Molecular Basis for Sesterterpene Diversity Produced by Plant Terpene Synthases

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ABSTRACT

Class I terpene synthase (TPS) generates bioactive terpenoids with diverse backbones. Sesterterpene synthase (sester-TPS, C25), a branch of class I TPSs, was recently identified in Brassicaceae. However, the catalytic mechanisms of sester-TPSs are not fully understood. Here, we first identified three nonclustered functional sester-TPSs (AtTPS06, AtTPS22, and AtTPS29) in Arabidopsis thaliana. AtTPS06 utilizes a type-B cyclization mechanism, whereas most other sester-TPSs produce various sesterterpene backbones via a type-A cyclization mechanism. We then determined the crystal structure of the AtTPS18–FSPP complex to explore the cyclization mechanism of plant sester-TPSs. We used structural comparisons and site-directed mutagenesis to further elucidate the mechanism: (1) mainly due to the outward shift of helix G, plant sester-TPSs have a larger catalytic pocket than do mono-, sesqui-, and di-TPSs to accommodate GFPP; (2) type-A sester-TPSs have more aromatic residues (five or six) in their catalytic pocket than classic TPSs (two or three), which also determines whether the type-A or type-B cyclization mechanism is active; and (3) the other residues responsible for product fidelity are determined by interconversion of AtTPS18 and its close homologs. Altogether, this study improves our understanding of the catalytic mechanism of plant sester-TPS, which ultimately enables the rational engineering of sesterterpenoids for future applications.

Keywords: terpenoid, terpene synthase, sesterterpene, cyclization mechanism, crystal structure

INTRODUCTION

Plants synthesize many terpenoids (including phytohormones) with diverse ecological functions to help them survive in their local environments. The structural diversity of plant terpenoids is largely determined by two consecutive, but separate, enzymes, namely, short-chain prenyltransferases and terpene synthases (TPSs) (Chen et al., 2011; Wang et al., 2016a). Plant TPSs can be divided...
Phylogenetic analyses show that eight subfamilies of plant TPSs are well established (TPS-a to TPS-h). Among them, the TPS-a subfamily (class I TPSs with an N-terminal inactive β domain and a C-terminal active α domain), which is common in flowering plants, provides the basis for the sub- and/or neofunctionalization of plant TPSs in this clade (Chen et al., 2011). There are 22 TPS-a members in the Arabidopsis genome (a total of 32 TPS genes), some of which have been demonstrated to be mono- (C10), sesqui- (C15), and di-TPS (C20) derivatives (Tholl and Osbourn, 2004). Recently, our group and Osbourn’s group functionally characterized novel PT-TPS-P450 gene clusters in Arabidopsis, and these genes showed unprecedented sester-TPS activity by utilizing geranylgeranyl pyrophosphate (GGPP) as a substrate (Wang et al., 2016a; Huang et al., 2016b). Phylogenetic analyses show that eight subfamilies of plant TPSs are well established (TPS-a to TPS-h). Among them, the TPS-a subfamily (class I TPSs with an N-terminal inactive β domain and a C-terminal active α domain), which is common in flowering plants, provides the basis for the sub- and/or neofunctionalization of plant TPSs in this clade (Chen et al., 2011). There are 22 TPS-a members in the Arabidopsis genome (a total of 32 TPS genes), some of which have been demonstrated to be mono- (C10), sesqui- (C15), and di-TPS (C20) derivatives (Tholl and Osbourn, 2004). Recently, our group and Osbourn’s group functionally characterized novel PT-TPS-P450 gene clusters in Arabidopsis, and these genes showed unprecedented sester-TPS activity by utilizing geranylgeranyl pyrophosphate (GGPP) as a substrate (Wang et al., 2016a; Huang et al., 2016b). Recently, our group and Osbourn’s group functionally characterized novel PT-TPS-P450 gene clusters in Arabidopsis, and these genes showed unprecedented sester-TPS activity by utilizing geranylgeranyl pyrophosphate (GGPP) as a substrate (Wang et al., 2016a; Huang et al., 2016b).
only a few studies on sester-TPSs from plants and fungi (Chiba et al., 2013; Matsuda et al., 2015, 2016; Ye et al., 2015; Chai et al., 2016; Bian et al., 2017; Huang et al., 2017, 2018; Shao et al., 2017). Two early-stage cyclization mechanisms of GFPP in the catalytic pocket of sester-TPSs have been proposed, namely, type A, C1-IV-V (cyclization between the C1 cation, the C14–C15 olefin [the fourth double bond, IV] and the C18–C19 olefin [the fifth double bond, V]) and type B, C1-III-IV (cyclization between the C1 cation, the C10–C11 olefin [the third double bond, III], and the C14–C15 olefin [the fourth double bond, IV]) (Minami et al., 2018; Sato et al., 2018). However, the amino acid(s) determining the difference between type-A sester-TPS and type-B sester-TPS remains largely unknown. Moreover, no sester-TPS from plants utilizing the type-B cyclization mechanism has been characterized to date. Although several routes to the formation of polycyclic sesterterpene backbones from GFPP have been proposed, all identified plant sester-TPSs share the 5/15 bicyclic intermediate (type-A route) (Huang et al., 2017, 2018; Shao et al., 2017). Recently, we demonstrated that single amino acids (Gly 328 in AtTPS18/AtTPS25, Gly325 in AtTPS19, and Pro328 in AtTPS30) are critical for the substrate specificity of sester-TPSs in Arabidopsis. Specifically, AtTPS18 G328W, AtTPS19 G325W, AtTPS25 G328W, and AtTPS30 P328W lost their sester-TPS activity, while mutants (AtTPS18G328W, AtTPS19G325W, and AtTPS25G328W) utilized geranyl pyrophosphate, farnesyl pyrophosphate (FPP), or geranyl-geranyl pyrophosphate (GGPP) as substrates (Chen et al., 2019). However, it is unclear how sester-TPSs accommodate GFPP conformers and direct the rearrangements and neutralization of C25 prenyl carbocation intermediates to produce diverse sesterterpenes.

In this study, we identified three additional sester-TPS encoding genes (AtTPS06, AtTPS22, and AtTPS29) that are not located in the GFPPS-TPS-P450 cluster in Arabidopsis. Interestingly, AtTPS06 catalyzed reactions through the 5/11 bicyclic intermediate (type-B route) based on the structure of the resulting sesterterpene. We further discovered the amino acids that control the product specificity/fidelity of plant sester-TPSs by determining the crystal structure of AtTPS18, and the key amino acids were

Figure 2. Proposed Cyclization Scheme for the Synthesis of Sesterterpene Backbones by Arabidopsis sester-TPSs.

The chemical structures elucidated in this study are indicated by red numbers. The products of AtTPS18 and its mutants are numbered C3–C7 according to their retention time in the GC–MS experiment carried out in this study (see below). All chemical structure elucidation data in this study are shown in Supplemental Tables 1 and 2 and Supplemental Figures 1–5. IM, intermediate. C3 ((+)-brassitetaene B), C6 ((+)-brassitetaene A), and C9 ((–)-caprutriene) were previously determined by Huang et al. (2017, 2018), whereas C7 ((–)-thalianatriene) and C8 ((–)-retigeranin B) were elucidated by Shao et al. (2017).
confirmed by site-directed mutagenesis. Our results provide a feasible pipeline for producing high-value or unnatural sesterterpenoids on a large scale using a synthetic biology strategy.

RESULTS AND DISCUSSION

Type-B Sester-TPSs, Possible Progenitors of Type-A Sester-TPSs, Are Discovered in Brassicaceae

Due to the huge functional divergence of plant TPSs (Chen et al., 2011; Zhou and Pichersky, 2020), a sufficient number of characterized sester-TPSs, together with the chemical structures of their products, is a prerequisite for understanding the catalytic mechanism of plant sester-TPSs. In an effort to discover new sester-TPSs from *A. thaliana*, we screened ten uncharacterized members (*AtTPS01*, *AtTPS06*, *AtTPS07*, *AtTPS09*, *AtTPS15*, *AtTPS16*, *AtTPS22*, *AtTPS26*, *AtTPS28*, and *AtTPS29*; we did not obtain the full-length cDNA of *AtTPS05*) in the TPS-a subfamily using an optimized *Escherichia coli* (E. coli) system (Figure 1A). A partial mevalonate (MVA) pathway (pMBIS construct [Martin et al., 2003]) was introduced into the *E. coli* system, and mevalonolactone was added to the culture medium to increase the production of the terpenoid of interest (Figure 1B). When coexpressed with *AtGFPPS2* (*At3g14550*), sesterterpenes (selective ion monitoring [SIM] mode for gas chromatography–mass spectrometry [GC–MS], *m/z* 340 for C25H40) were detected in transgenic *E. coli* strains harboring *AtTPS06*, *AtTPS22*, or *AtTPS29* (Figure 1C). No sester-TPS activity was observed for the other tested TPSs. Notably, *AtTPS01*, *AtTPS06*, *AtTPS07*, *AtTPS09*, *AtTPS15*, *AtTPS16*, and *AtTPS26* encoded di-TPSs (Supplemental Figures 6 and 7), and the chemical structures of these diterpenes merit further investigation. To determine their structures, we purified compound 1 from *AtTPS06* (0.26 mg) and compound 2 from *AtTPS29* (0.44 mg). One-dimensional and two-dimensional nuclear magnetic resonance (NMR) analysis of these compounds (*C1* and *C2*, respectively) revealed their planar structures and relative stereochemistries, and both compounds were found to be novel sesterterpenes (all chemical structure elucidation data in this study are shown in Supplemental Tables 1 and 2, and Supplemental Figures 1–5).

Figure 3. Structure of the AtTPS18–FSPP Complex.

(A) The overall structure of the AtTPS18–FSPP complex as a ribbon cartoon (NTD and CTD are shown in orange and cyan, respectively). The substrate analog, FSPP, is shown as a yellow stick model, and the two Mg2+ atoms are shown as magenta balls.

(B) Expansion of the substrate-binding pocket and the docking results showing the substrate binding of GFPP (electron static surface). FSPP and GFPP are shown as yellow and cyan stick models, respectively.

(C) Interaction of amino acid residues with FSPP in the binding site. The bound FSPP is depicted as a ball-and-stick model. The 24 residues in the active site of AtTPS18 are shown. Among them, Arg319, Arg321, Asp356, Asp357, Asp497, Asn500, Asp510, Thr504, Glu508, Arg511, and Glu513 were not considered for mutagenesis because they are conserved in the ionization of prenyl diphosphate substrates.

(D) Superimposition of the cartoon models of AtTPS18 (magenta), NTEAS (yellow, PDB: 5EAT), taxadiene synthase from *Taxus brevifolia* (tan, PDB: 3P5R), and limonene synthase from *Citrus sinensis* (blue, PDB: 5UV2) from a top-down perspective. Farnesyl hydroxyphosphonate (FHP) in NTEAS and GFPP in AtTPS18 are shown as yellow and cyan sticks, respectively. The positions of the I–J loop and helix G (G1 and G2) are indicated by red and blue circles, respectively.

(E) Close-up view of the helix G region of AtTPS18, 5EAT, 3P5R, and 5UV2 (for details, see the legend of D). The shift distance (Å) of helix G of AtTPS18, relative to that of other plant TPSs, was measured and labeled.

(F) Possible interactions between Asn493 (in helix H) and helix G in the AtTPS18–FSPP complex. Dashed lines indicate possible hydrogen bonds.
We did not detect expression was low in seedling roots and flowers were highly expressed in roots and siliques, whereas Ceroplastes floridensis which was initially isolated from the secretions of the insect of flocerene, a sesterterpene with an 11-6-5 fused ring system, (Naya et al., 1981). The 11-6-5 fused ring system of (Figures 1D and Supplemental Figure 2). Although 5-8-6-5 fused ring sesterterpenoids are shown in Figure 2. (Figures 1D and Supplemental Figure 2) and a–d have not yet been determined.

The chemical structures of C3–C7 are shown in Figure 2.

The sum of the areas of the chromatogram peaks (GC–MS with SIM mode, b The chemical structures of a–d have not yet been determined.

Consequently, we can hypothesize that it served as a progenitor of type-A sester-TPSs during the evolution of plants. AtTPS06 was initially characterized as a di-terpene synthase that converted GGPP to dolabelladienol and other unidentified diterpenes (Supplemental Figures 1D and Supplemental Figure 2; Wang et al., 2016b). The catalytic promiscuity of AtTPS06 (using GGPP and GFPP as substrates) suggests that it served as a progenitor of type-A sester-TPSs during the evolution of TPSs, at least in Brassicaceae (Weng, 2014).

qRT–PCR analyses revealed that both AtTPS22 and AtTPS29 were highly expressed in roots and siliques, whereas AtTPS06 expression was low in seedling roots and flowers (Supplemental Figure 8). We did not detect C1 and C2, the main C25 terpene products of AtTPS06 and AtTPS29, respectively, in any tested Arabidopsis tissues (100 mg [fresh weight] of roots and flowers) using GC–MS with the full-scan mode. However, we cannot simply conclude that C1 and C2 are not produced in Arabidopsis, given that the amount of C1 and C2 might be below the detection limit of GC–MS. Another possibility is that C1 and C2 might be further metabolized in planta.

### Plant Sester-TPSs Have a Larger Catalytic Pocket than Classic TPSs

To investigate the mechanism underlying the divergence of these Arabidopsis sester-TPSs (as they generated profiles different from that of GFPP), we recombinantly expressed truncated AtTPS18 (residues 52–605; the first 51 amino acids were predicted to be the plastidial signal peptides) in E. coli and purified the protein. The AtTPS18 complex with FSPP (farnesyl-diphosphate, an inert analog of FPP; Figure 3 and Supplemental Table 3) was crystallized, and its structure was determined at a resolution of 2.3 Å. The final model contained residues 71–605, and it was comprised of two compact α-helical domains, namely, the N-terminal domain (NTD) and the C-terminal catalytic domain (CTD) (Figure 3A). FSPP adopts an extension of 2.3 Å.

### Table 1. Product Distribution of AtTPS18 and Its Mutants.

| Mutations     | Product distribution (%) | Relative yield (-fold)c |
|---------------|---------------------------|-------------------------|
|               | a  | b  | C  | C3 | C4 | C5 | C6 | d  | C7 |                   |
| AtTPS18 (WT) | 0.15 | 0.01 | 2.8 | 0.6 | 1.3 | 3.2 | 0.8 | 6.1 | 85.1 | 1.0               |
| N493F        | –  | –  | 2.6 | 0.65 | 2.4 | 4.2 | 1.2 | 7.8 | 81.2 | 0.4               |
| Y465A        | 0.5 | 0.15 | 2.4 | 3.8 | 4.7 | 4.8 | 1.8 | 16.1 | 65.7 | 1.01              |
| G328P        | 0.34 | –  | 1.5 | 3.1 | 1.7 | 88.6 | 0.05 | 0.69 | 3.95 | 1.62              |
| G328A        | 0.72 | –  | 2.1 | 3.34 | 4.94 | 21.8 | 0.04 | 7.1 | 59.9 | 1.35              |
| G328S        | 0.04 | –  | 4.11 | 1.77 | 1.47 | 2.2 | 0.92 | 9.4 | 80.05 | 1.15             |
| G328C        | 0.3  | –  | 2.4 | 1.7 | 1.4 | 15.1 | 0.06 | 7.2 | 71.9 | 0.67              |
| G328L        | –  | –  | –  | 10.1 | 6.5 | 77.7 | –  | –  | 5.75 | 0.16              |
| G328N        | 2.3 | –  | –  | 16.9 | 0.19 | –  | 79.2 | –  | 1.35 | 0.08              |
| F496S        | 8.2 | 3.7 | –  | 9.5 | 40.1 | –  | 17.3 | 0.45 | 20.8 | 0.11              |
| F496M        | 1.6 | 0.25 | 1.7 | 8.1 | 10 | 5.2 | 4.5 | 5.2 | 63.6 | 0.49              |
| F496H        | 1.5 | –  | 0.2 | 13.2 | 45.9 | –  | 12.6 | 4.8 | 21.7 | 0.22              |
| F496A        | –  | 10.2 | –  | 9.3 | 28.3 | –  | 26.7 | 1.2 | 24.3 | 0.17              |
| F496C        | 7.9 | 6.8 | 0.2 | 9.7 | 37.0 | 0.55 | 10.7 | 2.7 | 24.3 | 0.14              |
| G328A/F496A  | 1.4 | 5.7 | 0.3 | 10.2 | 23.0 | 14.4 | 16.3 | 3.1 | 25.4 | 0.2               |

All product distributions were calculated from at least three independent experiments, whereas the results of F496S and F496A mutants were obtained from two replicates.

The structures of sesterterpenes a–d have not yet been determined.

The chemical structures of C3–C7 are shown in Figure 2.

The sum of the areas of the chromatogram peaks (GC–MS with SIM mode, m/z 340) of the main sesterterpenes produced by AtTPS18 and its mutants were calculated as the yield after normalization with an internal standard and culture density (OD600 value). The yield of AtTPS18 (wild type) was set to 1.0.

Notably, AtTPS06 is the first type-B sester-TPS identified in plants. Moreover, AtTPS06 is more similar to sesqui-TPSs than to sester-TPSs in the phylogenetic analysis (Figure 1A), which suggests that TPS06 and its close homologs are possible intermediate enzymes between mono-/-sesqui-/-di-TPSs and sester-TPSs in the evolution of plants. AtTPS06 was initially characterized as a di-terpene synthase that converted GGPP to dolabelladienol and other unidentified diterpenes (Supplemental Figures 6 and 7; Wang et al., 2016b). The catalytic promiscuity of AtTPS06 (using GGPP and GFPP as substrates) suggests that it served as a progenitor of type-A sester-TPSs during the evolution of TPSs, at least in Brassicaceae (Weng, 2014).

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synthesized GFPP and the soaking of apo-AtTPS18 crystals in this ligand were attempted, but none of the solved structures contained the GFPP substrate. We therefore used AutoDock to model the GFPP substrate in the AtTPS18 structure and found that GFPP fit well in the substrate-binding pocket and generally overlapped with the FSPP-binding site (Figure 3B). These results indicate that the substrate-binding pocket of AtTPS18 is very suitable for the binding of GFPP.

A comparison of the AtTPS18 structure with that of NtEAS (5-epi-aristolochene synthase, PDB: 5EAT [Starks et al., 1997]) showed a root-mean-square deviation value of 1.67 Å. AtTPS18 adopted a three-dimensional structure similar to that typically found in plant sesquiterpene synthases (Figure 3D). The major differences were found at the C-terminus. A superimposition of the structures of AtTPS18 and NtEAS revealed that all helices (helices A to K), except for helix G, H2, and the I–J loop (connecting helices I and J), fit well (Figure 3D). A comparison of the structures of other plant terpene synthases with the structure of AtTPS18 revealed that the I–J loop served as a lid, which was slightly separated from the top of the substrate-binding pocket (Figure 3D). In addition, the most obvious conformational change was the outward shift of helix G in AtTPS18, which created a larger catalytic pocket compared with those of other plant TPSs, thus allowing it to accommodate GFPP (Figure 3D and 3E). Compared with mono-TPS (limonene synthase, PDB: 5UV2 [Kumar et al., 2017]), sesqui-TPS (PDB: 5EAT [Starks et al., 1997]), di-TPS (taxadiene synthase, PDB: 3P5R [Koksal et al., 2011]), and AtTPS18 (obtained in this study), helix G in AtTPS18 shifted outward by 2.8 Å, 3.1 Å, and 3.0 Å, respectively (Figure 3E). We further measured the pocket volume of the three representative plant TPSs and AtTPS18 using CAVER Analyst 2.0 (Jurcik et al., 2018). The results showed that the volume of the pockets in terpene synthases (5UV2: 593.0 Å³; 5EAT: 801.9 Å³; 3P5R: 1285.5 Å³; AtTPS18: 1513.8 Å³) increased gradually with the chain length of the substrate (every isoprene unit occupied approximately 300 Å³, Supplemental Figure 9A–9D). Notably, AtTPS06 (type B; 1536.3 Å³) and AtTPS18 (type A) had similar pocket volumes, although they used different cyclization mechanisms (Supplemental Figure 9E).
Molecular Basis for Sesterterpene Diversity

Previous studies have clearly demonstrated that the amino acid residues in helix G affect the product specificity and capacity of TPS enzymes (Starks et al., 1997; Li et al., 2013). To further investigate the mechanism of the outward shift in helix G (the sequence alignment is shown in Supplemental Figure 10A and 10B), we carefully checked the possible interactions between the amino acids in the kink of helix G in the AtTPS18–FSPPP complex. As shown in Figure 3F, we found that the side chain of Asn 493 in helix H formed two hydrogen bonds with the main chain of residues Gly 468 and Met 469 in helix G, which probably resulted in the outward shift of helix G. Notably, position 493 was occupied by Asn in approximately 50% of active sester-TPSs (n = 18), whereas this position was always occupied by Ala in most mono-/sesqui-TPSs (n = 59; Supplemental Figure 10C). Because synthetic GFPP was in short supply, we used the E. coli system harboring AtGFPPS2 to test the product profiles of the mutated enzymes in this study (as shown in Figure 1B and 1C; for detailed calculations, see Methods). Western blotting (anti-myelin basic protein [MBP] tag) was used to ensure no significant difference in the protein solubility among the mutated sester-TPSs (Supplemental Figure 11). Site-directed mutagenesis and biochemical assays showed that mutation of the polar residue Asn 493 to the nonpolar residue Phe decreased sester-TPS activity by approximately 60% (Table 1). This might be due to the disruption of the hydrogen bonds contributed by Asn 493, thus resulting in an inward shift of the elastic helix G and shrinkage of the pocket volume. We also replaced the kink sequence in helix G of AtTPS18 (456TAGMDGY462), the kink sequence is underlined; Supplemental Figure 10A) with the kink sequence of 5EAS enzyme (60TTTYYLY467). This replacement might also reduce the volume of the catalytic pocket, thus inactivating the enzyme (Supplemental Data 1). These data confirm the importance of the enlarged catalytic pocket created by the outward shift of helix G in sester-TPS activity.

Type-A Sester-TPSs, Not Type-B Sester-TPSs, Have More Aromatic Residues in the Catalytic Pocket than Classic TPSs

In general, only two or three aromatic residues, which possibly stabilize reactive carbocation intermediates, are found in the classic TPS catalytic pocket (Christianson, 2017). However, the AtTPS18 structure clearly revealed that there were five aromatic residues (Tyr 462, Tyr 465, Phe 496, Phe 573, and Tyr 577) in the catalytic pocket (Figure 3C). Moreover, three positions (Tyr 462, Tyr 465, and Phe 496 in AtTPS18) were conserved and unique to type-A sester-TPSs (Figure 4A and Supplemental Figure 12), whereas these three positions were occupied by nonaromatic residues in most classic TPSs. We had previously demonstrated that position 328 (in AtTPS18) was occupied by Gly or Pro in plant sester-TPSs, whereas this position was occupied by an aromatic residue (always Trp) in most classic TPSs (Chen et al., 2019). Therefore, these residue features (Gly 328, Tyr 462, Tyr 465, and Phe 496) could be used to distinguish type-A sester-TPSs from classic plant TPSs (Supplemental Figure 12).

To test the effects of these aromatic residues on sester-TPS activity, we replaced the aromatic residues with Ala one by one. The results showed that most mutants (Y462A, F573A, and Y577A) completely lost their sester-TPS activity (Supplemental Data 1), implying their critical roles in stabilizing carbocation intermediates in catalysis. However, no change in total activity and product specificity was noted between the Y465A mutant and AtTPS18 (Table 1), although Tyr 465 was conserved across all characterized type-A sester-TPSs. We cannot presently arrive at a role for Tyr 465 in the catalysis of AtTPS18. The F496A mutant showed approximately 17% of the activity of AtTPS18 (Table 1). The effects of Phe 496 on sester-TPS product specificity were further investigated, as described in another section.

The discovery of type-B sester-TPS (AtTPS06) has made it possible to compare residues in the catalytic pocket of both types of sester-TPSs. Both residue features of type-A sester-TPSs could not be applied to type-B sester-TPSs, which are similar to classic TPSs, in that only two aromatic residues, Trp 335 and Tyr 584, were found in AtTPS06, and the two positions were obviously different (occupied by aromatic or nonaromatic) in AtTPS18 (Gly 328 and Phe 496) and AtTPS06 (Trp 335 and Ser 503; Figure 4A).

It has been proposed that the initial conformation of GFPP, which is fixed in the catalytic pocket of different sester-TPSs, determines the sesterterpene cyclization mechanism (type A or type B; shown in Figure 2 [Sato et al., 2018]). To test the effects of Trp 335 and Ser 503 on GFPP accommodation in type-B sester-TPSs, we replaced these two amino acids in AtTPS06 with Gly and Phe, both individually and simultaneously. All mutants (AtTPS06W335G, AtTPS06S503F, AtTPS06W335G/S503F, and AtTPS06W335G/S503F) completely lost their sester-TPS activity (Figure 4B), which suggested that both residues are required for correct GFPP accommodation in the catalytic pocket of AtTPS06, if not all plant type-B sester-TPSs. In addition, we exchanged the helix G region between AtTPS18 (456TAGMDGY462, including Tyr 462) and AtTPS06 (465STAGEVL469, including Leu 469), and this replacement also resulted in enzyme inactivity (Figure 4B). Similarly, all corresponding AtTPS18 mutants (with the replacement of the corresponding amino acid in AtTPS06), except AtTPS18F496 (no type-B sesterterpenes were found), were enzymatically inactive (Figure 4C and Table 1). These results indicate that, in addition to these two amino acids, there must be other amino acids controlling GFPP accommodation in type-B sester-TPSs. Cgr1961, isolated from Capsella grandiflora, was reported to be a nonclustered sester-TPS (Chen et al., 2019). Based on its sequence features (only two aromatic residues, Trp 302 and Ser 471, are found in Cgr1961; Figure 4A) and phylogenetic analyses, we propose that Cgr1961 is also a type-B sester-TPS; however, our findings will need to be validated in future structural elucidation studies of Cgr1961 products.

Controlling Product Specificity in AtTPS18 by Gly 328 and Phe 496

Based on the importance of Gly 328 and Phe 496 in type-A sester-TPS activity, as discussed above, we further investigated the effects of other residues at these two positions on the enzymatic properties of AtTPS18. We generated 12 mutants for the Gly 328 position of AtTPS18 from Gly 328 to 16 amino acids with hydrophobic side chains (Ala, Phe, Leu, and Trp), amino acids with positively charged side chains (Lys, His, and Arg), amino acids with negatively charged side chains (Asp), amino acids with polar
uncharged side chains (Ser and Asn), or other amino acids (Cys and Pro). The findings showed that exchanging Gly328 with a charged residue (Asp, Arg, His, or Lys) resulted in an enzyme that was inactive for the production of sesterterpenes, which suggested that a charged residue probably destabilizes the formation of the carbocation intermediate in the catalytic pocket (IM1 and IM2 in Figure 2). Moreover, G328F and G328W substitutions (both Phe and Trp have bulky side chains) also resulted in enzyme inactivity (Supplemental Data 1). By contrast, the G328P substitution increased the total production of sesterterpenes by at least 60% (n = 3; Table 1 and Supplemental Data 1).

Using the same strategy, we generated 11 mutants for the Phe496 position of AtTPS18 (replacing Phe496 with Ala, Ile, Leu, Met, Val, Lys, His, Asp, Ser, Cys, or Pro). Compared to AtTPS18, all mutants showed decreased sester-TPS activity. Interestingly, the F496A mutant retained 17.3% (n = 2) of the sester-TPS activity of AtTPS18; however, replacing the Phe496 position with Ala in other Arabidopsis sester-TPSs (AtTPS19F493A and AtTPS30F496A) resulted in the loss of sester-TPS activity (Supplemental Figure 13). Additionally, all Phe496 mutants showed decreased product specificity compared to AtTPS18, as no main product accounted for >50% of the total sesterterpene production (Table 1). These findings provide a starting point for creating new sester-TPSs using directed evolution strategies. Because multiple novel sesterterpene peaks were detected in the chromatograms of the reaction mixtures obtained with AtTPS18 mutants (Supplemental Figure 14), we scaled up the production of AtTPS18G328P and AtTPS18G328A/F496A using the E. coli system (Shao et al., 2017). Four compounds (C3 to C6) were purified and structurally elucidated by NMR spectroscopy (Supplemental Table 2; Supplemental Figures 4 and 5). Among them, C3 ((+)-brassitetraene B) and C6 ((+)-brassitetraene A) have been previously reported by Huang et al. (2018). C4 is an isomer of (2E)-α-cericereone with a different double-bond position (Qin et al., 2018).
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et al., 2016), whereas C5 (18S) is an epimer of C7 (18R, (-)-thalianatriene) (Shao et al., 2017). Notably, the ratio of C5/C7 isomers increased with the increase in the side-chain size of the amino acid at position 328 (Gly and Ser < Ala and Cys < Leu and Pro; Table 1). Moreover, AtTPS18G328N produced C6 as the main product, although with a lower yield than AtTPS18. It is possible that the side chain of the amino acid at position 328 is close to C19/C19 of the carbocation intermediate, allowing it to determine the conformation of the final product. For example, Asn328 might stabilize the formation of IM1 and facilitate the deprotonation of IM1 to produce C6 (Figure 2).

Thus far, there have been no reports on the catalytic parameters of sester-TPSs, probably because GFPP is not commercially available. We therefore compared the catalytic efficiency of plant sester-TPSs using GFPP with their ancestor enzyme, a mono-/sesqui-TPS from the TPS-a subfamily. Here, we only determined the catalytic efficiency of purified AtTPS18 and AtTPS18G328N due to insufficient synthetic GFPP. AtTPS18G328N showed much higher substrate affinity ($K_M = 1.28 \pm 0.49 \mu M$, $k_{cat} = 0.047 \pm 0.05 s^{-1}$, $n = 3$) than AtTPS18 ($K_M = 4.79 \pm 1.82 \mu M$, $k_{cat} = 0.11 \pm 0.017 s^{-1}$, $n = 3$). The catalytic efficiency of AtTPS18 fell within the range previously reported for mono-TPSs or sesqui-TPSs from plants (Chen et al., 2004; Tholl et al., 2005; Nagegowda et al., 2008; Wang et al., 2008).

Interconversion between AtTPS18 and Its Close Homologs Using Structure-Based Comparisons

Previous studies have reported that AtTPS18, AtTPS19, and Cru237 (a characterized sester-TPS from Capsella rubella) are similar to each other, sharing >80% identity at the protein level (Supplemental Figure 15; Shao et al., 2017; Huang et al., 2018; Chen et al., 2019). Moreover, the three sester-TPSs share common carbocation intermediates in the production of their own products (Figure 2) (Huang et al., 2017). Next, we attempted to interconvert their sester-TPS activities using the AtTPS18 structural information obtained in this study. Structural comparisons showed that there were three key amino acid pairs (Met352 in AtTPS18 versus Ile349 in AtTPS19; Asp431 in AtTPS18 versus Tyr428 in AtTPS19; Gly461 in AtTPS18 versus Asp458 in AtTPS19; Figure 5A). The results of site-directed mutagenesis and biochemical assays showed that a single mutant (M352I, D431Y, or G461D) did not notably change the product profile relative to wild-type AtTPS18. One of the double mutants (M352I/G461D) produced 34.5% C6 and 36.6% C7 (the yield was 0.12 when the yield of AtTPS19 was 0.36, $n = 2$; Supplemental Data 3). Moreover, one double mutant (I349M/D458G) produced 19.3% C8 and 14.9% C7 (the yield was 0.12 when the yield of AtTPS19 was set to 1.0, $n = 2$), and a triple mutant (I349M/Y428D/D458G) produced 1.7% C8 and 19.3% C7 (the yield was 0.36, $n = 2$).

In addition, we found that one amino acid (Val353 [hydrophobic side chain] in AtTPS18 versus Asn353 [polar side chain] in Cru237) is probably responsible for the product specificity between AtTPS18 and Cru237 (Figure 5C). The results of biochemical assays showed that the Cru237 mutant (N353V) showed complete AtTPS18 activity, whereas the corresponding AtTPS18 mutant (V353N) displayed inactivity (Figure 5D).

Equally important, there are still several open questions with regard to the catalytic mechanism of plant sester-TPSs, including the mutants generated in this study. For example, how are the carbocation intermediates controlled in the catalytic pocket of sester-TPSs to generate different products? Which residues are involved in the final deprotonation reaction? Computational-guided methods (e.g., quantum mechanical/molecular mechanical–molecular dynamic simulations), together with the structural features of AtTPS18, will help address these interesting questions (Zhang et al., 2020).

Concluding Remarks

Sester-TPSs, an evolutionarily new clade of the TPS-a subfamily, are not Brassicaceae-specific; instead, they are widely distributed in the plant kingdom (Liu et al., 2016). To date, approximately 140 sesterterpenoids have been identified; however, the chemical diversity of the plant sesterterpenes is expected to be much larger than that initially assumed due to the possible rearrangements of the C25 carbocation intermediate by sester-TPSs. More importantly, further studies are needed to investigate the physiological functions of sesterterpenoids from both plants and microbes (Wei et al., 2004; Luo et al., 2010). In this study, we revealed the general catalytic mechanism of plant sester-TPSs by combining chemical structure elucidation, enzyme structure analysis, and biochemical assays. The novel sesterterpenoids generated by the newly mutated sester-TPSs will significantly expand the diversity of terpenoids, which is important for further applications. As sester-TPSs have also been identified in microbes (Matsuda et al., 2015; Ye et al., 2015; Tian et al., 2017), whether microbe sester-TPSs (PT-TPS fusion proteins) utilize a catalytic mechanism similar to that of plant sester-TPSs is an interesting topic that should be addressed in future studies.

METHODS

Plant Materials, RNA Analysis, and Chemicals

A. thaliana plants (Columbia-0 ecotype) were grown in soil in a growth room or on half-strength solid Murashige–Skoog medium plates in a growth chamber at 22°C under a 16-h light/8-h dark photoperiod. RNA extraction, reverse transcription reactions, and gene cloning were performed as described previously (Chen et al., 2015).

All chemicals used in this study were purchased from Sigma-Aldrich (USA) except FSPP, which was purchased from Echelon Biosciences (USA). GFPP (1 mg) was chemically synthesized following a published protocol (Sato et al., 2013). All DNA constructs generated in this study were verified by sequencing.

Gene Cloning, Expression in E. coli, and Biochemical Assays

All truncated Arabidopsis TPS-a or mutated genes (signal peptide sequence was removed; for detailed primer information, see Supplemental Data 2) were cloned into the pMAL-c2x vector (New England Biolabs). An MBP tag was added at the C-terminus of each truncated TPS gene. AtGFPPS2 (At3g14550) was ligated into the pET-28a vector. To increase sesterterpene production, we cotransformed three plasmids, pMBIS-TC', containing five MVA genes (mevalonate kinase, phosphomevalonate kinase, mevalonate pyrophosphate decarboxylase, IPP isomerase, and FPP synthase [Martin et al., 2003], pET-Karl' -GFPPS2, and pMAL-AMP' -TPSa into the E. coli JM109 (DE3) strain. The transformed E. coli harboring pMBIS-TC', pET-Karl' -AtGFPPS2, and pMAL-AMP' (empty vector) were used as negative controls. The E. coli system (200 ml in Tris–borate [TB] buffer with the appropriate
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antibiotics) was induced by the addition of 0.25 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) and 10 mM mevalonolactone to an OD_{600} (optical density at a wavelength of 600 nm) of 0.6. After 48 h of induction at 16°C, the cells were harvested by centrifugation at 5000 g for 20 min. The harvested cells were resuspended in 12 ml of Tris–HCl buffer (pH 8.0) in a 50-ml conical tube and lysed with a probe sonicator. The resulting solution was then extracted with 20 ml of EtOAc by vortexing for 1 h. The EtOAc phase was collected and concentrated to 200 μl under flowing nitrogen. Sesterterpene analyses by triple-quadrupole GC–MS (GC-QQQ-MS; Agilent 7890B/7000C, SIM mode m/z 340) were performed as described previously (Chen et al., 2019). The sum of the areas of the chromatographic peaks of the main sesterterpene compounds produced by AtTPS18 and its mutants was calculated as the total yield after normalization to the internal standard and culture density (OD_{600} value).

The soluble recombinant sester-TPSs (AtTPS06, AtTPS18, AtTPS19, and Cru237) and their mutants in E. coli were quantitatively validated using western blotting with an anti-MBP monoclonal antibody (New England BioLabs). In brief, soluble crude proteins (20 ng) from each E. coli strain were separated by 8% SDS–PAGE, transferred onto a polyvinylidene fluoride membrane, and blotted with an anti-MBP antibody (1:50 000 dilution) following a published protocol (Chen et al., 2015).

To determine the catalytic efficiency of AtTPS18 and AtTPS18\textsuperscript{G328P}, we used 2 μg of purified recombinant protein in each biochemical assay (200-μl volume, buffer contained 10 mM Mg\textsuperscript{2+}, 2 mM dithiothreitol, and 50 mM Tris–HCl [pH 7.5]) and a range of GFPP concentrations (0, 1, 2, 5, 10, 20, 50, and 100 ng/μl). After 1 h of incubation at 30°C, the products were extracted twice with 200 μl of EtOAc. The extracted sesterterpenes were combined and analyzed by GC-QQQ-MS (MRM, 340.0 > 119.0) using the aforementioned program. The target sesterterpenes were quantified by comparing with standard curves that were generated with C5 (AtTPS18\textsuperscript{G328P} main product) and C7 (AtTPS18 main product). The K_{M} and k_{cat} of each protein were determined by extrapolating the values from Lineweaver–Burk plots. The enzymatic assays were carried out in triplicate.

**Purification of Sesterterpenes and Elucidation of Chemical Structures**

The E. coli JM109 (DE3) strain harboring pMAL-c2x-AtTPS06, AtTPS18 (and its mutants, AtTPS18\textsuperscript{G328P} for C5 preparation and TPS18\textsuperscript{G328A/F496A} for C3/C4/C6 preparation), or AtTPS29, pET-28a-AGFPFPS2, and pMBIS-TC\textsuperscript{p} plasmids (as mentioned above) was incubated in 1 l of TB medium containing 100 μg/ml of ampicillin (sodium salt), 20 μg/ml of tetracycline, and 50 μg/ml of kanamycin sulfate at 37°C and agitated at 130 rpm in a 2.1-L Erlenmeyer flask. When the cultures reached an OD_{600} of 0.4–0.5, 250 μl of 1 M IPTG and 8 ml of 50 mg/ml mevalonolactone were added in two allotments, followed by incubation for 48 h at 16°C/130 rpm. The cultured cells were harvested by centrifugation and lyophilized. The lyophilized cells from 8 l of culture media were extracted with acetone. The extract was concentrated, re-extracted with n-hexane, and repeatedly subjected to silica-gel column chromatography. The sesterterpenes were eluted with n-hexane to yield different amounts of pure C1 to C6 (all were obtained as colorless oils; for the NMR data of C1, C2, C4, and C5, see Supplemental Tables 1 and 2 and Supplemental Figures 2–5).

**Large-Scale AtTPS18 Production, Purification, and Crystallization**

AtTPS18 was inserted into the pET-Duet vector with a 6×His tag at the N-terminus. The plasmid was transformed into the E. coli BL21 (DE3) strain. The transformed bacterial cells were grown in Luria–Bertani medium containing ampicillin at 37°C and induced by 0.25 mM IPTG for 12 h at 18°C. The cells were harvested and resuspended in buffer A (20 mM Tris–HCl [pH 8.0] and 100 mM NaCl) supplemented with 1 mM phenylmethylsulfo-
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purification and structural elucidation of C3. Y.Z. and Hongwei Liu carried out the purification and structural elucidation of C4, C5, and C6. T. Shinada and T. Sato synthesized the GFPP. Q.C., J.L., Z.L., P.Z., and G.W. analyzed the data. G.W. wrote the manuscript with input from all authors.

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