Characterization of a Cytosolic Acyl-Activating Enzyme Catalyzing the Formation of 4-Methylvaleryl-CoA for Pogostone Biosynthesis in Pogostemon Cablin

Jing Chen, Lang Liu, Ying Wang, Zhengguo Li, Guodong Wang, George A. Kraus, Eran Pichersky and Haiyang Xu

1 School of Life Sciences, Chongqing University, Chongqing 401331, China
2 Center of Plant Functional Genomics, Institute of Advanced Interdisciplinary Studies, Chongqing University, Chongqing 401331, China
3 State Key Laboratory of Plant Genomics and National Center for Plant Gene Research, Institute of Genetics and Developmental Biology, Chinese Academy of Sciences, Beijing 100101, China
4 Department of Chemistry, Iowa State University, Ames, IA 50011, USA
5 Department of Molecular, Cellular, and Developmental Biology, University of Michigan, Ann Arbor, MI 48109, USA

*Corresponding author: E-mail, hyxu@cqu.edu.cn
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Pogostone, a compound with various pharmaceutical activities, is a major constituent of the essential oil preparation called Pogostemonis Herba, which is obtained from the plant Pogostemon cablin. The biosynthesis of pogostone has not been elucidated, but 4-methylvaleryl-CoA (4MVCoA) is a likely precursor. We analyzed the distribution of pogostone in P. cablin using gas chromatography-mass spectrometry (GC-MS) and found that pogostone accumulates at high levels in the main stems and leaves of young plants. A search for the acyl-activating enzyme (AAE) that catalyzes the formation of 4MVCoA from 4-methylvaleric acid was launched, using an RNAseq-based approach to identify 31 unigenes encoding putative AAEs including the PcAAE2, the transcript profile of which shows a strong positive correlation with the distribution pattern of pogostone. The protein encoded by PcAAE2 was biochemically characterized in vitro and shown to catalyze the formation of 4MVCoA from 4-methylvaleric acid. Phylogenetic analysis showed that PcAAE2 is closely related to other AAE proteins in P. cablin and other species that are localized to the peroxisomes. However, PcAAE2 lacks a peroxisome targeting sequence 1 (PTS1) and is localized in the cytosol.

Keywords: 4-Methylvaleric acid • 4-Methylvaleryl-CoA • Acyl-activating enzyme • Branched-chain fatty acid • Pogostemon cablin • Pogostone

Introduction

Pogostemonis Herba, commonly known as ‘Guang-Huo-Xiang’ in Chinese and patchouli in English, is produced from the dried aerial part of the plant Pogostemon cablin (Blanco) Benth. (Labiatae). As Pogostemonis Herba and its essential oil (patchouli oil) possess variety of pharmacological activities (Lu et al. 2011, He et al. 2013, Li et al. 2013, Wang et al. 2016), they have widely been used in many Asian countries as traditional Chinese medicine for the treatment of fatigue, summer heat, nausea, vomit, abdominal distension and other ailments since ancient time (Zhao et al. 2005, Chen et al. 2015, Hu et al. 2017). Patchouli oil has also been widely used for perfumes, cosmetics and food stuffs (Pattnaik et al. 1996, Lin et al. 2014, Yang et al. 2016). Pogostone (formula C_{17}H_{16}O_2, Fig. 1) is one of the major constituents of patchouli oil and is largely responsible for the intensive aromatic odor and, therefore, designated as a chemical marker for the quality control of patchouli oil (Swamy and Sinniah 2015, Yang et al. 2016). This compound has been demonstrated to exert various bioactive activities including anti-bacterial (Peng et al. 2014), anti-candida (Li et al. 2012), antifungal (Li et al. 2012, Luchesi et al. 2020), anticanter (Cao et al. 2017), insecticidal (Huang et al. 2014), anti-inflammatory (Li et al. 2014), immunosuppressive (Su et al. 2015) and gastroprotective (Chen et al. 2015) activities. The levels of pogostone in patchouli oil, mainly obtained from stems and leaves of Pogostemon plants, vary among samples collected from different geographic locations and at different periods (Luo et al. 2003, Li et al. 2004, Hu et al. 2006).

The biochemical steps involved in the biosynthesis of pogostone are currently unknown. It is likely that one or more polyketide synthases that use various starter molecules are involved (Fig. 1A). One obvious such starter is 4-methylvaleryl-CoA (4MVCoA), derived from the branched-chain fatty acid 4-methylvaleric acid (Fig. 1B). Acyl-CoA thioesters such as 4MVCoA are formed by members of the acyl-activating enzyme (AAE) superfamily that activate carboxylic acids through a...
Fig. 1 The proposed biosynthetic pathway of pogostone and its 4MVCoA precursor in *P. Cablin*. A, Proposed alternative pogostone biosynthetic pathways: 1) A pathway involving one type III polyketide synthase (PKSIII) that catalyzes the condensation of acetyl-CoA and malonyl-CoA to form 2-pyrone (4-Hydroxy-6-methyl-2-pyrone), which is then acylated by one typical acyltransferase using 4MVCoA as substrate to form an unstable intermediate, followed by spontaneous rearrangement to form pogostone. 2) A pathway involving one PKSIII that catalyzes the condensation of 4MVCoA and malonyl-CoA to form a tetraketide intermediate, which is then rearranged (spontaneously or by a cyclase) and cyclized by a cyclase to form pogostone. 3) A pathway involving one typical PKSIII that catalyzes the condensation of acetyl-CoA and malonyl-CoA to form a triketide intermediate, followed with condensation catalyzed by an unusual polyketide synthase with 4MVCoA as the extension unit to form a branched tetraketide intermediate, which is then rearranged by a cyclase to form pogostone. B, Proposed biosynthetic pathway of 4MVCoA. 4-Methylvaleric acid is derived from pyruvate through α-ketoacid elongation pathway, and catalyzed by an AAE to form 4MVCoA. PKSIII, type III polyketide synthase; AT, Acyltransferase; AAE. The carbon chain of the branched short-chain fatty acid precursor 4-methylvaleric acid and 4MVCoA is labeled in bold to show its incorporation into pogostone.
two-step mechanism. First, the carboxylic acid group is adenylated to form an enzyme-bound acyl-AMP intermediate by ATP pyrophosphorylisis, and this step is followed by the displacement of AMP with a CoA group to form the relevant acyl-CoA thioester (Shockey and Browse 2011) (Fig. 1B). One of the unifying features of this superfamily is the well-conserved 12 amino acid residue AMP-binding motif (PROSITE PS00455), which is used to identify putative genes encoding AAEs (Shockey et al. 2003). Besides carboxylic acids, various plant hormones are also conjugated to amino acids by members of this superfamily (Nobuta et al. 2007, Okrent et al. 2009).

We have begun identifying and characterizing the enzymes involved in the biosynthesis of pogostone. The combined analysis of transcriptome and metabolome of different tissues at different developmental stages has been effectively used to discover enzymes involved in the specialized metabolic pathway in many species such as pyrethrum (Xu et al. 2018, 2019). To facilitate the elucidation of the pogostone biosynthetic pathway, a transcriptome assembly of P. cablin was generated in this study from RNAseq analysis of seedling, root, stem and leaf harvested at different developmental stages. GC-MS analysis established that pogostone is largely present in seedlings, the main stems and top leaves of main stem of P. cablin at earlier developmental stage and showed a developmentally specific accumulation pattern in top leaves of the main stem of P. cablin.

Here, we report the identification and characterization of the enzyme catalyzing the formation of 4MVCoA. Through transcriptome analysis of P. cablin RNAseq database, we identified 31 putative AAE genes including PcAAE2, the transcript profile of which shows a positive correlation with the accumulation pattern of pogostone. Through in vitro enzymatic assays, we identified PcAAE2 as encoding 4MVCoA synthetase. We further showed that PcAAE2 is a cytosolic enzyme that may have evolved from an ancestral peroxisome-targeted AAE.

Results

Distribution of pogostone in P. cablin top leaves newly generated from the main stems shows a developmental pattern

The levels of pogostone in P. cablin seedlings, roots, main stems and top leaves from the main stems at different developmental stages (Fig. 2A) were determined by gas chromatography-mass spectrometry (GC-MS) analysis of methyl tert-butyl ether (MTBE)-extracted macerated tissues. This analysis showed that pogostone is largely produced in P. cablin seedlings and in the main stems and the top leaves of the main stem of plants younger than 7 weeks (Fig. 2B). Plants older than 7 weeks contained negligible amounts of pogostone compared to younger plants (Fig. 2B), which indicates that the distribution of pogostone in P. cablin newly generated leaves from the main stem is developmentally controlled (Fig. 2B). Besides pogostone, another compound related to pogostone (with one carbon shorter in the side chain and likely derived from isovaleryl-CoA),

Fig. 2 Distribution analysis of pogostone and pogostone B in different tissues of P. cablin at different developmental stages. A. Different developmental stages of P. cablin. B. Concentration of pogostone and pogostone B in different tissues of P. cablin at different developmental stages. The organic extracts of tissues were analyzed by GC-MS and quantification was achieved by normalization of the peaks to the internal standard tetracane and comparison with standard curves of authentic pogostone and pogostone B (Data are means ± 5D of four independent biological replicates). FW, Fresh weight. The tissues used in this study are described as follows: The seedling used in this study is 4-week-old, 5W-TLeaf, the top leaves from main stem of 5-week-old P. cablin plant; 5W-Stem, the main stems of 5-week-old P. cablin plant; 5W-Root, the roots of 5-week-old P. cablin plant; 6W-TLeaf, the top leaves from main stem of 6-week-old P. cablin plant; 6W-Stem, the main stems of 6-week-old P. cablin plant; 6W-Root, the roots of 6-week-old P. cablin plant; 7W-TLeaf, the top leaves from main stem of 7-week-old P. cablin plant; 7W-Stem, the main stems of 7-week-old P. cablin plant; 7W-Root, the leaves from main stem of 7-week-old P. cablin plant are beneath and most closest to top leaves; 8W-TLeaf, the top leaves from main stem of 8-week-old P. cablin plant; 8W-Stem, the main stems of 8-week-old P. cablin plant; 8W-Root, the leaves from main stem of 8-week-old P. cablin plant are beneath and most closest to top leaves; 9W-TLeaf, the top leaves from main stem of 9-week-old P. cablin plant; 9W-Stem, the leaves from main stem of 9-week-old P. cablin plant are beneath and most closest to top leaves. The top leaves used in this study are two symmetrical leaves with the main vein of around 2 cm. It is noteworthy that 7W-SLeaf is converted from 6W-TLeaf after 1 week, similar cases for 8W-SLeaf converted from 7W-TLeaf and 9W-SLeaf converted from 8W-TLeaf.
which we named pogostone B, was also detected in *P. cablin* plants (Supplementary Fig. S1). Pogostone B showed a very similar distribution pattern to that of pogostone in the tested tissues, albeit present at much lower levels (Fig. 2B).

**Identification of candidate genes responsible for 4MVCoA biosynthesis**

To identify the genes encoding the enzymes involved in the conversion of 4-methylvaleric acid to 4MVCoA, RNAseq libraries were constructed from six different *P. cablin* tissue samples including Seedling, SW-Root, SW-Stem, SW-Leaf, 8W-TLeaf and 8W-SLeaf from *P. cablin* plants (Fig. 2A). The corresponding transcriptome assemblies were then constructed and unigenes were annotated by comparison with genes on the UniProt database and using the BLASTX program. AAE genes of *P. cablin* were identified by using Arabidopsis representative AAE sequences of each of the seven clades of the AAE superfamily, which had been previously defined (Shockey and Browse 2011), to query our transcriptome assembly using TBlastN software. In addition, we searched the functional annotations of the unigene with keywords such as ‘AMP-binding protein’. Since we sought to search for the gene responsible for 4MVCoA biosynthesis from 4-methylvaleric acid, the unigenes encoding putative AAEs catalyzing the conjugation of plant hormones to amino acids were not included in this search. A total of 31 unigenes encoding putative AAEs, containing the conserved AMP-binding motif (PROSITE PS00455) of the AAE family, were identified and further analyzed (Table 1, Supplementary Tables S1 and S2).

Previous phylogenetic analysis has defined seven subfamilies of AMP-dependent synthetases and ligases in Arabidopsis (Shockey and Browse 2011). Phylogenetic analysis indicated that all the 31 *P. cablin* AAE unigenes group into the previously established six clades of the AAE superfamily (Supplementary Fig. S2). Several members of clade VI from *Arabidopsis thaliana* and *Humulus lupulus* have been previously functionally analyzed by in vitro biochemical assays. HICCL2 and HICCL4 from *H. lupulus* utilize various short-chain fatty acids and branched short-chain fatty acids as substrates (Xu et al. 2013). Both of these CCLs prefer branched short-chain fatty acids as substrates, with isovaleric acid and isobutyric acid as the preferred substrates for HICCL2 and HICCL4, respectively (Xu et al. 2013). The Arabidopsis gene At2G17650 encodes an enzyme that also prefers branched short-chain fatty acids as the substrates, with isovaleric acid serving as the most preferred substrate in *in vitro* enzymatic assays (Xu et al. 2013). Recently, identified HcAAE1 from *Hypericum calycinum* exhibits promiscuous substrate preference, with benzoic acid and several straight/branched short-chain fatty acids such as butyric acid and isobutyric acid as preferred substrates (Singh et al. 2020). These data suggest that there might also be one or more members among the *P. cablin* clade VI genes that encodes an enzyme with a preference for branched-chain fatty acids, including 4-methylvaleric acid. Therefore, we concentrated on the eight genes in the *P. cablin* clade VI, which we designated as PcAAE1–8, for further examination (Table 1 and Supplementary Table S1).

| Name   | BLASTP against Arabidopsis genome (functional annotation) | Identity (%) | Peptide length (amino acids) | Signal peptide prediction (using Target P 2.0 and PTS1 Predictor) |
|--------|----------------------------------------------------------|--------------|-----------------------------|---------------------------------------------------------------|
| PcAAE1 | AT2G17650, isovaleryl-CoA ligasea                         | 67           | 584                         | mTP (0.6691), cTP (0.0015), luTP (0.0005), PTS1 (−24.573)     |
| PcAAE2 | AT2G17650, isovaleryl-CoA ligasea                         | 64           | 543                         | mTP (0.0021), cTP (0.0296), luTP (0.0038), PTS1 (−25.047)     |
| PcAAE3 | AT3G16910, acetate/butyrate-CoA ligaseb                    | 78           | 568                         | mTP (0.0011), cTP (0.0083), luTP (0.0015), PTS1 (12.027)     |
| PcAAE4 | AT1G65890, AAE 12                                        | 61           | 590                         | mTP (0.0117), cTP (0.0144), luTP (0.0008), PTS1 (6.968)      |
| PcAAE5 | AT5G16340, AAE 6                                         | 63           | 553                         | mTP (0.01), cTP (0.1781), luTP (0.0198), PTS1 (6.992)        |
| PcAAE6 | AT5G16370, AAE 5                                         | 65           | 551                         | mTP (0.00), cTP (0.02914), luTP (0.0014), PTS1 (5.274)       |
| PcAAE7 | AT1G20560, butyryl-CoA ligasea                            | 70           | 555                         | mTP (0.004), cTP (0.006), luTP (0.0135), PTS1 (11.738)       |
| PcAAE8 | AT1G21540, AAE 9                                         | 65           | 609                         | mTP (0.0007), cTP (0.2914), luTP (0.0014), PTS1 (−35.162)   |

**Table 1** Bioinformatic analysis of the eight AAEs in the clade VI of AAE superfamily of *P. cablin*

*a*The in vitro enzymatic activities have been reported (Shockey et al. 2003, Xu et al. 2013). mTP, mitochondrial transit peptide; cTP, chloroplast transit peptide; luTP, thylakoid luminal transit peptide; PTS1, peroxisomal targeting signal 1. The websites of the PTS1 Predictor and TargetP 2.0 are http://mendel.imp.ac.at/pts1/ and http://www.cbs.dtu.dk/services/TargetP/, respectively.

**Tissue-specific/developmental expression patterns of PcAAE clade VI genes**

Analysis of expression patterns of the eight genes in *P. cablin* RNAseq database indicated that PcAAE2 expression shows the strongest positive correlation with the distribution pattern of pogostone in these tissues of all these eight genes (Supplementary Table S1; Fig. 2B). In particular, PcAAE2 transcript reads are much lower in older tissues (8W-TLeaf and 8W-SLeaf) than in seedling and younger plants (SW-Stem and SW-TLeaf). To more accurately test whether the transcript profiles of PcAAE1–8 in the clade VI of AAE superfamily are correlated with the distribution pattern of pogostone in different tissues of *P. cablin*, their expression patterns were further confirmed by quantitative reverse transcription polymerase chain reaction (RT-qPCR) in the same tissue samples used for distribution analysis of pogostone (Fig. 3). The expression patterns of PcAAE1–8 obtained by RT-qPCR analysis are mostly consistent with that in RNAseq database (Supplementary Table S1; Fig. 3). Again, the expression patterns of PcAAE2 in the tested samples show a strong positive correlation with the distribution pattern of pogostone: (i) PcAAE2 is highly expressed in the five
Fig. 3 RT-qPCR analysis of transcript levels of PcAAE1-8 in different tissues of *P. cablin* at different stages of development. The RT-qPCR data were expressed as relative expressions of PcAAE1-8 relative to *P. cablin* GAPDH (glyceraldehyde-3-phosphate dehydrogenase), Actin7 and Tubulin3 in each tissue sample. Data are means ± SD of four independent biological replicates. Significant difference among the expression in different tissues of *P. cablin* at different stages of development was tested using independent *t*-tests *P* < 0.05 [Two-tailed distribution; Two-sample unequal variance (heteroscedastic)]. Different letters above the bars mark the statistically significant groups after *t*-tests.

**PcAAE2 encode a 4MVal-CoA synthetase**

Sequence alignment showed that of the eight *P. cablin* clade VI proteins, PcAAE1 and PcAAE2 share the highest amino acid sequence identities with HICCL2 (66.0% and 66.2%, respectively) and HcAAE1 (74.6% and 68.9%, respectively) (Supplementary Fig. S3) and, thus, are the closest homologs to HICCL2 and HcAAE1, suggesting that they may have the similar
Table 2 Substrate Specificity of Recombinant PcAAE1, PcAAE2 and PcAAE3

| Substrate         | Structure | PcAAE1          | PcAAE2           | PcAAE3          |
|-------------------|-----------|-----------------|------------------|-----------------|
| Acetic acid       |           | N.D.            | 5 ± 1.2          | 47 ± 0.5        |
| Propionic acid    |           | 5 ± 0.2         | 6 ± 0.6          | 100* ± 2.2      |
| Butyric acid      |           | 69 ± 1.3        | 85 ± 5.4         | 74 ± 1.8        |
| Isobutyric acid   |           | N.D.            | 6 ± 0.7          | 42 ± 1.0        |
| Valeric acid      |           | 36 ± 0.6        | 89 ± 2.4         | 12 ± 0.1        |
| Isovaleric acid   |           | 100a ± 4.9      | 51 ± 2.4         | N.D.            |
| Hexanoic acid     |           | 11 ± 0.4        | 52 ± 3.5         | N.D.            |
| 4-Methylvaleric acid |      | 21 ± 0.6        | 100b ± 8.6       | N.D.            |
| Heptanoic acid    |           | N.D.            | 54 ± 3.6         | N.D.            |
| 5-Methylhexanoic acid |    | N.D.            | 51 ± 3.1         | N.D.            |
| Octanoic acid     |           | 9 ± 1.3         | 10 ± 0.4         | N.D.            |
| Benzoic acid      |           | N.D.            | 13 ± 0.3         | N.D.            |

These enzymatic assays were performed using the coupled enzymes assay (Koo et al. 2006, Chen et al. 2011). These activities were measured with substrate and CoA at concentration of 0.5 mM, respectively. Data are expressed as relative mean percentages of three independent biological replicates. N.D. represents ‘not detected’ or relative activity < 5%.

a 100% relative activity of PcAAE1 corresponds to 4.02 μmol min⁻¹ mg⁻¹ on isovaleric acid.
b 100% relative activity of PcAAE2 corresponds to 1.96 μmol min⁻¹ mg⁻¹ on 4-methylvaleric acid.
c 100% relative activity of PcAAE3 corresponds to 6.75 μmol min⁻¹ mg⁻¹ on propionic acid.

substrate preference toward branched-chain fatty acids. The combination of the positive correlation of expression patterns of PcAAE2 with the distribution pattern of pogostone and the predicted substrate preference of the protein encoded by PcAAE2 toward branched-chain fatty acids made it the most likely candidate gene encoding the enzyme responsible for the formation of 4MVCoA precursor of pogostone compared with the other seven genes encoding P. cablin clade VI proteins.

To test this hypothesis, we performed in vitro biochemical assays with all eight AAE proteins in the P. cablin clade VI of the family (Table 1). All proteins were produced in Escherichia coli with a fused N-terminal His6 tag and purified as soluble proteins. All the eight purified recombinant proteins were initially tested in the coupled enzyme assay with a variety of fatty acid substrates including straight short/medium-chain fatty acids and branched short/medium-chain fatty acids and benzoic acid at a 0.5 mM concentration of the acid substrate and 0.5 mM of CoA. Except for the three recombinant proteins PcAAE1~3, the other five recombinant proteins showed no detected enzymatic activities with any of these selected substrates.

Recombinant PcAAE1 utilizes isovaleric acid as the best substrate among the selected substrates and shows lower activities toward butyric acid, valeric acid and 4-methylvaleric acid (Table 2). Recombinant PcAAE2 shows high activities toward 4-methylvaleric acid, valeric acid and butyric acid and utilizes 4-methylvaleric acid as the most preferred substrate (Table 2). Recombinant PcAAE2 also accepts 5-methylhexanoic acid, heptanoic acid, hexanoic acid and isovaleric acid as substrates but at a lower rate (Table 2). Purified PcAAE3 accepts very short-chain fatty acids as substrates, including acetic acid, propanoic acid and butyric acid, with propionic acid as the preferred substrate (Table 2). Therefore, PcAAE1 and PcAAE2 are assigned as the probable isovaleryl-CoA synthetase and 4MVCoA synthetase, respectively, and may supply CoA ester precursors for pogostone and its analogue pogostone B biosynthesis. Thus, in-depth studies of the dependency of their enzymatic activities on incubation conditions and components were further performed. Highest activities of PcAAE1 and PcAAE2 with their respective preferred substrates were observed at pH 7.5 and 30°C. The activities of PcAAE1 and PcAAE2 were dependent on divalent cations with Mg²⁺ being preferred and increased by 1.7-fold and 0.8-fold, respectively, when the concentration of
Mg$^{2+}$ rose from 1 to 10 mM. Additional supplementation with dithiothreitol or a univalent cation at 2.5 mM including K$^+$ does not obviously increase their activities. These two enzymes were not stable at 4, −20 and −80°C and lost more activities at 4, −20°C than at −80°C for 24 h (Supplementary Fig. S4).

Since we aim to identify the AAEs responsible for the formation of 4MCCoA and isovaleryl-CoA, the predicted CoA ester precursors of pogostone and its analogue pogostone B, respectively, the enzymatic products of PcAAE1 with isovaleric acid as substrate and PcAAE2 with 4-methylvaleric acid and isovaleric acid as substrates were further verified with LC-QTOF-MS system (Fig. 4) and the kinetic properties of PcAAE1 with isovaleric acid as substrate and PcAAE2 with 4-methylvaleric acid and isovaleric acid as substrates were further determined (Table 3). The kinetic analysis revealed that PcAAE1 has a $K_m$ value of 12.9 ± 0.8 μM and a catalytic efficiency of 3.33 × 10⁵ s⁻¹M⁻¹ for isovaleric acid, and PcAAE2 has a $K_m$ value of 43.4 ± 5.1 μM and catalytic efficiency of 5.19 × 10⁴ s⁻¹M⁻¹ for 4-methylvaleric acid, while the $K_m$ value and catalytic efficiency for isovaleric acid are 216.9 ± 13.1 μM and 0.78 × 10⁴ s⁻¹M⁻¹, respectively (Table 3).

Considering the previously reported substrate promiscuity of AAEs such as GsAAE1, Ph4CL1 and HcAAE1 (Stout et al. 2012, Klemplien et al. 2012, Singh et al. 2020), we further determined the kinetic properties of PcAAE1 and PcAAE2 with substrates toward which PcAAE1 and PcAAE2 show relative activities higher than 40% in Table 2. The kinetic analysis further revealed that PcAAE2 also has low $K_m$ values with the substrates valeric acid, hexanoic acid and heptanoic acid besides 4-methylvaleric acid (Table 3). Although the $K_m$ value of PcAAE2 with hexanoic acid is about the same as that of PcAAE2 with 4-methylvaleric acid, the catalytic efficiency of PcAAE2 with hexanoic acid is only 44% of that of PcAAE2 with 4-methylvaleric acid (Table 3).

### Table 3 Kinetic Properties of Recombinant PcAAE1 and PcAAE2

| Enzyme  | Substrate          | $K_m$ (μM) | $K_{cat}$ (s⁻¹) | $K_{cat}/K_m$ (M⁻¹s⁻¹) |
|---------|--------------------|------------|----------------|-------------------------|
| PcAAE1  | Butyric acid       | 161.0 ± 5.0| 4.0 ± 0.2      | 24,741 ± 1276           |
|         | Isovaleric acid$^a$| 12.9 ± 0.8 | 4.1 ± 0.1      | 332,937 ± 7532          |
| PcAAE2  | Butyric acid       | 570.1 ± 86.5| 2.4 ± 0.1      | 4344 ± 90              |
|         | Valeric acid       | 53.9 ± 2.7 | 2.0 ± 0.02     | 37,288 ± 436           |
|         | Isovaleric acid$^b$| 216.9 ± 13.1| 1.7 ± 0.08    | 7759 ± 371             |
|         | Hexanoic acid      | 41.5 ± 2.4 | 1.0 ± 0.03     | 22,904 ± 842           |
|         | 4-Methylvaleric acid$^b$| 43.4 ± 5.1 | 2.3 ± 0.1     | 51,854 ± 2765          |
|         | Heptanoic acid     | 84.5 ± 7.1 | 1.1 ± 0.04     | 13,184 ± 433           |
|         | 5-Methylhexanoic acid | 175.5 ± 20.0 | 1.2 ± 0.05 | 6951 ± 312           |

These enzymatic assays were performed using the same coupled enzymes assay as Table 2. Data are presented as mean ± SD of three independent biological replicates.

$^a$The $K_m$ values of cosubstrates CoA and ATP were 28.2 and 89 μM, respectively.

$^b$The $K_m$ values of cosubstrates CoA and ATP were 150.8 and 540.2 μM, respectively.

$^c$The $K_m$ values of cosubstrates CoA and ATP were 40.2 and 160.5 μM, respectively.

Phylogenic relationships of PcAAE2 homologs
To gain further insight into the evolution of PcAAE2, we expanded the phylogenetic analysis of AAE clade VI to include 77 homologs from chlorophytes including _Chlamydomonas reinhardtii_ and _Volvox carteri_, Embryophyte, Tracheophyte and angiosperms. The phylogenetic tree was constructed using a maximum likelihood algorithm (Fig. 6). The proteins in the expanded clade VI were mainly grouped into four distinct subclades (Fig. 6). We designated the group that contains PcAAE1, PcAAE2 and PcAAE7 as subclade Vla, the group that contained PcAAE3 as subclade Vlb, the group that included PcAAE5, PcAAE6 and PcAAE8 as subclade Vlc, and the group that included PcAAE4 as subclade Vld (Fig. 6). Six homologs of the AAE clade VI...
Fig. 4 \textit{In vitro} analyses of \textit{PcAAE1} and \textit{PcAAE2} activities. \textbf{A}, LC-QTOF-MS analysis of products formed in reactions containing purified \textit{PcAAE1} or \textit{PcAAE2}, free CoA, ATP, MgCl\textsubscript{2}, isovaleric acid or 4-methylvaleric acid after 30 min of incubation. The extracted ion chromatograms in negative mode of \textit{m/z} 766.1079, 850.1654 and 864.1811 are shown for free CoA, isovaleryl-CoA and 4MVCoA, respectively. \textbf{B}, The mass spectrums in negative mode of standard CoA, standard isovaleryl-CoA and CoA esters (predicted isovaleryl-CoA and 4MVCoA) generated \textit{in vitro} by the action of \textit{PcAAE1} or \textit{PcAAE2} with the substrates of isovaleric acid or 4-methylvaleric acid. The singly [M-H]\textsuperscript{-} and doubly [M-2H]\textsuperscript{2-} charged pseudo-molecular ion are shown horizontally. The CoA ester from isovaleric acid by the action of \textit{PcAAE1} or \textit{PcAAE2} was identified as isovaleryl-CoA through the comparison of its retention time and mass spectrums with that of standard isovaleryl-CoA, while the CoA ester from 4-methylvaleric acid by the action of \textit{PcAAE2} was identified as 4MVCoA based on the exact mass and fragment pattern.

It is interesting that most of clade VI AAEs possess canonical PTS1 signals and are predicted to be targeted to the peroxisome by PTS1 predictor (Fig. 6). We also further investigated subcellular localization prediction of other selected genes using Targetp 2.0. 3 out of the 77 clade VI AAEs contain mitochondrial transit peptides at the N-terminus and are predicted to be located in mitochondria, while 15 out of these 77 AAEs contain no obvious signal peptides and are predicted to be located in cytoplasm (Fig. 6). In addition, all the six outgroups contain prototypical PTS1 signals with high prediction score, suggesting likely peroxisomal localization. \textit{PcAAE1} and \textit{PcAAE2} are grouped closely with \textit{HcAAE1}, \textit{HlCCL2} and At2G17650 in the subclade via (Fig. 6), which are consistent with the fact that they share very similar substrate preference toward branched/straight short-chain fatty acids.
mainly isolated from patchouli oil, which is usually produced by
et al. 2012 has been proved to exert many pharmaceutical activities (et al. 2004).
ous studies have indicated that pogostone, one of the major
ous odor, and more recently, this compound
thetic ability to blend with other essential oils
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lower level, showed very similar accumulation pattern to that of
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erated from main stems shows a developmental pattern
(Fig. 2). To begin elucidating the biosynthetic pathway of
ogostone, we searched for the enzyme that catalyzes the formation
of 4MVC0A, one of the precursors of pogostone. Transcriptome
proiling and searches of RNAseq databases obtained from
three different stages of developing P. cablin plants identified
1 AAE unigenes (Supplementary Table S1), with one unigene,
named PcAAE2, showing the strongest positive correlation with
the distribution pattern of pogostone (Supplementary Table
Figs. 2B and 3).

The purified recombinant PcAAE2 protein is able to ef-
citively catalyze the formation of 4MVC0A in vitro from
4-methylvaleric acid, with a \( K_m \) value of 43.4 \( \mu M \) and a cata-
ytic efficiency of \( 5.19 \times 10^3 \text{ s}^{-1} \text{M}^{-1} \) for 4-methylvaleric acid
(Table 3; Fig. 4). In addition to 4-methylvaleric acid, PcAAE2
is also able to effectively catalyze the formation of CoA esters
in vitro from valeric acid and hexanoic acid, with \( K_m \) values
of 53.9 and 41.5 \( \mu M \), respectively (Table 3), suggesting
substrate promiscuity, which has been commonly observed
for AAEs (Stout et al. 2012, Klempien et al. 2012, Xu et al.
2013, Singh et al. 2020). Despite the observation of
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In addition, PcAAE2 is also able to catalyze the formation
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which accounts for about 15% of that with 4-methylvaleric
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stone B compared with that of pogostone in tested P. cablin
tissues (Fig. 2), PcAAE2 may also supply isovaleryl-CoA pre-
cursor for pogostone B biosynthesis in cytoplasm, where the
polyketide pogostone and pogostone B are believed to be
synthesized.

Besides PcAAE2, PcAAE1, another member of clade VI AAEs
from P. cablin, also shows substrate preference toward branched
short-chain fatty acid (Table 2). The purified recombinant
PcAAE1 protein accepts isovaleric acid as the best-preferred
substrate and is able to effectively catalyze the formation
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isovaleric acid (Table 3; Fig. 4). Although the expression pat-
tern of PcAAE1 did not show an obvious positive correlation
with the accumulation pattern of pogostone B in the tested P. cablin
tissues (Supplementary Table S1; Figs. 2 and 3), the

**DISCUSSION**

**PcAAE2 catalyzes the formation of 4MVC0A precursor of pogostone in P. cablin**

The medicinal plant *P. cablin* is an important commercial source
of patchouli oil (Yang et al. 2016, van Beek and Joulain 2018).
Patchouli oil has the ability to blend with other essential oils
and give a strong base, lasting character, fixative properties and
aids in preventing evaporation, thus promoting tenacity; hence,
it is widely used in the manufacturing of perfume, cosmetics,
soaps, scents body lotions and detergents (Gubareva 2004, Kiso
et al. 2004, Swamy and Sinniah 2015, Hu et al. 2017). Previous
studies have indicated that pogostone, one of the major
chemical constituent of patchouli oil, is largely responsible for
its intense aromatic odor, and more recently, this compound
has been proved to exert many pharmaceutical activities (Li
et al. 2012, 2014, Yi et al. 2013, Su et al. 2015, 2017, Chen et al.
2016, Wang et al. 2016, Cao et al. 2017). At present, pogostone
is mainly isolated from patchouli oil, which is usually produced by
steam distillation of the shade dried leaves and stems (Swamy
and Sinniah 2015).

In patchouli oil, besides pogostone, a previous study also
detected its different analogues including pogostone B in much
smaller concentrations (Nakahara et al. 1975, Rijke et al. 1978,
van Beek and Joulain 2018), which is similar to that was
observed in the present study. Pogostone B, albeit at a much
lower level, showed very similar accumulation pattern to that of
pogostone in tested *P. cablin* tissues (Fig. 2B). The distribution
of both pogostone and pogostone B in *P. cablin* top leaves newly
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tissues (Supplementary Table S1; Figs. 2 and 3), the

![Subcellular localization of PcAAE1 and PcAAE2 in Arabidopsis leaf mesophyll protoplasts. GFP fusion proteins were visualized by laser confocal microscopy. A, Subcellular localization of PcAAE1 in Arabidopsis leaf mesophyll protoplasts. Both the first 30 amino acids (predicted mitochondrial transit peptide) of PcAAE1 fused C-terminal GFP and whole PcAAE1 fused N-terminal GFP were used for assay of subcellular localization of PcAAE1. Mitochondria were revealed by MitoTracker Red-staining. B, Subcellular localization of PcAAE2 in Arabidopsis leaf mesophyll protoplasts. Both PcAAE2 fused N-terminal and C-terminal GFP were used for assay of subcellular localization of PcAAE2. The signal of free GFP was used as control and chloroplasts were revealed by chlorophyll autofluorescence in pseudo-color blue. Scale bar = 10 \( \mu M \).](http://example.com/image.png)
**Fig. 6** Phylogenetic analyses of AAE clade VI homologs and PcAAE2 homologs from the plant kingdom. All proteins used for phylogenetic tree construction except PcAAE1 ~ 8 were obtained from the PHYTOZONE v12.1 site (https://phytozone.jgi.doe.gov/pz/portal.html), NCBI website (https://www.ncbi.nlm.nih.gov/) and TAIR website (https://www.arabidopsis.org/). The protein sequences of PcAAE1 ~ 8 were directly obtained from *P. cablin* transcriptome database. Sequences were aligned using ClustalW and tree were generated by the maximum likelihood algorithm using MEGA7.0 (Kumar et al. 2016). Branch point bootstrap values were calculated with 1000 replicates. The subcellular localization of each protein was analyzed using signal prediction algorithm including Targetp 2.0, SignalP 5.0 and The PTS1 predictor (http://www.cbs.dtu.dk/services/TargetP/, http://www.cbs.dtu.dk/services/SignalP/ and http://mendel.imp.ac.at/pts1/). The predicted signal peptide and prediction value were listed along with protein name and species. Peroxisome targeting sequence 1 (PTS1) was marked in blue color, while mitochondrial transit peptide (mTP) was marked in red color. None of chloroplast transit peptide was detected in all selected AAEs. The verified subcellular localizations of AAEs were labeled in green color (Supplementary Table S3). Black arrows indicate proteins of which the enzymatic activities and subcellular localizations were characterized in this paper. The subclade that PcAAE1, PcAAE2, HcAAE1, HICCL2 and At2g17650 were grouped into was indicated in black frame.
possibility of involvement of PcAAE1 in pogostone B biosynthesis cannot be completely ruled out. Subcellular localization assays indicated that PcAAE1 is a mitochondria-targeted protein (Fig. 5A). Therefore, it is mostly likely that PcAAE1 maintains the isovaleryl-CoA pool in mitochondria under certain conditions and plays a role in a dynamic balance between leucine degradation and the electron transport chain of mitochondria (Araujo et al. 2010).

**Cytosolic PcAAE2 may have evolved from an ancestral peroxisomal AAE**

The AAE gene family expanded and diversified as land plants evolved from their smaller, simpler algal ancestors (Shockey and Browse 2011). Each of the AAE gene family of *Chlamydomonas reinhardtii* and Volvox carteri, two unicellular green algae, contains one gene (Cr03g211745 for *Chlamydomonas reinhardtii* and Vc0013s0189 for *Volvox carteri*) only distally related to PcAAE2, and this gene was grouped out of the AAE clade VI subfamily in our phylogenetic tree (Fig. 6), suggesting that this clade VI is specific to multicellular land plants as previously reported (Shockey and Browse 2011).

Interestingly, most of clade VI AAEs in this phylogenetic analysis (46 out of 71) possess a predicted peroxisomal targeting signal 1 (PTS1) and are therefore predicted to be targeted to the peroxisomes (Fig. 6), while three clade VI AAEs possess predicted mitochondrial transit peptides and are predicted to be located in the mitochondria (Fig. 6). Fifteen clade VI AAEs possess no obvious predicted signal peptides and are predicted to be targeted to cytoplasm (Fig. 6). Each of PcAAE2’s closest homologs (Cr03g211745, Vc0013s0189, Pp3c25280, Pp3c96840, Sm409095, Sm440236, Sm236556 and Sm81559) from *Chlamydomonas reinhardtii* (a unicellular green algae), *Volvox carteri* (unicellular green algae), *Physcomitrella patens* (a moss) and *Selaginella moellendorfii* (an ancient vascular plant) contains a prototypical PTS1 sequence at the C-terminus with high prediction score (Fig. 6) and is therefore highly likely to be located in peroxisome where the catabolic process of β-oxidation occurs. These ancestor peroxisomal PcAAE2 homologs may play important roles in β-oxidation. PcAAE2 lacks a PTS1 and locates in cytoplasm where the polyketide pogostone is believed to be synthesized (Fig. 5B).

Since the possession of a PTS1 sequence is the ancestral trait in the AAE VI clade throughout the plant kingdom, and a PTS1 sequence is also present in the AAE6 clade (in PcAAE7, a protein that has 65% amino acid identity to PcAAE2), the recruitment of PcAAE2 for pogostone biosynthesis appears to have arisen through a duplication event of an ancestral peroxisomal protein and subsequent divergence in both subcellular localization and substrate specificity. Similar cases for a cytoplasm localized hexanoyl-CoA synthetase (GCAAE1) recruited to cannabinoid biosynthesis in *Cannabis sativa* trichomes and for two cytoplasm localized branched short-chain CoA ligases (HICCL2 and HICCL4) recruited to bitter acid biosynthesis in *H. lupulus* trichomes and for recently identified cytoplasm and peroxisome dually localized benzoyl-CoA ligase (HcAAE1) recruited to xanthone biosynthesis in *H. calycinum* have been already reported (Stout et al. 2012, Xu et al. 2013, Singh et al. 2020). A similar case may also have occurred for PcAAE1, which encodes a mitochondrial isovaleryl-CoA synthetase (Table 3; Fig. 5A), probably playing a role in a dynamic balance between leucine degradation and the electron transport chain of mitochondria through maintaining the pool of isovaleryl-CoA in mitochondria (Araujo et al. 2010). This conclusion could be confirmed by further experimental subcellular localization analysis of more clade VI genes from different plant species.

**Assessment of P. cablin PcAAE1 and PcAAE2 sequence properties**

AAEs belonging to the acyl-CoA synthetase subfamily are members of the so-called ANL superfamily, in which some recognized conserved signature motifs (A1–A10), common to all adenylation domains have been identified (Marahiel et al. 1997, Gulick 2009) and are also present in *P. cablin* PcAAE1 and PcAAE2 (Supplementary Fig. S3). AAEs exhibit similar overall folding patterns with large N-terminal and small C-terminal domains. Biochemical and structural studies demonstrate that AAE enzymes undergo large-scale domain movement to facilitate the two-step catalytic reaction, an adenylate-forming conformation during the first half-reaction and a thioester-forming conformation during the second step (Bar-Tana and Rose 1968, Gulick et al. 2003, Reger et al. 2008, Gulick 2009, Li and Nair 2015). The hinge residue that facilitates this movement is usually a part of the A8 motif (Gulick et al. 2003), which is conserved in *P. cablin* PcAAE1 and PcAAE2 (Supplementary Fig. S3). As in all adenylate forming enzyme, two conserved AMP-binding motifs (AMP1 and AMP2) (Weimar et al. 2002, Morgan-Kiss and Cronan 2004) and the catalytically relevant amino acid lysine (Reger et al. 2007, Hu et al. 2010, Li and Nair 2015) are also present in *P. cablin* PcAAE1 and PcAAE2 (Supplementary Fig. S3).

In addition, PcAAE1–3 from *P. cablin* and some other medium/short-chain acyl-CoA synthetases from *A. thaliana*, *C. sativa*, *H. lupulus*, *H. calycinum*, *P. cablin*, *Taxus media* have been shown to be active with a range of medium/short-chain fatty acids (Stout et al. 2012, Xu et al. 2013, Srividya et al. 2020, Singh et al. 2020). Phylogenetic analysis indicated that they fall into two clades (Fig. 7A). Clade I, which contains acyl-CoA synthetases preferring branched/straight short-chain fatty acids, is clearly distinct from clade II, which contains CsAAE3, TmAAE13, TmAAE15 and TmAAE16, all of which show high activities toward straight medium-chain fatty acids. Previous analysis of bacterial AAEs indicated an fatty acyl-CoA synthetase (FACS) signature motif appearing to contribute to the fatty acid-binding pocket and thus affects the preference for fatty acids of different chain lengths (Black et al. 1997, 2000). Sequence divergence across their FACS signature motifs indicated that they fall into two groups that are consistent with
that these amino acids may play a vital role in fatty acid chain length specificity. To verify this hypothesis, *in silico* homolog model based on the available crystal structures of AAEs such as Pt4CL1 and Nt4CL2 (Hu et al. 2010, Li and Nair 2015) should be required. Further extensive comparisons on the structures and gain-of-function mutations in this motif are needed for a deeper understanding of the substrate preferences for branch/straight chain and chain length of these enzymes.

### Metabolic engineering of pogostone biosynthesis

*P. cablin* (Lamiaceae) is a commercially important plant for its essential oil (patchouli oil). It is native to southeast Asia and is now cultivated widely in many tropical and subtropical regions (Wu et al. 2010, Xie et al. 2017), including China, Indonesia, the Philippines and Thailand for the extraction of essential oils from its leaves and stems, and especially as a natural source of patchouli alcohol and pogostone (Singh and Rao 2009, He et al. 2016, Swamy and Sinniah 2016). Different *P. cablin* cultivars exhibit significant differences in quality and bioactive components, as these factors are influenced by climate, soil nutrients and water in the different locations (Swamy et al. 2015). Based on the main components of essential oils, *P. cablin* in China can be divided into two chemotypes: pogostone type, with a high content of pogostone and low content of patchouli alcohol, and patchoulol type, which has a high content of patchouli alcohol but a low content of pogostone (Liu et al. 2002, Luo et al. 2003, Huang et al. 2016). Traditionally, the patchoulol-type cultivars are mainly used in the perfume industry, whereas the pogostone-type cultivars are considered as medicinal plants in China. However, the production of pogostone, likely the main effective compound in medicinal use, is extremely low due to the limitation of cultivation area (Hu et al. 2006, Wu et al. 2010). Due to the discovery of more and more pharmacological activities of pogostone, the pogostone-type cultivars are highly popular (Chen et al. 2021). However, their natural supplies cannot sufficiently satisfy the economic demand (Zhang et al. 2019).

Our discovery of *PcAAE2*, the enzyme that catalyzes the biosynthesis of 4MVCa, will serve as a step forward in the construction of high-yield pogostone *P. cablin* varieties. Increasing *PcAAE2* transcript levels or changing its developmental expression pattern could lead to further increase in the production of pogostone. Pogostone has received increased attention along with the discovery of its additional potential bioactivities and pharmacological properties such as antibacterial, anti-inflammatory, anticancer activities and others (Chen et al. 2015, 2016, Sun et al. 2016, Wang et al. 2016, Cao et al. 2017). The discovery of 4-MVCa synthetase could also facilitate the functional characterization of other enzymes such as polyketide synthases involved in the pogostone biosynthesis and, thus, will allow for the reconstruction of this pathway in the heterologous system using a synthetic biology approach to produce pogostone in large amounts.

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**Fig. 7** Analysis of the characterized medium/short chain acyl-CoA synthetase from *A. thaliana, C. sativa, H. lupulus, H. calycinum, P. cablin, T. media*. A. Maximum likelihood inference of the phylogenetic relationship of the characterized medium/short chain acyl-CoA synthetase from *C. sativa, H. lupulus, H. calycinum, P. cablin, T. media* (Supplementary Table S3). Sequences were aligned using ClustalW and tree were generated by the Maximum likelihood algorithm using MEGA7.0 (Kumar et al. 2016). Branch point bootstrap values were calculated with 1000 replicates. B. Alignment of the amino acid residues of the FACS signature motifs of the acyl-CoA synthetases. Alignment was conducted by ClustalW (https://embnet.vital-it.ch/software/ClustalW.html) and edited using GeneDoc software (https://genedoc.software.informer.com/2.7/).
Materials and Methods

Plant materials and chemicals

*P. cablin* seeds were germinated in soil, and plants were grown in a greenhouse with a 16 h light/8 h darkness photoperiod at 30°C. The various tissues samples including seedling, roots, main stems and leaves were collected from *P. cablin* plants. For RT-qPCR analysis, the harvested tissues were flash frozen in liquid nitrogen and stored at −80°C until use. *Arabidopsis* plants were grown in soil in a growth chamber with the photoperiod (12 h light/12 h dark at 23°C) under 60% humidity. The well-expanded leaves from 4-week-old *Arabidopsis* plant were selected for protoplast isolation for subcellular localization.

Pogostone and pogostone B standard were synthesized as described previously (Yu et al. 2018). Since standard 4MVeCoA is not available commercially, the CoA esters produced by *PcAAE2* from 4-methylvaleraldehyde were identified as 4MVeCoA based on the exact mass and fragment pattern by LC-QTOF-MS. MitoTracker Red used for mitochondria staining was purchased from ThermoFisher, and other chemicals were purchased from Sigma-Aldrich.

GC-MS analysis of pogostone and pogostone B from *P. cablin* seedling, roots, main stems and leaves from the main stem at different stages of development

*P. cablin* seedlings, roots, main stems and leaf tissues at different stages of development were collected and cut into small pieces, 100 mg of which were transferred into a tube containing 500 µl MTBE with 0.05 ng/µl tetradeucane as internal standard. The tube was vortexed for 3 min at maximum speed, incubated at room temperature for 1 h and then centrifuged for 3 min at the speed of 10,000 g/min. The MTBE phase was collected and analyzed by GC-MS. For GC-MS analysis, 1 µl aliquot of the sample was injected into Agilent GC-MSD (Agilent 7890B-5977B) system equipped with the DB-5MS column (30 m × 0.25 mm × 0.25 µm film thickness, Agilent, USA). The oven temperature was programmed as follows: initial temperature, 50°C for 3 min, 50 to 320°C at a rate of 10°C min⁻¹ and hold for 5 min. The identification of compounds was assigned by comparison of their retention times and mass fragmentations with corresponding authentic standards. The measurement of pogostone and pogostone analogues was performed by comparison to the corresponding authentic standard curves.

RNAseq analysis

Total RNA was extracted from seedling, root, main stem and leaf parts of *P. cablin* plants at different developmental stages using Total RNA Isolation Kit from Omega. The RNAseq library construction and transcriptome assembling were performed as described previously (Xu et al. 2018). The expression levels of all transcripts were estimated using the Trinity software, which estimated transcript abundance using the RSEM function (Li and Dewey 2011).

Identifying AAE genes in the *P. cablin* transcriptome

To find candidate unigenes encoding AAEs in our *P. cablin* transcriptome database, we constructed local Nucleotide Database using the software (BioEdit Sequence Alignment Editor, https://bioedit.software.informer.com/7.2/) and queried it with Arabidopsis representative AAEs from each of seven clades of AAE superfamily using the TBLASTN function. We also screened the entire annotated database for the functional annotations ‘AMP binding protein’ and ‘CoA ligase’ and added transcripts identified in this way to our list of candidates.

Quantitative reverse transcription polymerase chain reaction analysis of *PcAAE1~8* transcript levels

For quantitative real-time PCR analysis of transcripts in different tissues, total RNA was isolated using total RNA Isolation Kit from Omega with a DNA digestion step. RNA concentration and quality (260/280 and 260/230 ratios) were determined using the Thermo Scientific™ NanoDrop™ Lite Spectrophotometer. Reverse transcription polymerase chain reaction (RT-PCR) was prepared using High Capacity cDNA Reverse Transcription Kit from Thermo Fisher Scientific following the manufacturer’s instructions with minor modification. 3 μg total RNA was used for making cDNAs in a 20 µl RT-PCR reaction system containing 5 µM oligo(dT), which was obtained from Thermo Fisher Scientific (cat number: #50112). RT-qPCR analysis was performed using Power SYBR Green PCR master mix (Thermo Fisher) using Bio-Rad CFX Connect Real-Time PCR Detection System. Each reaction consisted of 10 µl Power SYBR Green PCR master mix, 2 µl cDNA and 1 µl of 5 µM forward and reverse primers in a total volume of 20 µl made up with sterile water. The thermal profile included, after a 2 min step at 95°C, 40 cycles of denaturation at 95°C for 5 s and annealing at 55°C for 30 s and amplification for 30 s. Melting curves were recorded by melting the amplified amplicon at 65−95°C in an increment increase of 0.5°C per cycle. *P. cablin* glyceraldehyde 3-phosphate dehydrogenase (PcGAPDH), PcActin7, and PcTubulin3 were used as internal reference genes. For *PcAAE1~8* and the three reference genes, single melting curve peaks were obtained. Primer specificities were evaluated by melt curve analysis and agarose gel electrophoresis. cDNAs of four biological replicates were analyzed. No-template (NTC) and no-reverse transcriptase controls (NRC) were used to rule out primer dimer formation and genomic DNA contamination, respectively.

The RT-qPCR data were expressed as relative expressions of *PcAAE1~8* relative to the three reference genes in each tissue. The relative expressions of *PcAAE1~8* in each tissue were calculated using 2 −ΔΔCq with the ΔCq calculated by subtracting Cq values of *PcAAE* genes from the geometric mean Cq values of the three reference genes in every tissue and the amplification efficiency (E) set as 2. The equation used in this process is defined by just one group of Cq values containing Cq for the gene of interest and geometric mean Cq for the three reference genes in tested tissues (Vandesompele et al. 2002). Stable expression of the three reference genes among all selected samples was confirmed before integrating their Cq values in normalization. Primers used in this study are designed using Primer 5.0 software and listed in Supplementary Table S4. Assay of unbiased RNA coverage of oligo (dT) and random primer for *PcAAE1~8* in the top leaf and main stem tissues of 6-week-old *P. cablin* plants was performed and the relevant results were supplied as Supplementary Fig. S5.

Isolation of His-tagged recombinant proteins of *PcAAE1~8*

The open reading frames of *PcAAE1~8* were obtained by RT-PCR from prepared cDNA of *P. cablin* seedlings and were introduced into the expression vector pET-28a (+) between the restriction sites of Ndel and BamHII using NEBuilder™ HiFi DNA Assembly Cloning Kit (NEB, Catalog number: E5520S), in each case generating a fusion gene that encoded a ‘tag’ of His6 residues at the N-terminus for expression in *Escherichia coli*. To obtain soluble proteins for expression in *E. coli*, a truncated open reading frame of *PcAAE1*, missing the first 30 codons, was obtained by RT-PCR from prepared cDNA of *P. cablin* seedlings and was introduced into the expression vector pET-28a(-) to generate a fusion gene that encoded a ‘tag’ of His6 residues at the N-terminus for expression in *E. coli*. Recombinant proteins of *PcAAE1~8* were expressed in *E. coli* strain BL21 (DE3) and purified using Ni-NTA agarose chromatography (Qiagen) as previously described (Xu et al. 2018).

Enzymatic assays of recombinant *PcAAE1~8*

Acyl-CoA synthetase activities of recombinant *PcAAE1~8* proteins were measured using the coupled enzymes assay (Koo et al. 2006, Chen et al. 2011). For substrate specificity assays, reaction mixtures (250 µl) contain 0.1 M Tris-HCl, pH 7.5, 2 mM DTT, 5 mM ATP, 10 mM MgCl₂, 0.5 mM CoA, 0.4 mM NADH, 0.5 mM fatty acid substrate, 1 mM phosphoenolpyruvate and four units each of myokinase, pyruvate kinase, and lactate dehydrogenase. The reaction was initiated by adding 1 µg of purified enzyme. Oxidation of NADH was monitored at 30°C for 15 mins at 340 nm (ε₃₄₀nm = 6220 M⁻¹ cm⁻¹) with a Beckman DU530 spectrophotometer. The enzyme concentration (1 µg) and incubation time (15 min) were selected to attain a linear reaction velocity during the assay period. To determine the kinetic parameters of recombinant *PcAAE2, a similar
protocol was followed. The $K_m$ value for fatty acid substrate was determined by fixing the concentration of CoA at 1 mM and ATP at 5 mM, whereas the $K_m$ value for CoA was determined by fixing the concentration of fatty acid substrate at 1 mM and ATP at 5 mM, and the $K_m$ value for ATP was determined by fixing the concentration of fatty acid substrate at 1 mM and CoA at 1 mM. $K_m$ and $k_{cat}$ values were calculated from initial rate data by using the hyperbolic regression analysis method in Hyper32 software (version 1.0.0, http://hyper32.software.informer.com/).

In addition, Acyl-CoA synthetase activity of recombinant PcAAE1 and PcAAE2 proteins was further measured using an LC-QTOF-MS-based method. Briefly, the reaction mixtures (250 µl) contain 0.1 M Tris-HCl, pH 7.5, 2 mM DTT, 5 mM ATP, 10 mM MgCl₂, 0.5 mM CoA, 0.5 mM fatty acid substrate. The reaction was initiated by adding 10 µl of purified PcAAE1 or PcAAE2 protein. Assays were stopped after 30 min by the addition of 250 µl methanol. The solution was filtered through a 0.45 µm filter prior to analysis by LC-QTOF-MS system (Agilent, 6545 LC/QTOF-MS) coupled with a C₁₈ column (ZORBAX RRHD Plus C₁₈, 3.5 µm x 50 mm, 1.8 µm) at 35°C. The gradient (solvent A, water + 0.1% formic acid; solvent B, methanol) program was set as follows at a flow of 0.5 ml min⁻¹: 0–0.5 min, a linear gradient from 5% to 40% of B; 0.5–2.5 min, 40% of B; 2.5–3.5 min, a linear gradient to 100% of B; 3.5–4.5 min, 100% of B; 4.5–6.6 min, a linear gradient to 100% of A and equilibration for 2 min before starting the next sample. The operating parameters were set as follows: capillary voltage, 3,500 V; nebulizer pressure, 35 pounds per square inch gauge; drying gas flow, 11 L min⁻¹; gas temperature, 320°C; sheath gas temperature, 350°C; sheath gas flow, 11 L min⁻¹ and the fragmentor, 175 V. Mass spectrum were acquired in negative mode, and lock mass correction was performed using standard hexakis (2,2,3,3-tetrafluoropropoxy) phosphazine and purine. The compounds were analyzed with the parameters as follows: 766,1079 (m/z) for CoA, 850,1654 (m/z) for isovaleryl-CoA and 864,1811 (m/z) for 4MVCoA.

Subcellular Localization of PcAAE1 and PcAAE2

The vector used for subcellular localization of analysis was pTF486. For the construction of these vectors expressing fusion proteins with GFP fused with C-terminus of the first 30 amino acids of PcAAE1 and whole PcAAE2, the open reading frames of the first 30 amino acids of PcAAE1 and whole PcAAE2 were amplified from P. cablin seedling cDNA, respectively, and inserted into pTF486 vector between restriction sites of Salt and BamHI using In-Fusion™ HD Cloning Kit (TaKaRa, Catalog number: 638910) following the protocol. For the construction of these vectors expressing fusion proteins with GFP fused with N-terminus of PcAAE1 and PcAAE2, respectively, the open reading frames of PcAAE1 and PcAAE2 and gene encoding GFP were amplified from Pc cbin seedling cDNA and pTF486 plasmid, respectively, the purified PCR product of either PcAAE1or PcAAE2 and gene encoding GFP and purified double enzyme digestion product of pTF486 plasmid between restriction sites of Salt and BamHI were assembled together using In-Fusion™ HD Cloning Kit (TaKaRa) following the protocol. Protoplasts were prepared from Arabidopsis leaves, and transformation and confocal microscopy were performed as described previously (Yoo et al. 2007).

Supplementary Data

Supplementary data are available at PCP online.

Data Availability

Data supporting the finding of this study are available within the article and its supplementary files. The sequences of the genes used in this study are available under GenBank accessions as follows: PcAAE1, MW413950; PcAAE2, MW413951; PcAAE3, MW413952; PcAAE4, MW413953; PcAAE5, MW413954; PcAAE6, MW413955; PcAAE7, MW413956; PcAAE8, MW413957; PcGAPDH, MW735976; PcActin7, MW735977; PcTubulin3, MW735978. The bioproject accession number for the RNAseq data is PRJNA713906.

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Author Contributions

H.X. designed the experiments; J.C., L.L., Y.W. and Z.L. conducted the experiments, analyzed data or provided material; J.C., G.W., E.P. and H.X. wrote the article; all authors edited the article.

Disclosures

The authors have no conflicts of interest to declare.

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