FUNCTIONAL IDENTIFICATION OF THE TERPENE SYNTHASE FAMILY INVOLVED IN DITERPENOID ALKALOIDS BIOSYNTHESIS IN ACONITUM CARMAICHELII

LIUYING MAO, BAOULONG JIN, LINGLI CHEN, MEI TIAN, RUI MA, BIWEI YIN, HAIYAN ZHANG, JUAN GUO, JINFU TANG, TONG CHEN, CHANGJIANGSHENG LAI, GUANGHONG CUI, LUQI HUANG

ABSTRACT: Aconitum carmichaelii is a high-value medicinal herb widely used across China, Japan, and other Asian countries. Aconitine-type diterpene alkaloids (DAs) are the characteristic compounds in Aconitum. Although six transcriptomes, based on short-read next generation sequencing technology, have been reported from the Aconitum species, the terpene synthase (TPS) corresponding to DAs biosynthesis remains unidentified. We apply a combination of Pacbio isoform sequencing and RNA sequencing to provide a comprehensive view of the A. carmichaelii transcriptome. Nineteen TPSs and five alternative splicing isoforms belonging to TPS-b, TPS-c, and TPS-e/f subfamilies were identified. In vitro enzyme reaction analysis functional identified two sesqui-TPSs and twelve diTPSs. Seven of the TPS-c subfamily genes reacted with GGPP to produce the intermediate ent-copalyl diphosphate. Five AcKSLs separately reacted with ent-CPP to produce ent-kaurene, ent-atiserene, and ent-13-epi-sandaracopimaradiene: a new diterpene found in Aconitum. AcTPSs gene expression in conjunction DAs content analysis in different tissues validated that ent-CPP is the sole precursor to all DAs biosynthesis, with AcKSL1, AcKSL2s and AcKSL3-1 responsible for C20 atisine and napelline type DAs biosynthesis, respectively. These data clarified the molecular basis for the C20-DAs biosynthetic pathway in A. carmichaelii and pave the way for further exploration of C19-DAs biosynthesis in the Aconitum species.
1. Introduction

*Aconitum carmichaelii* Debx. is a traditional herb belonging to Ranunculaceae. Its principal and lateral roots after processed, named “wu tou” and “fu zi” respectively, are its most commonly used components in traditional Chinese medicine (Fig. 1), and have been used for the treatment of pain, rheumatics, heart failure, colds, diarrhea, beriberi, and edema for 2000 years. Diterpenoid alkaloids (DAs) are believed to be the predominant bioactive compounds in *A. carmichaelii*, with over 100 isolated. C19-diterpenoid alkaloids, which contain unusual 6,7,5,6 carbon skeletons, are both the dominant bioactive and toxic constituents in *A. carmichaelii*. The most toxic compounds are aconitine, mesaconitine, and hypaconitine, characterized by acetyl and benzoyl esters (Fig. 1C), while alcohol amine-DAs are the primary bioactive compounds and exhibit reduced toxicity. In addition to the C19-type, several C18- and C20-DAs have also been found in *A. carmichaelii*, such as vilmorrianine D, songorine, napelline, atisine (Fig. 1C) and aconicarmissulfonine A: linked to significant analgesic activity.

DAs, whether C18, C19, or C20, are all presumed to be derived from the kaurane and atisane diterpenoid families. That is, C20-atisane and kaurane serve as biosynthetic precursors to C18- and C19-DAs, a mechanism proven by total synthesis. Thereby, the biosynthesis of DAs undergoes two principal phases. First, is the formation of the C20 diterpene skeleton by terpene synthase (TPS): the universal precursor geranylgeranyl pyrophosphate (GGPP) is cyclized by class II diterpene synthase (copalyl-diphosphate synthase, CPS) to produce ent-copalyl diphosphate (ent-CPP), then undergoes further cyclization or rearrangement by class I diterpene synthase (kaurene synthase-like, KSL) to form ent-kaurene or ent-atiserene. Second, is the insertion of the nitrogen into the mature diterpene scaffolds to form C20-DAs, e.g., atisine and napelline (Fig. 1C), which then undergo further development leading to the terminus C18- or C19-DAs. Historically, the specific enzymes and biosynthetic pathways in the plant have garnered substantially less interest than the total synthesis of these compounds: from 1963 to 2018, 24 different DAs have identified via total synthesis. One of the few key mechanistic findings has been that L-serine may serve as the nitrogen source of atisine-type DAs.

Recently, several RNA-seq approaches using the Illumina platform have analyzed the transcriptome of *A. carmichaelii* and *Aconitum heterophyllum*. Many candidate genes, such as terpenoid-related enzymes, monoxygenases, methyltransferases, and BAHD acyltransferases (named according to the first four identified enzymes of the family: BEAT, benzylalcohol O-acetyltransferase; AHCT, anthocyanin O-hydroxycinnamoyltransferase; HCBT, anthranilate N-hydroxycinnamoyl/benzoyltransferase; DAT, deacetylviolindole 4-O-acetyltransferase) related to DAs biosynthesis have been identified. However, the limitations of short-read sequencing results in the vast majority of isotigs not representing full-length cDNA sequences, and none of these candidates have yet been functionally identified through in vivo or in vitro analysis.

In addition to being initiated by class II diterpene synthase, diterpenes can also be initiated by class I diterpene synthase, such as taxa-4(5),11(12)-diene, casbene, and pseudolaratriene. Meanwhile, there are case reports that in addition to the TPS-c (e.g., CPS) and TPS-e/f (e.g., KSL) subfamily genes, the TPS-a and TPS-b subfamily genes have also been linked to diterpene biosynthesis. This suggests that C19-DAs backbones could be

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**Figure 1** The plant shows tissues used in this study for sequencing and the characteristic DAs in *A. carmichaelii*. The four-month-old plant (A) and roots (B) of *A. carmichaelii* in the greenhouse. C. The characteristic skeleton of DAs in *A. carmichaelii*. PR, principal root; LR, lateral root; FR, fibrous root; BPS, basal part of the stem; TL, Top leaf; CL, central leaf; LF, lower leaf; TS, Top stem; CS, central stem; LS, lower stem, NO, nodular like organs.
synthesized by unexpected enzymes. Accordingly, identifying all the TPS family genes from the whole transcriptome is a critical step in understanding the therapeutic agents of *A. carmichaelii*.

Single-molecule real-time (SMRT) sequencing provides reads > 10 kb and up to 60 kb (PacBio, Menlo Park, CA, USA 2016), enabling more complete and even full-length transcriptome data. \(^{27,28}\) Herein, we combined full-length isoform sequencing (Iso-seq) and RNA-seq technology to obtain a more reliable *A. carmichaelii* transcriptome. Combined with PCR cloning, we finally identified 19 TPS genes in *A. carmichaelii* and five alternative splicing isoforms. In *vivo* and *in vitro* assay identified 14 functional TPS genes, which establish biosynthetic routes to the diterpene scaffolds ent-atiserene, ent-kaurene, and ent-13-epi-sandaracopimaradiene, clarifying the molecular basis for *A. carmichaelii* DAs biosynthesis.

2. Materials and methods

2.1. Plant material

*A. carmichaelii* was acquired from Jiangyou, Sichuan Province, which is the genuine location for “wu tou” and “fu zi” material\(^ {29}\), in June 2018. The plant was grown in a greenhouse until February 2019. Then, 7 lateral roots from the same plant were divided and planted in separate flowerpots. These were then grown in a greenhouse at 25 ± 2 °C under a 16 h-light/8 h-dark cycle provided by a white fluorescent lamp (3000 lux). After 4 months growth, 3 plants with similar conditions were selected for the study. Different tissues were collected and immediately frozen in liquid nitrogen for further study.

2.2. RNA extraction

Total RNA was extracted using an RNA isolation kit (HuaYueYang Biotechnology, Beijing, China) following the manufacturer’s instruction. The integrity and concentration of total RNA was monitored using 1.0% agarose gel electrophoresis and NanoDrop ND-100 (NanoDrop Technologies, DE, USA). The electrophoreogram is shown in Supporting Information Fig. S1. The Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA) was further used to assess the quality of extracted RNA.

2.3. Iso-seq and RNA-seq library construction and sequencing

Due to the relatively high DAs content in the top leaves, principal roots, and lateral roots, to perform Iso-seq we mixed RNA from these three tissues together; concurrently, 13 tissues from three biological replicates were used for RNA-seq. The library construction for Iso-seq was described in Supporting Information Fig. S2. Briefly, mRNA was isolated from total RNA and converted into full-length cDNA using a SMARTer PCR cDNA Synthesis Kit (Clontech, CA, USA). Then, cDNA amplification was performed via an advantage 2 PCR kit (Clontech). After purification, size selection using a BluePippin Size selection system was performed (Sage Science, MA, USA) to enrich fragments longer than 4 kb. Then, an equal molar ratio mixed library of enriched fragments with those not processed by size selection was constructed using a SMRTbell Template Prep kit (Pacific Biosciences, CA, USA). The libraries’ quality was assessed by Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA) and Qubit fluorometer 2.0 (Life Technologies, CA, USA) before sequencing. To prepare the libraries for sequencing, sequencing primer was annealed and polymerase was added to the primer annealed template. Subsequently, the polymerase-bound template was bonded to MagBeads and sequenced on a PacBio RS II instrument (Novogene company, www.novogene.com).

RNA-seq was performed according to the standard protocol. Briefly, mRNA was converted into library templates using a TrueSeq RNA Sample Prep Kit v2 (Illumina, San Diego, CA, USA). The cDNA libraries were then sequenced from each end of the cDNA fragments on a HiSeq 2500 (Illumina).

2.4. Full-length transcript data processing and annotation

The SMRT-Analysis software package SMRTlink v7.0 (Pacific Biosciences) was used for Iso-seq data analysis. Briefly, low-quality reads (<50 bp) were first removed to obtain subreads. The subreads were then classified into circular consensus sequence (CCS) and non-CCS subreads using a ToFu pipeline (Pacific Biosciences). If the sequence contained a 3' and 5' primer or a poly (A) tail, the read was then used to obtain the full-length non-chimeric read (FLNC). Redundant FLNCs from same transcripts were discarded to form the consensus reads. The consensus reads were then polished with arrow software, and sequencing errors were further corrected using Illumina reads from the 13 different tissues via LoRDEC\(^ {30} \). Lastly, the redundant transcripts were removed via CD-HIT software\(^ {31} \). The resultant transcripts then underwent the standard annotation process using 7 databases: Nr, Nt, Pfam, KOG/COG, Swiss-prot, KEGG, and GO.

2.5. Cloning, sequence alignment, and phylogenetic analysis of TPS genes

1 μg of total RNA from 5 tissues were reverse transcribed into cDNA using the PrimerScript™ RT reagent kit with gDNA eraser (TaKaRa Corp., Dalian, China) according to the manufacturer’s instructions. TPS genes were then amplified with specific primers (Supporting Information Table S1) and cloned into a pET32 plasmid (Merck) with a seamless cloning kit (TransGen Biotech, Beijing, China). TPS alignment was represented using CLC Sequence viewer 7 (Qiagen, Denmark) software. The previously identified 51 diTPSs (Supporting Information Table S2) together with TPS from *Arabidopsis thaliana* were used to construct the maximum likelihood phylogenetic tree using MEGA 6.0\(^ {32} \).

2.6. In vitro assays

The functions of the AcTPSs were tested *via in vitro* assays as has been previously described\(^ {33} \). AcTPSs were analyzed alone or coupled with others, and an appropriate substrate supply. Briefly, recombinant plasmids were expressed in *Escherichia coli* Transetta (DE3). The crude proteins were then affinity-purified and individually assayed *via* GPP, FPP, GGPP, and CPP. CPP was obtained by incubating 200 μL of the purified ZmCPS2 enzymes with 20–50 μmol/L GGPP (Sigma) for 3 h at 30 °C. AcCPS function was estimated using the GGPP substrate; AcKSL function was assessed *via* the GGPP and CPP substrates; the other TPSs were tested with all possible substrates, respectively. Nine known enzymes, ZmCPS2 (Genbank: NM_001111787), IrCPS4 (KU180502), SmCPS1 (CCS), IrKSL4 (KX850633), SmKSL1 (EF635966), EpKSL4 (KP889108), OksKSL10 (DQ823355) and MsTPS1 (MH626616) were used as controls. Different enzymes were mixed with equal volume, typically 300 μL each, in a 2 mL vial. After incubation for 2–4 h at 30 °C, an equal volume of hexane was used to extract...
the terpene product. The hexane extracts were dried under a gentle stream of N2, and the residue suspended in 100 µL hexane. 1 µL was later subjected to GC–MS analysis.

2.7. Metabolic engineering of diterpenes in E. coli

A previously reported modular metabolic engineering system34 was used in our study. Briefly, the plasmid pIRS overexpressed IDI and DXR35, which are key genes from endogenous isoprenoid precursor pathway. The plasmid pGG-ZmCPS2 contained a pseudo-mature GGPS from Abies grandis36 and ZmCPS2/An2 from maize37, which have been designed to efficiently produce GGPP and CPP, respectively. These two plasmids, together with pET32-AcKSLs, were transfected into an E. coli C41 overexpression strain (Lucigen). The recombinant cultures were grown in 30mL TB medium (pH 7.0), with appropriate antibiotics, in 100 mL flasks. These cultures were grown at 37°C till OD600 reached 0.7. The temperature was then dropped to 16°C for 0.5 h prior to induction with 1 mmol/L isopropylthiogalactoside (IPTG), followed by supplementation with 40 mmol/L pyruvate and 1 mmol/L MgCl2. The induced cultures were grown for additional 48–72 h, then 3 mL of the cultures were used for extraction with an equal volume of hexane. The extraction process was the same in the in vitro assays.

2.8. Terpene product analysis by GC–MS chromatography

GC–MS analysis was carried on a Thermo TRACE 1310 gas chromatograph with a TQ8000 mass detector (Thermo Fisher Scientific) in electron ionization mode. A capillary column TR-5ms (30 mm × 0.25 mm ID; DF = 0.25 μm; Thermo Fisher Scientific) was used with a 1.0 mL/min helium flow rate. The temperature program was as follows: initial column oven temp 50°C, maintained for 2 min followed by a linear ramp at 40°C/min to 210°C; linear ramp at 5°C/min to 250°C; linear ramp at 40°C/min to 300°C, with a 5 min hold at 300°C. The ion trap temperature was 280°C. The terpenes were identified by a comparison of retention time and mass spectra to previously characterized enzymatic products or authentic standards.

2.9. DiTPS gene expression analysis

The expression of diTPS genes was obtained by two means: RNA-seq and qPCR analysis. For the RNA-seq, bowtie2 in RSEM software38 was used to map RNA-seq reads from the 13 tissues to the SMRT-based reference transcriptome. The read counts were then transformed to FPKM values to determine the gene expression levels of all identified genes. For qPCR analysis, 5 RNA extracts from the fibrous root, principal root, lateral root, leaf, and stem, respectively, were synthesized into cDNA by PrimerScript™ RT reagent kit with a gDNA eraser (TaKaRa Corp.). A SYBR Green kit (TaKaRa Corp.) for quantitative real-time polymerase chain reaction (qPCR) was applied to a ABI7500 real-time PCR detection system according to the manufacturer’s instructions. The primers sequences are listed in Table S1. Primer specificity was assessed by agarose gel and melting curve analysis. The results were normalized with the housekeeping gene Actin. Relative expression levels were calculated as the mean of three technical replicates of three biological replicates.

2.10. GenBank accessions

The full-length transcriptome reported in this paper has been deposited in China National Center for Bioinformation under accession number CRA003781, which is publicly accessible for all researchers at http://bgid.big.ac.cn/gsa. GenBank accession numbers for the functional terpene synthases described in this paper are AcCPS1 (MW478118), AcCPS2-1 (MW478119), AcCPS2-2 (MW478120), AcCPS2-3 (MW478121), AcKSL2-1 (MW478123), AcKSL2-2 (MW478124), AcKSL3-1 (MW478125), AcKSL3-2 (MW478126), AcTPS3 (MW478129) and AcTPS4 (MW478130). The sequences of different isoforms and E. coli codon optimized AcCPS1 were listed in Supporting Information isoform.seq.

3. Results

3.1. Full-length transcriptome sequencing using Iso-seq and RNA-seq platforms

Due to the relatively high DAs content in the top leaves, principal root, and lateral root1, we mixed RNA from these together tissues for Iso-seq. Concurrently, 13 tissues from differing spatial locations (Fig. 1A and B) were sequenced on an Illumina HiSeq 2000 platform to quantify gene/isoform expression levels and correct the Iso-seq reads. Utilizing 40.86 Gb sequenced data, we obtained 37, 182, 436 subreads (Supporting Information Table S3). The Iso-seq subreads’ length distribution was shown in Supporting Information Fig. S3. From the subreads we extracted 983,658 circular consensus sequence (CCS) reads. Subsequent trimming, assembly, filtering, classification, and clustering revealed that 52.95% were full-length reads, indicated by the detection of poly(A) or 3’ and 5’ sequences (Supporting Information Table S4). These were then used in the construction of full-length non-chimeric sequences (FLNCs), which ranged from 72 to 12,220 bp, with a 2023 bp average. The cDNA N50 values were determined to be 2612 bp. Of 520,860 full-length cDNAs, 48,240 polished, high-quality, isoform consensus reads were filtered out using arrow software. After isoform-level clustering, next generation short sequencing (Illumina) was performed for error correction, with redundant sequences removed via CD-HIT software, and 21,087 unigenes were generated for A. carmichaelii: average length 1991 bp, N50 value 2446 bp (Supporting Information Fig. S4). The N50 obtained in this study is 3-fold higher than previously reported (830 bp13 and 844 bp14), providing a novel opportunity to identify all TPS genes involved in DAs biosynthesis.

3.2. Identification of A. carmichaelii terpene synthases

To identify the A. carmichaelii TPS gene family, the above obtained unigenes were annotated using seven databases: Nr, Nt, Pfam, KOG/COG, Swiss-prot, KEGG, and GO (Supporting Information Fig. S5). We also mined the data using BLAST analysis against the enzymes like casbene and neocembrene synthases, which are responsible for macrocyclic or unusual skeleton formation22,23. In addition, for genes with unpredictable lengths we used BLAST analysis against the reported transcriptome database17. A total of 19 TPS unigenes were identified, of which 14 represented full-length sequences (Supporting Information Table S5). The remaining 5 represented alternative splicing events. Transcript13827 (1617 bp), transcript12676 (1443 bp) and transcript12546 (567 bp) had a 515 bp overlap (Supporting Information Fig. S6). The integration of transcript13827 and transcript12676 forms a common full-length CPS gene (average
2400 bp), annotated as AcCPS1. It was verified by polymerase chain reaction (PCR) amplification from the cDNA of fibrous root. At the same time, we obtained an isoform AcCPS1a, which had 98.70% sequence similarity with AcCPS1 (Supporting Information Fig. S7). We further used specific primers to clone the three alternative splicing isoforms. We got the expected PCR product of transcript13827 and transcript12676, while didn’t obtain any PCR product when using the specific primers of transcript12546. After sequenced the individual clones, we found nine of the transcript12676 clones were the same, which was nearly identical with AcCPS1 except the 11 nucleotides at the beginning of “ATG”. However, the condition was much more complexed for the clones from transcript13827. There were 37 nucleotide variations among the six clones and divided into three types. Four of them had the same sequence with AcCPS1a, clone-3-12 was identical with AcCPS1 except two nucleotide variations. Clone-4-12 was more different from the others. Sixteen nucleotides belonged to AcCPS1a, 8 nucleotides belonged to AcCPS1, and the other 13 nucleotides were different from both of them (Fig. S7).

Transcript11709 was 2247 bp, which was a little shorter than the common KSL genes (average 2400 bp). When blast against the reported transcriptome data we further got a full-length KSL gene with a 2454 bp open reading frame, which was annotated as AcKSL1. Further AcKSL1 and the transcript11709 were cloned from the cDNA of basal part of the stem. The process of cloning is not smooth for AcKSL1 (more than five times and from different organs); however, the cloning of transcript11709 was much easier. After sequencing, we got four isoforms of transcript11709. Clone-1-15 was identical with AcKSL1 besides the early termination. Clone-6-2 and clone-6-5 were extremely identical beside a three nucleotides insertion for clone-6-5 at the position of 123 bp, and they had about 97% sequence similarity with clone-1-15. Clone-1-8 is much more different from others, with about 94.7% sequence similarity with AcKSL1. At the same time, the stop codon of clone-1-8 mutated into “GGA”, which resulted in the absence of a complete open reading frame (Supporting Information Fig. S8).

At the process of cloning the other genes from cDNA, we further obtained two more isoforms of AcCPS2-3. They have more than 98.6% sequence similarity with AcCPS2-3 (Supporting Information Fig. S9). In summary, we cloned all the genes from various tissues except transcript19787 (AcTPS1). We further blast these TPS genes against the previously reported transcriptome13, it showed that nearly all genes except AcTPS4 could found corresponding unigenes with similarity higher than 92%.
(Supporting Information Table S7). However, all these unigenes didn’t represent the full-length cdDNAs except TR36313|c4_g1_i1, which corresponded to AcTPS2. Unigene TR48214|c4_g1_i1 had the highest similarity with AcKSL3-1 (98%), while when using AcKSL1 and AcKSL3-2 as the query, the result also to be TR48214|c4_g1_i1 with similarity of 92%. These results demonstrated that the Pacbio full-length platform enabled an in-depth investigation of the TPS gene families in A. carmichaelii. In order to make the result clear, the following analysis mainly focused on 16 TPS genes which had identical sequence with the transcriptome data (Table S6), and generally didn’t contain various isoforms cloned by PCR, unless otherwise specified.

3.3. Phylogenetic relationship of A. carmichaelii terpene synthases

We further categorized the 16 TPS genes by signature sequence motif analysis and phylogenetic comparison with TPS families from A. thaliana and S1 previously identified related functional diTPS (Fig. 2). The presence of the catalytic DXDD motif in combination with a close phylogenetic relationship to known TPSs of the TPS-c clade, illustrated that 4 TPS genes were linked to class II diTPS: AcCPS1 and AcCPS2-1 through AcCPS2-3 (Supporting Information Fig. S11). These 4 enzymes formed a single group, which differs from class II diTPS in other species (i.e., Salvia miltiorrhiza and Isodon rubescens) where class II diTPS usually form two distinct groups. AcCPS1 shares ~64% identity with AcCPS2-1 through AcCPS2-3, and the 3 AcCPS2s share >87% sequence identity. Due to the high consistency of AcCPS2s, we speculate that they may come from duplication in a single chromosome or from a similar allele of different chromosomes, given that A. carmichaelii has been reported to be a polyploidy plant. The close relationship found between CmCPS1 and CmCPS2, and other CPSs in S. miltiorrhiza and I. rubescens that have been linked to ent-CPP production indicates that AcCPS1 and AcCPS2s are ent-CPP synthases.

The 12 remaining candidate genes were designated as class I TPS, according to the presence of the conserved DDXXD motif (Fig. S11). Of these TPSs, 5 were clustered with diTPSs from the TPS-e/f clade: AcKSL1, AcKSL2-1, AcKSL2-2, AcKSL3-1, and AcKSL3-2 (Fig. 2). Their 5 respective enzymes were also clustered together. AcKSL1 shares ~63% identity with AcKSL2-1 and AcKSL2-2, and ~85% identity with AcKSL3-1 and AcKSL3-2. AcKSL2-1 and AcKSL2-2 share 98% sequence identity, while AcKSL3-1 and AcKSL3-2 share 92% sequence identity, again suggesting they are allelic variants or from tandem duplicates. The other 7 TPSs, AcCPS1 to AcCPS7, were determined to be within the TPS-b subfamily. Most of the characterized TPSs in this subfamily are monoterpane synthases or isoprene synthases. AcCPS1 and AcCPS2 share 98% identity, AcCPS5 and AcCPS7 share 87%, and the others share a substantially lower sequence identity (~40%). This clade included TwTPS27, whose function is corollary to diTPS and can convert copalyl diphosphate to miltiriadiene in Tripterygium wilfordii. These results indicate that all these TPS candidates have the potential to be involved in A. carmichaelii DAs biosynthesis.

3.4. Functional characterization of class II terpene synthases from A. carmichaelii

Based on the full-length transcriptome sequencing, we cloned all the AcCPSs from A. carmichaelii, including seven full-length
AcCPSs and two alternative splicing isoforms of AcCPS1 (Table S6). We then functionally characterized these AcCPSs through in vitro assays with recombinant proteins expressed via E. coli (Supporting Information Fig. S12). We first separately incubated the AcCPSs with GGPP to compare with the results from ent-CPP synthase ZmCPS2 of Zea mays 37. All enzymes except the two alternative splicing isoforms yielded a product with identical retention time and mass spectra to the product of ZmCPS2 after dephosphorylation (Fig. 3A and B), indicating that the primary product of these AcCPSs is CPP.

To further investigate the stereochemistry of the CPP products, these enzymes were coupled to stereospecific class I diterpene synthases, such as SmKSL133,43 (specific to normal-CPP) and IrKLS539 (specific to ent-CPP). Reactions where AcCPSs were coupled with IrKLS5 resulted in the formation of ent-kaurene, while no product was evolved when they were coupled with SmKSL1 (Fig. 3A). That all AcCPSs could produce ent-CPP indicates their involvement in A. carmichaelii DAs biosynthesis.

3.5. Functional characterization of class I terpene synthases from A. carmichaelii

We cloned 14 class I TPS genes for functional identification except AcTPS2, which included 5 KSLs: AcKSL1, AcKSL2-1, AcKSL2-2, AcKSL3-1 and AcKSL3-2, and 6 TPS-b subfamily genes: AcTPS1, AcTPS3 through AcTPS7, together with three alternative splicing isoforms of AcKSL1. Having identified only the ent-CPP activity of the 7 AcCPSs, we first identified the products of the 5 class I diTPS AcKSLs and three alternative splicing isoforms of AcKSL1 in combination with ent-CPP synthase (AcCPS2-1). When GGPP was used as the substrate, AcKSL2-1 and AcKSL2-2 turned ent-CPP into ent-atierase, a result identical to IrCPS4 in combination with IrKSL4 in I. rubescens; AcKSL3-1 produced ent-kaurene, identical to the product of IrCPS4 coupled with IrKLS539, and AcKSL3-2 coupled with AcCPS2-1 produced a product similar to ent-13-epi-sandaracopimaradie44 (Fig. 4). We further synthesized two known ent-sandaracopimaradie synthases, EpTPS23 from Euphorbia peplus44 and OsKSL10 from Oryza sativa45, along with sandaracopimaradie synthase (MsTPS1) from Mentha Spicata46 to verify the product stereochemistry of AcKSL3-2. When AcKSL3-2, EpTPS23, and OsKSL10 were respectively combined with AcCPS2-1, the product mass spectra of AcKSL3-2 revealed to be identical to the products of EpTPS23 and OsKSL10, though its retention time was ~0.8 min earlier (Fig. 4). While sandaracopimaradie, the product of MsTPS1 combined with normal-CPP synthase (SmCPS1 from S. miltiorrhiza33), had exactly the same retention time and mass spectra with the product of EpTPS23/OsKSL10. These results confirmed that the product of MsTPS1 with normal-CPP and EpTPS23/OsKSL10 with ent-CPP were enantiomers, while the product of AcKSL3-2 was the C-13 epimer due to the fact that they all originated from ent-CPP. Thus, the product of AcKSL3-2 was assigned to be ent-13-epi-sandaracopimaradie.

Due to the poor protein expression of AcKSL1 and its three alternative splicing isoforms in E. coli, we could not identify its...
function in vitro. Therefore, we used a highly efficient diterpene modular metabolic engineering system\textsuperscript{35,36} to further verify the function of all 8 of the KSLs. Three vectors, pIRS, pGG-ZmCPS2, and pET32-AcKSLs, were transfected into the C41 over-expressing strain of \textit{E. coli}. After incubation for 72 h the product was analyzed by GC–MS. The results of AcKSL2s and AcKSL3-1 were analogous to the above in vitro assays. AcKSL3-2 produced two other products in addition to \textit{ent}-13-\textit{epi}-sandaracopimaradiene: one was \textit{ent}-kaurene at 10.30 min based on the same retention time and mass spectra with the product of AcKSL3-1. The other product at 10.75 min showed to be an unidentified diterpene. \textit{Ent}-atiserene was produced by the \textit{E. coli} codon optimized AcKSL1, which had identical retention time and mass spectra with the products of AcKSL2s (Fig. 5). No obvious products were produced by the none codon optimized AcKSL1 and its three alternative splicing isoforms.

To investigate whether class I TPS genes have the capacity to produce diterpenes from GGPP, we first incubated all the recombinant class I TPS with GGPP. No product was formulated in these experiments (Supporting Information Fig. S13). We next incubated the 6 TPS-b enzymes with GGPP and AcCPS2-1, but again no products were formed. We thus conclude that in \textit{A. carmichaelii}, DAs are only produced by the combination of the TPS-c and TPSe/f subfamily genes. In order to identify the function of the TPS-b genes, we incubated them with FPP and GPP, respectively. Our results showed that two genes, AcTPS3 and AcTPS4, generated a product similar to farnesene with FPP, which was identified to be \textit{a}-farnesene with authentic standard \textit{b}-farnesene and \textit{b}-disabolene (Fig. S13), illustrating they are sesquiterpene synthases. The others showed no enzymatic activity in vitro assay though AcTPS1 and AcTPS6 had high quality of recombinant proteins (Fig. S12).

### 3.6. Gene expression patterns of \textit{A. carmichaelii} terpene synthases

In order to investigate the potential physiological roles of the candidate TPSs, we performed RNA sequencing across 13 tissues based on spatial location. Six were underground tissues: principal root, fibrous root, cortex, phloem and xylem from lateral root, and the basal part of the stem; and seven were aerial tissues: the top, central, and lower parts of the leaf and stem respectively, and the nodular like organs in the stem (Fig. 1A and B).

Thirteen tissues were able to be differentiated into two groups according to their spatial locations, with the aerial tissues forming one group and the underground tissues the other, illustrating that the expression of TPS genes indeed has a spatial difference (Fig. 6A). The 16 TPS genes formed 3 groups: group I included 3 AcCPS2s, AcKSL1, and AcKSL3-2, all of which had similar moderate expression levels in across all tissues; group II included 4 TPS-b subfamily genes (AcTPS1, AcTPS2, AcTPS3, and AcTPS6) and AcKSL3-1, which had relatively high expression levels in the aerial leaf and stem; and group III included AcKSL2-1, AcKSL2-2, AcCPS1 and 3 TPS-b subfamily genes (AcTPS4, AcTPS5, and AcTPS7), with relatively high expression in the underground tissues, especially for AcKSL2-1 and AcKSL2-2. Notably, AcCPS1 had specifically high expression levels in the fibrous root samples.

In order evaluate the expression levels from RNA-seq, we further evaluated the mRNA transcription levels of 4 diTPSs in the principal root, lateral root, fibrous root, stem, and leaves by qPCR (Fig. 6B). The results were in agreement with the RNA-seq data: AcCPS1 was specifically expressed in the fibrous root; AcCPS2-1 had relatively high expression levels in all tissues; AcKSL2-1 showed relatively high expression level in principal and lateral tissues, which was in line with the RNA-seq data.
root, but lower expression level in stem and leaves; and AcKSL1 had relative low expression levels in all the tissues.

4. Discussion

Diterpenoid alkaloids (DAs) are among the most well-known medicinal compounds of Aconitum. Most previous studies have focused on phytochemical composition and pharmaceutical usage of DAs, and significantly less research has gone into the molecular biology of DAs formation in Aconitum. To better understand the molecular basis for the complex biosynthesis of DAs; herein, we applied Pacbio full-length transcriptome in conjunction with RNA-seq and PCR cloning to mine for the TPS family and their potential roles, searching specifically for enzymes involved in DAs metabolism. Using this combination approach, we identified 19 TPS genes and five alternative splicing isoforms in A. carmichaelii. The diTPS, excepting the alternative splicing isoforms, were functionally identified using in vitro and in vivo diterpene metabolic engineering system. Using in vitro assays—with GPP, FPP, and GGPP as substrates—we eliminated the potential involvement of the TPS-b subfamily in DAs biosynthesis, and established biosynthetic routes to the diterpene scaffolds ent-atiserene, ent-kaurene, and ent-13-epi-sandaracopimaradiene (Fig. 7) based on 12 diTPS genes, clarifying the molecular basis for A. carmichaelii DAs biosynthesis.

The AcTPSs found in the TPS-c and TPS-e/f subfamilies cluster together, analogous to the diTPS found in T. wilfordii25, indicating a recent evolutionary event may responsible to the emergence of different classes of diterpene scaffolds. All of the class II diTPS genes proved to be ent-CPP synthase, which is accord with the common view that ent-CPP is the only intermediate involved in A. carmichaelii DAs biosynthesis10. Notably, AcCPS1 is specifically expressed in the fibrous root, as verified by RNA-seq and qPCR analysis, raising interesting questions about its function that necessitate further study. Due to their relatively high expression levels in underground tissues, AcKSL2-1 and AcKSL2-2 are presumed to be the key enzymes involved in DAs biosynthesis. If this is the case, the product of AcKSL2s, ent-atiserene, would be the only precursor for atisane-skeleton C_{20}-DAs, such as the atisine-, demudatine-, hetidine-, hetisine- or vakognanine-types, which is agreement with the previous hypothetical biosynthetic pathway based on chemical structure1. Besides AcKSL2s, AcKSL1 also identified to be an ent-atiserene synthase in A. carmichaelii. The relative low expression level of AcKSL1 in different tissues (Fig. 6) and the close phylogenetic relationship with ent-kaurene synthase (AcKSL3-1) (Fig. 2) illustrated AcKSL1 might be the ancestral enzyme for atisane-skeleton C_{20}-DAs biosynthesis, and AcKSL2s undergone a functional divergence from AcKSL1, which shed light on the evolutionary origin of C_{20}-DAs biosynthesis. The product of AcKSL3-1 was ent-kaurene, which would be the precursor for the biosynthesis of gibberellin and napelline-type C_{20}-DAs. The product of AcKSL3-2 was found to be ent-13-epi-sandaracopimaradiene, making it the first enzyme to be identified to synthesize this diterpene. Other ent-sandaracopimaradiene-like compound derivatives have been isolated from several species including rice45, Kaempferia galanga27, and Guarea rhophalocarpa48; however, it has not reported previously in Aconitum. On the other hand, beside ent-13-epi-sandaracopimaradiene, it also produced ent-kaurene and another unknown diterpene in the engineered E. coli. Further study into the chemistry and metabolic fate of these diterpenes in planta will be needed to fully elucidate its role. Collectively, AcCPS1, AcCPS2s, AcKSL1, AcKSL2s, and AcKSL3-1 appear to be responsible for all the known C_{20}-DAs biosynthesis in A. carmichaelii.

Many secondary metabolites are synthesized in highly specific tissues, such as tanshinones and forskolin in the roots. More specifically, tanshinones are synthesized in the root periderm of S. miltiorrhiza27,33 and forskolin is synthesized in the root cork of Coleus forskohlii68. Accordingly, the diTPS involved in their biosynthesis have specific, high expression levels in these organs27,33,48. In our recent study focused on DAs metabolites in different species, we found that despite the principal and lateral root being considered the primary medicinal components of A. carmichaelii, the characteristic DAs including aconitine, mesaconitine and hypeaconitine have a relatively high content in stem and leaves as well1. Indeed, when total DAs content was assessed, the top leaves contained the highest DAs content3. It is therefore difficult isolate which enzyme or substrate is responsible for the large array of C_{16}-DAs biosynthetic pathways of A. carmichaelii.
based on gene expression patterns and metabolite locations alone. Due to the high expression levels of AcKSL2s, we postulate that the atisine type C20-DAs are the main precursor for C19-DAs; however, further proof by RNA interference using microprojectile bombardment (gene gun) or virus-induced gene silencing technology to knock down AcKSLs gene expression in planta or through isotope labeling analysis is needed.

The Pacbio isoform sequencing offered information about alternative splicing transcripts 27,50. In terpene synthase, there were few reports about functional alternative splicing transcripts. In S. miltiorrhiza, the kaurene synthase like gene SmKSL1 and several genes involved in production of isoprenoid precursors were reported to exhibit alternative splicing without functional identification. In I. rubescens, the alternative splicing IrKLS3a was identified to alter the product outcome. IrKSL3 produced miltiradiene, while IrKSL3a produced isopimaradiene and miltiradiene 51. Here, we identified two alternative splicing transcripts of AcCPS1 and three of AcKSL1 in A. carmichaelii. However, none were identified to be functional genes. Compared with the full-length gene, clone-4-12 from transcript13827 lost a domain of CPS, while transcript12676 lost g and partial b domain52, thus it is reasonable for them to lose dTPS functions. However, we are surprised by the loss of function of AcKSL1 and its three alternative splicing transcripts in vitro. This was partly due to the low solubility of recombinant protein in prokaryotic expression, after codon optimized of E. coli for AcKSL1, we finally identified its function in the highly efficient diterpene modular metabolic engineering system 35,36, indicating we should attempt more platforms such as by transient expression of these genes in Nicotiana benthamiana 50 to characterize their functions.

In conclusion, by combining full-length Iso-seq and RNA-seq we obtained a robust A. carmichaelii transcriptome and identified all the TPS family genes involved in A. carmichaelii DAs biosynthesis. While not conclusive, our results highlight the atisine type C20-DAs as the predominate precursor of the medicinal C19-DAs, clarify the biosynthetic pathway for C20-DAs in A. carmichaelii, and pave the way for further exploration of C19-DAs biosynthesis pathway in Aconitum.

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Author contributions
Liuying Mao, Guanghong Cui, Juan Guo and Luqi Huang designed the research. Liuying Mao performed most of the experiment. Changjiangsheng Lai, Biwei Yin prepared the plant material for sequencing. Tong Chen analyzed the transcriptome data. Rui Ma, Haiyan Zhang performed the qPCR analysis. Baolong Jin, Lingli Chen and Mei Tian confirmed the catalytic analysis. Liuying Mao, Baolong Jin and Guanghong Cui wrote the manuscript. Juan Guo, Jinfu Tang revised the manuscript. All authors have read and approved the final manuscript.

Conflict of interest
The authors claim that the researchers in this study have no conflicts of interest.

Appendix A. Supporting information
Supporting information to this article can be found online at https://doi.org/10.1016/j.apsb.2021.04.008.

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