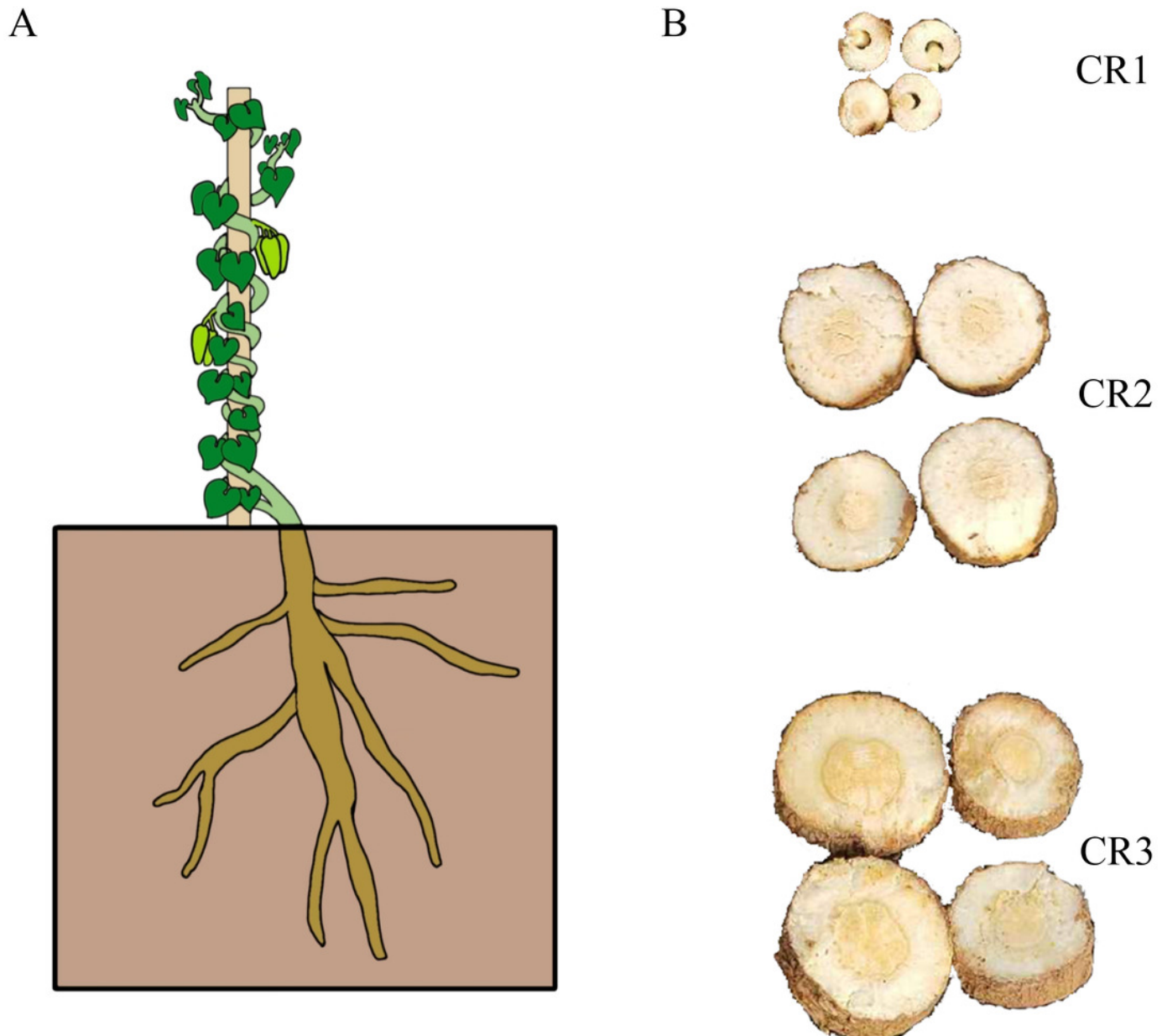
588 **Zou LQ, Wang CX, Kuang XJ, Li Y, Sun C. 2016.** Advance in flavonoids biosynthetic  
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590 DOI 10.4268/cjcm20162207.

591

# Figure 1

the sample used in this study

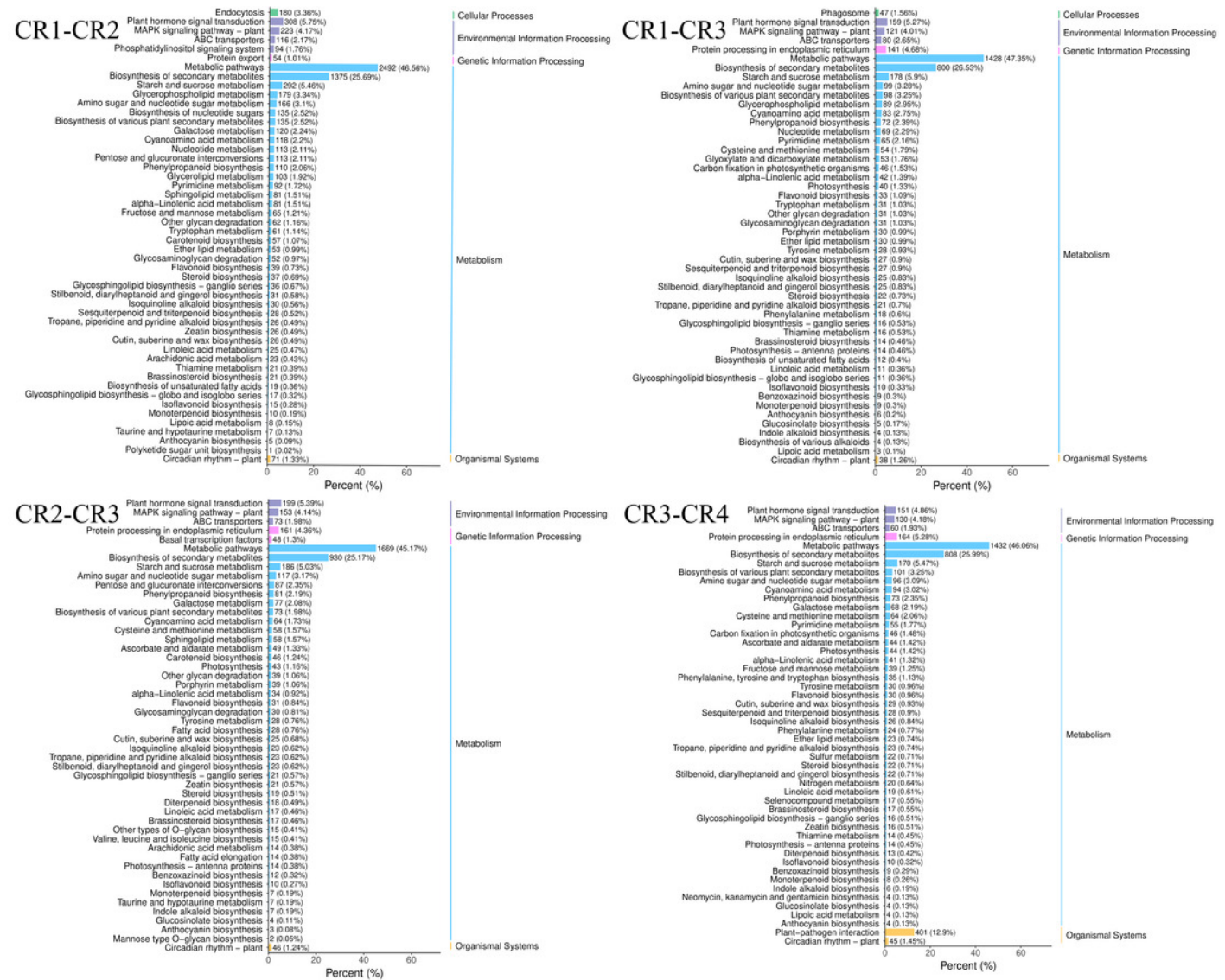
A, total plant of Dai-Bai-Jie. B, Root of cross-sections at different planting years. CR1: farmed for one year, CR2: farmed for two years, CR3, farmed for three years.



# Figure 2

The DEGs in the four groups were analyzed by KEGG metabolic pathway

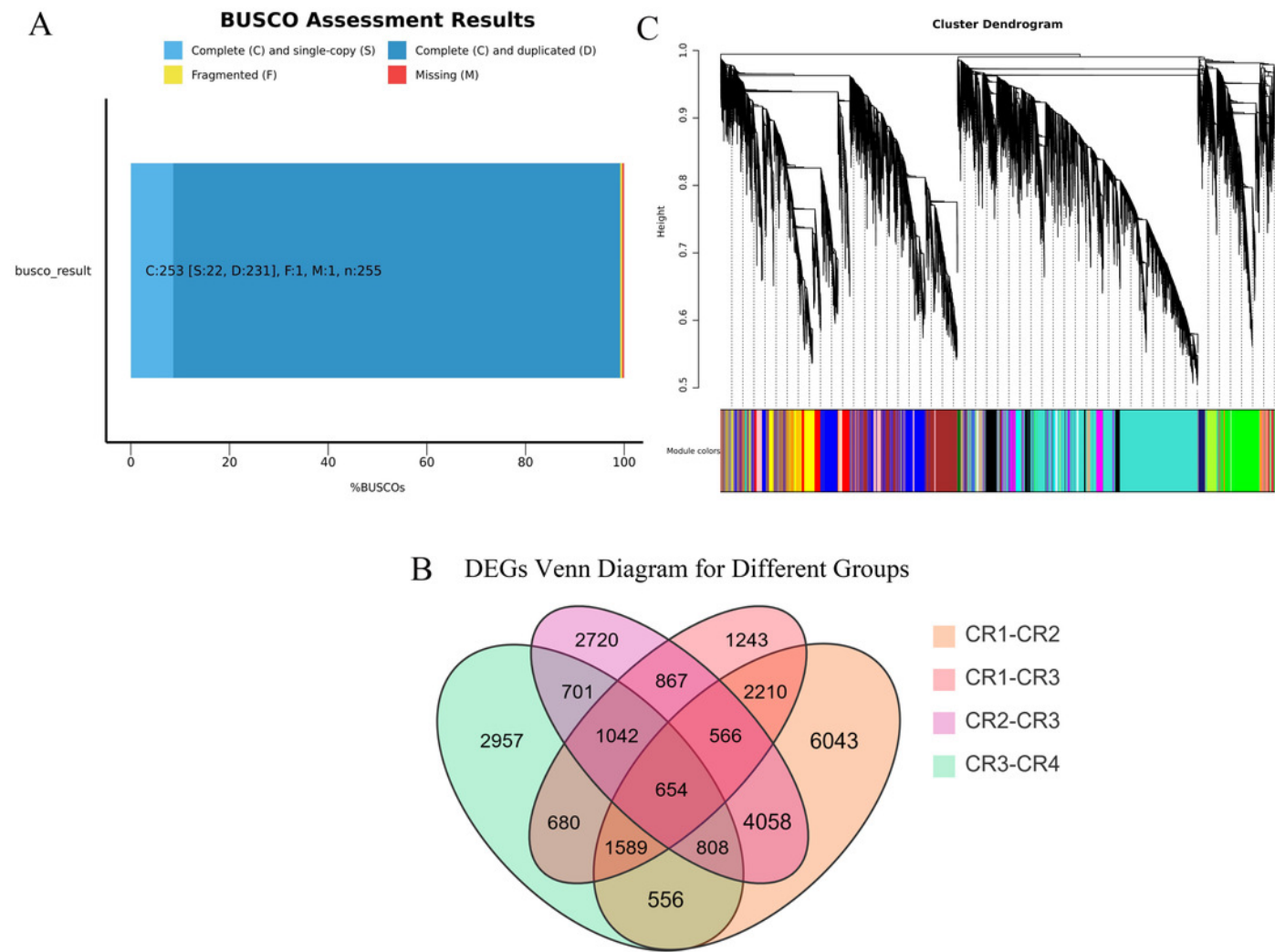
(A) CR1-CR2. (B) CR1-CR3. (C) CR2-CR3. (D) CR3-CR4.



# Figure 3

## Transcriptome analysis results

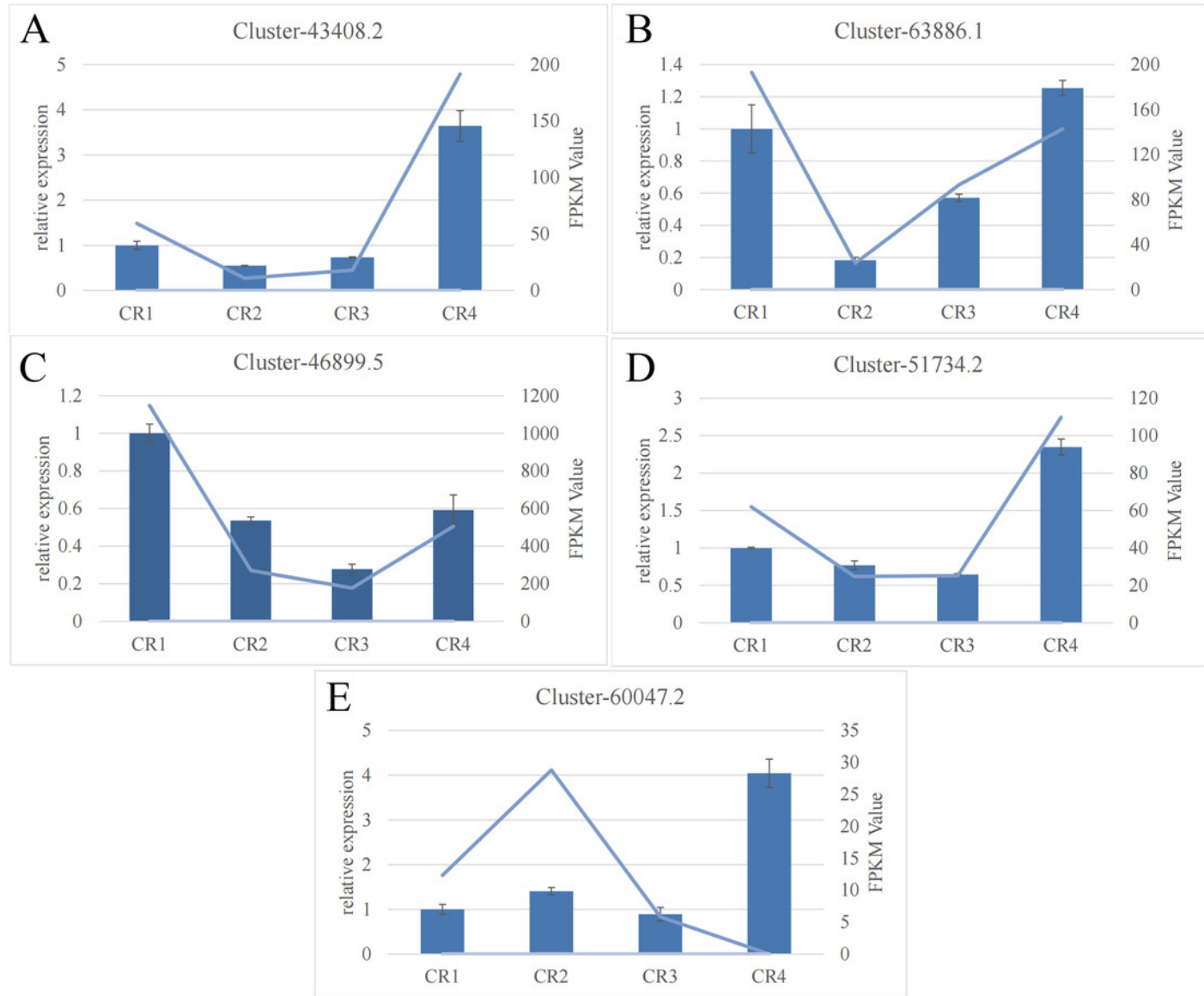
(A) BUSCO completeness assessments of the Dai-Bai-Jie transcriptome. (B) WGCNA clustering tree. (C) Venn Diagram representing the number of DEGs among four group sample.



## Figure 4

RNA-seq analysis of Dai-Bai-Jie and the qRT-PCR validation of five genes.

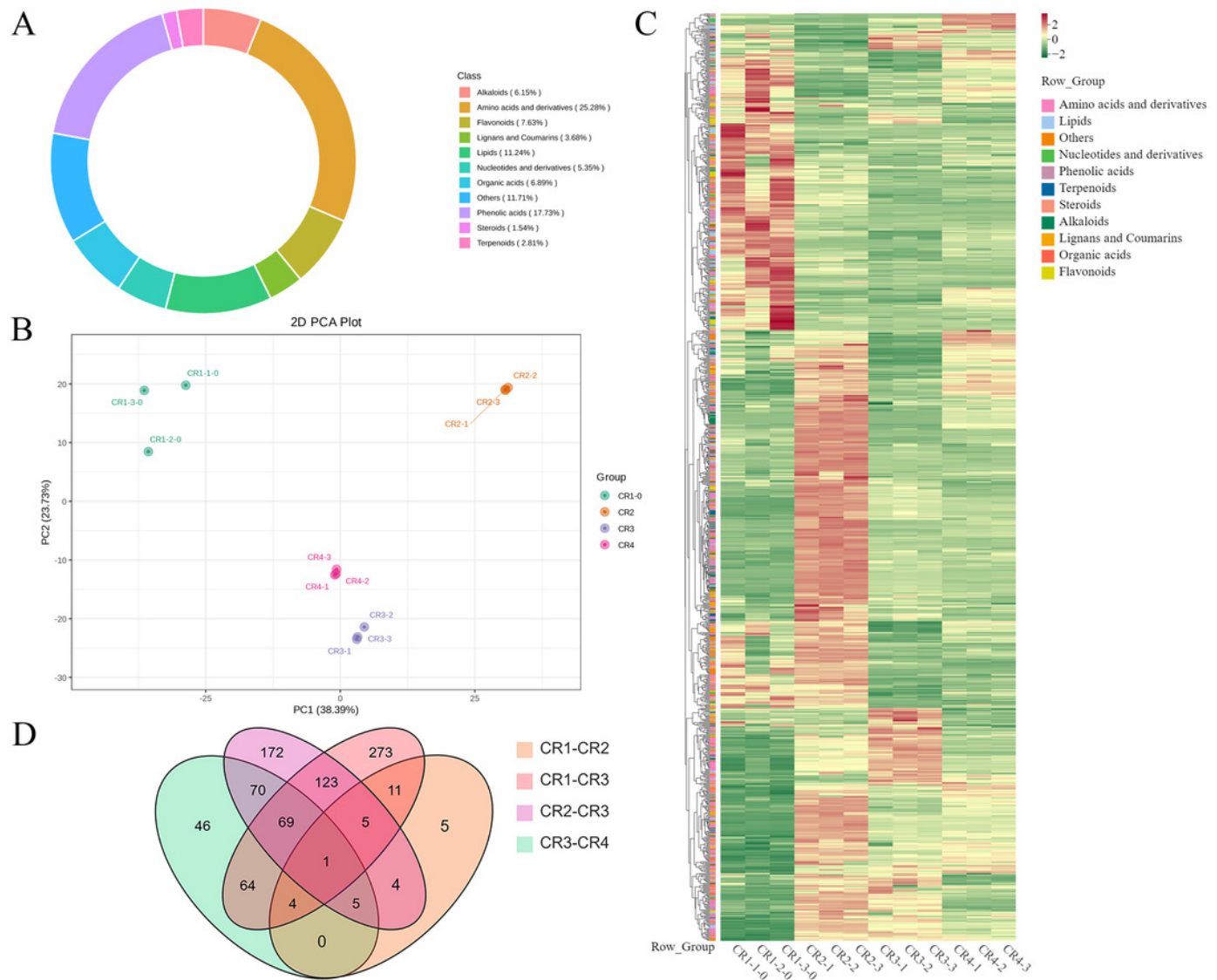
RNA-seq analysis of Dai-Bai-Jie and the qRT-PCR validation of five genes.



# Figure 5

## Metabolome analysis results

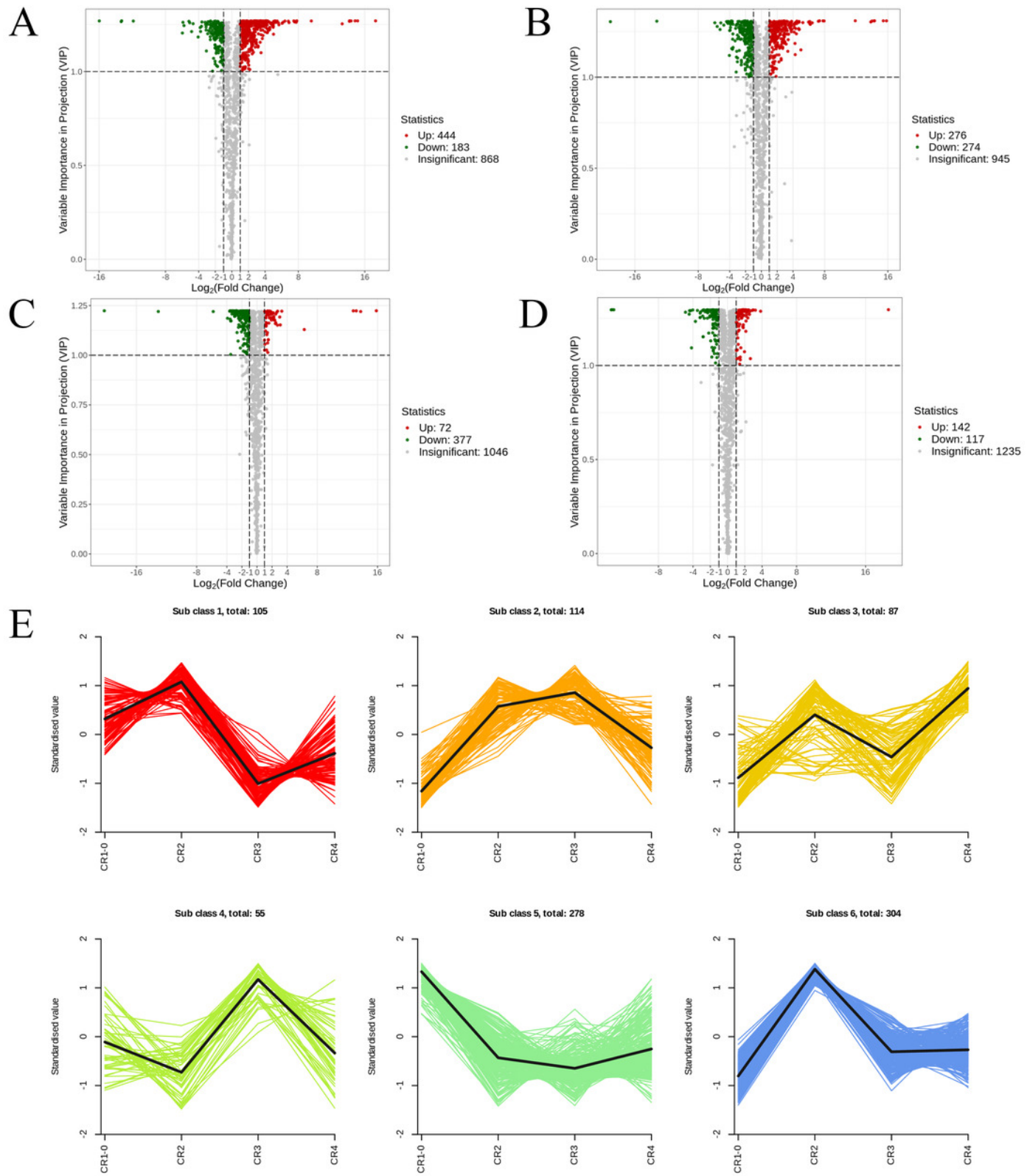
(A) Composition of metabolite in Dai-Bai-Jie. (B) PCA score plots for all samples. (C) Heat map of DAMs in four groups of samples. (D) Venn diagram of DAMs across groups.



## Figure 6

The volcano diagram and the k-means diagram of Metabolites.

(A) Volcano diagram of DAMs (CR1 vs. CR2). (B) Volcano diagram of DAMs (CR1 vs. CR3).  
(C) Volcano diagram of DAMs (CR2 vs. CR3). (D) Volcano diagram of DAMs (CR3 vs. CR4). (E)  
The K-means analysis of all Metabolites. The black line in the figure represents the average  
pattern of all Metabolites in each class, and different colors represent different trend.

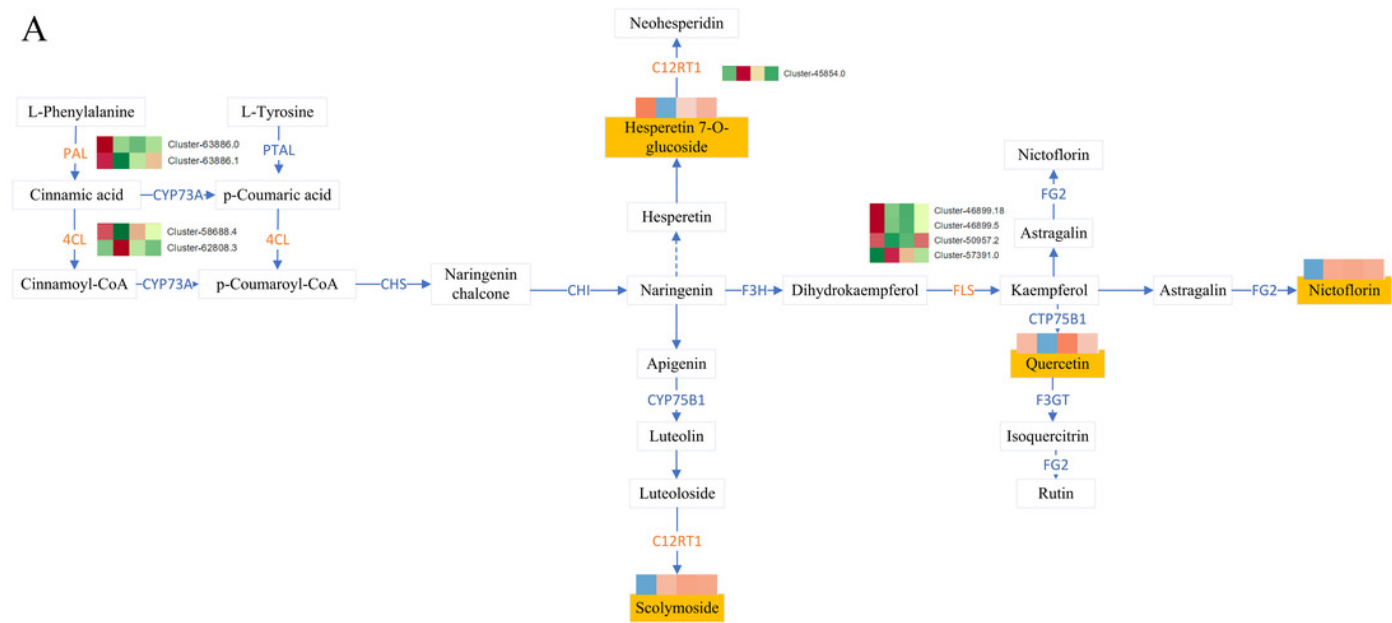


## Figure 7

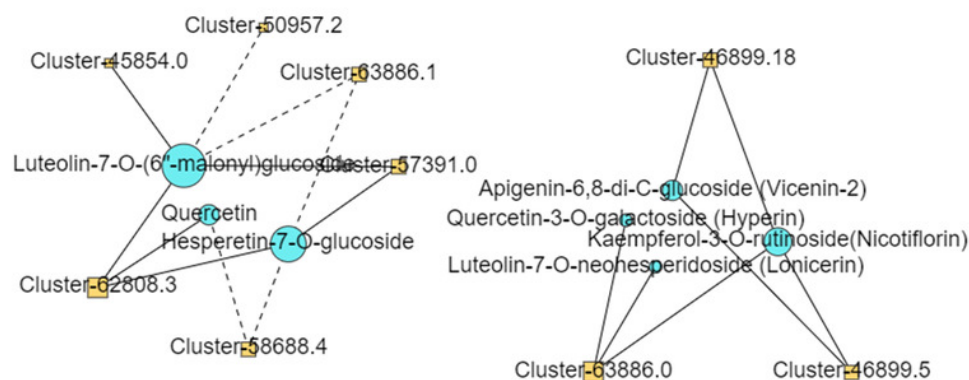
## Flavonoid synthesis pathway and Network diagram

(A) Flavonoid synthesis pathway. (B) Network diagram of flavonoids and differential genes.

A



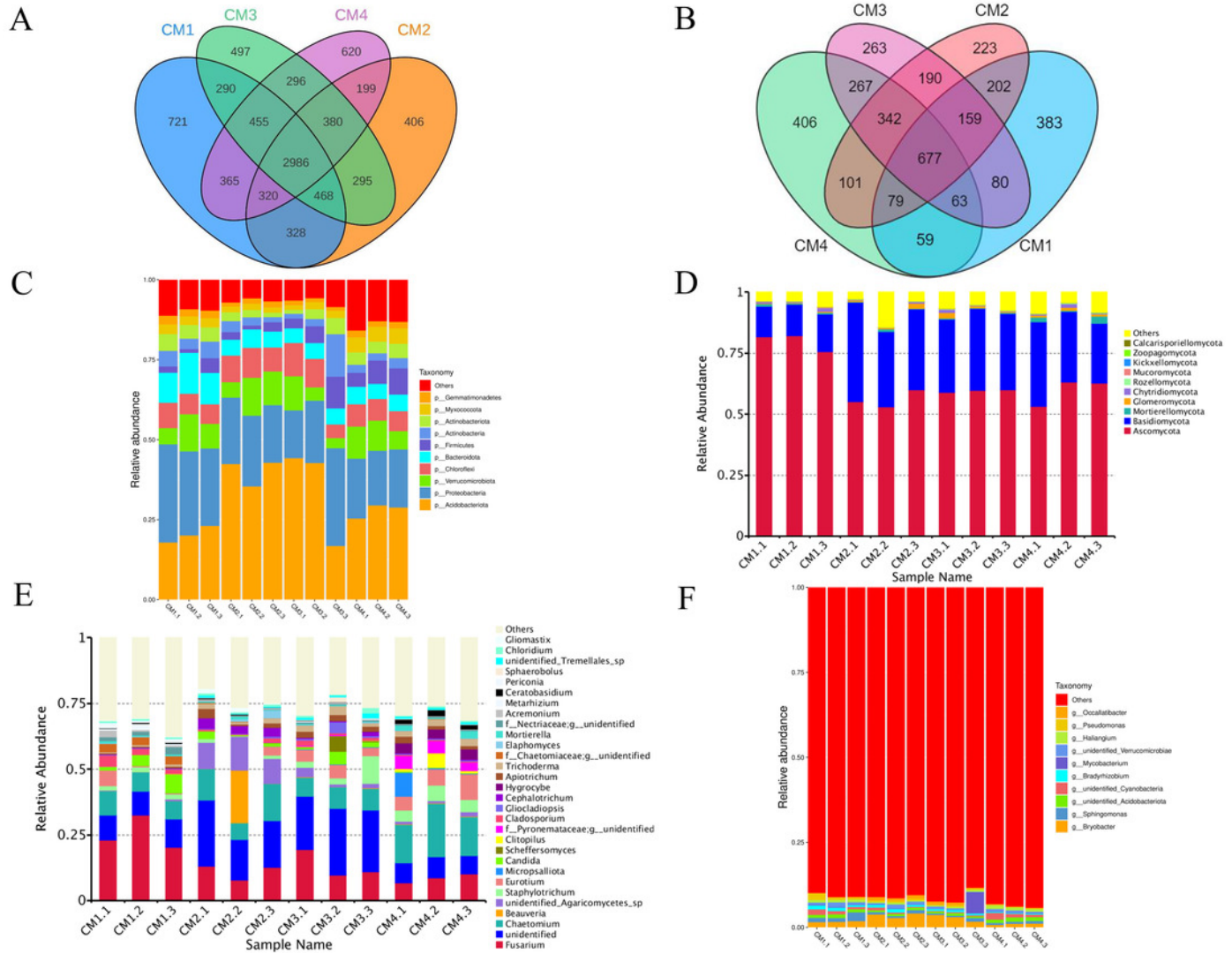
B



# Figure 8

Venn diagram and the relative abundance of phylum and genus among CM1, CM2, CM3, and CM4 in rhizosphere soil of Dai-Bai-Jie.

(A) Venn diagram of bacterial. (B) Venn diagram of fungus. (C) relative abundance of bacterial phylum. (D)relative abundance of fungal phylum. (E) relative abundance of bacterial genus. (F)relative abundance of fungal genus.



**Table 1**(on next page)

primer of Five genes

1  
2

Table 1 primer of Five genes

Gene	sequence (5'-3')	product size
Cluster-43408.2	F: TGATGAATGGGAAGCCCCGAG	175bp
FLS	R: TAGCGGTCCCTGTTTGGCTT	
Cluster-46899.5	F: AGCCCTTGAAGAATTGGTTGT	114bp
FLS	R: ATCTCTTGTAAAGGCCGATCAA	
Cluster-51734.2		166bp
CYP73A	F: GGACCTGGCTAAGGAAGTGT R: TGTGAAGAAAGGCACCGTCA	
Cluster-60047.2	F: GCATCCGTGGCGATCAAATC	179bp
4CL	R: TGCCACTTGGAACCCTTGT	
Cluster-63886.1	F: CATGCCCTCCTCAACAAACGA	171bp
PAL	R: GGACCTGCACTCCTGATCC	
GAPDH	F: GGCATTGTCGAGGGTCTCAT R: CCGGTGCTGCTGGAAATAAT	131bp

3

**Table 2**(on next page)

Diversity index of microbial communities in roots soils

1  
2  
3

4 Table 2 Diversity index of microbial communities in roots soils (mean  $\pm$  SD, n = 3)

	Sample	Shannon	Chao1	ACE	Goods_coverage
16s	CM1	10.275 $\pm$ 0.133	4844.719 $\pm$ 755.638	4912.204 $\pm$ 744.152	0.972 $\pm$ 0.006
	CM2	9.546 $\pm$ 0.128	4449.788 $\pm$ 173.103	4563.577 $\pm$ 259.590	0.972 $\pm$ 0.002
	CM3	9.507 $\pm$ 0.159	4580.545 $\pm$ 122.909	4696.261 $\pm$ 115.363	0.972 $\pm$ 0.002
	CM4	10.039 $\pm$ 0.067	4857.131 $\pm$ 150.655	4936.003 $\pm$ 163.463	0.971 $\pm$ 0.002
ITS	CM1	5.894 $\pm$ 0.317	1207.431 $\pm$ 60.164	1227.965 $\pm$ 58.934	0.997 $\pm$ 0.001
	CM2	5.267 $\pm$ 0.459	1152.941 $\pm$ 259.092	1185.874 $\pm$ 266.003	0.997 $\pm$ 0.001
	CM3	5.722 $\pm$ 0.276	1408.72 $\pm$ 171.792	1445.749 $\pm$ 164.948	0.996 $\pm$ 0.001
	CM4	6.320 $\pm$ 0.133	1424.764 $\pm$ 70.520	1449.286 $\pm$ 80.612	0.996 $\pm$ 0.001