

1 Running title: Diterpene synthases in *Isodon rubescens*

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22 L.D., H.Z. carried the experiments; J.T., Y.S. and H.L. coordinated the preparation of the
23 article; all authors contributed to the analysis of the collected data and writing of the article.

Functional diversification of kaurene synthase-like genes

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Summary:

The Lamiaceae family contains numerous diterpenoids, offering a valuable model system for the study of diterpenoid chemical diversity. In this study we report that the *ent*-kaurene diterpenoids in *Isodon rubescens* have the largest diTPS gene family identified in Lamiaceae. Three genes are predicted to be involved in *ent*-kaurene diterpenoid synthesis while another two KSL genes are functional enzymes utilizing *ent*-CPP as a substrate. We also showed the presence of a normal-CPP mediated biosynthesis pathway in *I. rubescens* and that three KSL genes possessing different domain architectures are involved in this pathway. This study provides new information regarding the understanding of diterpenoid chemical diversity and this compounds' evolution with in the Lamiaceae family.

48 **ABSTRACT**

49 *Ent*-kaurene diterpenoids are the largest group of known *Isodon* diterpenoids. Among them,
50 oridonin is accumulated in the leaves, and is the most frequently studied compound because of its
51 anti-tumor and anti-bacterial activities. We have identified five copalyl diphosphate synthase (CPS)
52 and six kaurene synthase-like (KSL) genes by transcriptome profiling of *I. rubescens* leaves. An *in*
53 *vitro* assay assigns ten of them to five different diterpene biosynthesis pathways, except IrCPS3
54 that has a mutation in the catalytic motif. The Lamiaceae-specific clade genes (IrCPS1 and IrCPS2)
55 synthesize the intermediate copalyl diphosphate (normal-CPP), while IrCPS4 and IrCPS5
56 synthesize the intermediate *ent*-copalyl diphosphate (*ent*-CPP). IrKSL2, IrKSL4 and IrKSL5 react
57 with *ent*-CPP to produce an *ent*-isopimaradiene-like compound, *ent*-atiserene and *ent*-kaurene,
58 respectively. Correspondingly, the Lamiaceae-specific clade genes IrKSL1 or IrKSL3 combined
59 with normal-CPP led to the formation of miltiradiene. The compound then underwent
60 aromatization and oxidization with a cytochrome P450 (CYP76AH30) forming two related
61 compounds, abietatriene and ferruginol, which were detected in the root bark. IrKSL6 reacts with
62 normal-CPP to produce isopimaradiene. IrKSL3 and IrKSL6 have the $\gamma\beta\alpha$ tri-domain structure, as
63 these proteins tend to possess the bi-domain structure of IrKSL1, highlighting the evolutionary
64 history of KSL gene domain loss and further elucidating chemical diversity evolution from a
65 macroevolutionary stance in Lamiaceae.

66 INTRODUCTION

67 The mint family (Lamiaceae) is the sixth largest family of flowering plants and one of the most
68 economically valuable natural products for foods and toiletries. The Lamiaceae family produces a
69 unique class of terpenoids called diterpenoids. The most common diterpenoids in Lamiaceae are
70 abietanes, labdanes, neoclerodanes, primaranes and *ent*-kauranes. These are taxa-specific
71 compounds that have been found in six of the seven subfamilies (Harley et al., 2004). The
72 diterpenoid skeleton, abietane, has the highest occurrence and is the most widespread among
73 Lamiaceae (Vestri Alvarenga et al., 2001), but is characteristic of the Nepetoideae subfamily.
74 *Ent*-kauranes are characteristic of the *Isodon* genus (Sun et al., 2006) and labdanes are
75 characteristic of many genera in the subfamily Lamioideae (Harley et al., 2004). These diterpenoids
76 fall into the labdane-related superfamily, which evolved from the same machinery involved in
77 gibberellin (GA) metabolism (Peters, 2010). Biosynthesis is initiated by two sequential diterpene
78 synthases (diTPS) that catalyze cyclization and/or rearrangement of the general acyclic precursor
79 (*E,E,E*)-geranylgeranyl diphosphate (GGPP) to form different requisite skeletal structures (Zi et al.,
80 2014). The first step is a protonation-initiated cyclization of GGPP by the class II diTPS copalyl
81 diphosphate synthase (CPS) that leads to the formation of a bicyclic labdadienyl/copalyl
82 diphosphate (CPP) with a specific stereochemistry (*ent*, *syn*, normal / (+)). CPP is further cyclized
83 and/or rearranged in a diphosphate ionization-initiated reaction catalyzed by a class I diTPS,
84 kaurene synthase-like cyclase (KSL), which is usually specific to a particular CPP stereoisomer.

85 The diterpenoid biosynthesis has been explored in nine species in Lamiaceae, including *Salvia*
86 *miltiorrhiza* (Gao et al., 2009; Cui et al., 2015; Su et al., 2016), *S. sclarea* (Caniard et al., 2012;
87 Schalk et al., 2012), *Isodon eriocalyx* (Li et al., 2012), *Coleus forskohlii* (Pateraki et al., 2014),
88 *Rosmarinus officinalis* (Bruckner et al., 2014), *S. fruticosa* (Božić et al., 2015), *S. pomifera* (Triikka et
89 al., 2015), and *S. divinorum* (Pelot et al., 2017) from the biggest subfamily Nepetoideae, and
90 *Marrubium vulgare* (Zerbe et al., 2014) from the subfamily Lamioideae. These enzymes have the
91 ability to produce abietane, labdane, clerodane, and *ent*-kaurane diterpenes via an enzymatic
92 hydrolysis catalyzed reaction (Gao et al., 2009; Cui et al., 2015; Caniard et al., 2012; Schalk et al.,
93 2012; Li et al., 2012; Pateraki et al., 2014; Zerbe et al., 2014; Bruckner et al., 2014; Božić et al.,
94 2015; Triikka et al., 2015; Pelot et al., 2017). Based on the stereochemistry of the intermediate that
95 is initially cyclized by CPS, these enzymes can be divided into two groups. One group produces an

intermediate of normal copalyl diphosphate (CPP), known as normal-CPP, which has been found to be the precursor of a large array of diterpenoids with high economic and medicinal value, including tanshinone (Gao et al., 2009; Cui et al., 2015), sclareol (Caniard et al., 2012; Schalk et al., 2012), and forskolin (Pateraki et al., 2014). The other group produces the stereoisomeric intermediate of *ent*-CPP that is involved in GA biosynthesis as well as the synthesis of labdane or *ent*-kaurane in *S. miltiorrhiza* and *I. eriocalyx* (Li et al., 2012; Cui et al., 2015). Phylogenetic analysis has shown that these enzymes are grouped in accordance with their functions except the newly identified SdCPS1 and SdKSL1 from *S. divinorum* (Pelot et al., 2017), that is, those involved in normal-CPP biosynthesis form a sister group to those involved in *ent*-CPP biosynthesis, and positive selection has played an important role in this functional divergence (Cui et al., 2015). A similar divergence has also been observed in class I KSL enzymes that are responsible for the subsequent transformation of normal-CPP synthase products. Many of these enzymes have lost the characteristic 'internal/ γ ' domain that is responsible for metabolite specialization, a deletion event found in KSLs in Lamiaceae (Hillwig et al., 2011; Caniard et al., 2012; Schalk et al., 2012; Bruckner et al., 2014; Pateraki et al., 2014; Zerbe et al., 2014; Cui et al., 2015; Božić et al., 2015; Triikka et al., 2015). A recent work has shown the plasticity of class I KSLs, such as SsScS from *S. sclarea*, that can react with any available CPS product (Andersen-Ranberg et al., 2016; Jia et al., 2016). These studies demonstrate the value of Lamiaceae as a useful system to study diterpenoid chemical diversity evolution. While most recent works focus on specific biosynthesis pathways in given species, little is known about the diversified diterpenoids' evolutionary origin.

Here we report a large diTPS gene family from *I. rubescens* (Tribe: Ocimeae, Subfamily: Nepetoideae, Fig. 1A) composed of many *ent*-kaurane diterpenoids (Sun et al., 2006). One of these, oridonin (Fig. 1B), is found in the leaves of many *Isodon* plants and is considered to be a potential cancer chemoprevention agent with broad spectrum anti-tumor and anti-bacterial activities *in vitro* and *in vivo* (Zhou et al., 2007). In contrast to the numerous *ent*-kaurane diterpenoids, rubesanolides A-E, with only five abietane-related compounds, has been isolated from the leaves of *I. rubescens* (Zou et al., 2011, 2012). Rubesanolide D (Fig. 1C) has shown inhibition activity against biofilm formation of the dental bacterium *Streptococcus mutans* (Zou et al., 2012). Although seven diTPS genes have been reported from an *I. rubescens* transcriptome sequencing project (Zerbe et al., 2013), none of them were identified. Here we report 11 diTPS genes from a transcriptome

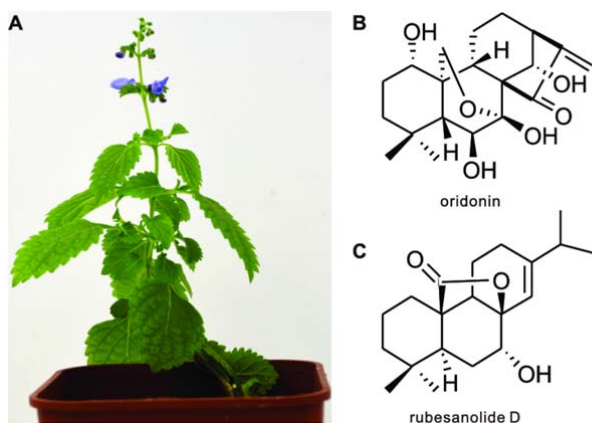


Figure 1. The major medicinally active diterpenoids in *I. rubescens*. A, The plant of *I. rubescens*. B, The typical *ent*-kaurene diterpenoid of oridonin. C, The typical abietane diterpenoid of rubesanolide D.

126 sequencing project. We identified *IrCPS4* that is predicted to be involved in oridonin biosynthesis,
 127 three KSL genes involved in *ent*-CPP interactions, and a normal-CPP mediated multiradiene and
 128 isopimaradiene biosynthesis pathway. The three KSL genes bearing different domain structures are
 129 involved in the normal-CPP mediated biosynthesis, revealing information regarding the
 130 evolutionary origin of the γ domain loss and further shedding light on the macroevolution of
 131 diterpenoid chemical diversity in Lamiaceae.
 132

133 Results

134 Transcriptome sequencing and *de novo* assembly

135 A non-normalized complementary DNA (cDNA) library was prepared for transcriptome
136 sequencing using the Illumina Hiseq2500 platform from leaf tissues of *I. rubescens* containing
137 the highest oridonin content (Committee CP, 2015). A total of 27,248,946 reads were obtained. The
138 Trinity *de novo* assembler (Grabherr et al., 2011) found 60,483 contigs with an average of 886 bp;
139 contig length varied from 201 to 12,118 bp (Supplemental Table S1).

140 Annotation of diTPS family in *I. rubescens*

141 All contigs were aligned to the GenBank non-redundant protein sequence (NR) database using
142 BlastX with e-values $< 1e^{-5}$. Twenty-nine unigenes were annotated as copalyl diphosphate
143 synthase or kaurene synthase-like cyclase genes (Supplemental Table S2). After manual
144 assembly using known diTPS sequences from *I. eriocalyx*, *S. miltiorrhiza*, and a previous
145 reported *I. rubescens* transcriptome sequencing dataset (Zerbe et al., 2013), we found five
146 putative diTPS with full length cDNA, annotated as *IrCPS1*, *IrCPS3*, *IrCPS4*, *IrKSL1* and
147 *IrKSL3* due to their homology to diTPS in *S. miltiorrhiza* (Supplemental Fig. S1). Together with
148 5'- and 3'-rapid amplification of cDNA ends (RACE), six other full length diTPS were identified
149 and annotated as *IrCPS2*, *IrCPS5*, *IrKSL2*, *IrKSL4*, *IrKSL5* and *IrKSL6*. In total we established
150 five CPS and six KSL genes from *I. rubescens* (Supplemental Table S3) which comprises 28
151 unigenes (does not include c40624_g1_i1). Among these 11 genes, three of them, *IrCPS1*,
152 *IrKSL1* and *IrKSL3* have corresponding full length sequence in the reported dataset (Zerbe et al.,
153 2013) with 98%~99% identities. Four of them, *IrCPS3*, *IrCPS4*, *IrCPS5* and *IrKSL6* have
154 corresponding sequences with 99% identity, and the other four genes have sequences with
155 identities lower than 93% (Supplemental Table S4).

156 The deduced amino acid sequences of the four putative CPS genes (*IrCPS1*, *IrCPS2*, *IrCPS4* and
157 *IrCPS5*) contain the '(D,E)XDD' motif typical of CPS enzymes (Prisic et al., 2004), while the six
158 putative KSL genes have the 'DDXXD' motif that is characteristic of KSL enzymes (Christianson,
159 2006) (Supplemental Fig. S1). In *IrCPS3*, the second Asp of the DXDD motif (corresponding to
160 D379 in AtCPS from *Arabidopsis thaliana*) is substituted by Asn (Fig. 2), The Asp amino acid has
161 been reported to be essential in forming hydrogen bonded "proton wires" during catalysis (Koksal
162 et al., 2014).

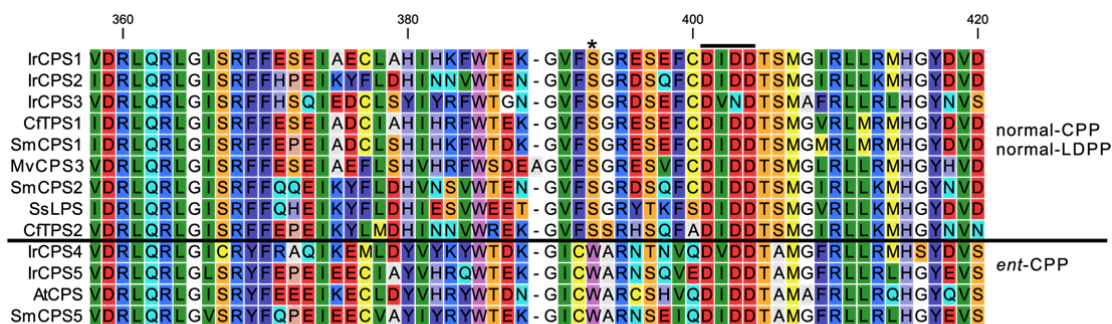


Figure 2. Alignment of the conserved DXDD motif and the positive selection site (Ser-362 / Trp-364) of CPS from *I. rubescens* and other characterized CPSs. The DXDD motif is labeled above the alignment with a solid line, and the conserved Ser or Trp position is marked with an asterisk. The second Asp of the DXDD motif is substituted by Asn in IrCPS3. The enzymes are separated by the horizontal line according to normal-CPP/LDPP or *ent*-CPP stereochemistry. The CPS in the upper section (including IrCPS1 and IrCPS2 from this study) were characterized as being involved in normal-CPP/LDPP formation while those in the lower section (including IrCPS4 and IrCPS5 from this study) were characterized to be involved in *ent*-CPP production. CftPS1 and CftPS2 are from *Coleus forskohlii*, SmCPS1, SmCPS2, SmCPS5 from *Salvia miltiorrhiza*, MvCPS3 from *Marrubium vulgare*, SsLPS from *S. sclarea*, and AtCPS from *Arabidopsis thaliana*.

163 The positive selection site Ser-362 (SmCPS1 in normal-CPP synthase) / Trp-364 (SmCPS5 in
164 *ent*-CPP synthase) has shown to be essential for formation of normal-CPP versus *ent*-CPP (Cui et
165 al., 2015). In *I. rubescens*, IrCPS1 and SmCPS2 have a Ser (S) at the 373 and 367 position, as do
166 the other six CPS from Lamiaceae (Fig. 2). This is necessary for proper enzymatic function to
167 produce normal -CPP or normal-LDPP using GGPP as a substrate (Schalk et al., 2012; Pateraki
168 et al., 2014; Zerbe et al., 2014; Cui et al., 2015). The other two genes, IrCPS4 and IrCPS5, have a
169 Trp (W) in this position (Fig. 2), similar to SmCPS5 from *S. miltiorrhiza* and AtCPS, whose
170 product is *ent*-CPP when using GGPP as a substrate (Cui et al., 2015).

171 **Phylogenetic relationship of diTPS within Lamiaceae**

172 A maximum likelihood (ML) tree was constructed with 96 diTPS enzymes with known functions
173 (Supplemental Table S5). The diTPS from *I. rubescens* cluster with other genes from Lamiaceae,
174 forming four clades. Two clades contain CPS and KSL that are involved in *ent*-CPP or its
175 hydroxylated LDPP-mediated biosynthesis while the other two clades contain CPS and KSL that
176 are involved in specialized normal-CPP mediated biosynthesis (Fig. 3).

177 IrCPS1, IrCPS2, and IrCPS3 are found in the Lamiaceae-specialized CPS clade (Fig. 3),
178 which also contains CftPS1/CftPS2 from *C. forskohlii* (Pateraki et al., 2014), SmCPS1/SmCPS2

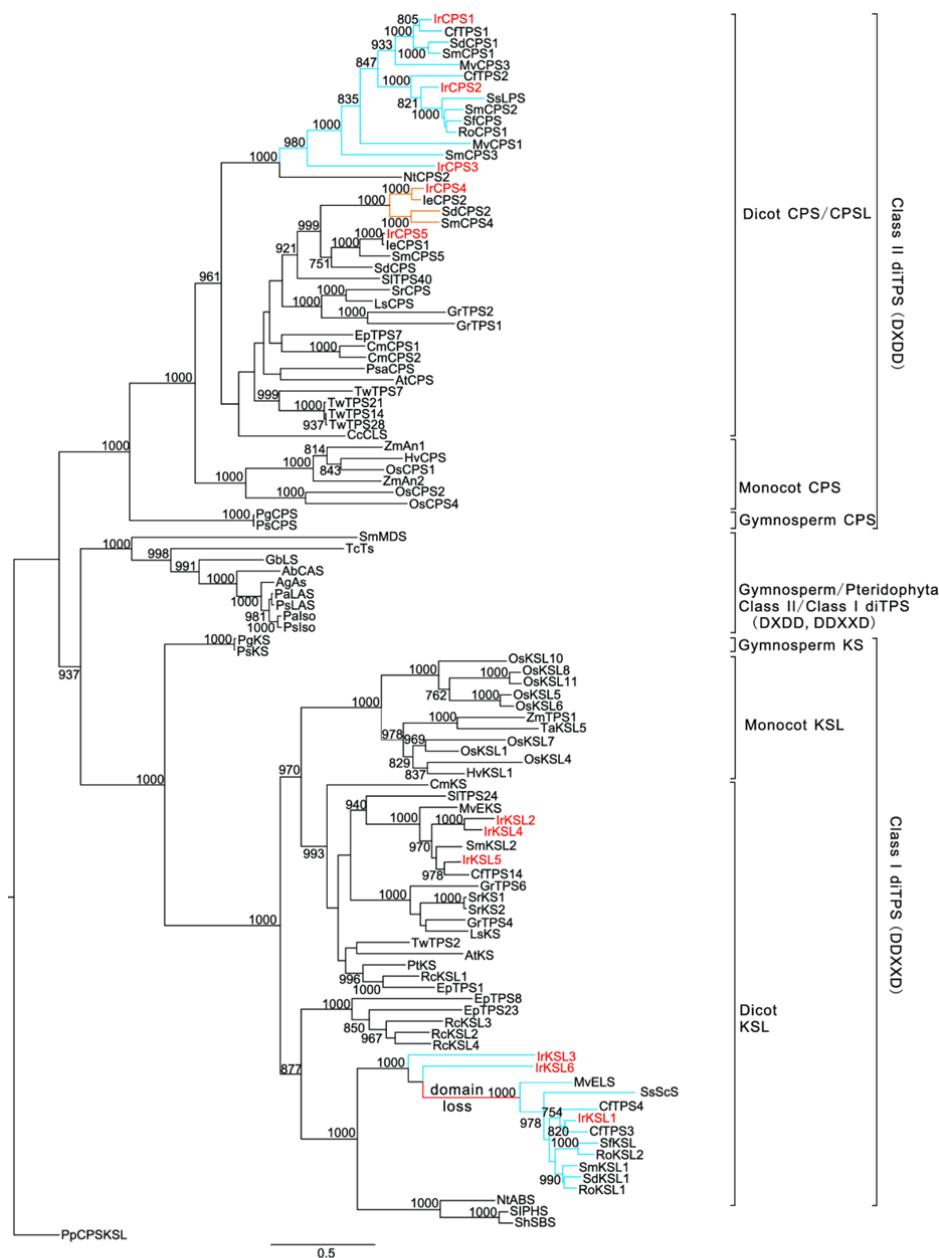


Figure 3. Phylogeny of *I. rubescens* diterpene synthases. The maximum likelihood tree illustrates the phylogenetic relationship of *I. rubescens* diterpene synthases with 96 representative characterized diTPS (Supplemental Table S5). Numbers on branches indicate the bootstrap percentage values calculated from 1000 bootstrap replicates. *Physcomitrella patens* CPS/kaurene (PpCPSKS) was used as an outgroup. Blue lines show diTPS genes from Lamiaceae involved in normal-CPP/LDPP-mediated diterpenoid metabolism and red lines show the loss of the N-terminal γ domain found in eudicot and monocot KSLs. Yellow lines show genes from Lamiaceae involved in the specialized *ent*-CPP/LDPP-related diterpenoid metabolism. Red-marked enzymes show diTPS from *I. rubescens* in this study.

- 179 from *S. miltiorrhiza* (Cui et al., 2015), MvCPS1/MvCPS3 from *Marrubium vulgare* (Zerbe et al.,
 180 2014), RoCPS1 from *R. officinalis* (Bruckner et al., 2014), SsLPS from *S. sclarea* (Schalk et al.,
 181 2012), SfCPS from *S. fruticosa* (Božić et al., 2015) and SdCPS1 from *S. divinorum* (Pelot et al., 2017).

182 The enzymes mentioned above have catalytic activity with GGPP to form normal-CPP and the
183 stereochemically related normal-LDPP (Schalk et al., 2012; Bruckner et al., 2014; Pateraki et al.,
184 2014; Zerbe et al., 2014; Cui et al., 2015; Božić et al., 2015) with the exception of SdCPS1 (Pelot et
185 al., 2017). IrCPS4 and IrCPS5 are found in the *ent*-CPP synthase clade. IrCPS4 is closely related to
186 an *ent*-CPS from *I. eriocalyx* (IeCPS2), which is predicted to be involved in the biosynthesis of
187 *Isodon ent*-kaurene diterpenoids (Li et al., 2012). IrCPS5 is closely related to IeCPS1 and SmCPS5
188 that are involved in the GA biosynthetic pathway (Fig. 3).

189 IrKSL1 is grouped in the specialized Lamiaceae KSL clade, having the less common $\beta\alpha$
190 bi-domain architecture through the loss of the γ domain at the N-terminus (Fig. 3). These enzymes
191 combine with the Lamiaceae-specialized CPS clade to form intermediates like miltiradiene,
192 (13R)-manoyl oxide and sclareol which are precursors to a large array of pharmaceutic compounds
193 (Schalk et al., 2012; Bruckner et al., 2014; Pateraki et al., 2014; Zerbe et al., 2014; Cui et al., 2015;
194 Božić et al., 2015). It is interesting to note that IrKSL3 and IrKSL6 form a monophyletic clade with
195 IrKSL1 despite the fact that they have the common $\gamma\beta\alpha$ tri-domain structures. IrKSL2, IrKSL4 and
196 IrKSL5 belong to the KSLs that are involved in the *ent*-CPP mediated biosynthetic pathway that
197 catalyzes reactions with *ent*-CPP and *ent*-LDPP (Fig. 3).

198 **Functional analysis of *ent*-CPP mediated diterpene synthesis**

199 Consistent with its closer relationship to other dicot CPS and KSL genes involved in *ent*-CPP
200 mediated GA or other related biosynthesis, IrCPS4 and IrCPS5 were found to produce *ent*-CPP.
201 We first incubated IrCPS4 and IrCPS5 with GGPP alone for comparison with a known enzyme
202 SmCPS5 (*ent*-CPP synthase) from *S. miltiorrhiza*. IrCPS4 and IrCPS5 yielded a diterpene with
203 identical retention time and mass spectrum to the product of SmCPS5 after dephosphorylation,
204 (Supplemental Fig. S2A-B) illustrating that the primary product of IrCPS4 and IrCPS5 is copalyl
205 diphosphate. The stereochemistry of IrCPS4 and IrCPS5 was further investigated by reaction
206 assays of each protein with SmKSL1 (specific to normal-CPP) and AtKS (specific to *ent*-CPP).
207 Reactions with IrCPS4 or IrCPS5 coupled with AtKS resulted in the formation of *ent*-kaurene,
208 while no product was obtained with SmKSL1 (Supplemental Fig. S2A). This illustrates that IrCPS4
209 and IrCPS5 are *ent*-copalyl diphosphate synthase and are predicted to be involved in oridonin or
210 GA biosynthesis in *I. rubescens*.

211 The catalytic activity of IrKSL2, IrKSL4 and IrKSL5 were each analyzed by incubation with

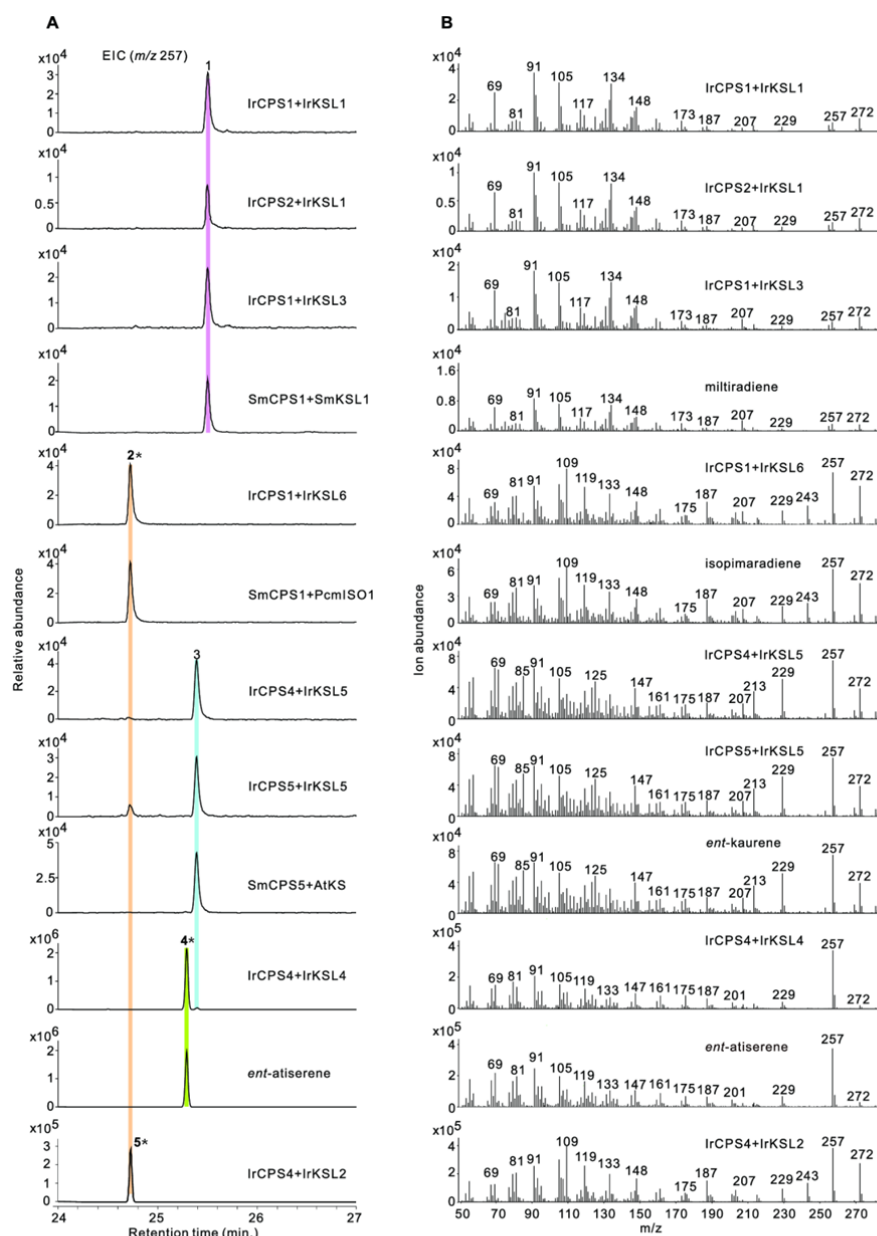


Figure 4. GC-MS analysis of *in vitro* assays with *I. rubescens* diTPS on a Cyclodex- β GC column. A, Extracted ion chromatograms (EIC) of *m/z* 257 of *in vitro* assays with IrCPS coupled with different IrKSL. The characterized SmCPS5 (*ent*-CPP synthase from *S. miltiorrhiza*), SmCPS1 (normal-CPP synthase from *S. miltiorrhiza*), SmKSL1 (multiradiene synthase from *S. miltiorrhiza*), AtKS (*ent*-kaurene synthase from *A. thaliana*) and PcmISO1 (pimaradiene from *Pinus banksiana*) were used as positive controls, along with the corresponding authentic standard *ent*-atiserene. 1. multiradiene; 2. isopimaradiene; 3. *ent*-kaurene; 4. *ent*-atiserene; 5. *ent*-isopimaradiene like compound. * indicates the unexpected enzyme products identified in this study. B, Corresponding mass spectra of recombinant enzyme assay products and authentic standard.

212 IrCPS4 that was identified to produce *ent*-CPP. All enzymes were found to react with *ent*-CPP.
 213 Specifically, IrKSL2 generated a product with identical EI mass spectrum to isopimaradiene
 214 (isopimara-7,15-diene, Hall et al., 2013), IrKSL4 generated a main product of *ent*-atiserene (95%),

215 and a minor product of *ent*-kaurene (5%), while IrKSL5 produced *ent*-kaurene (Fig. 4). At the same
216 time, these enzymes also produced the same product when combined with IrCPS5 (Supplemental
217 Fig. S3).

218 Many enzymes require copalyl diphosphate (normal-CPP) as a substrate including
219 Isopimara-7,15-diene synthase (PaTPS-Iso) from *Picea abies* (Martin et al., 2004) and the later
220 identified PbmISO1 and PcmISO1 from *Pinus banksiana* and *P. contorta* (Hall et al., 2013). Thus,
221 the stereochemistry of the IrKSL2 product, isopimaradiene, was further compared with PcmISO1.
222 As expected, isopimaradiene was produced when combining PcmISO1 with SmCPS1 (normal-CPP)
223 and has identical retention time with the product of IrCPS4 and IrKSL2 (Fig. 4). No product was
224 observed when combining PcmISO1 with IrCPS4. Therefore, the product of IrCPS4 and IrKSL2
225 appears to have a different stereochemistry compared to isopimaradiene, however, its absolute
226 configuration needs further identification.

227 **Functional analysis of normal-CPP-mediated diterpene synthesis**

228 Subsequently, we analyzed the function of the Lamiaceae-specialized clade gene products IrCPS1,
229 IrCPS2, IrKSL1, IrKSL3 and IrKSL6 *in vitro*. IrCPS1 and IrCPS2, in combination with IrKSL1 or
230 IrKSL3, formed miltiradiene as identified by comparison to the product of the combined activity of
231 normal-CPP synthase (SmCPS1) and miltiradiene synthase (SmKSL1) from *S. miltiorrhiza* (Fig. 4).
232 No product was observed when combining IrCPS1 and IrCPS2 with AtKS (specific to *ent*-CPP)
233 providing further evidence the product of IrCPS1 and IrCPS2 was normal-CPP.

234 An unexpected result was observed when combining IrCPS1 or IrCPS2 with IrKSL6. The
235 product has the same retention time and EI mass spectrum as the product of PcmISO1 and SmCPS1
236 (normal-CPP) (Fig.4; Supplemental Fig. S3). Thus, IrKSL6 can react with normal-CPP to produce
237 isopimaradiene in *I. rubescens*.

238 **Expression analysis of the identified diTPS**

239 To understand the physiological roles of the candidate diTPS in *I. rubescens*, we used quantitative
240 reverse transcription PCR (qRT-PCR) to evaluate mRNA transcript levels in various organs,
241 including root, stem, leaf, and flower. Transcripts of *IrCPS1* and *IrKSL1* were detected at
242 extremely high levels in the root (Fig. 5), a similar finding to the expression patterns of *SmCPS1*
243 and *SmKSL1* in *S. miltiorrhiza* (Cui et al., 2015). *IrCPS2*, *IrCPS5*, *IrKLS5* and *IrKSL6* were found
244 to have relatively ubiquitous expression in all sampled organ tissues. *IrCPS4*, *IrKSL2*, *IrKSL3* and

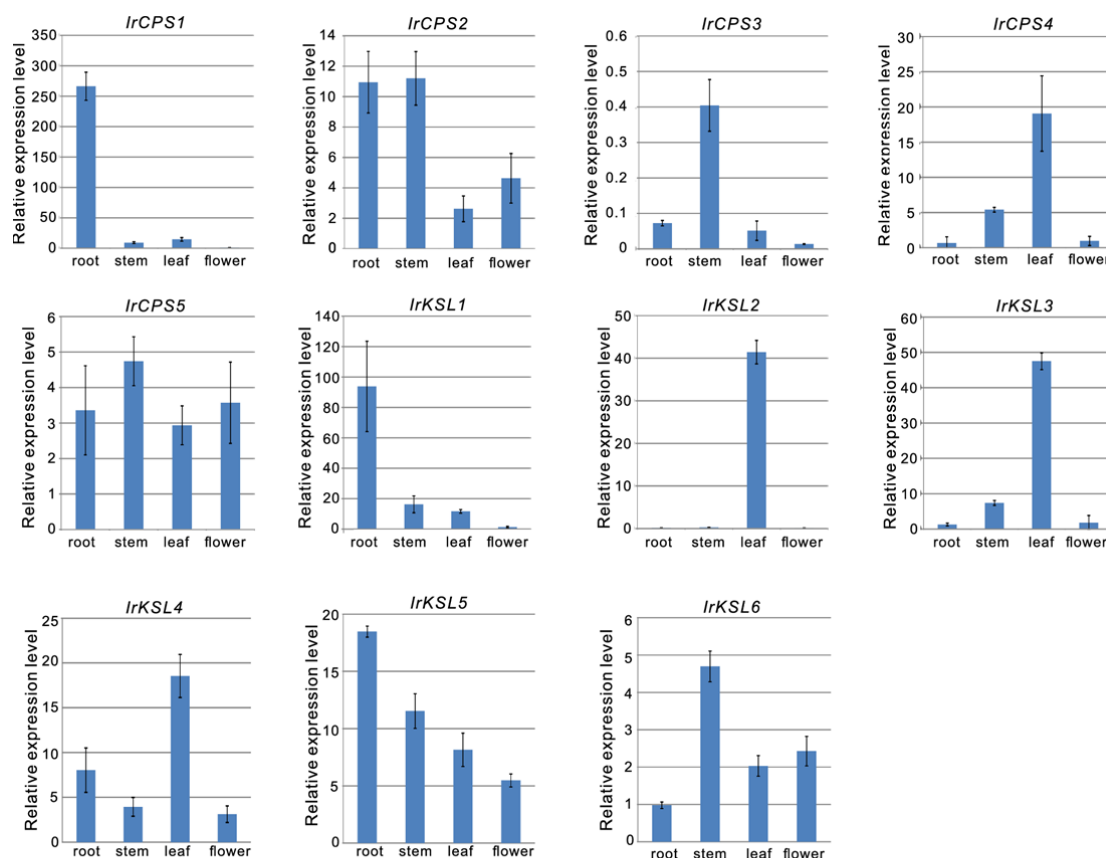


Figure 5. The mRNA expression levels of the 11 candidate diTPS in root, stem, leaf and flower tissues from *I. rubescens*. The expression level was normalized to that of *actin*. The error bars show the SDs from mean value ($n = 3$ experiments).

245 *IrKSL4* were highly expressed in the leaf tissues. *IrCPS3* showed relative lower transcription levels
 246 in all organs, which was also similar to expression levels in *SmCPS3* of *S. miltiorrhiza* (Cui et al.,
 247 2015) (Fig. 5).

248 In order to confirm that abietane diterpenoids (like miltiradiene produced by *IrCPS1* and
 249 *IrKSL1/KSL3*) exist in *I. rubescens*, different organs including root (periderm and cortex), stem,
 250 leaf and flower were extracted with hexane and analyzed by GC-MS. Miltiradiene, together with
 251 abietatriene and ferruginol (two compounds related to miltiradiene biosynthesis), were detected in
 252 the periderm of the root (Fig. 6). No abietane diterpenoids were found in other organs
 253 (Supplemental Fig. S4).

254 Oridonin is found in plant leaves and is a principle constituent of *I. rubescens*. This compound
 255 accumulates in the leaves at concentrations as high as 0.25 % of the dry weight (Committee CP,
 256 2015). Thus, the highly expressed levels of *IrCPS4* in the leaves illustrate that it may be

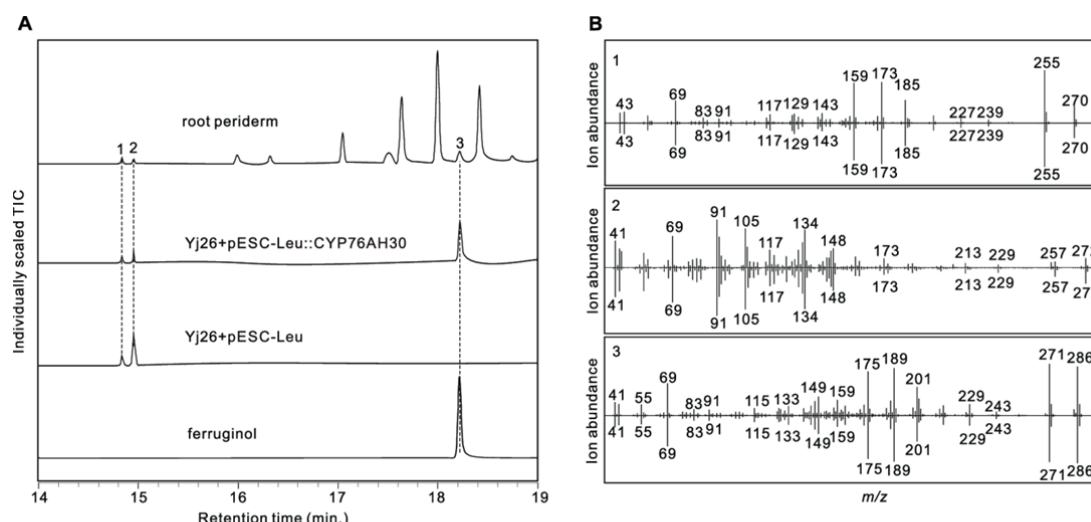


Figure 6. GC-MS detection of abietane diterpenoids in the periderm of the root and identification of CYP76AH30 in *I. rubescens* on a TR-5ms capillary column. A, Total ion chromatograms (TIC) of abietane diterpenoids in the hexane extract of the root periderm and the ferruginol production from strain YJ26 transformed with the plasmid pESC-Leu::CYP76AH30. Strain YJ26 with the plasmid pESC-Leu was used as a control, along with the authentic standard of ferruginol. B, Mass spectra of (1) abietatriene, (2) miltiradiene and (3) ferruginol from the root periderm (upper halves) and yeast (lower halves).

responsible for oridonin biosynthesis in *I. rubescens*. Other diterpenes, including *ent*-atiserene, *ent*-isopimaradiene like compound and isopimaradiene were not found in the organs studied.

Identification of CYP76AH30 as ferruginol synthase in *I. rubescens*

The conversion of miltiradiene to ferruginol have been elucidated in *S. miltiorrhiza* (Guo et al., 2013), *R. officinalis* (Zi and Peters, 2013; Ignea et al., 2016), and *S. fruticosa* (Božić et al., 2015; Scheler et al., 2016), where a 76AH sub-family member plays an important role. It is interesting to note that CYP76AH1 from *S. miltiorrhiza* is characterized as ferruginol synthase, while CYP76AH22-24 from *R. officinalis* and *S. fruticosa* has the ability to produce both ferruginol and 11-hydroxyferruginol. Mutagenesis analysis revealed that reciprocal activity changes from ferruginol synthase to hydroxyferruginol synthase only depend on three amino acids (and vice versa) (Scheler et al., 2016). When using CYP76AH1 from *S. miltiorrhiza* as a query sequence, one full-length homolog called CYP76AH30 was found in our transcriptome data. It shares 80.5% amino acid sequence identity with CYP76AH1, and 76.1%~ 77.6% identities with CYP76AH22-24. The three residues, 301E, 303N and 479L of CYP76AH30 are different from both CYP76AH1 and CYP76AH22-24 (Supplemental Fig. S5).

The role of the CYP76AH30 protein was identified by transforming the yeast strain, YJ26 (Zhou

273 et al., 2012) with a plasmid expressing the protein. This yeast strain harbors modules that express a
274 GGPP synthase and farnesyl diphosphate synthase fusion, a SmCPS1 and SmKSL1 fusion, and a
275 truncated hydroxy-3-methylglutaryl coenzymes A reductase (Zhou et al., 2012; Guo et al., 2013).
276 The pESC yeast epitope tagging vector harboring CYP76AH30 (pESC-Leu::CYP76AH30) under
277 the control of the GAL1 promoter was transformed into YJ26. After 48h of fermentation, we
278 detected ferruginol from the strain transformed with CYP76AH30 (Fig. 6). Thus, our results show
279 that CYP76AH30 is a ferruginol synthase and responsible for ferruginol biosynthesis in *I.*
280 *rubescens*.
281

282 Discussion

283 Here, we identified a large diTPS family in *I. rubescens* containing five CPS and six KSL
284 genes. Ten of these genes were functionally characterized *in vitro*, representing the most
285 extensive diTPS gene family found in Lamiaceae. Three of these genes are predicted to be
286 involved in *ent*-kaurene diterpenoids, such as oridonin and the requisite of GA metabolism, while
287 two other genes were also identified as KSLs functioning with *ent*-CPP as substrates. We also
288 showed that there is a normal-CPP mediated biosynthesis pathway that exists in *I. rubescens* and
289 that three KSL genes with different domain architectures are involved in this pathway,
290 highlighting the role of KSL domain loss in the evolution of chemical diversification in
291 diterpenoid biosynthesis.

292 *I. rubescens* contains a functionally diverse diTPS family

293 The intricate nature of diterpene metabolism (Fig. 7) is more complex than previously reported
294 in *I. rubescens* (Sun et al., 2006). Despite containing four shared functional CPS genes, *I.*
295 *rubescens* has four additional KSL genes compared to the two functional KSL genes identified in *S.*
296 *miltiorrhiza* (Cui et al., 2015).

297 Similar to *S. miltiorrhiza*, the two CPS (IrCPS1 and IrCPS2) and one KSL (IrKSL1) gene
298 products of *I. rubescens* are closely related to the Lamiaceae specialized CPS and KSL, which have
299 the same enzymatic activity as GGPP to produce normal-CPP and miltiradiene (Fig. 4). The
300 transcription pattern of *IrCPS1* and *IrKSL1* was similar to those involved in tanshinone
301 biosynthesis in *S. miltiorrhiza* and was mainly transcribed in the root periderm (Cui et al., 2015).
302 Our *in vitro* results show the identification of miltiradiene together with abietatriene and ferruginol
303 in the root periderm and further highlight the role of IrCPS1 in the miltiradiene mediated
304 biosynthesis (Fig. 7). In *I. rubescens*, IrKSL3 also has the ability to react with normal-CPP to
305 produce miltiradiene. IrKSL3 is highly expressed in leaves, suggesting its role in the biosynthesis
306 of abietane diterpenoids such as rubesanolides A-E. However, miltiradiene was not detected in the
307 aerial parts and its metabolic fate *in planta* calls for further study.

308 Phylogenetic and sequence analysis suggested that two genes, *IrCPS4* and *IrCPS5*, had the
309 ability to produce *ent*-CPP, the precursor of *ent*-kaurene diterpenoids. Unlike the relatively
310 ubiquitous expression patterns of *IrCPS55*, *IrCPS4* exhibited high transcription levels in leaf
311 tissues (Fig. 5), suggesting its role in specialized diterpene metabolism, such as oridonin

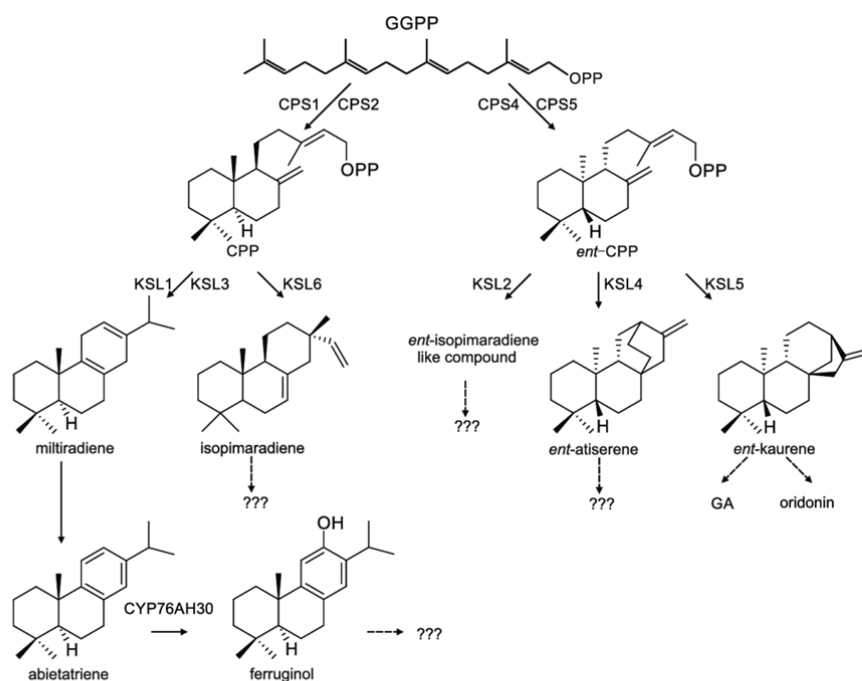


Figure 7. The proposed diterpenoid biosynthesis in *I. rubescens*. Two sequential diterpene synthases CPS and KSL, and corresponding reactions are indicated, along with the possible downstream natural products. Dashed arrows indicate multiple enzymatic reactions.

312 biosynthesis. The close phylogenetic relationship of IrCPS4 with IeCPS2 from *I. eriocalyx*,
 313 SmCPS4 from *S. miltiorrhiza* and the newly identified clerodienyl diphosphate synthase SdCPS2
 314 from *S. divinorum* (Pelot et al., 2017) highlights a Lamiaceae-specific lineage (Fig. 3), responsible
 315 for specialized *ent*-kaurene or labdane diterpenoid formation in Lamiaceae plants. The role of
 316 IrCPS4 in oridonin biosynthesis should further be confirmed by RNA interference or virus induced
 317 gene silencing *in planta*.

318 Three other KSL genes with unexpected enzymatic activities were identified in *I. rubescens*. The
 319 IrKSL2 gene product was found to be an *ent*-isopimaradiene-like molecule. Other
 320 *ent*-isopimaradiene-like compound derivatives have been isolated from *Calceolaria foliosa* (Chamy
 321 et al., 1989) and *I. lophanthoides* subsp. *Gerardiana* (Sun et al., 2006). The IrKSL4 gene product
 322 was found to be an *ent*-atiserene compound. A similar *ent*-atiserene gene product has been
 323 identified as a minor product (~4%) of the RcKSL4 protein from castor bean (*Ricinus communis*)
 324 (Jackson et al., 2014). Likewise, the product of IrKSL6, isopimaradiene, has not been found in
 325 *Isodon* species previously. Diterpenoid gene products from the leaves of *I. rubescens* revealed in
 326 this study provide a foundation for future *in planta* studies in the chemistry and metabolic fate of
 327 these molecules.

328 **Macroevolutionary of diterpenoids in Lamiaceae**

329 Since secondary metabolites play vital roles in defense or as signaling compounds, their occurrence
330 reflects adaptations and life strategies embedded in a given species. The identification of diTPS
331 genes in *S. miltiorrhiza* and *I. rubescens* offer an opportunity to elucidate diterpenoid biosynthesis
332 on a macroevolutionary level. The similar number and function of CPS enzyme in *S. miltiorrhiza*
333 and *I. rubescens* shows that these genes may originate from a common ancestral origin. The
334 resulting speciation of *S. miltiorrhiza* and *I. rubescens* led to the different evolution patterns of
335 diterpenoids. As a result, we speculate that because the fleshy root of *S. miltiorrhiza* is often under
336 herbivory attack (Chen et al., 2008) and survival has been based on the steady adaptation of
337 evolving molecules to increase the complexity of their chemical arsenal. This is evident in the root
338 production of miltiradiene, and the array of tanshinone compounds (Cui et al., 2015). The
339 undershrub of *I. rubescens* is also attacked by herbivores and the increasing complexity of the leaf
340 chemical arsenal, including oridonin and other *ent*-kaurene-derived compounds, is further
341 elucidated in the growing number of diterpene biosynthetic pathways able to produce different
342 complex chemical mixtures (Fig. 7). Diterpenoid pathways in Lamiaceae appear to have several
343 instances of multifunctional ancestral pathways including the production of *ent*-kaurene generated
344 by the same biosynthetic pathway as the universal hormone GA (Zi et al., 2014) and the production
345 of miltiradiene involved in the more chemically elaborate mixture of normal-CPP mediated
346 diterpenoids (Gao et al., 2009; Zerbe et al., 2014; Brückner et al., 2014; Pateraki et al., 2014).

347 **The evolution of domain loss in Lamiaceae**

348 To our knowledge, among KSL enzymes retaining the tri-domain ($\gamma\beta\alpha$) structure in Lamiaceae,
349 IrKSL3 and IrKSL6 have the ability to react with normal-CPP (Fig. 4). It's most likely that the
350 tri-domain enzymes (IrKSL3 and IrKSL6) belong to an ancestral KSL clade that has the loss of γ
351 domain, a widespread occurrence in the Lamiaceae (Gao et al., 2009; Caniard et al., 2012; Li et al.,
352 2012; Schalk et al., 2012; Bruckner et al., 2014; Pateraki et al., 2014; Božić et al., 2015; Cui et al.,
353 2015; Triikka et al., 2015; Su et al., 2016). Therefore, these sequences from *Isodon* represent a step in
354 the evolution of these Lamiaceae KSL with a domain loss. It is possible that those bi-domain KSLs
355 evolved independently in Lamiaceae on at least three separate occasions (Fig. 8). One instance is
356 the evolution of the tri-domain KSLs from an ancestral KSL that produced *ent*-kaurene for GA
357 biosynthesis and arose from an early gene duplication and neofunctionalization during Angiosperm

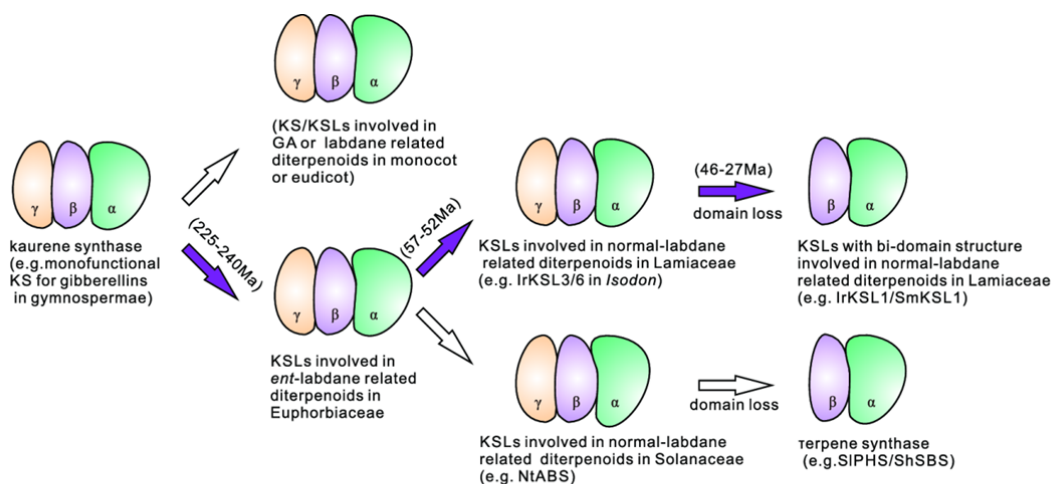


Figure 8. Proposed process for the evolution of bi-domain terpene synthases in Lamiaceae. The purple arrows show the proposed evolutionary steps from the ancient tri-domain kaurene synthase to the bi-domain KSLs in Lamiaceae.

evolution, prior to the dicot-monocot split (Brückner et al., 2014; Fig. 3 and 8) at about 240-225 million years ago (Ma) (Zeng et al., 2014). The second possibility is the emergence of a three-domain KSL described here (IrKSL3 and IrKSL6) in Lamiaceae along with the radiation of the subfamily Nepetoideae at the Paleocene/Eocene boundary (57-52 Ma) (Drew and Sytsma, 2012). The third option is the occurrence of $\beta\alpha$ bi-domain KSLs along with the rise of the tribe Menthae in Subfamily Nepetoideae during the mid-Eocene (46 Ma) (Drew and Sytsma, 2012). Since the genus *Isodon* arose 27 Ma (Yu et al., 2014), the existence of both tri-domain (i.e. IrKSL3 and IrKSL6) and bi-domain (i.e. IrKSL1) KSLs in *I. rubescens* may demonstrate that the bi-domain KSLs originated between 46 Ma to 27 Ma.

It is interesting to note that the recent emergence enzymes MvELS in *Marrubium vulgare* (12.6 Ma) (Roy and Lindqvist, 2015) and SsScS in *S. sclarea* (10 Ma) (Drew and Sytsma, 2012) exhibit more diversified enzymatic activity than other KSLs (Zerbe et al., 2014; Schalk et al., 2012; Andersen-Ranberg et al., 2016; Jia et al., 2016). This alludes to the fast evolution of KSL enzymes in Lamiaceae. To date, all of the identified $\beta\alpha$ bi-domain KSL enzymes have dual substrate activities (Caniard et al., 2012; Bruckner et al., 2014; Zerbe et al., 2014; Andersen-Ranberg et al., 2016; Jia et al., 2016), highlighting their roles in contributing to the diversity of diterpenoids in Lamiaceae. The proposed intermediate steps in KSL protein evolution studied here can be uncovered when additional genera are explored at the sequence level. Next generation sequencing technologies allow this at an accelerated pace and should provide valuable information on the

377 evolution of enzymes (not just diterpene synthases) within or across plant families.

378

379 **MATERIALS AND METHODS**

380 **Plant Materials**

381 Seeds of *I. rubescens* were from Henan Province, given by Prof Chen Suiqing. Plants were grown
382 in a controlled environment chamber, maintained at 25 °C under a 16 h light/8 h dark photoperiod.

383 **RNA extraction, cDNA library preparation, and transcript analyses by RNA-seq**

384 The leaves of five-month old plants were used for RNA-seq analysis. RNA was extracted
385 (HuaYueYang biotechnology, Beijing) and shipped to the GENEWIZ Company (www.genewiz.com)
386 for library construction and RAN-seq. The RNA quality was checked using an Agilent 2100 with
387 Agilent Eukaryote Total RNA Nano Kit. The RNA-seq library was prepared using the TruSeq
388 RNA Sample Prep Kit for Illumina starting with 1µg of total RNA. The library was purified on
389 Beckman AMPure XP beads. The barcoded RNA-seq library was assessed by qRT-PCR using the
390 Library Quantification kit. The size range of the final cDNA libraries was determined on an Agilent
391 bioanalyzer DNA7500 DNA chip (Agilent Technologies). The cDNA libraries were sequenced on
392 one lane for 151 cycles from each end of the cDNA fragments on a HiSeq2500 using a TruSeq SBS
393 sequencing kit v3-HS (Illumina). The sequence images were transformed to bcl files with the Real
394 Time Analysis 1.17.21.2 Illumina software, which were multiplexed to fastq files with CASAVA
395 version 1.8.2. The quality-scores line in fastq files processed with Casava1.8.2 use an ASCII offset
396 of 33 for presentation in the Sanger format.

397 **Transcriptome assembly and annotation**

398 The Trinity package version trinityrnaseq-r2013-02-25 (Grabherr *et al.*, 2011) was utilized to
399 assemble the transcriptome from the RNA-seq data. Prior to assembly, adaptor sequences were
400 trimmed from both ends of the sequence and so were any low quality bases (<30 phred score).
401 Sequences of 50 nucleotides and higher were utilized for the Trinity assembly, which was
402 performed according to guidelines in the published protocols (Grabherr *et al.*, 2011), and on the
403 Trinity website (<https://github.com/trinity-rnaseq/trinityrnaseq/wiki>). Individual isotigs were
404 annotated by searching the NCBI non-redundant protein sequence (nr) database using BLASTX
405 software with default parameters. Isotig functions were assigned based on the annotation associated
406 with the top hit that satisfied the following criteria: (i) $\geq 30\%$ sequence identity; (ii) $\geq 30\%$

407 alignment coverage of either the query or subject sequences; and (iii) with BLAST e-values < E-5.
408 Manually assemblies were carried out with Bioedit and DNAMAN.

409 **Cloning of the full length diTPS genes**

410 The total RNA used for RNA-seq was further used to verify the full length of diTPS obtained from
411 transcriptome assembly and clone the other diTPS by rapid amplification of cDNA ends (RACE).
412 5'-RACE and 3'-RACE was performed using the SMARTerTM RACE DNA Amplification kit
413 (Clontech Laboratories Inc., Mountain View, CA, USA) according to the manufacturer's protocol.
414 All the primers for RACE-PCR were shown in Supplemental Table S6. The full length of diTPS
415 genes were then cloned into pMD19-T (Takara, Dalian, China) vector for sequencing.

416 **DiTPS Gene Expression Analysis**

417 *I. rubescens* flowering plants were used to determine diTPS expression levels. Samples were
418 harvested and immediately frozen in liquid nitrogen. RNA was extracted (HuaYueYang
419 Biotechnology, Beijing) and RNA integrity and quality checked by denaturing gel electrophoresis.
420 1~5 µg of total RNA was reverse transcribed into cDNA using the PrimerScriptTM RT reagent kit
421 with gDNA eraser (TaKaRa Corp., Dalian, China), according to the manufacturer's instructions.
422 The synthesized cDNA was then diluted 10-fold and 1 µL of this diluted template with 0.4 µL of
423 ROX Reference Dye II (TaKaRa) were used for subsequent qRT-PCR analysis with a total PCR
424 reaction volume of 20 µL. The primers sequences are shown in Supplemental Table S6. qRT-PCR
425 was performed with a SYBR Green kit (TaKaRa Corp., Dalian, China) on ABI7500 real-time PCR
426 detection system. All reactions were performed using the following PCR conditions: initial
427 denaturation step of 95 °C for 10 min, followed by 40 cycles for 95 °C for 5 s, 60 °C for 34 s, with
428 a final melting stage from 60 °C to 95 °C. Primer specificity was assessed by agarose gel and
429 melting curve analysis. The results were normalized with housekeeping gene *actin*. Relative
430 expression levels were calculated as the mean of three technical replicates of 3 biological
431 replicates.

432 **Phylogenetic analysis**

433 The previously identified 96 diTPSs (Supplemental Table S5) were used to construct the
434 phylogenetic tree follow the method described previously (Cui et al., 2015).

435 **In Vitro Assays**

436 For *in vitro* functional assays, the full coding sequence of diTPS obtained from RNA-seq data

437 (*IrCPS1*, *IrCPS3*, *IrCPS4*, *IrKSL1*) was proved by PCR and further synthesized with codon
438 optimization for *Escherichia coli* (GENEWIZ biotech Co.) and cloned into pET32 plasmid (Merck).
439 The full coding sequence of other seven diTPS obtained from RACE (*IrCPS2*, *IrCPS3*, *IrCPS5*,
440 *IrKSL2*, *IrKSL3*, *IrKSL4*, *IrKSL5* and *IrKSL6*) were digested with specific restriction enzyme site
441 (Supplemental Table S6) and subcloned into the expression plasmid pET32a. The recombinant
442 diTPS proteins were expressed in *E. coli* Tuner, affinity-purified and assayed with GGPP as the
443 substrate.

444 *SmCPS1* (GenBank: KC814639), *SmCPS5* (KC814642), *SmKSL1* (ABV08817) from *S.*
445 *miltiorrhiza*, *AtKS* (AAC39443) from *A. thaliana*, *PcmISO1* (JQ240314) from *Pinus banksiana*
446 were used in this study. The expression and purification of the recombinant proteins were
447 performed as described previously (Cui et al., 2015). In short, the constructs were transformed into
448 Tuner (DE3) or TransB (DE3) competent cells. Three to five positive colonies were cultured in LB
449 medium with 50 mg·L⁻¹ carbenicillin, and 0.1~0.4 mM isopropyl β-D-thiogalactopyranoside (IPTG)
450 was added to induce the expression of the protein. Subsequently, cell pellets were collected and
451 re-suspended in assay buffer (50 mM phosphate pH 7.4, 10% glycerol, 2 mM DTT, 10 mM MgCl₂)
452 and sonicated six times for 10 sec, on ice. Lysate from the samples was centrifuged at 12000 x g
453 for 20 min at 4 °C. The proteins were purified with the nickel-nitrilotriacetic acid agarose beads
454 followed the previous described method (Cui et al., 2015).

455 The conversion of GGPP to CPP was carried out by incubating 200 μL of the purified enzymes
456 with 20-50 μM GGPP (Sigma) for 2-4 h at 30 °C. Assay mixtures were hydrolysed
457 (dephosphorylated) with 75 U bacterial alkaline phosphatase at pH 8 for 16 h at 37 °C to produce
458 hexane-soluble products. GGPP was converted to kaurene, miltiradiene and other diterpenes by
459 mixing 250 μL of purified recombinant CPS and KSL proteins with 20-50 μM of GGPP, and
460 incubated for 2-4 h at 30 °C. Assay mixtures were extracted three times with an equal volume of
461 hexane. The hexane fractions were pooled, evaporated under nitrogen and resuspended in 50 μL of
462 hexane or derivatized with 80 μL N-methyl-N-(trimethylsilyl) trifluoroacetamide (MSTFA) at 80 °C
463 for 40 minutes, and then analyzed by GC-MS.

464 **Identification of CYP76AH30 using the Engineering Yeast**

465 Full-length cDNA of CYP76AH30 were cloned from the leaves using primers shown in
466 supplemental Table S6. The open reading frame of CYP76AH30 was subcloned into the yeast

467 epitope-tagged vector pESC-Leu using BamHI and SalI, yielding pESC-Leu::CYP76AH30. The
468 plasmid was transformed into the mitreriadiene production strain YJ26, using lithium
469 acetate/single-stranded carrier DNA/polyethylene glycol transformation method (Daniel Gietz & Woods,
470 2002; Zhou et al., 2012). Transformants were selected on yeast nitrogen base without amino acids
471 (YNB) medium containing 20 g l⁻¹ glucose and grown at 30°C for 48 h. The recombinant yeast
472 strains were grown in YNB medium containing 2% glucose (YNB/glucose) at 30°C, shaking at 250
473 rpm, for 48 h, then transferred to 50 ml YNB/glucose medium in 250 ml flasks and grown to an
474 initial OD₆₀₀ of 0.05, and cultivated an additional 12-16 h to reach logarithmic phase. Cells were
475 centrifuged and washed twice with sterile water to remove any residual glucose. The cells were
476 then resuspended in 50 ml YNB medium containing 2% galactose (YNB/gal) for induction, and
477 grown for 30-72 h to produce diterpenoids. Yeast cultures were extracted three times by
478 ultrasonication with an equal volume of hexane. After separation, the organic extract was
479 concentrated under vacuum, and the residue resuspended in hexane for GC-MS analysis.

480 **Diterpene analysis using GC-MS**

481 Plant material was lyophilized for 48 h and 100 mg of the lyophilized powder extracted with 2 mL
482 hexane. The extracts were sonicated twice for 15 min and centrifuged (3000 × g) for 10 min. The
483 supernatant was evaporated under nitrogen, resuspended in 50 µL of hexane and analyzed by
484 GC-MS.

485 Two GC-MS systems were used in this study. Plant material and the entire *in vitro* assay
486 product were first carried out using a Trace 1310 series GC with a TSQ8000 MS detector
487 (Thermo Fisher Scientific Co. Ltd.). A 1 µL portion of extract was injected in splitless mode onto
488 the column. Chromatographic separation was performed on a TR-5ms capillary column (30 m ×
489 0.25 mm ID; DF = 0.25 µm; Thermo Fisher Scientific Co. Ltd.). Helium was used as the carrier
490 gas at a constant flow rate of 1 mL/min through the column. The injector temperature was set at
491 280 °C and the injector temperature was 280 °C. The oven program was as follows: 50 °C for 2
492 min, linear ramp at a rate of 20 °C·min⁻¹ to 200 °C, followed with a linear ramp at a rate of
493 5 °C·min⁻¹ to 300 °C, held at 300 °C for 10 min. The transfer line temperature was 280 °C.

494 The other GC-MS system is Agilent GC-MS 7890B connected to an Agilent 7000B triple
495 quadruple MS with electron impact ionization. A 1 µL portion of the extract was injected in
496 splitless mode onto the column. The column used was a chiral column Agilent Cyclodex-β (30 m x

0.25 mm i.d., 0.25- μ m film thickness). Helium was used as the carrier gas for GC at a flow rate of 1.0 ml·min⁻¹. The injector temperature was 250 °C. The oven program was as follows: 50 °C for 2 min, linear ramp at a rate of 8 °C·min⁻¹ to 230 °C, then held at 230 °C for 8 min. The transfer line temperature was 280 °C.

Identification of *ent*-kaurene, multiradiene and isopimaradiene were achieved by comparison to the previously defined enzymes. *Ent*-atiserene and ferruginol were identified by comparison to authentic standard.

The Illumina-derived nucleotide sequences reported in this paper has been submitted to the short read archive (SRA) at NCBI with the accession number of GSE96954. Nucleotide sequences of characterized enzymes can be found in the GenBank/EMBL data libraries with the following accession numbers: IrCPS1 (KU180499), IrCPS2 (KU180500), IrCPS3 (KU180501), IrCPS4 (KU180502), IrCPS5 (KU180503), IrKSL1 (KU180504), IrKSL2 (KU180505), IrKSL3 (KU180506), IrKSL4 (KX580633), IrKSL5 (KX580634), IrKSL6 (KX580635), Ir76AH30 (KX580636).

510

Supplemental Data

Supplemental Figure S1. Amino acid alignment of diterpene synthase in *I. rubescens* and *S. miltiorrhiza*.

Supplemental Figure S2. GC-MS analysis of *in vitro* assays with *I. rubescens* IrCPS4 and IrCPS5

Supplemental Figure S3. GC-MS analysis of different clade of IrKSL with the corresponding IrCPS on a TR-5ms capillary column.

Supplemental Figure S4. GC-MS detection of abietane diterpenoids in different organs

Supplemental Figure S5. Amino acid alignment of CYP76AH30 with CYP76AH1 and CYP76AH22-24

Supplemental Table S1. The number and length of all of the unigenes obtained by RNA-Seq

Supplemental Table S2. Unigenes annotated as copalyl diphosphate synthase and kaurene synthase genes

Supplemental Table S3. Information of diTPS gene family in *I. rubescens*

Supplemental Table S4. Blast result of genes identified from this paper to the reference transcriptome sequencing dataset

Supplemental Table S5. List of plant diTPS involved in this paper

527 **Supplemental Table S6.** Primers used in this study

528

529 **Figure legend**

530 **Figure 1.** The major medicinally active diterpenoids in *I. rubescens*. A, The plant of *I. rubescens*.
531 B, The typical *ent*-kaurene diterpenoid of oridonin. C, The typical abietane diterpenoid of
532 rubesanolide D.

533 **Figure 2.** Alignment of the conserved DXDD motif and the positive selection site (Ser-362 /
534 Trp-364) of CPS from *I. rubescens* and other characterized CPSs. The DXDD motif is labeled
535 above the alignment with a solid line, and the conserved Ser or Trp position is marked with an
536 asterisk. The second Asp of the DXDD motif is substituted by Asn in IrCPS3. The enzymes are
537 separated by the horizontal line according to normal-CPP/LDPP or *ent*-CPP stereochemistry. The
538 CPS in the upper section (including IrCPS1 and IrCPS2 from this study) were characterized as
539 being involved in normal-CPP/LDPP formation while those in the lower section (including
540 IrCPS4 and IrCPS5 from this study) were characterized to be involved in *ent*-CPP production.
541 CfTPS1 and CfTPS2 are from *Coleus forskohlii*, SmCPS1, SmCPS2, SmCPS5 from *Salvia*
542 *miltiorrhiza*, MvCPS3 from *Marrubium vulgare*, SsLPS from *S. sclarea*, and AtCPS from
543 *Arabidopsis thaliana*.

544 **Figure 3.** Phylogeny of *I. rubescens* diterpene synthases. The maximum likelihood tree illustrates
545 the phylogenetic relationship of *I. rubescens* diterpene synthases with 96 representative
546 characterized diTPS (Supplemental Table S5). Numbers on branches indicate the bootstrap
547 percentage values calculated from 1000 bootstrap replicates. *Physcomitrella patens* CPS/kaurene
548 (PpCPSKS) was used as an outgroup. Blue lines show diTPS genes from Lamiaceae involved in
549 normal-CPP/LDPP-mediated diterpenoid metabolism and red lines show the loss of the
550 N-terminal γ domain found in eudicot and monocot KSLs. Yellow lines show genes from
551 Lamiaceae involved in the specialized *ent*-CPP/LDPP-related diterpenoid metabolism.
552 Red-marked enzymes show diTPS from *I. rubescens* in this study.

553 **Figure 4.** GC-MS analysis of *in vitro* assays with *I. rubescens* diTPS on a Cyclodex- β GC
554 column. A, Extracted ion chromatograms (EIC) of *m/z* 257 of *in vitro* assays with IrCPS coupled
555 with different IrKSL. The characterized SmCPS5 (*ent*-CPP synthase from *S. miltiorrhiza*),
556 SmCPS1 (normal-CPP synthase from *S. miltiorrhiza*), SmKSL1 (miltiradiene synthase from *S.*
557 *miltiorrhiza*), AtKS (*ent*-kaurene synthase from *A. thaliana*) and PcmISO1 (pimaradiene from
558 *Pinus banksiana*) were used as positive controls, along with the corresponding authentic standard
559 *ent*-atiserene. 1. miltiradiene; 2. isopimaradiene; 3. *ent*-kaurene; 4. *ent*-atiserene; 5.
560 *ent*-isopimaradiene like compound. * indicates the unexpected enzyme products identified in this
561 study. B, Corresponding mass spectra of recombinant enzyme assay products and authentic
562 standard.

563 **Figure 5.** The mRNA expression levels of the 11 candidate diTPS in root, stem, leaf and flower
564 tissues from *I. rubescens*. The expression level was normalized to that of *actin*. The error bars
565 show the SDs from mean value ($n = 3$ experiments).

566 **Figure 6.** GC-MS detection of abietane diterpenoids in the periderm of the root and identification
567 of CYP76AH30 in *I. rubescens* on a TR-5ms capillary column. A, Total ion chromatograms (TIC)
568 of abietane diterpenoids in the hexane extract of the root periderm and the ferruginol production
569 from strain YJ26 transformed with the plasmid pESC-Leu::CYP76AH30. Strain YJ26 with the

570 plasmid pESC-Leu was used as a control, along with the authentic standard of ferruginol. B,
571 Mass spectra of (1) abietatriene, (2) miltiradiene and (3) ferruginol from the root periderm (upper
572 halves) and yeast (lower halves).

573 **Figure 7.** The proposed diterpenoid biosynthesis in *I. rubescens*. Two sequential diterpene
574 synthases CPS and KSL, and corresponding reactions are indicated, along with the possible
575 downstream natural products. Dashed arrows indicate multiple enzymatic reactions.

576 **Figure 8.** Proposed process for the evolution of bi-domain terpene synthases in Lamiaceae. The
577 purple arrows show the proposed evolutionary steps from the ancient tri-domain kaurene
578 synthase to the bi-domain KSLs in Lamiaceae.

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Parsed Citations

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