Genetics

Dual activities of ACC synthase: Novel clues regarding the molecular evolution of ACS genes

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Ethylene plays profound roles in plant development. The rate-limiting enzyme of ethylene biosynthesis is 1-aminocyclopropane-1-carboxylate (ACC) synthase (ACS), which is generally believed to be a single-activity enzyme evolving from aspartate aminotransferases. Here, we demonstrate that, in addition to catalyzing the conversion of S-adenosyl-methionine to the ethylene precursor ACC, genuine ACSs widely have C-S lyase activity. Two N-terminal motifs, including a glutamine residue, are essential for conferring ACS activity to ACS-like proteins. Motif and activity analyses of ACS-like proteins from plants at different evolutionary stages suggest that the ACC-dependent pathway is uniquely developed in seed plants. A putative catalytic mechanism for the dual activities of ACSs is proposed on the basis of the crystal structure and biochemical data. These findings not only expand our current understanding of ACS functions but also provide novel insights into the evolutionary origin of ACS genes.

Introduction

During the evolution from aquatic algae to terrestrial gymnosperms and angiosperms, plants underwent tremendous and extremely complex changes to adapt to the challenges of the terrestrial environment. To survive and reproduce, plants must change their growth, development, and metabolism accordingly. Gaseous ethylene is a very ancient phytohormone that exerts profound effects on many aspects of plant growth and development. It is also indispensable for plants to deal with a range of biotic and abiotic stresses (1, 2). Although the emergence of ethylene as a phytohormone is considered an important bridge between the changing environment and plant developmental adaptation (3), the origin and evolutionary history of the ethylene biosynthesis pathway in plants remain largely unknown.

In seed plants, the biosynthesis of ethylene mainly includes three important catalytic reactions (4). Methionine is first converted to S-adenosyl-methionine (SAM), which is further converted to 1-aminocyclopropane-1-carboxylic acid (ACC) by ACC synthase (ACS), and finally, ACC is oxidized to ethylene by ACC oxidase. Among these, the conversion from SAM to ACC represents the first committed and rate-limiting step of ethylene biosynthesis. The ethylene biosynthesis pathway of seed plants is often called the ACC-dependent pathway. The ACS proteins in seed plants are encoded by a multigene family. All ACS proteins contain a conserved aspartate aminotransferase (AAT)–like domain and belong to the α superfamily of pyridoxal-5′-phosphate (PLP)–dependent enzymes (5, 6).

The efficiency and level of ethylene biosynthesis in seed plants are generally higher than that in nonseed plants (7–9). Unlike the well-characterized ethylene biosynthesis pathway in seed plants, the ethylene synthesis routes in nonseed plants remain elusive. Some early studies indicated that exogenous ACC treatment can promote ethylene production in some uniconnate green algae (10–12) and multicellular charophytes (1), mosses (13), and ferns (14). However, other studies found that although ethylene emissions can be detected in the major groups of nonseed plants, exogenous ACC cannot be converted into ethylene, providing evidence for a non-ACC-dependent ethylene biosynthesis pathway in lower plants (7, 8, 15–17). In addition to these contrary findings, although the ancestry of ACS homologous genes can be traced back to the algal genome, biochemical and molecular biological analyses of ACS-like proteins in nonseed plants are very rare due to a lack of techniques in the early days. Our group previously cloned the only two ACS-like genes (PpACLs) in the moss Physcomitrella patens genome and found that neither of their encoded proteins had ACS activity (9). Of particular note, PpACL1 actually functions as a C-S lyase (9). It is still unclear when during evolution the ACC-dependent pathway emerged and how plants acquired the genes encoding ACS enzymes that can catalyze the conversion of SAM to ACC in the “Yang cycle” (4), which are termed genuine ACSs herein.

In addition to ACS, the α superfamily of PLP-dependent enzymes also includes aminotransferases and carbon-sulfur lyases (C-S lyases) (18). A previous bioinformatics-based study proposed that ACS genes might originate from plant-ACS–like genes that come from AA Tase genes (19). In particular, another study reported that an apple ACS protein, MdACS1, exhibits extremely low aminotransferase activity in addition to ACS activity (20). Therefore, it is generally believed that plant ACSs are evolutionarily related to aminotransferases. However, neither of the above studies included C-S lyases that belong to the same superfamily. The possibility of an evolutionary relationship between ACSs and C-S lyases has long been ignored.

Here, starting with Arabidopsis ACS7 (AtACS7), we found that, in addition to catalyzing the formation of the ethylene precursor ACC, genuine ACSs widely have C-S lyase activity. Two N-terminal motifs and a glutamine residue were found to be essential for conferring ACS activity to ACS-like proteins during the evolutionary process.
RESULTS
AtACS7 has ACS and Cβ-S lyase dual enzymatic activities
AtACS7 is one of the genuine ACS enzymes of ethylene biosynthesis in Arabidopsis, catalyzing the conversion of SAM to ACC through α,β-elimination (Fig. 1A, top). However, when using purified AtACS7 as a control for the in vitro Cβ-S lyase activity assay of PpACL1, we unexpectedly found that AtACS7 could also catalyze the cleavage of the Cβ-S bond of the substrate l-cystine and convert it into thiocysteine, ammonia, and pyruvate (Fig. 1A, bottom).

Analysis using an amino acid analyzer revealed the production of NH₄⁺ in both PpACL1- and AtACS7-catalyzed reactions but not in the two negative control reactions (Fig. 1B). The generation of pyruvate was examined using the 2,4-dinitrobenzene-hydrazine method. Reddish-brown colorations of 2,4-dinitrobenzene-hydrazone generated from pyruvate and 2,4-dinitrobenzene were only found in the two negative control reactions (Fig. 1B). The generation of Cβ-S bond of the substrate l-cystine and conversion into thiocysteine, ammonia, and pyruvate were unexpectedly found that AtACS7 can also catalyze the cleavage of the Cβ-S bond of the substrate l-cystine and convert it into thiocysteine, ammonia, and pyruvate (Fig. 1A, bottom).

Characterization of the Cβ-S lyase activity of AtACS7
As a PLP-dependent enzyme, it is well established that ACS requires PLP as the cofactor for its catalysis. We then tested the effects of PLP and aminoethoxyvinylglycine (AVG), a competitive inhibitor of PLP-dependent enzymes, on the Cβ-S lyase activity of purified AtACS7. The Cβ-S lyase activity of AtACS7 was also PLP dependent and could be totally suppressed by adding AVG to the reaction mixture to a final concentration of 5 μM (Fig. 1, H and I).

When using l-cystine as a substrate, AtACS7 achieved its maximal Cβ-S lyase activity at approximately 30°C (Fig. 1J), while the optimum pH was slightly basic at approximately 7.4 (Fig. 1K). Under optimal temperature and pH conditions, the enzymatic activity of AtACS7 peaked when the substrate concentration was approximately 4 mM. The maximum reaction rate reached approximately 79 μmol pyruvate mg⁻¹ min⁻¹, and the Kₘ value was determined to be 1.5 mM (Fig. 1L).

The structures of AtACS7 and MdACS1 are nearly identical
To address the structural basis that may support its dual enzyme activity, we determined the crystal structure of AtACS7 in complex with pyridoxal 5’-phosphate-γ-aminoethoxyvinylglycine (PPG) at 2.20-Å resolution [Protein Data Bank (PDB) code: 7DLW; statistics are summarized in table S1]. There are four AtACS7-PPG complexes that make up two dimers in each asymmetric unit. The PPG molecules showed good electron density, except for the flexible aminoethoxyvinyl groups (fig. S2A). The two dimers (chain A and B dimer and chain C and D dimer) exhibit almost identical structures, with a root mean square deviation (RMSD) of 0.76 Å, and both have the classic AAT-like folding (Fig. 2A). We used chain C and D dimer as reference for the following discussion. The structures of the two subunits are nearly the same, with an RMSD of 0.44 Å.

For the active site (Fig. 2B), Asn²¹⁷ and Tyr²⁴⁸ both form hydrogen bonds with the O3 of PPG, and Asp²⁴⁵ forms a hydrogen bond with the N1 of PPG. Arg²⁹³ and Arg²¹⁹ form salt bridges with the carboxyl group and phosphate group of PPG, respectively. Tyr¹⁶⁰ exhibits parallel stacking over the pyridine ring of PPG, while the key catalytic residue Lys²⁸⁵ is located at the opposite side of the PPG from Tyr¹⁶⁰. In addition to the above residues that belong to the same AtACS7 subunit, the side chain of Gln⁹⁸ from the other subunit is protruding near the PPG molecule (Fig. 2B).

Since apple MdACS1 has been suggested to have ACS and very low aminotransferase dual activities (20), we subsequently compared the complex structure of AtACS7-PPG with the complex structure of MdACS1-PPG (PDB code: 1M7Y) (22). The overall and the active site structures of the two ACSs are almost identical (Fig. 2, C and D, and fig. S1). Together, these results showed that AtACS7 has both ACS and Cβ-S lyase activities in vitro.

To further examine whether the dual enzymatic activities of AtACS7 exist in plant cells, we measured the ethylene emissions and pyruvate contents in the AtACS7-overexpressing transgenic Arabidopsis [lines AtACS7-ox-3-4 and AtACS7-ox-7-5, as used in our previous study (21)]. Besides the elevated ethylene production and the triple response phenotype, the higher levels of accumulated AtACS7 protein resulted in a significant increase in the content of pyruvate in both of the transgenic lines (Fig. 1, E to G). These results demonstrated that AtACS7 also has in planta Cβ-S lyase activity.

The Cβ-S lyase activity of ACSs is a common phenomenon
Since structural similarity generally indicates similarity of protein function, the extreme structural similarity between AtACS7 and MdACS1 persuaded us to purify the MdACS1 protein and examine whether it also has Cβ-S lyase activity. As expected, MdACS1 indeed had both ACS and Cβ-S lyase activities in vitro, although both were lower than AtACS7 (Fig. 3A).

ACS proteins are divided into three main groups based on their C-terminal sequences (23). AtACS7 is the only type III ACS in Arabidopsis, while MdACS1 belongs to the type II ACSs. We further examined the activities of different types of ACS proteins, such as GmACS7-like in soybean, OsACS5 and OsACS1 in rice, SslACS4 in tomato, and AtACS6, AtACS8, and AtACS11 in Arabidopsis. The results showed that all of them had ACS and Cβ-S lyase dual enzymatic activities (Fig. 3, B to H), suggesting that dual enzymatic activity may be a common feature of all types of ACSs in higher plants.

Q98 plays a substantial role in ACS activity
The Cβ-S lyase activities of ACS proteins in higher plants implied that ACSs have a close evolutionary relationship with Cβ-S lyases. It was previously reported that ACS proteins contain seven conserved domains (namely, boxes 1 to 7) (24). We swapped each of the seven boxes of Arabidopsis AtACS7 with P. patens PpACL1 and examined the enzymatic activities of these purified recombinant proteins. Among them, a recombinant named AtACS7-R6, in which the BOX2 of AtACS7 was replaced by PpACL1-BOX2, exhibited Cβ-S lyase activity but no ACS activity in the in vitro assays (Fig. 4A and fig. S3).

We subsequently determined the 2.95-Å resolution crystal structure of the AtACS7-R6 mutant in complex with PPG (PDB code: 7DLY; statistics summarized in table S1 and fig. S2B). The asymmetric unit contains one AtACS7-R6 dimer, and each AtACS7-R6 monomer forms a complex with the PPG molecule. The structures of these two subunits are nearly the same, with an RMSD of 0.12 Å. Compared with the wild-type AtACS7 dimer, the structure of the AtACS7-R6 mutant overall does not change much (with an RMSD of only 0.59 Å), except that the region replaced by PpACL1 BOX2 and adjacent region (residues 91 to 104) become disordered (Fig. 4B). Such a change...
Fig. 1. AtACS7 has ACS and C\(_2\)-S lyase dual enzymatic activities both in vitro and in planta. (A) Chemical reactions catalyzed by ACSs (top) or C\(_2\)-S lyases (bottom).

(B) Chromatograms of NH\(_4\)\(^+\) produced from the C\(_2\)-S bond cleavage of L-cystine, which is catalyzed by either PpACL1 or AtACS7 in vitro. (C) In vitro pyruvate production assays of PpACL1- or AtACS7-containing C\(_2\)-S lyase reaction systems. (D) In vitro ACS activity assays of PpACL1 and AtACS7. (E to G) The prominent accumulation of AtACS7 protein resulted in a strong triple-response phenotype, an elevated ethylene production, and a significant increase in the content of pyruvate in the etiolated seedlings of two independent AtACS7-overexpressing lines (AtACS7-ox-3-4 and AtACS7-ox-7-5). (H) The C\(_2\)-S lyase activity of purified AtACS7 was PLP dependent. (I) Exogenous AVG totally suppressed the C\(_2\)-S lyase activity of purified AtACS7. (J and K) Determination of the optimal temperature and pH for C\(_2\)-S lyase activity of purified AtACS7. (L) Estimation of kinetic parameters of C\(_2\)-S lyase activity of purified AtACS7 under the optimal temperature and pH conditions. The C\(_2\)-S lyase activities were measured by the generation of pyruvate using L-cystine as substrate. Data represent means ± SE (n ≥ 3, biological replicates). The number of biological replicates for each experiment is indicated in Materials and Methods. Asterisks indicate statistically significant differences based on Student’s t test (α = 0.01). For negative control (NC), blank buffer was used instead of purified proteins.
Insights into the catalytic mechanism of ACSs with dual enzymatic activities

In addition to AtACS7\textsuperscript{Q98A}, we also examined the activity of the AtACS7\textsuperscript{N217A} and AtACS7\textsuperscript{D245N} mutants. The N217A mutation abolished both the ACS and C\textsubscript{\(\beta\)}-S lyase activities of AtACS7 in vitro (Fig. 4A). However, the AtACS7\textsuperscript{D245N} mutant showed almost no C\textsubscript{\(\beta\)}-S lyase activity but still had ACS activity, a completely opposite phenotype to the AtACS7-R6 and AtACS7\textsuperscript{Q98A} mutants (Fig. 4A). This observation provided insights into a novel catalytic mechanism for the ACS and C\textsubscript{\(\beta\)}-S lyase dual activities of ACS proteins (Fig. 5).

For all PLP-dependent AAT-like enzymes, the aldehyde group of PLP first forms a Schiff base linkage with an amine acid substrate (l-cysteine for C\textsubscript{\(\beta\)}-S lyase activity or SAM for ACS activity) under the mediation of the lysine residue corresponding to K285 of AtACS7, thereby generating an external aldimine (26–28). In the C\textsubscript{\(\beta\)}-S lyase activity of AtACS7, on the basis of the bacterial C\textsubscript{\(\beta\)}-S lyases (26, 27), the C\textsubscript{\(\alpha\)} proton is extracted by K285, producing a quinonoid intermediate. The proton is then transferred to the S\textsubscript{\(\gamma\)} atom of the quinonoid intermediate, which breaks the bond between C\textsubscript{\(\beta\)} and S\textsubscript{\(\gamma\)}, resulting in the release of thiol. The remaining aldimine is later hydrolyzed to produce pyruvate and ammonia. During this process, the formation of a quinonoid intermediate is critical for the catalysis. The quinonoid formation requires the protonation of pyridine nitrogen by D245. Therefore, the D245N mutation leads to the deprotonation of the pyridine nitrogen (28), resulting in the abrogation of C\textsubscript{\(\beta\)}-S lyase activity, which is consistent with the results of the C\textsubscript{\(\beta\)}-S lyase activity assay (Fig. 4A).

Interestingly, the AtACS7\textsuperscript{D245N} mutant still retained considerable, albeit reduced, ACS activity (Fig. 4A). This is not entirely consistent with previous suggestions that the catalytic process of ACS activity also requires a quinonoid intermediate (29, 30), implying that the ACS activity is less dependent on the quinonoid intermediate than C\textsubscript{\(\beta\)}-S lyase activity. We proposed that the C\textsubscript{\(\gamma\)}-S bond of SAM in the external aldimine intermediate can be broken before C\textsubscript{\(\alpha\)} deprotonation. In this case, with the C\textsubscript{\(\gamma\)}-enzyme covalent intermediate, when C\textsubscript{\(\alpha\)} is deprotonated, the \(\alpha\)-carbanion may covalently link to C\textsubscript{\(\gamma\)} immediately to form ACC, without forming the quinonoid intermediate. Thus, the formation of a quinonoid intermediate is no longer necessary (Fig. 5). In this process, some residue other than K285 (possibly Y160) (29, 30) is required to break the C\textsubscript{\(\gamma\)}-S bond. Therefore, compared

### Phagelogenetic analysis was further carried out using sequences of ACSs and a series of aminotransferases and C-S lyases with AAT-like folding from a wide variety of organisms, such as humans, mice, yeast, bacteria, protozoa, mosquitoes, barley, Arabidopsis, tomato, and apple (table S2). The results showed that only functional ACSs had the glutamine residue corresponding to Q98, further supporting the importance of this residue for ACS activity (Fig. 4E).

To estimate the role of Q98, SAM-aldimine was docked to the wild-type AtACS7 structure using the program AUTODOCK (25). The result showed that the amide group of Q98 directly approached the sulfur atom of SAM (Fig. 4F). Considering that no reaction between the amide group and sulfonium ion has been reported, we propose that Q98 functions to modulate the SAM conformation.

### Fig. 2. Structure of AtACS7. (A) Illustration of AtACS7 dimer. (B) The active site of AtACS7. (C) Superposition of the AtACS7 dimer and MdACS1 dimer. (D) Comparison of the active sites of AtACS7 and MdACS1. In all the panels, the two AtACS7 subunits are colored yellow (chain C) and cyan (chain D), and the PPG molecules of AtACS7 are colored violet. The contents from MdACS1 are colored white for (C) and (D). See also table S1.
with the Cβ-S lyase activity, which only requires K285 for covalent bond breaking or bonding, the catalytic process of ACS activity is more complicated, supporting that it is evolutionarily more recent than Cβ-S lyase activity.

Proposed structural model of ACS-like proteins with ACS activity

To identify the motifs required for ACS activity, we performed web-based Multiple Expectation-maximum for Motif Elicitation (MEME) analysis using ACSs from *Arabidopsis* or ACSs that were functionally confirmed by ourselves (Fig. 3) with Cβ-S lyase PpACL1 and *Arabidopsis* aminotransferases AtACS10 and AtACS12 as controls. The results revealed highly conserved motifs among all sequences in the C terminus and variable motifs in the N terminus (Fig. 6A and table S3). Compared with those of PpACL1 and the two amino transferases, two N-terminal motifs and seven C-terminal motifs that are spatially conserved in a specific order were named ACS motifs 1 to 9. The motif that is only present in the N terminus of two amino-transferases AtACS10 and AtACS12 was named the AAT motif. Considering the key glutamine residue mentioned above, all nine ACS motifs, but excluding the AAT motif, and a glutamine residue corresponding to Q98 of AtACS7 in the second ACS motif were proposed to be collectively required for ACS activity (Fig. 6B).

Validation of the proposed ACS model

Compared to AtACS7, the Cβ-S lyase PpACL1 has seven C-terminal ACS motifs but lacks the two N-terminal ones (Fig. 6A). We then swapped the N-terminal domain of PpACL1 with that of AtACS7 so that...
Fig. 4. The Q98 residue plays a substantial role in conferring ACS activity. (A) In vitro C₅-S lyase and ACS activity assays of purified AtACS7 mutants. The activities of wild-type AtACS7 were regarded as 100%. Data represent means ± SE (n ≥ 3, biological replicates). The number of biological replicates for each protein is indicated in Materials and Methods. (B) Superimposition of the overall structure of AtACS7-R6 and wild-type AtACS7. The two AtACS7-R6 subunits are colored yellow and cyan, respectively, and the PPG molecules are colored violet. Wild-type AtACS7 is colored white, except for the 91-to-104 region, which is colored red. (C) Comparison of the active sites of AtACS7-R6 (colored) and wild-type AtACS7 (white). (D) Determination of in planta C₅-S lyase and ACS activities of the AtACS7 Q98A mutant by measuring the contents of pyruvate or ACC in the Agrobacterium-infiltrated tobacco leaves as described in Materials and Methods. Free eYFP was injected as an NC, while wild-type AtACS7 was used as a positive control. Data represent means ± SE (n = 6, biological replicates). Black asterisks indicate statistically significant differences in pyruvate contents, while a red asterisk indicates statistically significant differences in ACC contents compared with the eYFP control based on Student’s t test (α = 0.05). (E) Phylogenetic analysis of functional ACS proteins, ACS-like proteins, aminotransferases, and C₅-S lyases from a wide variety of organisms. The phylogenetic tree was constructed using the neighbor-joining method in MEGA X software. Numbers at each interior branch indicate the bootstrap values of 1000 replicates. The bar indicates a genetic distance of 0.2 cM. Detailed organisms and locus numbers or PDB IDs of all protein sequences are listed in table S2. All functional ACS proteins form a separate clade (red) containing the glutamine residue corresponding to Q98 of AtACS7. (F) Docking for SAM at the AtACS7 active site.
that the recombinant protein N7-PpACL1 met our proposed model (Fig. 6C). As expected, subsequent in vitro enzymatic analysis revealed the ACS activity of N7-PpACL1 (Fig. 6, D and E). The results confirmed the model we proposed and, more importantly, demonstrated that the N terminus plays a pivotal role in conferring ACS activity to ACS-like proteins.

To further verify the model, we bacterially expressed 21 ACS-like proteins from plant species ranging from chlorophytes to angiosperms (fig. S5) and determined their ACS activities by measuring ACC contents in the supernatants of the transformed bacterial cultures as described in Materials and Methods (31). As shown in Fig. 7A and figs. S6A and S7, significant increases in ACC contents were detected in the supernatants of bacteria overexpressing ACS-like proteins that contain all nine conserved ACS motifs and the key glutamine residue corresponding to Q98 of AtACS7, for instance, Gm.01G003900.1, Gm.07G128000.1, and Gm.08G030100.1 of *Glycine max*; AmTr_v1.0_scaffold00111.98 of *Amborella trichopoda*; MA_103524g0010 of *Picea abies*; PITA_24974 of *Pinus taeda*; and Gb_12852 and Gb_38571 of *Ginkgo biloba*. In contrast, ACC contents were not detected in the bacterial cultures overexpressing the rest of the ACS-like proteins that lack any of the nine ACS motifs and/or the glutamine residue (Fig. 7A and figs. S6A and S7), further supporting the standard ACS model we proposed (Fig. 6B). These included AmTr_v1.0_scaffold00069.217 of *A. trichopoda*, MA_66897g0010 of *P. abies*, PITA_38831 of *P. taeda*, Gb_22779 of *G. biloba*, and all the tested ACS-like proteins from nonseed plant species such as lycophytes, ferns, liverworts, mosses, charophytes, and chlorophytes (Fig. 7A). The only exception was AtACS1, which met the model requirements but did not show ACS activity, as AtACS1 lacks the asparagine corresponding to N217 in AtACS7. This residue interacts with PLP and plays an essential role in all AAT-like enzymes [fig. S7 and (24)]. The ACS activities of several ACS-like proteins were double checked using purified recombinant GST- or His-fusion proteins to further confirm the results (fig. S6B).

In addition to these unreported ACS-like proteins, we also performed motif analysis to the ACS-like proteins whose activities have already been reported in the literature (9, 24, 32–43). As shown in Fig. 7B and fig. S7, all genuine ACSs meet the ACS model requirements. Those that did not show ACS activities failed to have the nine ACS motifs and/or the key Q98 residue. We also measured the Cβ-S lyase activity of ACS-like proteins listed in Fig. 7A using crude protein extracts (fig. S8A) or purified proteins (fig. S8B). We found that, except for the genuine ACSs, all tested ACS-like proteins having in vitro Cβ-S lyase activity did not meet the ACS model requirements (Fig. 7A and fig. S8), indicating that the nine motifs and the key glutamine residue are not entirely necessary for Cβ-S lyase activity.
DISCUSSION

Although it is generally believed that all plant ACSs originated from AATase-derived plant-ACS–like proteins (19, 20), our results suggested that ACSs may have a closer evolutionary relationship with Cβ-S lyases than aminotransferases. This was mainly supported by two lines of evidence: (i) Genuine ACSs with aminotransferase activity are very rare in seed plants. So far, the only example of ACC synthase with aminotransferase activity is apple MdACS1 (20). However, here we demonstrated that seed plant ACSs widely have considerable Cβ-S lyase activity. (ii) Neither of the two ACS-like proteins in

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Fig. 6. The acquisition of two N-terminal motifs is a key event for ACS activity. (A) MEME analysis was performed using Arabidopsis ACS proteins and the functionally confirmed ACS sequences from soybean, rice, and tomato in addition to P. patens Cβ-S lyase PpACL1 and Arabidopsis aminotransferases AtACS10 and AtACS12 as controls. The results revealed nine motifs (ACS motifs 1 to 9) that are collectively required in a specific order for ACS activity. The motif present only in the two Arabidopsis aminotransferases AtACS10 and AtACS12 was named AAT. (B) Proposed structural model of the genuine ACS proteins contains nine conserved ACS motifs and the glutamine residue corresponding to Q98 of AtACS7 (indicated with a star and an arrow) in the second ACS motif. (C) Schematic diagram of N-terminal substitution between AtACS7 and PpACL1. The N7-PpACL1 recombinant protein is composed of the N-terminal sequence of AtACS7 and the C-terminal sequence of PpACL1. The purified recombinant protein N7-PpACL1 had no Cβ-S lyase activity in vitro (D) but gained in vitro ACS activity (E). The empty vector pET28a and Cβ-S lyase PpACL1 were used as controls. Data represent means ± SE (n = 3, biological replicates). See also table S3.
Fig. 7. Validation of the proposed ACS model. (A) Validation of the proposed ACS model using ACS-like proteins from plant species ranging from chlorophytes to angiosperms. (B) Validation of the proposed ACS model using functionally confirmed ACS proteins in the literatures. The plus symbols (+) represent the presence of identical ACS motifs or Q98 residue and/or exhibiting ACS or Cβ-S lyase activity, while the minus symbols (−) represent the absence of identical motifs or Q98 residue and/or no ACS or Cβ-S lyase activity detected, respectively. Data shown for ACS activity determination were based on at least four biological replicates, while those for Cβ-S lyase activity measurements were from at least three biological replicates. Raw data are presented in figs. S5 to S8. * indicates the special issue of AtACS1, which meets the model requirements but lacks a key asparagine residue as discussed in the text.
the moss *P. patens* exhibits ACS activity, and PpACL1 functions as a Cβ-S lyase (9). Combined with the fact that several nonseed plant ACS–like proteins displayed Cβ-S lyase activity (Fig. 7A), it is tempting to postulate that Cβ-S lyase activity may be quite ancient and widely present in plants of different taxonomic categories. More notably, the Cβ-S lyase PpACL1 could gain ACS activity when its N terminus was replaced by that of AtACS7 (Fig. 6, C to E), confirming that ACSs are closely related to Cβ-S lyases. These findings shed new light on the molecular evolution of ACSs.

On the basis of the crystal structures and MEME analyses, we proposed that nine ACS motifs and a specific glutamine residue corresponding to Q98 of AtACS7 in the second ACS motif are required for genuine ACSs. While most of the tested ACS–like proteins from chlorophylls to angiosperms have the seven conserved ACS motifs at the C terminus, only those with a complete assembly of the nine motifs showed ACS activity, suggesting that the two ACS motifs on the N terminus are prerequisites for ACS-like proteins to gain ACS activity during evolution (Fig. 7). This also explained why the recombinant protein N7-PpACL1 could acquire ACS activity. The “natural” acquisition of the two N-terminal motifs occurred following the appearance of gymnosperm species during plant evolution (Fig. 7A). This suggests that genuine ACS activity may be unique to seed plant species.

It is controversial whether the ACS-like proteins in *Marchantia polymorpha* have ACS activity. There are two ACS-like genes, Mapoly0034s0060.1 and Mapoly0001s0058.1, in the *Marchantia* genome. Li *et al.* (44) proposed that the two *Marchantia* ACS homologs, they named as MpcACS1 and MpcACS2, have ACS activity based on measurements of changes of ACC contents in the *Marchantia* knockout mutants and MpcACSL- or MpcACSL-expressing yeast cells. However, Katayose *et al.* (45) recently found that knockout of one or both MpcACSL-like genes did not significantly change ACC contents and ethylene production in the *Marchantia* mutants. Our results here support the findings of Katayose *et al.:* No ACC production was detected in *Escherichia coli* cell cultures expressing either of the two *Marchantia* ACS–like genes (Fig. 7A and fig. S6A). In addition, purified Mapoly0034s0060.1 or Mapoly0001s0058.1 protein failed to convert SAM to ACC in our in vitro activity assays (fig. S6B). In contrast, we found that both Mapoly0034s0060.1 and Mapoly0001s0058.1 have Cβ-S lyase activity (fig. S8B). Furthermore, both ACS-like proteins in *Marchantia* lack the key glutamine residue corresponding to the Q98 of AtACS7 (Fig. 7A and fig. S7). This residue is critical for maintaining the conformation of SAM for catalysis (Fig. 4, C and F). Mutation of this amino acid reduced the enzyme activity by more than 93% (Fig. 4, A and D). In addition, the *Marchantia* ACS–like protein Mapoly0001s0058.1 (named as MpcACSL in 44) lacks ACS motif 2, one of the nine motifs collectively required for ACS activity (Fig. 7A). Therefore, the two ACS-like proteins in *Marchantia* do not seem to be genuine ACSs as defined here.

Endogenous ACC is detected in wild-type liverwort *M. polymorpha* (44, 45). It also exists in moss and ferns (7, 13). However, in this study, we found that the ACS–like proteins of liverwort *M. polymorpha* and ferns *Salvinia cucullata* and *Azolla filiculoides* do not have ACS activity (Fig. 7A and fig. S6). Hence, it is reasonable to speculate that there might be alternative pathways for ACC biosynthesis in species lacking functional, genuine ACSs as defined here.

ACS activities detected in the in vitro assays and in an *E. coli* expression system are generally consistent with those in the in planta studies (Figs. 1 and 4 and fig. S6B) (32–34, 45). However, they are not the same as demonstrating the existence or absence of physiological activities in plants. Besides, there are also potential issues with the in planta transient expression system in tobacco. For example, expression levels of the ACS-like genes in transiently transformed tobacco leaves might be different from the endogenous levels expressed in a certain organ or at a certain developmental stage of plants, especially of nonseed plant species. Likewise, functions of the ACS-like genes of nonseed plants might be different when expressed in seed plant tobacco. Therefore, in planta functions of ACS-like genes and the proposed ACS model should be further tested using ACS-like gene knockout mutants and overexpressing lines of organisms belonging to different taxonomic classifications.

ACSs are widely known as rate-limiting enzymes in the ethylene biosynthesis pathway. Here, we showed that this enzyme also has Cβ-S lyase activity and produces pyruvate when using l-cysteine as a substrate. Although only the in vivo Cβ-S lyase activity of AtACS7 was measured here (Figs. 1G and 4D), our in vitro experiments demonstrated that the Cβ-S lyase activity of ACS or ACS-like proteins simply requires PLP and cysteine, both of which exist in plants. We speculate that the in vitro Cβ-S lyase activities of the tested ACS-like proteins should reflect their Cβ-S lyase activity in vivo. However, whether a given ACS-like protein has in vivo Cβ-S lyase activity needs further verification.

Pyruvate is a common metabolite that is an end product of glycolysis and an energy substrate for the mitochondrial Krebs cycle. It is also well known for its protective properties against stressful conditions in both animal and plant cells (46, 47). In addition, another product of the Cβ-S lyase activity of ACSs, thioicysteine, is prone to breakage of the disulfide bond under reducing conditions and production of hydrogen sulfide, a multifunctional signaling molecule that participates in almost all aspects of plant life (48). Together, the discovery of the ACS and Cβ-S lyase dual enzymatic activities of ACSs in seed plants not only improves the general knowledge of the diversity of Cβ-S lyases but also greatly expands the current understanding of the biological functions of ACS proteins. However, it is also possible that ACS activity is the main function of the modern ACS and that Cβ-S lyase activity is a residual activity. Whether it has significant biological functions awaits further studies.

**MATERIALS AND METHODS**

Identification of ACS homologs

To identify ACS homologs from the sequenced genomes of land plants, lycophytes and ferns, liverworts, charophytes, and chlorophytes listed in Fig. 7, all ACS proteins of the model plant *Arabidopsis thaliana* were used to construct a hidden Markov model (HMM) profile. Subsequently, HMMER (http://hmmer.org/download.html) searches against the above-mentioned plant proteomes (table S4) were performed. All putative ACS proteins sequences obtained were evaluated and confirmed using the NCBI Conserved Domain Database (https://ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi). Transcript sequences of the confirmed ACS protein sequences were retrieved on the basis of the locus numbers.

Expression and purification of ACS homologous proteins

ACS homologous genes were either cloned from the cDNA of the corresponding plant species or chemically synthesized at the Beijing Genomics Institute. Fragments in TA cloning vectors were then
transferred into pET-28a or pGEX-6p-1 expression vector by the routine digestion and ligation method. The recombinant plasmids were then sequenced for confirmation and transformed into expression host E. coli BL21 [Rosetta 2 (DE3) pLysS]. Construction details for each gene are presented in table S5. Recombinant proteins were purified by affinity chromatography using HisTrap FF columns or Glutathione Sepharose 4 Fast Flow (GE Healthcare) columns according to the manufacturer’s instructions.

Crystallization and structure determination

HisTrap affinity chromatography–purified AtACS7 or AtACS7-R6 mutant protein was further purified with ion exchange (Hitrap Q HP) and gel filtration (Superdex 200) chromatography. The purified protein was concentrated to 10 mg/ml in 20 mM tris (pH 8.0) and 150 mM NaCl. Before crystallization, 1 mM PLP and AVG were supplemented to the protein solutions. Using the hanging drop vapor diffusion method, AtACS7 was crystallized in 0.1 M tris (pH 8.6), 23% (w/v) polyethylene glycol (PEG)−3350, and the AtACS7-R6 mutant was crystallized in 0.1 M Hapes (pH 7.6), 6% (w/v) PEG−10000. Diffraction data were collected at SSRF (Shanghai Synchrotron Radiation Facility) synchrotron and processed with the HKL2000 (49) program. The structure of apple ACS (PDB code: 3P1U) was used as the searching model for the structure of AtACS7-R6 mutant. Both structure models were refined using COOT (51) and PHENIX.

ACS activity assay

The in vitro ACS activity assay using purified ACS-like proteins with His or GST tag was performed as described previously (9) with modifications. Briefly, 20 μg of the purified protein was pipetted into 20-ml gas chromatography (GC) vials containing 460 μl of ACS assay buffer [50 mM EPPS [N-(2-Hydroxyethyl)piperazine-N′- (3-propanesulfonic acid)] (Sigma-Aldrich) (pH 8.5), 10 μM PLP (Sigma-Aldrich), and 2 mM dithiothreitol] and 20 μl of 10 mM SAM (Sigma-Aldrich). The mixture was made to a total volume of 500 μl and then incubated at 30°C for 30 min. The amount of ACC was determined by the method in (52). Briefly, when the reaction was terminated, 18 drops of fresh cold mixture of 10% NaClO and saturated NaOH (1:2, v/v) was added to convert the formed ACC to ethylene. Vials were capped immediately and incubated on ice for 5 min. Ethylene emission was measured using GC (Agilent 7890A). The ACS activity was calculated as the amount of ACC converted from SAM per minute and per microgram protein according to an ACC standard curve. The protein concentration in each assay was determined by the Bradford method. Data shown for purified GmACS7-like OsACS5, OsACS1, SIACS4, and AtACS6 in Fig. 2; AtACS7-R6, AtACS7Q98A, AtACS7N217A, and AtACS7D245N in Fig. 4A; N7-PpACL1 in Fig. 6E; and PITA_24974 in fig. S6B were results of three biological replicates. Data shown for purified GmACS7-like OsACS5, and PITA_24974 in fig. S6B were from six biological replicates.

Purification of some ACS-like proteins using available conventional methods is difficult. Li et al. (31) reported that ACS overexpressed in BL21 E. coli cells could convert the endogenous, highly prevalent SAM in the bacterial cells into ACC and subsequently secrete it into the growth medium. Following their method, we determined ACS activities of ACS-like proteins by measuring the contents of ACC accumulated in the growth medium of bacteria transformed with ACS-like genes. In brief, BL21 strains harboring ACS-like genes were grown in the LB liquid medium and then induced by isopropyl-β-d-thiogalactopyranoside (IPTG). After induction for 20 hours at 16°C, accumulations of these proteins were visualized on SDS–polyacrylamide gel electrophoresis gel. The cell cultures containing ACS-like protein with similar accumulation levels of AtACS7 were centrifuged at 10,000g. Aliquots of the supernatants were transferred to 20-ml GC vials and sealed with parafilm. ACC contents were indirectly determined by measuring ethylene product, as described in (52). Data shown for Glyma.01G039001.0, Glyma.07G128000.1, Glyma.08G031000.1, AmTr_v1.0_scaffold00111.98, MA_103524g0010, PITA_24974, Gb_12852, Gb_38571, and Mapoly0034s0060.1.p were results from three biological replicates. Data shown for AmTr_v1.0_scaffold00069.217, MA_66897g0010, Gb_22779, Mapoly001s0058.1.p, 16049, and Cre06.g306400.t1.2 were from four biological replicates. Data for PITA_38831, Sacu_v1.1_s0086.g018403, Azf_S0335.g065524, 75495, GBSM10000679.1.p, and GBSM1015261.1.p1 were based on results of five biological replicates.

To measure the contents of ACC in Agrobacterium-infiltrated tobacco (Nicotiana benthamiana) leaves, Agrobacterium tumefaciens strain GV3101 harboring 35S: eYFP, 35S:AtACS7Q98A-eYFP, or 35S:AtACS7-eYFP fusion gene was mixed with the strain containing P19 silencing suppressor and injected into the epidermis of the leaves of 5-week-old tobacco plants, as described previously (53). For each targeted leaf, half (divided by the midrib) was infiltrated with the eYFP-only control culture (35S:eYFP) and the other half with an AtACS7 construct culture (35S:AtACS7Q98A-eYFP or 35S:AtACS7-eYFP). After being incubated in the dark for 48 hours, five to six infiltrated leaves were pooled. ACC contents were measured as described in (54). Briefly, leaves were grounded in 95% ethanol and incubated at 85°C for 20 min. After centrifugation at 10,000g for 15 min at 4°C, the supernatant was collected and mixed with 85% ethanol by vortexing. The mixture was incubated at 70°C for 30 min and then centrifuged again at 10,000g for 15 min at 4°C. The supernatant was dried in a SpeedVac, and the residues were then resuspended in 1 ml of ddH2O for ACC quantification, as described above. Data shown here were from six biological replicates.

Ethylene emission measurements

Ethylene emissions from 3-day-old etiolated Arabidopsis seedlings of the wild-type and two individual 35S:AtACS7-eGFP transgenic lines were measured as described (55). Briefly, Arabidopsis etiolated seedlings were incubated in 12-ml vials with 3-ml liquid 1/2 MS (Murashige and Skoog) medium and grown in plant growth chamber for 24 hours (22/19°C) in darkness after being sealed with caps. Ethylene accumulated in the vials were measured by GC (Agilent 7890A), and the rate of ethylene production was expressed as nanoliter per gram of seedlings (fresh weight) per 24 hour. All experiments were performed in three biological replicates.

Determination of C3′-S lyase activity and kinetics

In vitro C3′-S lyase activity was assayed using purified ACS proteins with a His or GST tag as described (9). Briefly, 100 μg of purified enzyme was added to 300 μl of reaction buffer (100 μM PLP, 4 mM L-cystine, and 75 mM potassium phosphate). The mixture was incubated at 30°C for 30 min before chloroform was added to denature
proteins. The mixture was then centrifuged at 12,000 rpm for 10 min at 4°C. To quantify the pyruvate product, the supernatant recovered after centrifugation was mixed with 2,4-dinitrophenylhydrazine [0.1% (w/v) in 2 M HCl], and the reaction was stopped by adding 1.5 M NaOH. Then, the pyruvate content was determined by measuring the absorbance at 520 nm and comparing to a standard curve. To measure the reduction of l-cystine and production of NH₄⁺, supernatant of the reaction mix mentioned above was purified by suction filtration (Luer syringe filter, PES (Polyethersulfone) 0.22 μm; syringe 2.5 ml) and analyzed using an amino acid analyzer (MembraPure A300, GmbH) following the manufacturer’s instructions. Data shown for the in vitro C₅-S lyase activity of purified GmACS7-like, OsACS5, OsACS1, SlACS4, and AtACS6 in Fig. 2; AtACS7-R6, AtACS7⁷⁶⁹⁸A, AtACS7ⁿ²¹⁷A, and AtACS7¹²¹⁵⁵ in Fig. 4A; and N7-PpACL1 in Fig. 6D were from three biological replicates. Data shown for the in vitro C₅-S lyase activity of purified ACS7 with different tags in Fig. 1D, and MdACS1 and AtACS11 in Fig. 2 were from four biological replicates. Data shown for the in vitro C₅-S lyase activity of purified AtACS8 were from five biological replicates. Data shown for the in vitro C₅-S lyase activity of purified OsACS6 in Fig. 2D, and Sacu_v1.1_s0086.g018403, Mapoly0001s0058.1.p, and Mapoly0034s0060.1.p in fig. S8B were results of six biological replicates. To test the inhibitory effect of AVG, 5 μM of the inhibitor was included in the reaction mixtures. Data shown in Fig. 11 are the results of four biological replicates. The kinetic parameters Kₘ and Vₘₐₓ of AtACS7 C₅-S lyase were calculated using the Michaelis-Menten equation. Data presented in Fig. 1 (J to L) are results of three biological replicates. Because purification of some ACS-like proteins is difficult, we measured C₅-S lyase activities of these proteins using crude protein extracts as described in (56). Data shown for Glyma.07G128000.1, MA_103524g0010, PITA_24974, PITA_38831, Gb_12852, Gb_38571, Gb_22779, and Cre06.g306400.t1.2 are results from three biological replicates. Data shown for AmTr_v1.0_scaffold0111.98, AmTr_v1.0_scaffold0069.217, MA_66897g0010, 75495, and 16049 were results from four biological replicates. Data shown for Glyma.08G030100.1 were from five biological replicates. Data shown for Glyma.01G039000.1, Azhi_S3353_g065524, GBSM01015261.1.p1, and GBSM01000679.1.p were from six biological replicates.

In planta C₅-S lyase activity of AtACS7⁷⁶⁹⁸A was determined by measuring the content of pyruvic acid in Agrobacterium-infiltrated tobacco (N. benthamiana) leaves. The same infiltration extracts prepared for ACC content measurements were used to determine the pyruvate level in each treatment. Measurements of pyruvate contents were performed following the manufacturer’s instructions (BC2000, Solarbio Life Sciences). Data shown here were based on the results of six biological replicates. For 35S:AtACS7⁺gfp stable transgenic lines and their wild-type control, 3-day-old etiolated seedlings were harvested, and pyruvate contents were quantified using the same kit (BC2000, Solarbio Life Sciences). The results presented were from three biological replicates.

**Phylogenetic analysis**

Protein sequences of aminotransferases, C₅-S lyases, and ACSs from a wide variety of organisms such as human, mouse, yeast, bacteria, protozoa, mosquito, barley, Arabidopsis, tomato, and apple (table S2) were aligned using MUSCLE (Multiple Sequence Alignment, https://ebi.ac.uk/Tools/msa/muscle/). The phylogenetic tree was generated using the neighbor-joining method in MEGA X software (57). Numbers at each interior branch indicate the bootstrap values of 1000 replicates. The bar indicates a genetic distance of 0.2 cM.

**Motif identification**

Motifs of the ACS proteins were identified statistically by the web-based MEME motif finder (http://meme-suite.org/tools/meme) (58) with motif length set as 6 to 50 and number of motifs 25. The MAST (Motif Alignment and Search Tool) program (http://meme-suite.org/tools/mast) (59) was used to search protein motifs of all ACS proteins; the ACS-like and AAT proteins were also included in the analysis.

**SUPPLEMENTARY MATERIALS**

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