Abstract: A reinvestigation of the linalool synthase from Chryseobacterium polytrichastri uncovered its diterpene synthase activity, yielding polytrichastrene A and polytrichastrol A with new skeletons, besides known wanju-2,5-diene and thunbergol. The enzyme mechanism was investigated by isotopic labeling experiments and DFT calculations to explain an unusual ethyl group formation. Rationally designed exchanges of active site residues showed major functional switches, resulting for I66F in the production of five more new compounds, including polytrichastrene B and polytrichastrol B, while A87T, A192V and the double exchange A87T, A192V gave a product shift towards wanju-2,5-diene.

Terpenoid biosynthesis starts with the cyclization of an acyclic oligoprenyl diphosphate into a structurally complex terpene hydrocarbon, which can be followed by several functionalizations to introduce bioactivity. The cyclization step is catalyzed by a single enzyme, a terpene synthase (TPS), and proceeds by substrate ionization through abstraction of diphosphate (type I TPSs) or protonation (type II) to initiate a cationic cascade reaction. Among bacterial type I TPSs, several sesquiterpene synthases (STPSs) and diterpene synthases (DTPSs) have been reported, while monoterpenes synthases (MTPSs) such as 1,8-cineol synthase are rare. The sestermobaraene synthase recently discovered from S. moharaensis is the only known bacterial type I sesterterpene synthase (StTPS).

While the overall structures of type I TPSs are highly conserved and exhibit a similar α-helical fold as avian farnesyl diphosphate synthase, their amino acid sequences can be very different which challenges a sequence-based function prediction. It has been suggested that initial cyclization modes can be predicted from phylogenetic analyses and an algorithm for sequence based function predictions has been implemented into antiSMASH. Computational approaches to predict the product from an enzyme homology model by fitting cationic intermediates into the active site, or to distinguish between the functions of enzymes as MTPS, STPS or DTPS have been developed. A phylogenetic tree constructed from the amino acid sequences of 3278 bacterial TPS homologs reveals a scattered distribution of MTPSs, STPSs, DTPSs and StTPSs and of cyclization modes, demonstrating that a function prediction from amino acid sequences of bacterial TPSs is difficult. Therefore, we turned our attention to a rational approach based on the crystal structure of selina-4(15),7(11)-diene synthase from S. pristinaespiralis (SdS) in complex with the substrate surrogate 2,3-dihydrofarnesyl diphosphate (DHFP). For TPSs using large substrates (geranylgeranyl diphosphate, GPP for StTPSs or geranylgeranyl diphosphate, GGPP for DTPSs) the active site cavity should offer a larger space than for TPSs using small substrates (farnesyl diphosphate, FPP for STPSs or geranyl diphosphate, GPP for MTPSs).

Several non-polar active site residues enclose DHFP in the SdS structure that define the substrate conformation and the available space within the active site (Figure 1). Their van der Waals volumes can be calculated according to Abraham and co-workers (Table S1). A summary of the 13 residues located in the analogous positions of other charac-

Figure 1. Active site of SdS showing several non-polar residues enclosing the substrate surrogate DHFP (brown). Black labels are for foreground residues, grey labels for background residues. Green spheres represent Mg²⁺ cations.
polytrichastri diphosphate (DMAPP) and (phosphate buffer.\textsuperscript{[20]} Analysis of its active site residues gave activities towards different substrates in a mixed Tris/phosphate buffer.\textsuperscript{[20]} The enzyme originally described as CpLS was reassigned as \textit{Chryseobacterium polytrichastri} (CpLS), based on the observed linalool synthase (CpLS), with an average decrease of 58 Å$^3$ between two consecutive TPS groups. This value is similar to the increase of the substrate size of 84 Å$^3$ for each isoprene unit (Table S1), suggesting that the active site space becomes larger according to the requirements of the substrate. We have recently described a TPS from \textit{Chryseobacterium polytrichastri} as linalool synthase (CpLS), based on the observed activities towards different substrates in a mixed Tris/phosphate buffer.\textsuperscript{[20]}

Analysis of its active site residues gave a $\Sigma V_{\text{obs}} = 550$ Å$^3$, which is the smallest value among all TPSs (red entry in Table S2) and did not seem to fit for a MTPS, but would rather suggest a function as DTPS or StTPS.

![Image](50x466 to 293x546)

Figure 2. Added $V_{\text{obs}} (\Sigma V_{\text{obs}}$) of TPS active site residues. Bold bars indicate spanned regions, thin bars show the mean ± SD for each group of TPSs.

Additional experiments with DMAPP, (E)- and (Z)-(4-\textsuperscript{13}C,4-\textsuperscript{2}H)IPP.\textsuperscript{[25]} GGPPs and CpPS (Figures S29 and S30) confirmed the absolute configurations of 1 and 2.

The cyclization mechanism by CpPS requires GGPP isomerization to GLPP to explain the 2Z double bond in 1–3. Following Arigoni’s mechanistic model for cadalanes,\textsuperscript{[26]} this isomerization proceeds through (S)-GLPP by syn-allylic transposition of diphosphate with a specific stereochemical fate for the 1-pro-R and the 1-pro-S hydrogen of GGPP, ending up in the 1E and 1Z positions of GLPP, respectively. After vinyl group rotation around C2-C3 the 1.1-cyclization continues with attack of C14 at C1 from the Si face (top, anti- S$_2$-$^2$ reaction) to install the correct configuration at C14 in A, which also turns H$_1$ to the bottom. This hydrogen is now closer to the cation at C15 than H$_1$ and shifts into the iPr group in B, as confirmed experimentally with (R)- and (S)-(1-\textsuperscript{2}H)GGPP (Figure S31). For the enantiomers of 1–4, the migration of H$_1$ into the iPr group would be expected and, thus the migration of H$_1$ versus H$_4$ can give evidence for the absolute configurations of terpenes, as demonstrated previously for other sesqui- and diterpenes.\textsuperscript{[22,25–27]} The optical rotation of 4 from CpPS ([\alpha]$_D$\textsuperscript{25} = $-35.0$, $c = 0.1$, CHCl$_3$) discloses that this is the same enantiomer as in \textit{Pseudotsuga menziesii} ([\alpha]$_D$ = $+74.4$, $c = 6.3$, CHCl$_3$).\textsuperscript{[30]}

Cation B is a first branching point and can be attacked by water to yield 4, while the steps from B to F1 towards 3 (shown in blue) were supported by previous labeling experiments.\textsuperscript{[22]} Along these steps two 1,2-hydride migrations from C1 to D1 and from E1 to F1 occur that were both supported for I with (11-\textsuperscript{13}C,10-\textsuperscript{2}H)GGPP and (7-\textsuperscript{13}C,6-\textsuperscript{2}H)GGPP enzymatically prepared from (3-\textsuperscript{13}C,2-\textsuperscript{2}H)IPP\textsuperscript{[28]} and (3-\textsuperscript{13}C,2-\textsuperscript{2}H)IPP\textsuperscript{[29]} with IPP and GGPPs. In these experiments a deuterium atom migrates to the labeled carbon, resulting in a slightly upfield shifted triplet in the \textsuperscript{13}C-NMR spectrum (Figures S32 and S33). The steps from F1 towards 1 may include a dyotropic rearrangement\textsuperscript{[18]} to H (orange arrow) to explain the E1 group formation. Stereoselective deuterations introduced at C12 using DMAPP plus (E)- or (Z)-(4-\textsuperscript{13}C,4-\textsuperscript{2}H)IPP and GGPPs revealed the specific shift of the 12-\textsuperscript{13}C in the 12-pro-R hydrogen, while the usage of (3-\textsuperscript{13}C,4,4-\textsuperscript{2}H$_2$)IPP\textsuperscript{[31]} demonstrated its migration to C11 (Figure S34). A subsequent Wagner–Meerwein rearrangement (WMR) leads to J. Deprotonation to I with cyclopropanation proceeds with loss of the 5-pro-R hydrogen, as demonstrated by stereoselective deuterium labeling introduced with (R)- and (S)-(1-\textsuperscript{13}C,1-\textsuperscript{2}H)IPP (Figure S35). Another WMR from J results in K, the precursor of 2 that is formed by quenching with water. The overall mechanism towards 1–4 is also in line with \textsuperscript{13}C-labeling experiments, in which each of the twenty carbons of GGPP were labeled individually (Figures S36–S55).

For a deeper understanding of the role of active site residues in TPS catalysis (Figure S56), several CpPS variants targeting positions homologous to SdS active site residues were constructed (Tables S7 and S8). The purified enzyme variants were adjusted to same concentration (Figure S57), followed by conversion of GGPP and GC/MS analysis of the products (Figures S58 and S59, Table S9). The position equivalent to I66 in CpPS ($=\text{F55}$ in SdS, Figure 1) is in many TPSs and especially DTPSs occupied by an aromatic...
residue (Table S2). Indeed, the I66F variant gave an enhanced diterpene production (195 ± 43 % of wildtype level, Figure 3) and a strongly changed product profile, with only low amounts of 1, but no 2 and slightly increased 3 and 4. In addition, several new compounds were obtained. A preparative scale incubation allowed the isolation and structure elucidation by NMR spectroscopy (Figures S61–S100, Tables S10–S14) of wanju-2,6-diene (5), wanju-2-en-6α-ol (6), polytrichastrol B (7), bonn-2-en-11α-ol (8) and polytrichastrene B (9). Their absolute configurations were deter-
mined by analogous stereoselective deuterium labeling experiments as described above for 1–4, giving uniform results for all nine compounds (Figures S101–S107).

Alcohol 8 exhibits the skeleton of bonnadiene that is a side product of CwWS. The enantiomer is known from bonnadiene synthase (BdS) from Allokutzneria albata. All compounds fit well into the biosynthesis mechanism (Scheme 1), as 5 can be explained by deprotonation of E1, 6 can arise by water addition to F1, and 7 can be formed from K by another cyclization to L and capture with water. Only compounds 8 and 9 seem to follow a pathway that branches out early at B by 1,6-cyclization to C2, 1,2-hydride shift to D2 and 6,10-cyclization to E2, the precursor to 8 by water quenching (red pathway). A 1,4-hydride shift from F2 to M, 2,14-cyclization and deprotonation can lead to 9. Notably, F2 can also be linked to F1 through WMR (purple), so that the cyclization cascade along B, the red branch to F2, the purple link to F1 and the downstream reactions to L contain all intermediates in one direct line to explain all products by simple deprotonation or attack of water (only 9 requires a short side branch via M and N). The labeling experiments performed here cannot distinguish between the blue and the red pathway, because both pathways contain analogous elementary steps, so that all substrate atoms end up in the same positions.

To distinguish between these pathways, DFT calculations were performed (Table S15, Figure S108). The 1,3-hydride shift from A to B proceeds smoothly, ending in a B conformer that allows both cyclization reactions to C1 and C2 through similar transition state (TS) barriers (Scheme 1). The steps from C1 to F1 (blue) are all associated with low TS barriers or even barrierless, while the step from C2 to D2 has a fairly high barrier (+18.42 kcal mol\(^{-1}\)). The conversions from D2 to F2 are more feasible, and also the purple link between F1 and F2 can be passed in both directions. This analysis suggests that the blue pathway may be preferred for the formation of 1–7, and also 9 can be reached via this pathway and the purple link to F2, as the 1,4-hydride transfer to M can be realized computationally. Ring closure of M to N was only possible with assistance of a base (MeNH\(_2\)) in contact with Me20 to suppress spontaneous ring opening of N to M, ultimately leading to 9. An alternative two-step process with 1,3- and 1,2-hydride transfer is geometrically not possible, similar to 1,3-hydride transfers in guaiane biosynthesis that are rare and only allowed for the few stereoisomers with suitable geometry. Compound 8 may be the only case for which the red path is entered from B, because the step from F2 to E2 has an even higher TS than the C2 to D2 conversion. The high TS barriers to reach E2 from any side may explain the observation of 8 only as a minor compound in the 166F variant.

Based on the calculations the concerted F1 to H dyotropic rearrangement discussed above seems not possible (+37.77 kcal mol\(^{-1}\)), but assistance by a catalytic base, for which MeNH\(_2\) was selected as a model (initial attempts with water or NH\(_3\) failed), allows a stepwise process with low barriers through deprotonation of H12 and fragmentation to G followed by back transfer of the same proton to C11 and ring closure to H.

The SdS positions I75, L78, L79 and F297 are also often occupied by aromatic residues in DTPSs. In the corresponding positions of CpPS small (Gly, Ala) or non-polar (Met) residues are found that were exchanged to obtain the A87F, G90F, A91F, and M308W variants. G90F showed a decreased activity (23 ± 2 %) with main product 3, while A87F and A91F were inactive, presumably because of steric interference with GGPP. The M308W exchange did not yield a soluble protein, suggesting a major impact on the enzyme structure.

Side product 3 of CpPS is the main product of CwWS, and these enzymes differ in their active site residues in only two positions: A87 and A192 in CpPS are taken by T86 and V191 in CwWS (Table S2). The importance of active site residues to control the product profile of a TPS is impressively demonstrated by the A87T, A192V and—with two simultaneous exchanges—the A87T,A192V enzyme variants, all of which gave 3 as main product. The A192V variant showed an increased activity (176 ± 8 %), but also the single A87T and the double A87T,A192V exchange did not disturb substrate acceptance (117 ± 6 % and 108 ± 13 % activity, respectively).

In summary, we have characterized a bacterial DTPS that converts GGPP into diterpenes with novel skeletons. An unusual ethyl group formation was studied in detail by isotopic labeling experiments and DFT calculations, revealing a two-step rather than a concerted dyotropic rearrangement likely assisted by an active site base that needs to be identified by future structural work ideally in conjunction with QM/MM simulations. We have also identified 13 active site residues that are promising candidates for functional switches within TPSs, which allows for a rational design of engineered biocatalysts in future work.

Acknowledgements

This work was funded by the DFG (DI1536/7-2). Open access funding enabled and organized by Projekt DEAL.
Conflict of Interest

The authors declare no conflict of interest.

Keywords: biosynthesis · enzyme mechanisms · isotopes · mutagenesis · terpenes

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