

REVIEW

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The role and mechanisms of canonical and non-canonical tailoring enzymes in bacterial terpenoid biosynthesis

Yuya Kakumu, ^{ab} Ayesha Ahmed Chaudhri ^{ab} and Eric J. N. Helfrich ^{abc}

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Terpenoids represent the largest and structurally most diverse class of natural products. According to textbook knowledge, this diversity arises from a two-step biosynthetic process: first, terpene cyclases generate a vast array of mono- and polycyclic hydrocarbon scaffolds with multiple stereocenters from a limited set of achiral precursors, a process extensively studied over the past two decades. Subsequently, tailoring enzymes further modify these complex scaffolds through regio- and stereocontrolled oxidation and other functionalization reactions, a topic of increasing interest in recent years. The resulting highly functionalized terpenoids exhibit a broad spectrum of unique biological activities, making them promising candidates for drug development. Recent advances in genome sequencing technologies along with the development and application of sophisticated genome mining tools have revealed bacteria as a largely untapped resource for the discovery of complex terpenoids. Functional characterization of a limited number of bacterial terpenoid biosynthetic pathways, combined with in-depth mechanistic studies of key enzymes, has begun to reveal the versatility of bacterial enzymatic processes involved in terpenoid modification. In this review, we examine the various tailoring reactions leading to complex bacterial terpenoids. We first discuss canonical terpene-modifying enzymes, that catalyze the functionalization of unactivated C–H bonds, incorporation of diverse functional groups, and oxidative and non-oxidative rearrangements. We then explore non-canonical terpene-modifying enzymes that facilitate oxidative rearrangement, cyclization, isomerization, and dimerization reactions. The increasing number of characterized tailoring enzymes that participate in terpene hydrocarbon scaffold formation, rather than merely decorating pre-formed scaffolds suggests that a re-evaluation of the traditional two-phase model for terpenoid biosynthesis might be warranted. Finally, we address the potential and challenges of mining bacterial genomes to identify terpene biosynthetic gene clusters and expand the bacterial terpene biosynthetic and chemical space.

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^aInstitute for Molecular Bio Science, Goethe University Frankfurt, Max-von-Laue Strasse 9, 60438 Frankfurt am Main, Germany. E-mail: eric.helfrich@bio.uni-frankfurt.de

^bLOEWE Center for Translational Biodiversity Genomics (TBG), Senckenberganlage 25, 60325 Frankfurt am Main, Germany

^cSenckenberg Gesellschaft für Naturforschung, Senckenberganlage 25, 60325 Frankfurt am Main, Germany



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Yuya Kakumu

Yuya Kakumu received his BSc (2018) and MSc (2020) from Gifu University, Japan. Following a short period in industry, he returned to Gifu University as a research associate. Supported by a JSPS research fellowship, he earned his PhD from Gifu University under the tutelage of Prof. Tohru Mitsunaga in 2023. His doctoral studies focused on the metabolomics-based discovery, structural elucidation, and biological activities of plant-derived terpenoids. He is currently a postdoctoral researcher at Goethe University Frankfurt, Germany, working with Prof. Eric J. N. Helfrich. His research is centered around the genome mining-based identification and characterization of bacterial terpenoid biosynthetic pathways.

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Ayesha Ahmed Chaudhri

Ayesha Ahmed Chaudhri received her BSc from Punjab University and M.Phil. (2020) from Quaid-i-Azam University, Pakistan. She is currently pursuing her PhD at Goethe University Frankfurt, Germany, in the research group of Prof. Eric J. N. Helfrich, supported by an HEC/DAAD fellowship. Her research involves the genome mining-based discovery and functional characterization of non-canonical bacterial terpenoid biosynthetic gene clusters, with an emphasis on tailoring reactions involved in the biosynthesis of complex terpenoids.



1. Introduction

Terpenoids are ubiquitously biosynthesized in nature as primary or secondary metabolites. Their physiological roles range from coenzymes (*e.g.*, Q10) and vitamins (A, D, and K) to hormones (*e.g.*, steroids), pigments (*e.g.*, carotenoids), and defensive metabolites (*e.g.*, momilactones). Terpenoids, with about 100 000 characterized compounds (Dictionary of Natural Products),¹ represent the largest and structurally most diverse class of natural products. Terpenoids can be categorized into more than 400 structural families.¹ When taking diastereomers and natural product hybrids (*i.e.*, meroterpenoids) consisting of a terpenoid component and a substructure that is derived from other natural product classes (*e.g.*, polyketides and alkaloids) into consideration, the number of identified compounds rises to 180 000 that have been reported from all domains of life including plants, marine invertebrates, fungi, bacteria, and archaea.² Many terpenoids feature highly oxygenated sp^3 -rich carbon scaffolds with multiple stereocenters. These three-dimensionally distinct terpenoids frequently show potent biological activities and selectively interact with their molecular target.^{3,4} In these interactions, hydrogen bonds between highly oxygenated terpenoids and their targeted molecules play a crucial role. Several complex terpenoids have been exploited as drugs in human and veterinary medicine, agrochemicals, and chemical probes for biological studies. Notable examples include the anti-cancer agent taxol,⁵ the antibiotic pleuromutilin,⁶ the insecticide azadirachtin,⁷ and the phosphoinositide 3-kinase inhibitor wortmannin.⁸ Consequently, the biosyntheses of these complex terpenoids has garnered attention which resulted in the meticulous characterization of a variety of tailoring enzymes that catalyze unprecedented biosynthetic transformations. Moreover, complex terpenoids have served as a source of inspiration for the development of multi-step organic synthesis. Most notable is the recent progress in the development of efficient strategies for the rigorous

stereocontrolled functionalization of seemingly undistinguishable aliphatic C–H bonds.⁹

Despite their vast structural diversity, all terpenoids share a common biosynthetic origin.^{10,11} The biosynthesis of terpenoids begins with the formation of simple C₅ isoprene units, isopentenyl pyrophosphate (IPP) and dimethylallyl pyrophosphate (DMAPP), which are synthesized *via* two distinct pathways: the mevalonate (MVA) pathway¹² or the methylerythritol phosphate (MEP) pathway.¹³ Textbook knowledge describes terpenoid biosynthesis as a “two-step” process comprising a backbone assembly and a hydrocarbon scaffold modification phase (Fig. 1).¹⁴

In the first phase, the isoprene units are condensed in a head-to-tail fashion by oligoprenyl synthases to produce the methyl-branched acyclic and achiral polyene pyrophosphates, geranyl (GPP, C₁₀), farnesyl (FPP, C₁₅), geranylgeranyl (GGPP, C₂₀), geranylgeranyl (GFPP, C₂₅), hexaprenyl (HexPP, C₃₀), and heptaprenyl pyrophosphates (HepPP, C₃₅).^{15,16} Additionally, FPP and GGPP are condensed in a head-to-head fashion to yield squalene (C₃₀) and phytoene (C₄₀), respectively.¹⁷ The linear polyenes then undergo a series of carbocation-driven cyclization and rearrangement reactions catalyzed by two canonical types of terpene cyclases (TCs)^{10,18–20} or non-canonical TCs.^{21,22} Notably, this enzymatic process results in the formation of numerous mono- or polycyclic hydrocarbon scaffolds with multiple stereocenters.²³ For the biosynthesis of meroterpenes, the oligoprenyl pyrophosphates are transferred by prenyltransferases (PTs) onto nonterpenyl molecules such as polyketides and amino acid-derived compounds. The resulting hybrid compounds often undergo cyclization catalyzed by TCs.²⁴

In the second phase, (mero)terpenes undergo extensive modifications by oxidation of double bonds and unactivated C–H bonds, oxidative rearrangements, and incorporation of various functional groups.^{14,25} These modification reactions result in a myriad of structurally distinct terpenoids. Enzymes responsible for the decoration of terpene hydrocarbon scaffolds are primarily oxidoreductases and transferases, such as cytochrome P450 monooxygenases (P450s), nonheme iron α -ketoglutarate (Fe/ α KG)-dependent oxygenases, flavin-dependent monooxygenases (FMOs), NAD-dependent short-chain dehydrogenases/reductases (SDRs), methyltransferases (MTs), acetyltransferases (ATs), and glycosyltransferases (GTs). While most tailoring enzymes are chemo-, regio-, and stereoselective, a subset of specialized tailoring enzymes are multifunctional or promiscuous, leading to the production of a range of structurally similar terpenoids from the same pathway.^{26–30} Among these enzymes, P450s are particularly noteworthy for their ability to introduce various modifications to the terpene scaffold, such as hydroxylations, multi-electron oxidations, epoxidations, and oxidative structural rearrangements.³¹

Complex terpenoids have predominantly been isolated from eukaryotes.² Bacteria, on the other hand, have for the longest time been regarded as incapable of producing complex terpenoids.¹¹ However, with the advancements in analytical techniques, highly complex terpenoids have been discovered from bacterial origins, albeit fewer in number compared to those from eukaryotic origin.³² These bacterial terpenoids often



Eric J. N. Helfrich

Eric Helfrich earned his BSc (2010) from the University of Bonn and his MSc (2013) from the University of Jena, Germany. He pursued his PhD (2017) at ETH Zurich, Switzerland, under the supervision of Prof. Jörn Piel. He received an SNSF Fellowship to conduct postdoctoral research in the lab of Prof. Jon Clardy at Harvard Medical School, Boston, USA. In 2020, he was appointed Professor of Natural Product Genomics at Goethe University

Frankfurt, Germany. His research group develops genome mining tools to uncover overlooked biosynthetic gene clusters and investigates non-canonical biosynthetic pathways and enzymatic transformations in terpenoid, peptide and alkaloid biosynthesis.

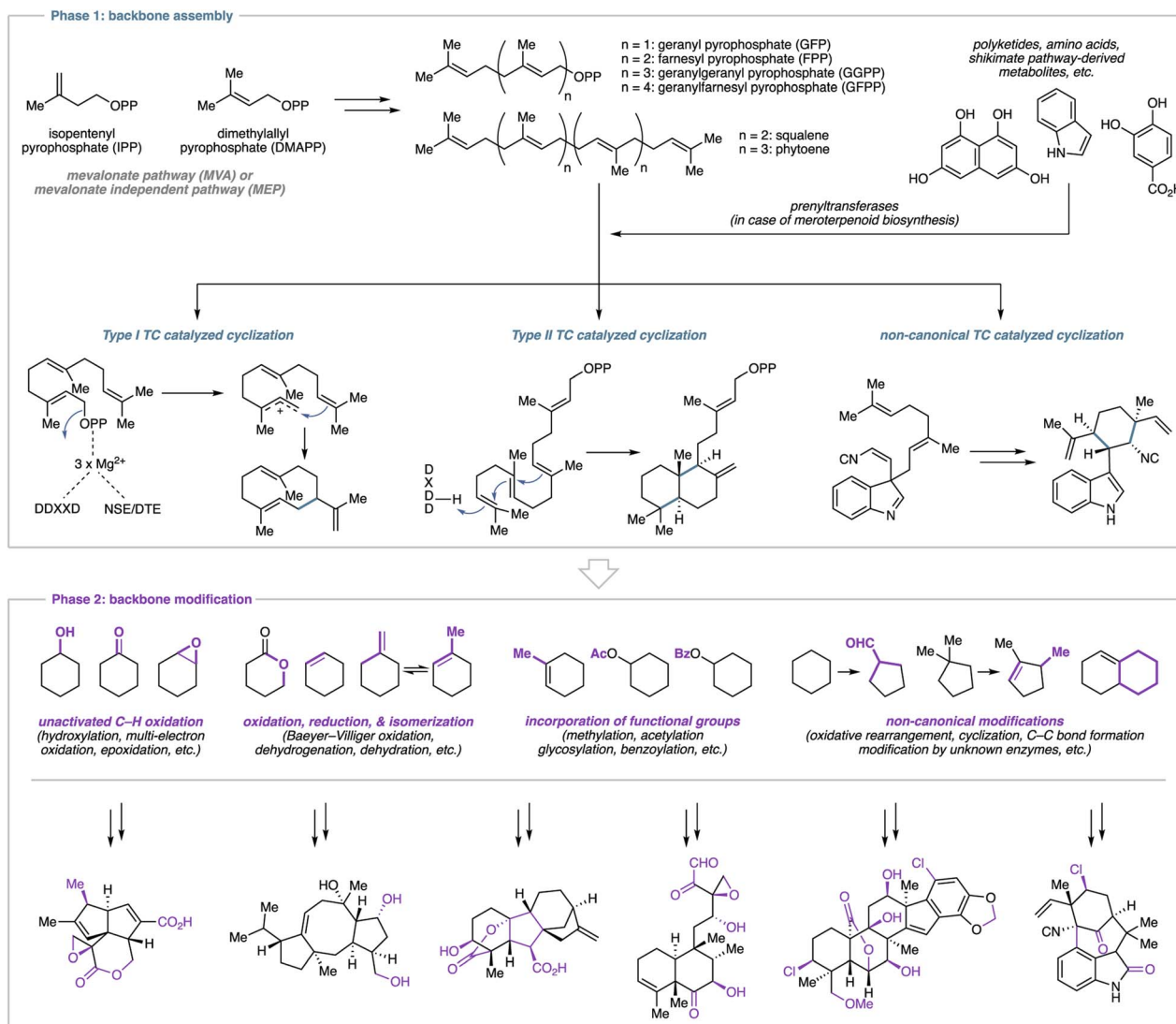


Fig. 1 Schematic overview of the two-phase terpenoid biosynthetic process. Phase 1: terpene hydrocarbon backbone assembly, phase 2: extensive modification of terpene backbone by tailoring enzymes.

exhibit remarkable biological activities and have the potential to serve as lead compounds for therapeutic applications. Examples of complex terpenoids with promising bioactivities include the immunosuppressant brasiliocardin A (**1**),³³ the selective fatty acid synthase inhibitor platensimycin (**2**),³⁴ the glyceraldehyde-3-phosphate dehydrogenase (GAPDH) inhibitor pentalenolactone (**3**),³⁵ and persicamidine C (**4**) which shows anti-SARS-CoV-2 activity (Fig. 2).³⁶ The ever increasing number of publicly available genome sequences in the post-genomic era, along with the development of sophisticated genome mining tools, have revealed that terpenoid biosynthetic gene clusters (BGCs) are widely distributed in bacterial genomes.^{37,38} Genes encoding TCs in bacterial genomes are frequently co-localized with multiple genes encoding tailoring enzymes, particularly P450s.^{37,38} As a result, bacteria are increasingly regarded as an almost untapped resource for the discovery of novel complex terpenoids.³² Despite the immense biosynthetic

potential to produce complex terpenoids as indicated by genome mining studies, the small number of characterized bacterial terpenoids suggests that most bacterial terpenoid BGCs are silent under standard laboratory culture conditions.

Over the past two decades, molecular biology techniques, such as heterologous expression, have emerged as powerful tools for the production of bacterial terpenoids and the characterization of the corresponding biosynthetic pathways. These techniques have not only enabled the identification of numerous highly functionalized terpenoids but also showcased the diverse enzymatic repertoire of tailoring enzymes involved in their biosynthesis. The vast majority of terpene hydrocarbon scaffold modifications involves functionalization of unactivated C–H bonds. However, a subset of tailoring enzymes acts as catalysts for unprecedented transformations, significantly modifying terpene frameworks and resulting in distinct terpenoid scaffolds. This growing body of knowledge on bacterial



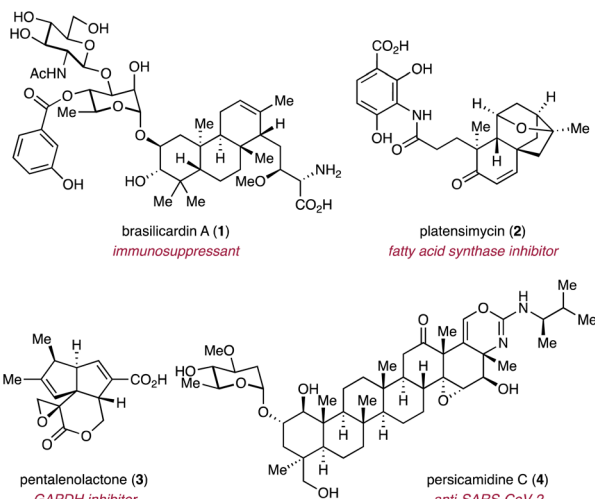


Fig. 2 Representative examples of complex bacterial terpenoids with promising biological activities.

terpenoid biosynthesis prompts us to review the broad spectrum of characterized tailoring reactions that we divided into canonical and non-canonical modifications.

Canonical modifications involve archetypical functionalizations catalyzed by oxidoreductases and transferases that operate *via* well-characterized catalytic mechanisms, such as oxygen rebound of iron-dependent enzymes and the methyl transfer reactions to nucleophiles by *S*-adenosyl-L-methionine (SAM)-dependent MTs. The reaction outcomes are highly conserved within enzyme families, as evidenced by numerous characterized examples. Canonical modifications of the terpene hydrocarbon scaffolds can encompass, but are not limited to, hydroxylations, epoxidations, four or six electron oxidations, dehydrogenations, methylations, glycosylations, benzoylations, and Baeyer–Villiger (BV) oxidations.

Non-canonical modifications, on the other hand, result from tailoring enzymes that employ previously uncharacterized reaction mechanisms to catalyze unprecedented biochemical transformations. These enzymatic activities often arise from neofunctionalization, enabling enzymes to drive key biosynthetic steps in shaping unique skeletal architectures of terpenoids. Examples of non-canonical modifications include oxidative rearrangements, cyclization, dimerization, and non-oxidative cyclization reactions. Furthermore, non-canonical modifications can also encompass the functionalization by members of known enzyme families with unprecedented catalytic activities and enzymes annotated as hypothetical proteins. The catalytic mechanisms underlying non-canonical tailoring enzymes are typically absent from textbooks, as these reactions have only recently been described with a limited number of characterized examples. To date, the majority of terpene biosynthetic studies have focused on the TC-catalyzed cyclization reactions. These canonical and non-canonical cyclizations have been covered in excellent review articles and are hence not addressed in this review.^{10,18–22}

In this review, we will begin with a brief description of a selection of hydrocarbon scaffolds that are biosynthesized

across kingdom borders, albeit with different modification patterns and as a result of convergent evolution. The following sections will delve into recent advancements in our understanding of canonical and non-canonical modifications of the terpene hydrocarbon backbone catalyzed by tailoring enzymes in bacterial terpenoid biosynthetic pathways. Each section will focus on a family of tailoring enzymes and will be further subdivided based on individual reaction types catalyzed by members of the enzyme family. Rather than reviewing the biosynthetic models of each bacterial terpenoid family separately, we have structured our review to focus on specific families of tailoring enzymes. This approach allows us to highlight similarities and differences between tailoring reactions and pathways, providing a more cohesive and informative narrative. We provide a brief overview of the biosynthetic model for each terpenoid family upon its first mention before delving into detailed descriptions of the respective tailoring reaction. We believe this structure offers a more insightful perspective compared to the traditional approach of reviewing terpenoid families individually. Our review does not include proteins with previously unknown functions involved in the cyclization phase, *e.g.*, Stig cyclases participating in cyanobacterial halapindole biosynthesis³⁹ or Pyr4-family transmembrane cyclases involved in meroterpenoid production.^{40,41} Furthermore, we will discuss the potential and limitation of genome mining strategies for the identification of highly oxidized bacterial terpenoids to further expand bacterial terpene biosynthetic and chemical space in the future.

2. Structural similarity and divergence of bacterial terpenoids and terpenoids of other origin

Terpenoids produced by bacteria often fall into the same structural families as those found in other kingdom of life.³² Since the number of oligoprenyl precursors is limited, it is not surprising that the same hydrocarbon scaffolds are produced across kingdom borders. Despite the commonality of terpene hydrocarbon backbones, most bacterial terpenoids exhibit distinct oxidation patterns when compared to eukaryote-derived terpenoids. However, in certain cases (*e.g.*, the gibberellins), the same modification patterns originate from different pathways as a result of convergent evolution. This section will introduce examples of bioactive terpenoids where the distinct modification patterns observed for bacterial and non-bacterial terpenoids are responsible for, in many cases, non-overlapping bioactivities. These drastic changes in bioactivity between members of a scaffold family highlight the importance of tailoring reactions during the biosynthesis of complex terpenoids.

2.1. Gibberellins

Gibberellins are norditerpenoid phytohormones that are widely found in plants, plant-associated fungi, and bacteria.⁴² In case of plant-associated microorganisms, gibberellins are utilized for the suppression of the plant host's immune response or the



promotion of nodule formation.⁴² Gibberellins that function as phytohormones—GA₁, GA₃, GA₄, and GA₇ (5–8)—share the 6/5/6/5-tetracyclic *ent*-gibberellane scaffold with a C-3 β hydroxy group, C-6 carboxylic acid, and γ -lactone bridge (Fig. 3A).^{43,44} The C-3 β hydroxy functionality is crucial for the promotion of plant growth and development.^{42,43} Plant pathogenetic bacteria such as *Xanthomonas oryzae* pv. *oryzicola* produce 7.^{45–47} In contrast, most nodule-forming symbionts, such as *Rhizobium* spp., do not produce plant growth-promoting gibberellins and instead produce GA₉ (9) that does not act as a phytohormone,^{46–48} although some rhizobia produce 7.⁴⁹ The difference in gibberellin production between pathogenic and symbiotic bacteria is hypothesized to be related to their respective types of relationships with plants.⁴² Plant pathogenetic bacteria might strategically produce 7 to compromise the plant host's immune system to facilitate infection.^{45,46} Symbiotic bacteria, on the other hand, delegate the C-3 β hydroxylation step to their plant hosts so that plants can balance the production of gibberellins.⁴⁷ This strategy allows symbionts to support plant growth without suppressing the plant's immune system.⁵⁰

2.2. Fusicoccane diterpenoids

Fusicoccane diterpenoids are produced by fungi, bacteria, and plants.⁵¹ The family features a 5/8/5-tricyclic dicyclopenta[*a,d*] cyclooctane ring system with characteristic oxidation and modification patterns that are dependent on the producing organism.^{51,52} Phytopathogenic fungus-derived fusicoccin A (10) and cotylenin A (11) feature a tetra-oxygenated terpene scaffold with a reverse-isoprenylated glycosyl group (Fig. 3B).^{53–56} Both compounds act as molecular glues that stabilize protein–protein interactions in the eukaryotic 14-3-3 protein family.⁵⁷ On the other hand, cyclooctatin (12),^{58,59} found in various *Streptomyces* spp.,^{60–63} exhibits a different oxidation pattern compared to the fungus-derived fusicoccane-type diterpenoids such as cotylenol (13) which is an aglycone of 11 (Fig. 3B).^{64,65} Interestingly, 12 is a potent lysophospholipase inhibitor⁵⁸ and hybrids of 12 and a polyketide from *Streptomyces violascens*, e.g., fusicomycin B (14), suppress migration and invasion of human hepatocarcinoma cells through the inhibition of matrix metalloproteases.⁶⁰ The biological activities of fusicoccane diterpenoids largely rely on the oxidation patterns and the incorporation of other building blocks.

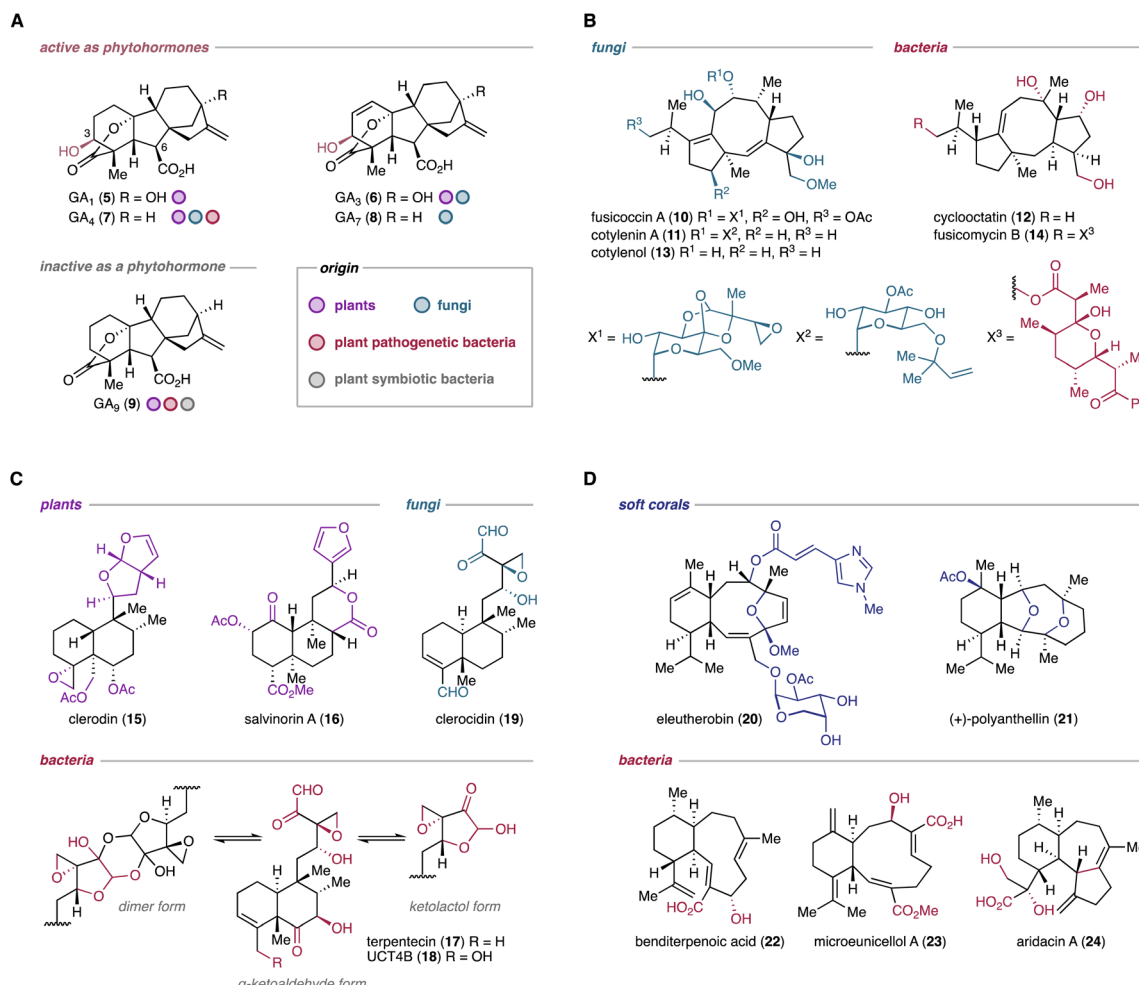


Fig. 3 Structural similarity and difference of diterpenoids from bacterial and eukaryotic origins. (A) Gibberellin norditerpenoids that are active or inactive as phytohormones from different taxonomic origins. (B) Selected fusicoccane diterpenoids from bacteria and fungi. (C) Selected clerodane diterpenoids from plants, fungi, and bacteria. (D) Selected eunicellane diterpenoids from soft corals and bacteria.



2.3. Clerodane diterpenoids

Clerodane diterpenoids are mainly found in plants, where they typically feature a branched carbon chain connected to a decalin core, often forming heterocycles such as furan and furanone rings.⁶⁶ Furthermore, the stereochemical configuration of the decalin ring is divided into eight types, namely four *neo*-clerodane types and four *ent*-*neo*-clerodane types, providing a wide range of structural diversity.⁶⁶ Representative examples from plants include the insecticide clerodin (15)^{67–70} and the potent selective κ-opioid receptor agonist salvinorin A (16).^{71,72} In contrast, actinobacteria are capable of producing a small family of clerodane-type diterpenoids, such as terpentecin (17)^{73,74} and UCT4B (18) (Fig. 3C).^{75,76} Bacterial clerodanes feature a highly oxygenated alkyl chain that exists in equilibrium between the α-ketoformyl aldehyde, hemiacetal, and dioxane dimer forms in solution.⁷⁷ Interestingly, fungi produce the highly similar diterpenoid clerocidin (19) that features a slightly different oxidation pattern at the decalin ring.^{77,78} These microbial diterpenoids act as topoisomerase II inhibitors that prevent DNA religation and result in DNA cleavage.^{75,79–81} This activity is significantly influenced by the presence of the unique oxygenated alkyl chain, wherein a strained epoxide adjacent to a ketolactol residue alkylates guanine bases.^{82,83}

2.4. Eunicellane diterpenoids

Historically, eunicellane diterpenoids have been almost exclusively identified from soft corals.^{84–86} Extensive isolation and structure elucidation efforts have shown that most of the coral-derived eunicellane diterpenoids feature a 6/10-*cis*-bicyclic ring system, which typically contains a transannular ether bridge.^{84–86} These diterpenoids often exhibit remarkable biological activities, such as the taxol-like microtubule stabilization activity of eleutherobin (20)^{87,88} and the anti-migration and anti-invasion activities of (+)-polyanthelin A (21).⁸⁹ Only recently, bacteria have been shown to produce various eunicellane diterpenoids with either *cis*- or *trans*-fused bicyclic scaffolds.⁹⁰ The oxidation patterns of bacterial eunicellanes are distinct from their coral-derived counterparts. It is noteworthy that all previously identified bacterial terpenoids lack the ether bridge on the 10-membered ring, although the number of identified compounds remains relatively small.⁹⁰ Instead, almost all bacterial eunicellanes feature fully oxidized allyl methyls. Notable examples include the antibacterial benditerpenoic acid (22),⁹¹ the antiproliferative microeunicellol A (23),⁹² and aridaicin A (24) with weak cytotoxicity against cancer cells (Fig. 3D).⁹³

2.5. Other terpenoids

In addition to the above-described examples, plant-like oxygenated eudesmane sesquiterpenoids have been identified from several *Streptomyces* spp.^{94–99} Furthermore, among structurally diverse bacterial carotenoids (>300 from ProCarDB),¹⁰⁰ some compounds such as β-carotene and astaxanthin are also produced by eukaryotes.¹⁰¹ Actinomycetes produce (iso) pimarane-type norditerpenoids that lack one of the *gem*-methyls, which is a feature almost exclusively found in (iso) pimaranes of bacterial origin.^{94,102–104}

The structural diversity of bacterial terpenoids and their unique oxidation patterns reflect the versatile biosynthetic capabilities of bacteria, demonstrating their potential to produce novel bioactive compounds albeit retaining the common frameworks observed across kingdom borders. Moreover, examples like the gibberellins showcase that some biosynthetic pathways have convergently evolved to produce structurally identical terpenoids using different strategies across the tree of life.

3. Modification of terpene hydrocarbon scaffolds by canonical tailoring enzymes in bacteria

3.1. Oxidation by cytochrome P450s

P450s are heme-thiolate-containing monooxygenases that play a crucial role in the oxidative transformation of members of various natural product classes.¹⁰⁵ They are predominantly known to catalyze hydroxylations of unactivated C–H bonds, epoxidations of alkenes, and four- and six-electron oxidations, the formation of aldehydes, ketones, and carboxylic acids, as well as biaryl couplings.^{106–108} Most P450s require redox partners, *e.g.*, a ferredoxin (Fd) and a ferredoxin reductase (FDR) in bacteria, to obtain single electrons for the reduction of the heme iron species during the catalytic cycle.¹⁰⁵ Although the catalytic mechanism of P450s has been extensively reviewed,^{106–109} we briefly describe the most crucial steps of its archetypical catalytic cycle using C–H hydroxylation reactions through the classical oxygen rebound mechanism as an example. The purpose of this brief introduction is to compare the catalytic cycle of canonical modifications to that of non-canonical modifications, such as skeletal rearrangements (*vide infra*). In the oxygen rebound mechanism (Fig. 4), a highly electrophilic oxoiron(IV) porphyrin cation radical (Compound I) abstracts a hydrogen atom from the substrate to generate a substrate radical. At the same time, Compound I is converted to a hydroxoiron(IV) intermediate (Compound II). The rebound of the hydroxyl radical from Compound II onto the substrate radical yields the hydroxylated substrate, returning Compound II to the incipient ferric state of the catalytic cycle. The formation of aldehydes, ketones, and carboxylic acids can be explained by multiple rounds of C–H hydroxylation to form diols coupled with H₂O elimination. The epoxidation of olefinic bonds is also realized *via* the radical rebound mechanism (Fig. 4).¹⁰⁶ This almost concerted reaction begins with coupling of an olefin and Compound I to yield the heme iron-substrate radical complex, with the components connected by a C–O bond. The resulting iron alkoxy radical intermediate immediately forms a second C–O bond through the addition of an iron(IV) oxyl radical to the alkyl radical, leading to the formation of an epoxide.¹⁰⁶

P450s serve as the archetypical biocatalyst for oxidative functionalization in terpenoid biosynthetic pathways.³¹ In bacterial genomes, most core biosynthetic genes encoding TCs and oligoprenyl synthases are clustered with one or multiple genes encoding P450s that are likely involved in the decoration



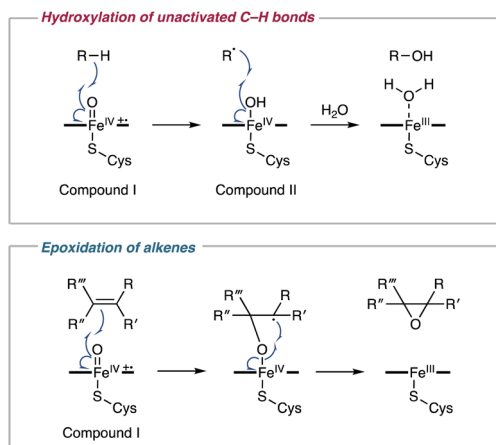


Fig. 4 Consensus oxygen rebound mechanism of the P450-catalyzed hydroxylation of unactivated C–H bonds and epoxidation of alkenes.

of the terpene hydrocarbon scaffold.^{37,38} The vast majority of these P450s encoded in bacterial terpenoid BGCs have not yet been functionally characterized. However, several studies have demonstrated the diverse catalytic activities of P450s involved in bacterial terpenoid biosynthesis.

3.1.1. Hydroxylation. Hydroxylation reactions in terpenoid biosynthetic pathways have a key role in generating a wide range of oxidation patterns in hydrocarbon backbones, affecting physicochemical and biological properties of the scaffold.¹¹⁰ Moreover, the installed hydroxy groups frequently serve as handles for the transfer of non-terpenyl building blocks by transferases. Alternatively, the oxidases further oxidize the installed hydroxy groups to aldehyde or carboxylic acid functionalities, respectively.¹¹¹ Leveraging the well-established heterologous expression systems that produce terpene precursors in high titers, P450s encoded in terpene BGCs have been shown to hydroxylate unactivated 1°, 2°, and 3° carbons of sesqui- and diterpene scaffolds to produce various products.^{112,113}

3.1.1.1. Cyclooctatin. Cyclooctatin (12) was isolated from *Streptomyces melanoporofaceins* MI614-43F2.^{58,59} The corresponding BGC (*cot*) is comprised of four genes encoding a GGPP synthase (GGPPS), a TC, and two P450s (CotB3, CotB4).¹¹⁴ Heterologous expression of the *cot* BGC-encoded genes revealed that CotB3 installs a hydroxy group at C-5 of cyclooctat-9-en-7-ol (25) to produce cyclooctat-9-ene-5,7-diol (26). Subsequently, CotB4 hydroxylates C-18 of 26 to yield 12 (Fig. 5).¹¹⁴ These two P450s catalyze regio- and stereospecific reactions with the assistance of an endogenous redox system of the heterologous host. Further studies revealed that the catalytic efficiencies of CotB3 and CotB4 are influenced by the redox system.¹¹⁵ Specifically, the AfR/Afx redox system identified from *Streptomyces afghaniensis* shows increased compatibility with CotB3 and CotB4 than the *Pseudomonas putida*-derived PdR/Pdx system. Heterologous expression of CotB3 and CotB4 combined with AfR/Afx results in higher production titers of 12 (15 mg L⁻¹) compared to the native producer (0.35 mg mL⁻¹).¹¹⁵ Additionally, substrate scopes of CotB3 and CotB4 were

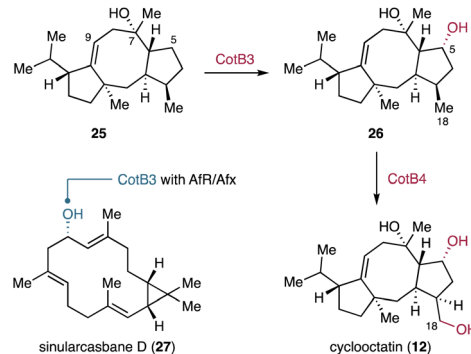


Fig. 5 Sequential hydroxylations in the cyclooctatin biosynthetic pathway by P450s and substrate promiscuous activity of the P450 CotB3 in dependence of redox partners.

explored using an *Escherichia coli* heterologous host harboring genes encoding diterpene TCs and AfR/Afx.¹¹⁵ These combinatorial biosynthetic studies revealed that CotB3 is also capable of hydroxylating a casbane-type diterpene backbone produced by the plant-derived TC *JcCSH*, leading to the formation of sinularcasbane D (27) (Fig. 5).¹¹⁵ Homologous BGCs of the *cot* BGC have been identified in the genomes of at least 72 actinomycetes.¹¹⁶ *In vitro* mechanistic investigations of CotB2 homologs revealed that ScCotB2 from *Streptomyces collinus* Tü365 converts GGPP to collinodiene (cyclooct-5,7-diene) as the main product, along with 25.¹¹⁶ Collinodiene may also serve as a substrate for the P450 homologs of CotB3 and CotB4 in *S. collinus*.

3.1.1.2. Phenalinolactones. Phenalinolactone A (28) is an *anti-anti-syn*-fused tricyclic perhydrophenanthrene diterpenoid which is decorated with a methyl pyrrolate, a dihydroxy furanone, and a methyl L-amicetose moiety.¹¹⁷ The *pla* BGC identified in the genome of *Streptomyces* sp. Tü6071 spans 42 kbp and harbors four P450 genes (*plaO2*, *plaO3*, *plaO4*, *plaO5*).¹¹⁸ Targeted inactivation of phenalinolactone biosynthetic genes and structure determination of the resulting intermediates indicated the functions of PlaO3, PLaO4, and PlaO5 (Fig. 6).^{118,119} PlaO3 first catalyzes hydroxylation of the terminal methyl C-19 at the A-ring of the dihydroxyfuranone-bearing PL HS6 (29) to form 30. The installed hydroxy group is further decorated by the acyltransferase PlaP2 to yield 31. Subsequently, PlaO4 converts 31 to 32 by installing a hydroxy group at C-20. The resulting OH-20 is used for the condensation with L-amicetose which is transferred by the glycosyltransferase PlaA6, leading to the formation of 33. Afterwards, the acyltransferase PlaV acetylates at OH-3 to produce phenalinolactone D (34). Finally, PlaO5 hydroxylates C-1 of 34 to yield 28. The remaining PlaO2 is not essential for the biosynthesis of 28.¹¹⁹

3.1.1.3. Brasilicardins. Brasilicardin A (1), an *anti-syn-anti*-fused tricyclic diterpenoid glycoside, is a promising immunosuppressive drug candidate.^{33,120,121} The mechanism of action of 1 is distinct from that of currently used immunosuppressants such as FK506, cyclosporin, and phingolimod.¹²² Structure-activity relationship studies revealed that the functional groups at C-2, composed of L-rhamnose, N-acetylglucosamine, and 3-hydroxybenzoate, are vital for the immunosuppressive activity



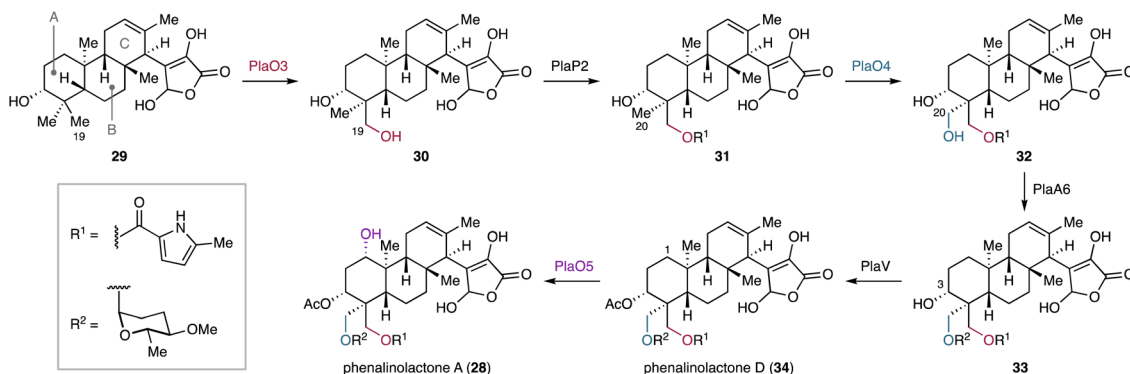


Fig. 6 Proposed tailoring steps catalyzed by the three P450s, PlaO3, PlaO4, and PlaO5 in the biosynthesis of phenalinolactone A.

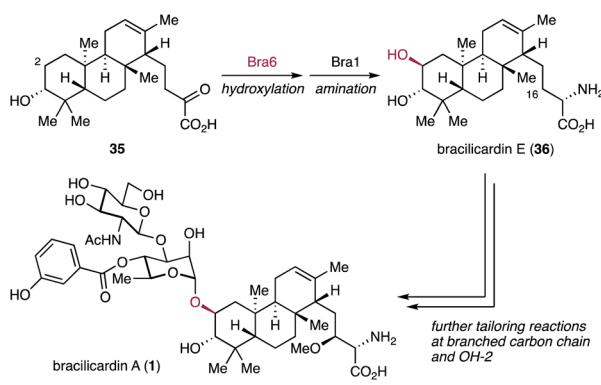
of **1**.^{121,123} Functional analyses of the *bra* gene products from *Nocardioides* strains showed that the P450 Bra6 catalyzes the C-2 β hydroxylation of the perhydrophenanthrene intermediate **35** (Fig. 7).^{124,125} The concomitant transamination of the α -ketoacid catalyzed by the aminotransferase Bra1 results in the production of brasiliocardin E (**36**). The aglycone **36** undergoes methylation at C-16, catalyzed by the Fe/ α KG-dependent oxygenase Bra0 and the methyltransferase Bra11, followed by extensive functionalization at OH-2 by transferases derived from the *bra* BGC, ultimately yielding **1**.^{124,125}

3.1.1.4. Other terpenoids. In addition to the above mentioned examples, C-H hydroxylation by P450s are found in biosynthetic pathways of various bacterial terpenoids, e.g., **37–43** (Fig. 8).^{112,113,126} Notably, CryP derived from *Cryophilus cryophilus* CGMCC 4.1710 installs a hydroxy group on a bridgehead carbon to yield cryophilain (**39**). Although hydroxy bridgeheads are frequently found in complex terpenoids, CryP stands out as one of the handful examples that is biochemically validated to oxidize a bridgehead carbon in a stereospecific manner.¹¹³

3.1.2. Hydroxylation sets up spontaneous downstream reactions. P450-catalyzed hydroxylation reactions sometimes set up spontaneous downstream reactions.

3.1.2.1. Platensimycin. Platensimycin (**2**) is an antibiotic that selectively inhibits the β -ketoacyl-(acyl-carrier-protein) synthase I (FabF) in bacterial fatty acid biosynthesis.³⁴ It also selectively inhibits the mammalian fatty acid synthase and acts as a potent

suppressor of hepatic *de novo* lipogenesis.¹²⁷ In comparison to the structurally related *ent*-*seco*-atisane diterpenoid platencin (**44**), an inhibitor of both FabF and β -ketoacyl-(acyl-carrier-protein) synthase III (FabH),¹²⁸ **2** harbors an ether bond between the C and D rings of the *ent*-kaurane derived carbon skeleton, which likely contributes to its selective inhibition of FabF.¹²⁹ Inactivation of candidate biosynthetic genes and *in vitro* biochemical characterization experiments revealed the biosynthetic origin of the tetrahydrofuran ring in **2**.^{130,131} The P450 PtmO5 stereoselectively hydroxylates C-11 β of 16 α -hydroxy-*ent*-kauranoic acid **45** (Fig. 9).¹³⁰ The resulting diol **46** undergoes nonenzymatic dehydroxylation of the tertiary alcohol OH-16, forming the tertiary cation **IM1**. The cation **IM1** is subsequently quenched by the C-11 hydroxy group which results in the formation of 11,16-epoxy-*ent*-kauranoic acid **47**.^{130,131} The C-11 hydroxy group installed by PtmO5 is positioned in close proximity to the generated carbocation, leading to a spontaneous intramolecular S_N1 reaction.¹³⁰ The intramolecular ether formation of **46** is only observed under strong acidic conditions *in vitro*. Therefore, the possibility that PtmO5 or other enzymes encoded in the *ptm* BGC are involved in the ether formation *in vivo* cannot be excluded. After the formation of the ether bridge, **47** undergoes various modifications, including B-ring oxidation (Chapter 3.2.3), A-ring cleavage (Chapter 4.2.5), β -oxidation, and non-terpenyl building block incorporation (Chapter 3.3.4), to yield **2**.¹³²



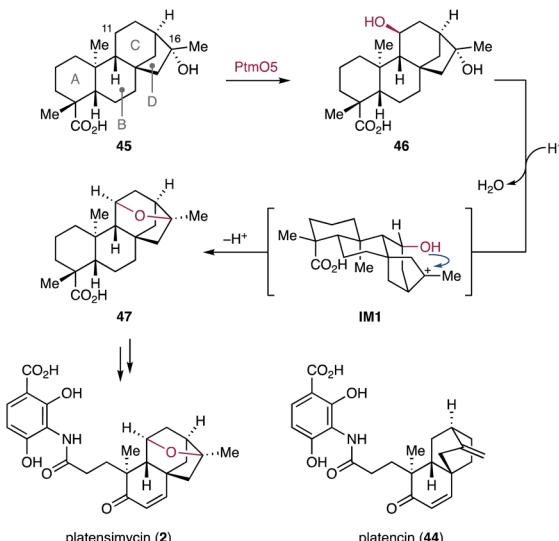


Fig. 9 P450-mediated hydroxylation and nonenzymatic ether bond formation in platensimycin biosynthesis.

3.1.3. Multi-electron oxidations. Terpene scaffold modifying P450s frequently catalyze two or three sequential oxidations to yield ketones and carboxylic acids from methylenes and methyls, respectively.

3.1.3.1. Pentalenolactone. The GAPDH inhibitor pentalenolactone (3) is a sesquiterpenoid lactone that features a densely oxidized 5/5/6/3-tetracyclic ring framework.¹³³ The *ptl* gene cluster for the biosynthesis of pentalenolactone-type sesquiterpenoids was identified in the genome of *Streptomyces avermitilis* MA-4680.¹³⁴ The hydrocarbon backbone of 3, namely pentalenene (48), is biosynthesized by the TC PtIA from FPP.¹³⁴ Decoration of the scaffold **48** begins with a multi-electron oxidation of the C-13 methyl by the P450 PtII (Fig. 10A).¹³⁵ *In vitro* experiments showed that PtII catalyzes the oxidation of **48** to yield the alcohol **49**, the aldehyde **50**, and trace amounts of the carboxylic acid **51**.¹³⁵ Consequently, PtII was proposed to catalyze a two-step oxidation that forms the aldehyde from the allylic methyl at C-13. Given the low efficient conversion of **48** into **51**, it remains unclear whether PtII or another oxygenase, with appropriate redox partners, catalyzes the oxidation of the aldehyde in **50** to the carboxylic acid in **51**.¹³⁵

3.1.3.2. Albaflavenone. Another prominent example of P450-catalyzed multi-electron oxidations is the CYP170A1-mediated ketone formation during albaflavenone (52) biosynthesis in *Streptomyces coelicolor* A3(2) (Fig. 10B).¹³⁶ CYP170A1 performs two sequential allylic oxidations at the C-5 methylene to yield **52**. Interestingly, the CYP170A1-catalyzed ketone formation in *epi*-isoizaene (**53**) proceeds through the formation of the epimeric albaflavenols (**54**), both of which are converted to **52**.¹³⁶

3.1.3.3. Venezuelaene B. Venezuelaene B (**55**) is a fragrant diterpenoid. The *ven* BGC has been identified in the genome of *Streptomyces venezuelae* ATCC 15439 (Fig. 10C).¹³⁷ This compound features an unprecedented 6/5/5/7-tetracyclic ring system with a ketone as the lone oxygenated functionality. Among *ven* gene products, the P450 VenC catalyzes the

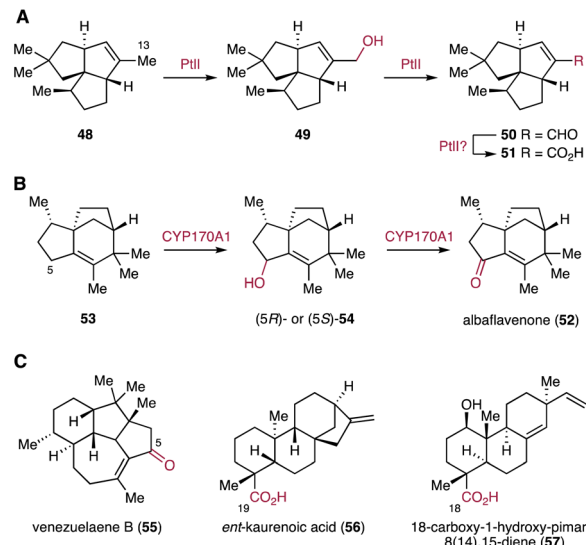


Fig. 10 Examples of P450-catalyzed sequential oxidations in bacterial terpenoid biosynthetic pathways. (A) PtII-catalyzed carboxylation of pentalenolactone biosynthetic pathways. (B) Ketone formation in albaflavenone catalyzed by CYP170A1. (C) Other examples of terpenoids with ketones or carboxylic acids installed by P450s.

formation of the ketone by four-electron oxidation. Interestingly, another BGC-encoded P450, VenB, with 65% sequence homology to VenC shows no enzymatic activity using either **55** or the unmodified hydrocarbon scaffold as a substrate.¹³⁷

3.1.3.4. Gibberellins. In the biosynthetic pathway of bacterial gibberellins, CYP117 catalyzes a six-electron oxidation of the C-19 methyl in *ent*-kaurene to yield *ent*-kaurenoic acid (**56**) (Fig. 10C).^{48,138}

3.1.3.5. Pimarane-type diterpenoids. The *asp* BGC that encodes type I and type II TCs (*aspT1*, *aspT2*) and two P450s (*aspP1*, *aspP2*) was identified from the genome of *Actinomadura* sp. NAK0032.¹¹² Heterologous expression studies in the *S. albus* J1074M chassis revealed that the terpene scaffold pimara-8(14),15-diene is first hydroxylated by Asp2 to give a terpene alcohol. The methyl C-18 of the resulting alcohol is then sequentially oxidized by Asp1 to the carboxylic acid (**57**).¹¹²

3.1.4. Multi-site oxidation. Compared to monofunctional P450s that modify a single site of the terpene scaffold, some P450s are capable of oxidizing terpene hydrocarbon scaffolds on multiple sites. Some of the P450s even catalyze different reactions at distinct sites of the same molecule.¹³⁹ Enzymatic activities of these multifunctional P450s result in the production of various terpenoids with different oxidation patterns at the same time.¹⁴⁰

3.1.4.1. Cylabdan. Cylabdan are hybrids of a labdane diterpenoid and an acyl cysteine linked via a thioether bond.^{141–143} These compounds have been isolated from *Streptomyces cylabdanicus* K04-0144. They enhance the activity of imipenem against methicillin-resistant *Staphylococcus aureus*.¹⁴⁴ Genome mining identified the *cld* BGC comprised of four genes encoding a GGPPS, type I and II TCs, and a P450.¹⁴⁵ Subsequent heterologous expression of the *cld* genes in



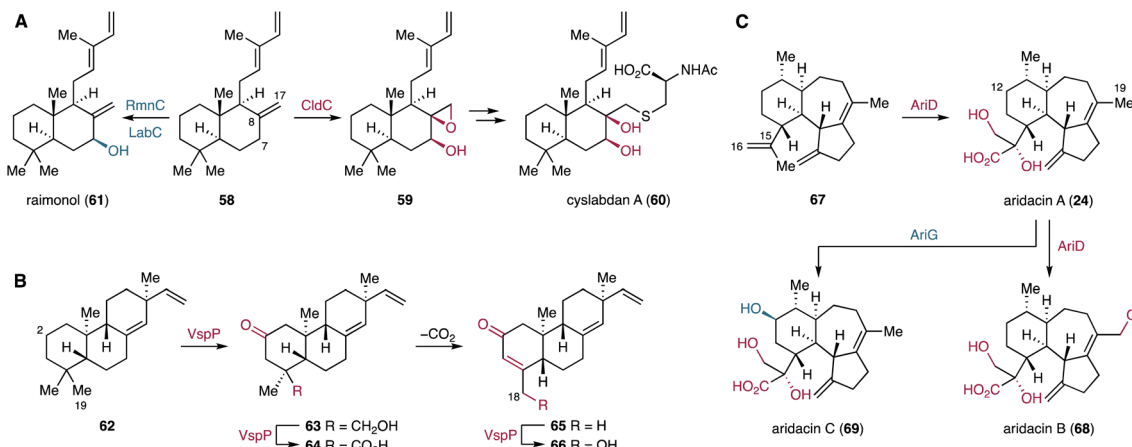


Fig. 11 Representative examples of bacteria terpenoid functionalization by multifunctional P450s. Proposed biosynthesis of cyslabdan A and raimonol (A), *ent*-sandaracopimaradiene derived diterpenoids (B), and aridacins (C).

Streptomyces avermitilis SUKA22 indicated that the P450 CldC catalyzes hydroxylation at C-7 and epoxidation at the C-8/C-17 olefin of labda-8(17)-12,14-triene (58) to yield 59 (Fig. 11A). The epoxide of 59 is opened by nucleophilic addition of mycothiol. The resulting mycothiol derivative is then hydrolyzed by a mycothiol-S-conjugate amidase to yield cyslabdan A (60). The *rnn* and *lab* BGCs were discovered as homologous BGCs of the *cld* BGC in the genomes of *Streptomyces anulatus* GM95 and *Streptomyces* sp. KIB 015, respectively.^{145,146} Interestingly, the P450s RmnC and LabC, homologs of the multifunctional CldC, are monofunctional and only install a hydroxy group at C-7 β of 58 to form raimonol (61) (Fig. 11A).^{145,146}

3.1.4.2. Nor-*ent*-sandaracopimaradiene diterpenoids. The *nor-ent*-sandaracopimaradiene diterpenoids have been identified through heterologous expression of the *vsp* BGC encoded in the genome of *Verrucospora* sp. NA02020.¹¹² Functional characterization of the *vsp* BGC revealed that the P450 VspP is a versatile oxygenase involved in all oxidative modifications of the *ent*-sandaracopimaradiene scaffold (62).¹¹² Heterologous expression of the type I and II TC genes (*vspT1* and *vspT2*) of the *vsp* BGC in *S. albus* J1074M afforded 62.¹¹² The expression of *vspP* alongside *vspT1* and *vspT2* in *S. albus* led to the production of several oxidized diterpenoids, including 63–66 (Fig. 11B).¹¹² Based on the characterized products, VspP was proposed to be a multifunctional oxygenase that catalyzes ketone formation at C-2 and six electron oxidations of the methyl C-19 in 62, resulting in the formation of 64.¹¹² Furthermore, VspP decarboxylates C-19 of 64 and subsequently hydroxylates C-18 to form 66.

3.1.4.3. Aridacins. The recent discovery and biosynthetic studies of aridacins, eunicellane-type diterpenoids from *Amycolaptosis arida* CGMCC 4.5579, led to the identification of one of the highest oxidizing P450s, AriD, to date.⁹³ AriD forms a glyceric acid moiety from an isopropenyl residue of the 6/7/5-tricyclic arida-3,6,15-triene scaffold (67) to produce aridacin A (24) (Fig. 11C). The formation of the glyceric acid moiety was proposed to result from carboxylation of the C-17 methyl and epoxidation of the C-15 alkene, followed by epoxide ring opening through the addition of H₂O.⁹³ Additionally, AriD is

also capable of hydroxylating the C-19 methyl in 24 to form aridacin B (68). Another P450, AriG, catalyzes the hydroxylation of the methylene C-12 in 24, yielding aridacin C (69).⁹³ Although the catalytic activity of AriD likely follows the classic oxygen rebound mechanism, the incorporation of at least four oxygens in the terpene hydrocarbon backbone is quite rare.

3.2. Oxidation by other oxidoreductases

In bacterial terpenoid biosynthesis, the vast majority of enzymes that decorate the hydrocarbon scaffold during the modification phase are P450s. However, many large bacterial terpenoid BGCs also harbor genes encoding other oxidoreductases, such as FMOs, Fe/αKG-dependent oxygenases, and SDRs. These oxidoreductases are frequently utilized in the biosynthesis of highly oxidized terpenoids.

3.2.1. Flavin-dependent monooxygenases. One commonly observed FMO-catalyzed reaction in terpenoid biosynthetic pathways is the epoxidation of terminal olefins in oligoprenyl precursors, which serve as substrates for type II TCs in the cyclization phase. In addition to their role in terpene backbone cyclization, FMOs also catalyze various late-stage oxidative functionalizations of terpene scaffolds.

3.2.1.1. Sesquisabinene-type sesquiterpenoids. The recently identified *net* BGC in *Streptomyces netropsis* DSM 40259 harbors genes encoding a type I TC (*netT*), a P450 (*netP*), an acetyltransferase (*netA*), and a FMO (*netO*).¹¹² Based on the functional characterization of the *net* BGC by heterologous expression, NetO epoxidizes the terminal olefin C-10/C-11 of the hydrocarbon scaffold 7-*epi-cis*-sesquisabinene hydrate (70) generated by NetT (Fig. 12).¹¹² The stereoselectivity of the NetO-catalyzed epoxidation remains undetermined, as the products isolated from the co-expression of *netT* and *netO* are diastereomeric diols resulting from epoxide hydrolysis of 71 in the heterologous host.¹¹² The epoxidation of the dimethyl allyl group by NetO resembles the epoxidation of FMOs acting on oligoprenyl pyrophosphates and squalene to set up type II TC-catalyzed cyclization reactions. In contrast, NetO is a tailoring enzyme that epoxidizes the cyclized terpene scaffold 70. The P450 NetP



is proposed to convert **71** into the ketone-containing intermediate **72**. Alternatively, it has been suggested that the enzyme order can be reversed: **70** might be first converted to the ketone-containing intermediate **73** by NetP, followed by the NetO catalyzed epoxidation to yield **72** (Fig. 12).¹¹²

3.2.2. Bayer–Villiger monooxygenases. Baeyer–Villiger (BV) oxidations are reactions that convert a carbonyl to an ester or a lactone through the formation of a Criegee intermediate and its rearrangement.¹⁴⁷ In synthetic organic chemistry, BV reactions are widely applied for ring expansions and lactone formation, such as in steroid synthesis.¹⁴⁸ Nature widely employs Baeyer–Villiger monooxygenases (BVMOs) that mainly belong to the FMO enzyme family for late-stage tailoring reactions in terpenoid biosynthesis or in terpenoid degradation pathways.¹⁴⁹ However, there is, at present, only one biochemically verified example that employs a BVMO in a bacterial terpenoid biosynthetic pathway, *i.e.*, in the biosynthesis of the pentalenolactone family of sesquiterpenoids.^{150,151}

3.2.2.1. Pentalenolactones. Pentalenolactone (**3**) and neopentalenoketolactone (**74**) are representatives of highly oxygenated sesquiterpenoids isolated from various *Streptomyces* sp.^{133,150} The structure of **3** bears an epoxidized δ -lactone moiety that is likely derived from the methyl cyclopentane ring of 1-deoxypentalenoic acid (**51**) through ketone formation followed by BV oxidation. The biosynthetic transformations from **51** to pentalenolactone F (**75**) and **74** were thoroughly characterized using a combination of *in vitro* biochemical assays of tailoring enzymes, gene inactivation, and gene overexpression studies as well as single X-ray crystallography of the Fe/ α KG-dependent oxygenase PtlH.^{150–154} BGCs, named *ptl*, *pnt*, and *pen*, were characterized from the genomes of *S. avermitilis*, *Streptomyces arenae* Tü 469, and *Streptomyces exfoliatus* UC5319, respectively.^{134,151} After the production of **51** through a six electron oxidation (Fig. 10A), PtlH (PntH, PenH) hydroxylates the methylene C-11 to form 1-deoxy-11 β -hydroxypentalenic acid (**76**). The alcohol **76** is subsequently converted to 1-deoxy-11-oxopentalenic acid (**77**) by the SDR PtlF (PntF, PenF) (Fig. 13).^{152–154} The BVMO PntE (PenE) then catalyzes the BV oxidation of **77** to afford pentalenolactone D (**78**).¹⁵¹ The orthologous BVMO PtlE unexpectedly shows the opposite regioselectivity of BV oxidation on the ketone of **77**, resulting in the production of neopentalenolactone D (**79**) (Fig. 13).¹⁵⁰ Since

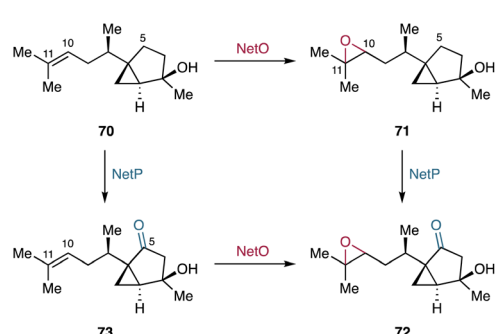


Fig. 12 Oxidative functionalization of sesquisabinene hydrate by the FMO NetO and the P450 NetP.

the BVMOs involved in pentalenolactone and neopentalenoketolactone biosynthesis share high sequence identity, crucial amino acid residues that dictate the distinct regioselectivity of PntE (PenE) and PtlE remain unknown.¹⁵¹ Nevertheless, it is the distinct regioselectivity of these BVs that is the bases for the diverging pathways of the pentalenolactones and neopentalenoketolactones.¹⁵¹

3.2.3. Nonheme iron α -ketoglutarate-dependent oxygenases. Fe/ α KG-dependent oxygenases are a large family of enzymes that use high-valent oxoiron(IV) intermediates.¹⁰⁸ The consensus catalytic mechanism of Fe/ α KG-dependent oxygenases is reminiscent of that of P450s.¹⁰⁸ The activated oxoiron(IV) species facilitates the homolysis of unactivated C–H bonds, resulting in the formation of a substrate radical and a hydroxoiron(III) species. The rebound of hydroxoiron(III) to the substrate radical follows to produce the hydroxylated substrate. Their product scope ranges from the hydroxylation of unactivated C–H bonds, to desaturation, halogenation, and oxidative rearrangement reactions.²⁷

3.2.3.1. Pentalenolactone. In the biosynthetic pathways of the pentalenolactone family of sesquiterpenoids, pentalenolactone D (**78**) and neopentalenolactone D (**79**) are structurally diversified by multi-step oxidizations catalyzed by the multi-functional Fe/ α KG-dependent oxygenases PntD (PenD) and

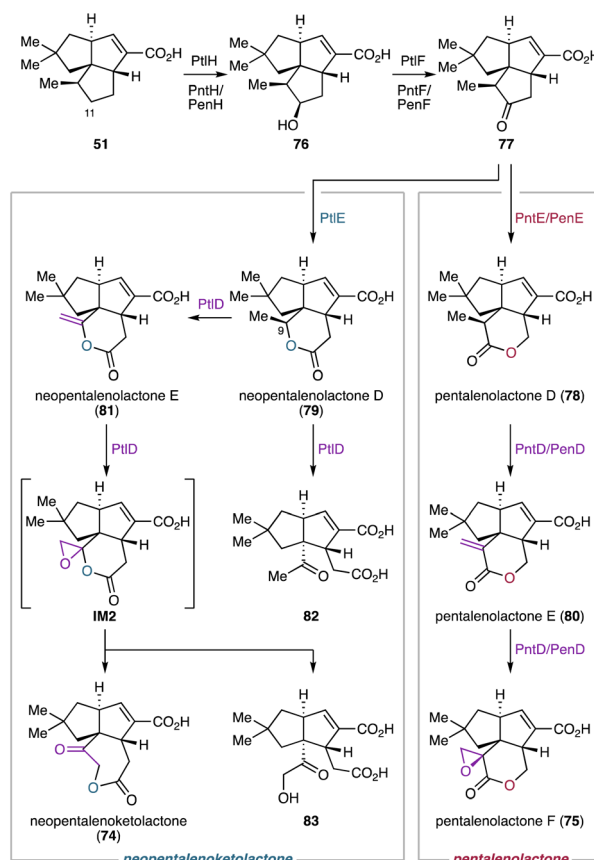


Fig. 13 Biosynthetic pathways of pentalenolactone F and neopentalenoketolactone branched by BVMO-mediated lactone formation.



PtID, respectively (Fig. 13).¹⁵¹ In these pathways, PntD (PenD) desaturates **78** to form pentalenolactone E (**80**) and epoxidizes the generated *exo*-methylene to produce **75**.¹⁵¹ *In vitro* biochemical studies using $H_2^{18}O$ or $^{18}O_2$ and site-directed mutagenesis of PtID confirmed its remarkable multifunctionality in the structural diversification of neopentaketolenolactones.^{155,156} PtID converts **79** into either neopentalenolactone E (**81**) by dehydrogenation or the ketoacid product **82** by hydroxylation-mediated δ -lactone cleavage. Notably, in the context of the PtID-catalyzed desaturation reaction, a hydrogen at C-9 is abstracted by the iron(IV)-oxo species and the subsequent electron transfer yields a carbocation intermediate, which is stabilized by the π -cation interaction with the active site residue Tyr113 in PtID.¹⁵⁶ The carbocation intermediate is then converted to the oxocarbenium ion, facilitating the deprotonation of H-10 by the basic nature of Lys288 in PtID to give **81**.¹⁵⁶ The *exo*-methylene **81** is epoxidized by PtID to yield the epoxyketal **IM2**, which immediately undergoes a spontaneous rearrangement reaction to form the ketoester **74** or a hydrolytic reaction to yield the δ -lactone-opened product **83**.¹⁵⁵ The remarkable functionality of PtID in catalyzing different reactions with various substrates gives rise to the complex and branched biosynthetic pathways of neopentalenolactone-related sesquiterpenoids.

3.2.3.2. Hapalindoles. In bacterial terpenoid biosynthesis, a representative example of Fe/ α KG-dependent oxygenases is the enantioselective halogenation of hapalindoles. The family of hapalindoles is composed of structurally diverse terpenoid indole alkaloids produced from stigonematalean cyanobacteria *Fischerella* spp. and *Hapalosiphon* spp.³⁹ The widespread occurrence of C-13 chlorinated hapalindoles suggests that chlorination is likely an early functionalization step in the oxidative maturation of hapalindole alkaloids (Fig. 14). The *wel* and *amb* BGCs for welwitindolinones and ambiguines have been identified in the genomes of *Hapalosiphon welwitschii* UTEX B1830 and *Fischerella ambiguia* UTEX1903, respectively. Each BGC harbors genes encoding five nonheme iron-dependent oxygenases including a Fe/ α KG-dependent oxygenase (*welO5* and *ambO5*).^{157,158} *In vitro* enzymatic assays demonstrated that WelO5 stereospecifically converts 12-*epi*-fischerindole U (**84**) and 12-*epi*-hapalindole C (**85**) to the chlorinated products, 12-*epi*-fischerindole G (**86**) and 12-*epi*-hapalindole E (**87**), respectively (Fig. 14).^{157,159} In addition to the chlorination, *in vitro* enzymatic assays showed that WelO5 is also capable of incorporating bromine in **84**.¹⁶⁰ The *apo* and *holo* structures of WelO5 provide insights into the catalytic mechanism of the stereoselective halogenation that requires ligand rearrangement.¹⁶¹ Mutating active site residues in WelO5 indicated that a single residue (Ser189) plays a key role in determining the selectivity of the halogenation reaction through regulating the position of an oxygenic group in the Fe(IV)-oxo intermediate by hydrogen bonding.¹⁶¹ In contrast to WelO5, the homologous AmbO5 (79% id) chlorinates the C-13 position of various premature hapalindole scaffolds.¹⁶² AmbO5 converts structurally distinct hapalindoles **84**, **85**, and **88–90** into the C-13 chlorinated molecules **86**, **87**, and **91–93**, respectively (Fig. 14). Sequence comparison of WelO5 and AmbO5 and

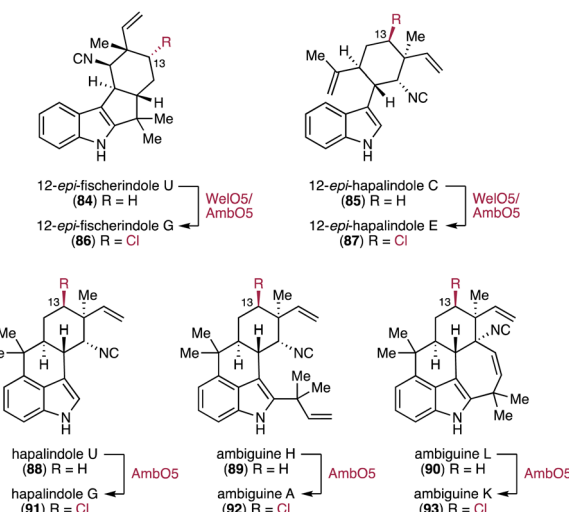


Fig. 14 Site-selective halogenation by Fe/ α KG-dependent halogenases in the biosynthesis of hapalindoles.

mutagenesis of WelO5's C-terminal residues revealed that the distinct C-terminal 18 amino acids determine the substrate scope.¹⁶²

3.2.4. Short-chain dehydrogenases/reductases. SDRs are a family of NAD(H)- or NAD(P)H-dependent oxidoreductases that have a highly varied substrate scope including alcohols, steroids, and aromatic compounds.¹⁶³ Their catalytic activities range from hydroxylation, dehydrogenation, to carbonyl reduction, and isomerization, as shown in steroid metabolism and terpenoid biosynthesis.^{164–166}

3.2.4.1. Platensimycin and platencin. In the biosynthesis of platensimycin (**2**) and platencin (**44**), stereospecific hydroxylation through cryptic carbonylation is orchestrated by the redundant α KG-dependent oxygenase pair PtmO3/PtmO6 and SDR family enzymes PtmO8 and PtmO1, a pair of NAD⁺/NADPH-dependent dehydrogenases.¹⁶⁷ Targeted gene inactivation and *in vitro* enzyme assays revealed that the C-7 oxidation and epimerization are initiated by the Fe/ α KG-dependent oxygenases PtmO3/PtmO6 that convert **47** and **94** to **95** and **96**, respectively, that each harbor a C-7 β hydroxy group (Fig. 15A).¹⁶⁷ The C-7 β hydroxy groups of **95** and **96** are oxidized by the SDR PtmO8 to the C-7 carbonyls **97** and **98**. Subsequently, **97** and **98** are reduced by PtmO1 to afford the C-7 α hydroxy **99** and **100**, respectively, resulting in a net epimerization of the C-7 hydroxy group.¹⁶⁷ The discovery of a three-enzyme cascade responsible for stereospecific hydroxylation, oxidation, and reduction at C-7 explains the origin of the C-7 oxidation patterns commonly found in *ent*-kaurane and *ent*-atisane-derived diterpenoids.¹⁶⁷

3.2.4.2. Gibberellins. The characteristic carbon backbone of the gibberellins is derived from an *ent*-kaurane scaffold through contraction of the B-ring (*vide infra*). In the bacterial gibberellin biosynthetic pathway, the P450 CYP114 from *Sinorhizobium fredii* NGR234 converts **56** to GA₁₂-aldehyde (**101**) along with a low amount of GA₁₂ (**102**) (Fig. 15B).⁴⁸ Heterologous expression and gene inactivation experiments have demonstrated that the SDR_{GA} efficiently catalyzes the conversion of the aldehyde



101 to the carboxylic acid **102** (Fig. 15B).⁴⁸ The unique tandem enzymatic conversion in bacterial gibberellin biosynthesis differs from that in plants and fungi, where P450s catalyze both the ring contraction and the complete conversion of the extruded C-7 methylene into a carboxylic acid moiety.⁴⁸

3.3. Functionalization by transferases

Transferase-encoding genes are mostly found in BGCs associated with highly functionalized terpenoids where they further decorate terpene scaffolds by installing functional groups including methyls, sugars, and aromatic compounds onto functional groups installed during the decoration phase. In most cases, these BGCs are relatively large (>10 kb) because the genes encoding enzymes for the biosynthesis of the building blocks to be transferred typically cluster with genes encoding terpene synthases, transferases, and oxygenases that install the handles for building block attachment.

3.3.1. Phenalinolactone A. Perhydrophenanthrene diterpenoids receive several peripheral decorations of the scaffold catalyzed by a series of transferases through their biosynthetic pathways. In phenalinolactone A (**28**), two MTs PlaP5 and PlaM1 are proposed to catalyze *C*-methylation of pyrrole carboxylic acid and *O*-methylation of L-amicetose, respectively.¹¹⁸ Then, these building blocks are transferred onto the diterpene scaffold by the AT PlaP2 and the GT PlaA6, respectively (Fig. 16).¹¹⁸

3.3.2. Brasilicardin A. The tailoring steps *en route* to brasiliocardin A (**1**) include the methoxylation of OH-16 by the MT Bra11, incorporation of L-rhamnose by the GT Bra10, and transfer of 3-hydroxybenzoic acid by the AT Bra8 (Fig. 16).^{124,125} The transferase responsible for the incorporation of *N*-

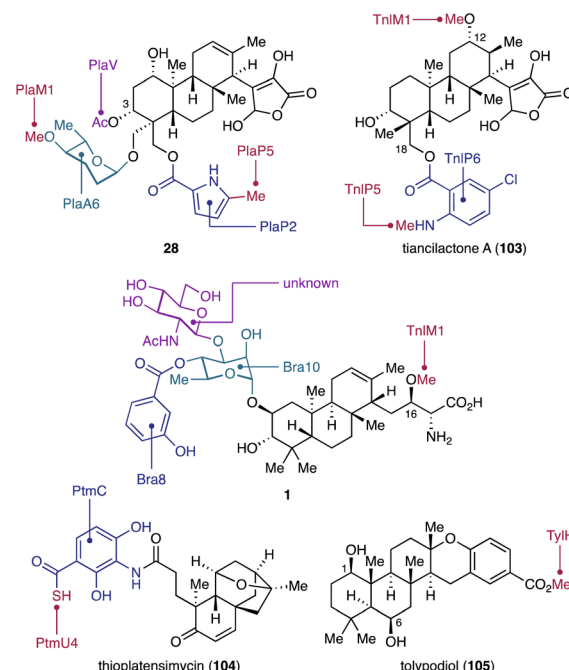


Fig. 16 Selected bacterial terpenoids that are modified by the incorporation of non-terpenyl building blocks.

acetylglucosamine remains unknown because of the absence of candidate genes in the *bra* BGC.^{124,125}

3.3.3. Tiancillactone A. Tiancillactones, analogs of phenalinolactones, have been identified by mining the genome of *Streptomyces* spp. CB03234 and CB03238 strains.¹⁶⁸ Compared to **28**, the carbon scaffold of tiancillactone A (**103**) is decorated by a chloroanthralinate moiety at C-18 and a methoxy moiety at C-12. The *anti-anti-syn*-fused perhydrophenanthrene scaffold of **28** is generated by the type II TC TnlT2 that does not harbor the catalytic DXDD motif typically conserved in type II TCs.¹⁶⁸ Chloroanthralinic acid is biosynthesized by the tryptophan halogenase TnlP3, tryptophan dioxygenase TnlP2, and two endogenous enzymes, a formamidase and a kynureinase. The resulting building block is activated by the adenosine triphosphate-dependent synthase TnlP4 and then transferred onto the peptidyl carrier protein TnlP1. The hydrolase TnlP6 is proposed to incorporate chloroanthralinate at OH-18.¹⁶⁸ Gene inactivation experiments indicate that the *N*-methylation of chlorine anthralinic acid is catalyzed by the MT TnlP5, although its timing in the tiancillactone biosynthetic pathways is unknown.¹⁶⁸ In the last step *en route* to **103**, the MT TnlM1 installs a methyl group at OH-12 (Fig. 16).¹⁶⁸

3.3.4. Thioplatensimycin. Thioplatensimycin (**104**) is a benzenecarbothioic acid-bearing intermediate of **2** (Fig. 16), which has been isolated from *S. platensis* SB12029.¹⁶⁹ For the biosynthesis of benzenecarbothioic acid as a building block, 3-amino-4-dihydroxybenzoic acid (AHBA) is initially synthesized from aspartate 4-semialdehyde and dihydroxyacetone phosphate by the 2-amino-4,5-dihydroxy-6-oxo-7-(phosphonoxy) heptanoate synthase PtmB1 and the AHBA synthase PtmB2.¹⁷⁰ AHBA is subsequently hydroxylated by the FMO PtmB3 to yield

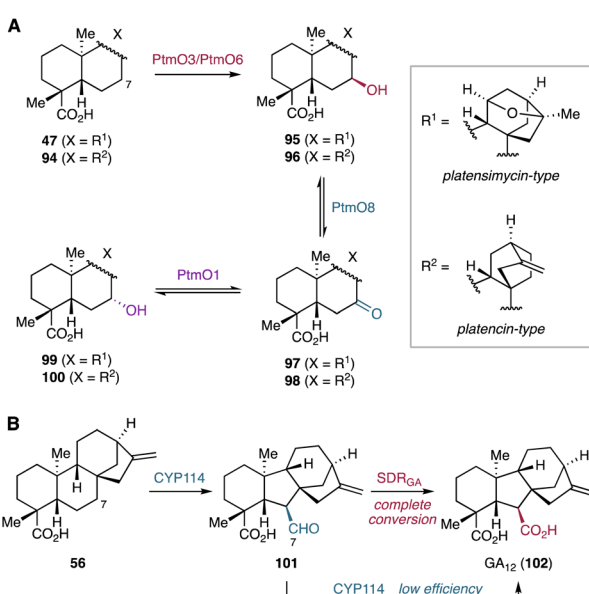


Fig. 15 B-ring functionalization of *ent*-kaurane and *ent*-atisane diterpenoids by oxidoreductases. (A) Cryptic carbonylation and hydroxy epimerization in the platensimycin and platencin biosynthetic pathways. (B) SDR-assisted conversion of an aldehyde into a carboxylic acid moiety during bacterial gibberellin biosynthesis.



3-amino-2,4-dihydroxybenzoic acid (ADHBA).¹⁷⁰ Gene inactivation and *in vitro* enzymatic studies verified that 3-amino-2,4-dihydroxythiobenzoic acid (ADHBSH) is synthesized from ADHBA by the sulfur transfer protein PtmU4.¹⁷¹ The N-AT PtmC catalyzes condensation of ADHBSH with platenicyl-CoA, a C₁₇ CoA thioester derived from the 4,5-*seco*-*ent*-kaurane diterpenoid, resulting in the production of **104**. The thiocarboxylic acid residue in **104** is non-enzymatically hydrolyzed to the carboxylic acid moiety in **2**.^{171–173} PtmC exhibits a broad substrate flexibility for benzoic acid-type building blocks.¹⁷² Inspired by the promiscuity of PtmC, mutasynthesis of platenimycin variants has been performed using a series of aminobenzoic acids.¹⁷² Furthermore, modeling studies and sequence similarity network analysis of arylamine N-ATs suggested that PtmC is an unusual arylamine N-AT that has likely evolved from xenobiotic N-ATs to accommodate the structurally bulky platenicyl-CoA.¹⁷³

3.3.5. Tolypodiol. Tolypodiol (**105**) is an anti-inflammatory meroditerpenoid isolated from the cyanobacterium *Brasilonema* sp. HT-58-2.¹⁷⁴ Based on retrobiosynthetic analysis and BLAST searches of chorismate lyase homologs that might be involved in the biosynthesis of the *p*-hydroxybenzoic acid component, the putative tolypodiol BGC (*tyl*) has been identified and subsequently verified through heterologous expression of the *tyl* genes in *Anabaena* sp. UTEX 257.¹⁷⁵ The *tyl* BGC includes a gene encoding the non-canonical Pyr4-type transmembrane TC (*tylF*).¹⁷⁵ For the decoration of the pentacyclic scaffold derived from GGPP and *p*-hydroxybenzoic acid, the P450 TylJ and the Rieske oxygenase TylG were proposed to install hydroxy groups at C-1 and C-6.¹⁷⁵ *In vitro* enzymatic assays revealed that the MT TylH catalyzes a methylation at the aryl carboxylic acid group of desmethyltolypodiol to form **105**.¹⁷⁵

3.4. Modification of terpene precursors

Modifications introduced by tailoring enzymes do not always occur after cyclization of the oligoprenyl precursors. In some cases, MTs and P450s act as modifying enzymes involved in either the assembly of unusual oligoprenyl pyrophosphates or the construction of novel hydrocarbon backbones through cyclization reactions (see Chapters 4.1.5 and 4.2.7).^{93,176–179}

3.4.1. 2-Methylisoborneol. Several actinomycetes, cyanobacteria, and myxobacteria produce the musty or earthly smelling homoterpene 2-methylisoborneol (**106**).^{180–184} The structure of **106** resembles that of the plant terpenoid camphor, but the presence of an additional methyl group at C-2 does not follow the isoprene rule. Labeling studies showed that the C-2 methyl originates from SAM that is incorporated into GPP.^{182,185} A candidate BGC only constituting TC and MT encoding genes was identified by genome mining and verified to be responsible for the production of **106** by heterologous expression.¹⁷⁶ Based on *in vitro* studies, the MT SCO7701 from *S. coelicolor* A3(2) has been shown to convert GPP into (2*E*)-2-methyl GPP (**107**) that is accepted as a substrate by the BGC-encoded TC (Fig. 17A).¹⁷⁷

3.4.2. Benzastatins. In contrast to SCO7701 that installs the methyl at C-2 of GPP, the MT BezA found in the biosynthetic pathways of benzastatins catalyzes methylation at the C-6

position of GPP (Fig. 17A).¹⁷⁹ BezA is substrate-specific and produces 6-methyl GPP (**108**), which is further oxidized by the P450 BezC. The resulting 10-hydroxy-6-methyl GPP (**109**) is transferred onto *p*-aminobenzoic acid by the PT BezF and further functionalized in downstream tailoring reactions to form benzastatins (see Chapter 4.1.4).¹⁷⁹

3.4.3. KS-505a. KS-505a (**110**),¹⁸⁶ also known as longestin,¹⁸⁷ from *Streptomyces argenteolus* A-2 has been identified as a cyclic nucleoside phosphodiesterase inhibitor.^{188–190} This unusual merotetraterpenoid features an unusual methyl branched octacyclic carbon skeleton fused with a benzoic acid moiety. Feeding experiments using isotope-labelled SAM showed that the branched methyls at C-1 and C-12 originate from SAM.¹⁸⁷ The corresponding *lon* BGC harbors three genes encoding MTs,¹⁹¹ one of which, *lon23*, is homologous to the GGPP MT-

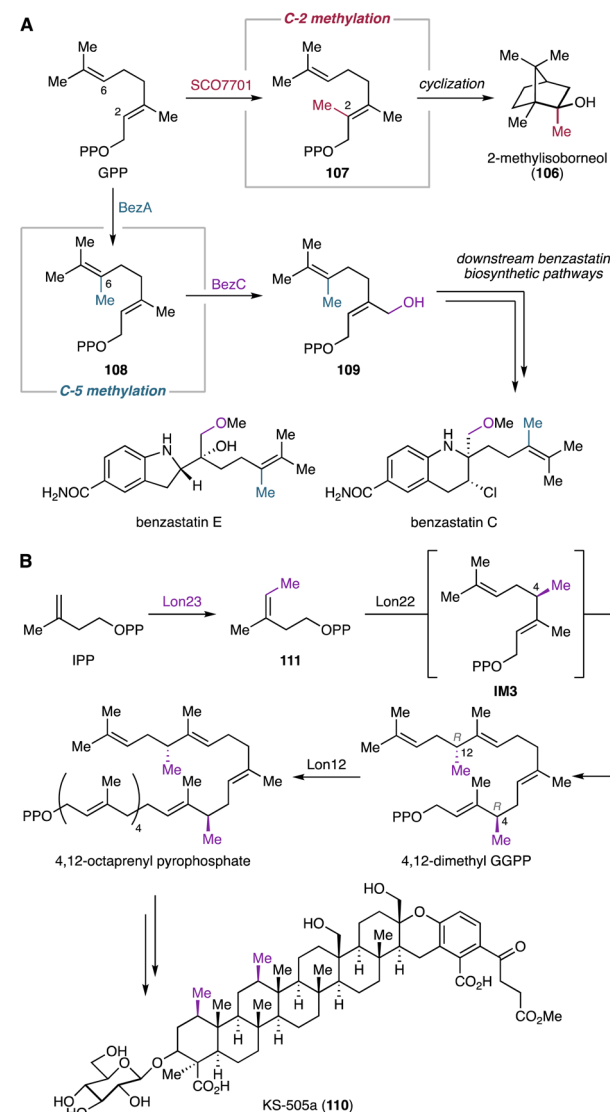


Fig. 17 Early-stage modification of oligoprenyl pyrophosphates by MTs before the formation of cyclic terpene hydrocarbon backbones. (A) Biosynthetic models of 2-methylisoborneol and benzastatins. (B) Biosynthetic pathway of KS-505a.



encoding gene *SCO7701* (38% id) from *S. coelicolor* A3(2).¹⁷⁸ *In vitro* enzymatic studies showed Lon23 to be a (3*Z*)-3-methyl IPP (**111**) synthase that utilizes IPP as a substrate (Fig. 17B). The unusual prenyl building block **111** is utilized for the subsequent stereospecific condensation reaction with IPP catalyzed by the 4,12-dimethyl GGPP synthase Lon22. This reaction proceeds through the formation of 4-methyl GPP (**IM3**) and the subsequent head-to-tail condensation of two **IM3** molecules to yield 4,12-dimethyl GGPP. The resulting 4,12-dimethyl GGPP is further elongated by the 4,12-dimethyl octaprenyl pyrophosphate synthase Lon12.¹⁷⁸ Notably, these polyprenyl pyrophosphate synthases are also capable of synthesizing desmethyl or monomethyl polyprenyl pyrophosphates.^{178,191}

4. Modification of terpene hydrocarbon scaffolds by non-canonical tailoring enzymes in bacteria

4.1. Non-canonical modifications by P450s

4.1.1. Oxidative methyl migration. Methyl migration is an important rearrangement reaction involved in terpene hydrocarbon backbone construction. TC-catalyzed carbocation-driven cascade reactions frequently proceed through non-oxidative Wagner–Meerwein-type 1,2-methyl shifts *en route* to diverse terpene scaffolds.²⁰ In contrast, oxidative methyl migrations at later stages are rare.

4.1.1.1. Pentalenolactone. Biosynthetic studies on pentalenolactone (**3**) revealed an oxidative rearrangement from **75** to **3** catalyzed by the P450 PntM (PenM).^{192,193} Typically, neopentyl radicals generated through P450 catalysis do not cause skeletal rearrangements.¹⁹⁴ Given the structures of shunt products biosynthesized in trace amounts from pentalenolactone-producing *Streptomyces* spp.,^{195,196} the PntM-catalyzed oxidative methyl migration was proposed to proceed through the formation of a neopentyl cation (Fig. 18).¹⁹² Specifically, the neopentyl radical **IM4** generated by H-1 *Si* hydrogen abstraction undergoes a rapid electron transfer from **IM4** to the highly oxidizing heme

Fe^{III}–OH radical cation to produce the neopentyl cation **IM5**. This cation facilitates the *syn*-1,2-methyl shift and the antarafacial deprotonation of H-3*α*, resulting in the formation of the final product **3** that features the *vic*-dimethylcyclopentene ring (Fig. 18).¹⁹²

X-ray structure analyses of the wild-type and single amino acid mutated PntM variants imply that a complex network of noncovalent interactions tightly regulates the positioning of both substrates and products so that the *Si* face of C-1 is located close to the heme iron species.¹⁹³ The appropriate orientation of **75** is further retained by steric hindrance caused by the C-2 methyl and C-7 olefin adjacent to the C-1 *Si* face. The typical oxygen rebound of P450s requires the rapid rotation of the Fe-bound OH group formed after hydrogen abstraction, and the rate constant for oxygen rebound is $>10^{10}$ – 10^{11} s⁻¹.^{197–199} In contrast, the thermodynamically unfavorable electron transfer to the heme Fe^{III}–OH radical species has a significantly slower rate ($<10^8$ – 10^9 s⁻¹).²⁰⁰ However, in the case of a radical center with β-branched alkyl groups such as the neopentyl radical, oxygen rebound is extremely slow because of the steric hindrance and thus the electron transfer is compatible with oxygen rebound.^{200,201} The naturally occurring steric barrier in **75** due to the presence of the C-2 *gem*-dimethyls and H-7 olefinic proton makes the kinetically insignificant electron transfer favorable, resulting in the dominant production of **3**.¹⁹³ In addition, quantum mechanical/molecular mechanics (QM/MM) calculations support that the electron transfer process for the formation of the C-1 cation **IM5** from **IM4** is favorable compared to the oxygen rebound mechanism because the overlap between the π^* orbital of Fe–OH and the σ_z^2 orbital of the C-1 radical in **IM4** is blocked by the presence of H-2.²⁰² The migration reaction is likely terminated by a H-3*α* abstraction, wherein a water molecule that shows hydrogen bonding with the carboxylic acid of porphyrin acts as Brønsted base.^{193,202}

4.1.2. Semipinacol rearrangement. Pinacol rearrangements are acid-induced rearrangement reactions of *vic*-diols that form ketones or aldehydes by 1,2-alkyl migrations.²⁰³ In analogy, semipinacol rearrangements are reactions related to pinacol rearrangements that mechanistically share the electrophilic reactive species at a vicinal position of an oxygenated carbon and the 1,2-migration to terminate the reaction.²⁰⁴ These pinacol and semipinacol rearrangements are historically thought to be key modifications to facilitate late-stage ring contractions, ring expansions, and spirocyclization reactions *en route* to highly modified terpenoids.^{205–208} Semipinacol-type rearrangements have been proposed in multiple biosynthetic models but the corresponding biosynthetic enzymes have not been characterized. Inspired by the rearrangement reactions in nature, several biomimetic total syntheses of complex terpenoids have been elegantly achieved employing (semi)pinacol rearrangements.^{208–211}

4.1.2.1. Gibberellins. A prominent example of a semipinacol rearrangement can be found in the biosynthesis of gibberellins. Its characteristic 6/5/6/5-tetracyclic carbon framework is derived from the contraction of the B-ring of the *ent*-kaurene-type hydrocarbon skeleton.²¹² Isotope-labeling studies of fungal and plant gibberellin biosynthesis revealed that the

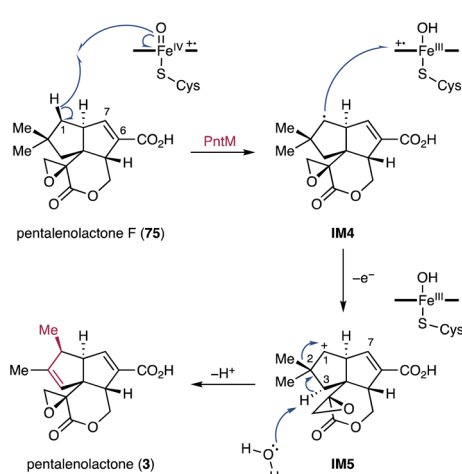


Fig. 18 PntM-catalyzed oxidative methyl migration in the pentalenolactone biosynthetic pathway.



semipinacol rearrangement-based B-ring contraction proceeds through the abstraction of H-6 β in *ent*-7 β -hydroxykaurenoic acid (112), forming the radical intermediate IM6.²¹³ A subsequent single electron transfer from IM6 to the hydroxoiron(IV) species generates the secondary cation IM7.²¹³ This intermediate undergoes an 1,2-alkyl migration onto the C-6 carbocation, ultimately leading to the formation of GA₁₂-aldehyde (101) (Fig. 19).²¹³⁻²¹⁷ The P450s responsible for this rearrangement reaction have been identified from both plants and fungi.⁴² Furthermore, recent studies on the bacterial gibberellin biosynthetic pathways provided insights into the enzymatic structure–function relationships underlying the semipinacol rearrangement-based ring contraction reaction.²¹⁸

Functional characterization of biosynthetic enzymes encoded in the gibberellin BGC in *S. fredii* revealed that CYP114 not only catalyzes the hydroxylation at C-7 of *ent*-kaurenoic acid (56), but also the B-ring contraction of 56 to produce 101 and trace amounts of GA₁₂ (102).⁴⁸ The ring contraction reaction catalyzed by CYP114 requires the redox partner Fd_{GA} which is encoded in the gibberellin BGC.⁴⁸ In the absence of Fd_{GA}, the enzymatic activity of CYP114 is limited to C-7 β hydroxylation with the help of an endogenous Fd from the heterologous host.⁴⁸ The relationships between CYP114 and Fd_{GA} are unique compared to their counterparts from plants and fungi, where the P450s do not require a dedicated redox partner and rather use ubiquitous cytochrome P450 reductases for their ring contraction activity.²¹⁹⁻²²¹ Furthermore, the enzymatic capability of plant- and fungi-derived P450s (CYP88A/CYP68A) encompass the full oxidation of the extruded C-7 into a carboxylic acid moiety. The fungal P450, CYP68A, exhibits an additional hydroxylation activity at C-3 β of 102 to yield GA₁₄ (113) (Fig. 19).⁴² Nevertheless, the substrate and the unique rearrangement reaction of the P450s are conserved beyond kingdom borders. In light of the structural rearrangements observed in plant and fungal gibberellin biosynthetic pathways, the contraction reaction of the B-ring was hypothesized to be a semipinacol rearrangement.⁴⁸ Feeding experiments of *ent*-kaurenal and *ent*-kaurenoic acid methyl ester revealed that the

C-19 carboxylic acid anchimerically assists in stabilizing the C-6 carbocation of IM7 that is presumably generated during the semipinacol-type rearrangement.⁴⁸ Further isotope labeling studies supported the CYP114-catalyzed semipinacol rearrangement extruding C-7 from the B-ring of 112 *via* H-6 β abstraction and 1,2-alkyl migration (Fig. 19).²²²

Recently, the reaction mechanism of the CYP114-catalyzed ring contraction has been elucidated.²¹⁸ *In vitro* enzymatic assays and *in vivo* feeding experiments indicated that *EtCYP114* from *Erwinia tracheiphila* converts 56 to 101 along with 112 and 102 in the presence of the redox partner *EtFd_{GA}* from *E. tracheiphila* or *BjFd_{GA}* from *Bradyrhizobium japonicum*. However, *EtCYP114* loses its ability to contract the B-ring when incubated with spinach Fd, which results exclusively in the formation of 112. While *EtCYP114* is able to accommodate both 56 and 112, its ring contraction activity is limited to using 56 as a substrate.^{47,218,222} Structural analysis of *EtCYP114* bound to 56 showed the positioning of H-7 β (3.7 Å) and H-6 β (3.9 Å) in close proximity to the heme iron. The crystal structure of *EtCYP114* shows a notable lack of ionic interactions at the C-18 carboxylic acid, which provides the anchimeric effect for the C-6 carbocation of IM7. This interaction strongly supports the carbocation-mediated semipinacol rearrangement reaction (Fig. 19), although an unusual radical mediated rearrangement cannot be ruled out.²¹⁸ Furthermore, *EtCYP114* lacks the highly conserved acid–alcohol motif (e.g., Asp–Thr) that facilitates sequential protonation of the heme iron species in P450s.^{218,223} When the acidic amino acid residue is introduced into *EtCYP114*, the enzyme completely loses its catalytic ability to contract the ring.²¹⁸ These results along with the *apo* structure of *EtCYP114*:A261D suggested that complementation of the acid–alcohol motif in *EtCYP114* may alter the hydrogen-bonding network within the active site, potentially disrupting its interaction with the dedicated redox partner Fd_{GA}.²¹⁸ Consequently, the ring contraction reaction catalyzed by CYP114 requires both the characteristic redox partner and the absence of the acidic amino acid residue typically conserved in other P450s.^{48,218}

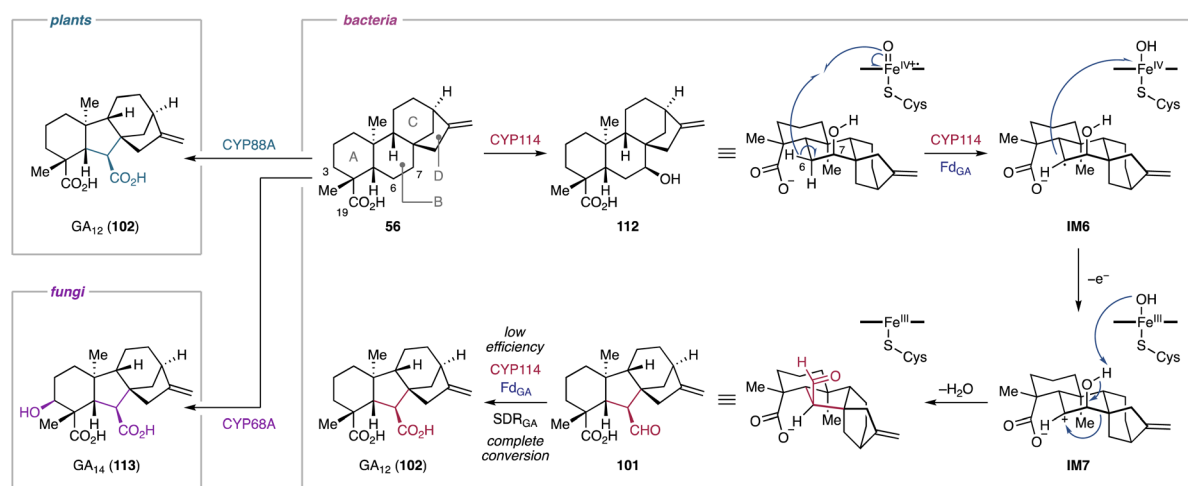


Fig. 19 Semipinacol rearrangement-based ring contraction in the gibberellin biosynthetic pathways in bacteria, plants, and fungi.



4.1.3. Lactonization. Lactone moieties in terpenoids frequently play a pivotal role in the biological activities of the corresponding products, as demonstrated by plant and fungal lactone-containing terpenoids, *e.g.*, artemisinin and mycophenolic acid.^{224,225} Despite its importance, lactone formation in terpenoid biosynthesis remains largely uncharacterized, with only a handful of biochemically validated examples.^{150,226–229} Most cases involve BVMO-catalyzed ring expansion reactions.

4.1.3.1. Gibberellins. Many gibberellin family norditerpenoids possess a γ -lactone bridge at the A-ring.²¹² The formation of the lactone bridge with the concurrent loss of C-20 in **102** or **113** is catalyzed by the Fe/ α KG-dependent oxygenase GA20ox in plants or the P450 CYP68B in fungi.⁴² In a similar manner, CYP112 in bacteria is capable of converting **102** into GA₉ (**9**).⁴⁸

In vitro enzymatic assays and ¹⁸O₂ labeling studies were conducted to gain insights into the CYP112-catalyzed C-20 dissociation and γ -lactone formation.²³⁰ *Et*CYP112 converts **102** into **9** by decarboxylation of C-20 in the presence of spinach Fd, *E. tracheiphila* derived *Et*FdR, and NADPH. The mixed origin of the γ -lactone oxygen from molecular oxygen and the C-18 carboxylic acid indicates that CYP112 is a multi-functional enzyme that catalyzes demethylation and γ -lactone formation *via* two proposed diverging routes (Fig. 20).²³⁰ Initially, CYP112 hydroxylates the terminal methyl C-20 of **102** to form 20-hydroxy-GA₁₂ (**114**), followed by dehydration to yield the δ -lactone-harboring GA₁₅ (**115**). Subsequently, **114** and **115** each follow two sequential oxidation reactions along separate routes (routes A and B) (Fig. 20). In route A, CYP112 first converts **114** into the *gem*-diol form (**116**) of GA₂₄ through the

oxygen rebound mechanism. Furthermore, H-20 of **116** is abstracted by Compound I of CYP112, generating the radical intermediate **IM8**. Compound II then abstracts the hydrogen of the C-18 carboxylic acid in **IM8**. The resulting diradical undergoes intramolecular radical coupling, yielding the *gem*-diol cyclic anhydride **117** of GA₂₅. In route B, **115** undergoes two rounds of hydroxylation at C-20 to form **117** through the formation of the lactol form (**118**) of GA₂₄. The key intermediate **117**, convergently produced from both proposed routes, is further oxidized by CYP112 to obtain **9**. Specifically, the hydrogens of the C-20 *gem*-diol are sequentially abstracted by the heme iron species, which induces a radical-based rearrangement of **IM9** to form the carbonate diradical **IM10**. Subsequent intramolecular radical coupling results in the formation of **9** through decarboxylation at C-20 (Fig. 20).²³⁰ The unusual diradical-mediated C-C bond scission and lactonization in bacterial gibberellin biosynthesis are remarkable for their similarity to the corresponding reactions in plant and fungal-derived gibberellin biosynthetic pathways.^{213,231} Despite the involvement of evolutionarily independent oxygenase families (P450s *vs.* α KG-dependent oxygenases) or P450 classes (cytosolic *vs.* membrane-bound), these enzymes showcase that tailoring enzymes for the biosynthesis of terpenoids have converged to utilize similar mechanistic routes across different branches of life.²³⁰

4.1.4. Nitrene transfer-mediated heterocyclization. In the archetypical catalytic cycle of P450s (Chapter 3.1), Compound I (Fe oxene) serves as the key oxidizing intermediate for oxene transfer reactions.²³² Inspired by the structural similarities

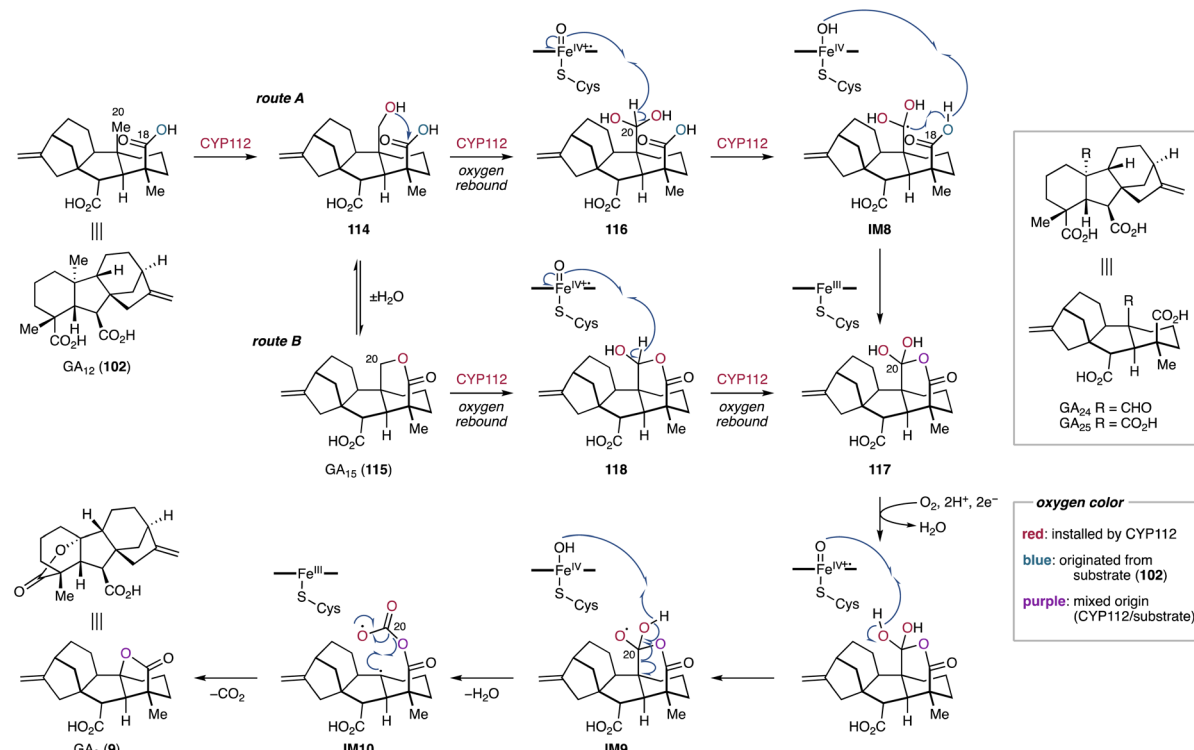


Fig. 20 CYP112-catalyzed C-20 elimination and γ -lactone formation in the biosynthetic pathway of bacterial gibberellins.



between Fe oxene, Fe carbene, and Fe nitrene species, extensive studies have focused on protein engineering of native cytochrome P450s, such as P450_{BM3} from *Bacillus megaterium*, to harness engineered variants for carbene and nitrene transfer reactions, including cyclopropanation, aziridination, and C–H amination.²³³ However, so far, naturally occurring P450-catalyzed carbene or nitrene transfer reactions have only been observed in one case, *i.e.*, in the biosynthesis of benzastatins.¹⁷⁹

4.1.4.1. Benzastatins. Benzastatins are *p*-aminobenzoic acid-derived meroterpenoids with neural cell protective and antiviral activities.^{234–237} Most of benzastatins bear tetrahydroquinoline or indoline scaffolds which result from the cyclization of a geranyl moiety. The benzastatin BGC and its homologs have been identified in the genomes of *Streptomyces* sp. RI18 and several other actinobacteria.¹⁷⁹ Heterologous expression of *bez* genes in *Streptomyces lividans* and *in vitro* enzymatic assays have led to the characterization of the benzastatin biosynthetic pathway.¹⁷⁹ The pathway is initiated by the *N*-hydroxylase *BezJ* and the AT *BezG*, which sequentially catalyze *N*-hydroxylation and *O*-acetylation of *p*-aminobenzoic acid (119), respectively, to form *p*-*N*-acetoxybenzoic acid (120). The PT *BezF* then incorporates modified GPPs 108 or 109 into 120, resulting in the formation of geranyl *p*-*N*-acetoxybenzoic acid (121), a *bona fide* intermediate of the cyclized benzastatins.¹⁷⁹ The highly unstable 121 is nonenzymatically converted into the indoline 122 and the tetrahydroquinoline 123 with low efficiency, as well as the dihydrobenzo[*b*]azepine shunt product 124 (Fig. 21).¹⁷⁹ The production of 122–124 likely involves the nonenzymatic generation of the arylnitrenium ion IM11 through hydrolysis of the *N*-acetoxy moiety. This highly electrophilic ion reacts with the C-9/C-10 olefin to generate an azirinium ion, which is hydrolyzed to yield 122 and 123. Alternatively, the arylnitrenium ion IM11

reacts with the C-17 methyl of the geranyl moiety to form 124.¹⁷⁹ More importantly, the P450 *BezE* has been identified as the heterocyclization catalyst that mediates the formation of the benzastatins (122, 125). During the cyclization reaction, *BezE* does not require a redox partner for its catalytic cycle, indicating a nonoxidative reaction mechanism.¹⁷⁹

Mechanistically, the *BezE*-catalyzed nonoxidative heterocyclization reaction is proposed to employ a nitrene transfer mechanism (Fig. 21).¹⁷⁹ In case of aziridination catalyzed by P450 variants engineered from P450_{BM3}, the nitrene transfer begins with the acquisition of N₂ from an azide-containing compound to form an iron nitrenoid intermediate, which further reacts with olefins to yield aziridines.²³⁸ Instead of using an azide to form the iron nitrenoid,²³⁸ *BezE* utilizes the *N*-acetoxy group as nitrene source.^{179,239} The heme-iron nitrenoid (IM12) of *BezE* reacts with an olefin of the geranyl moiety through nitrene transfer, resulting in aziridine formation (IM13, IM14). Subsequently, the highly strained aziridine ring is opened by nucleophilic attack of a hydroxide or a chloride to form indoline 122 or tetrahydroquinoline 125, respectively.¹⁷⁹ The chlorine in 125 is nonenzymatically substituted to afford 123 and its C-9 diastereomer. The higher yield of 123 compared to its diastereomer suggested that the nonenzymatic conversion proceeds *via* an S_N1 reaction rather than an S_N2 reaction.¹⁷⁹ This preference is likely due to the steric hindrance of the dimethylpentenyl moiety at C-10.¹⁷⁹ *BezE* is the first example of a native P450 that catalyzes nitrene transfer. The *N*-acetoxylation likely facilitates bypassing of the conventional P450 catalytic cycle, which typically involves oxene transfer.²⁴⁰ This unusual catalytic mechanism enables *BezE* to act without the need for electron supplementation from redox partners.¹⁷⁹ Consequently, *BezE* is classified as a P450 nitrene transferase.¹⁷⁹

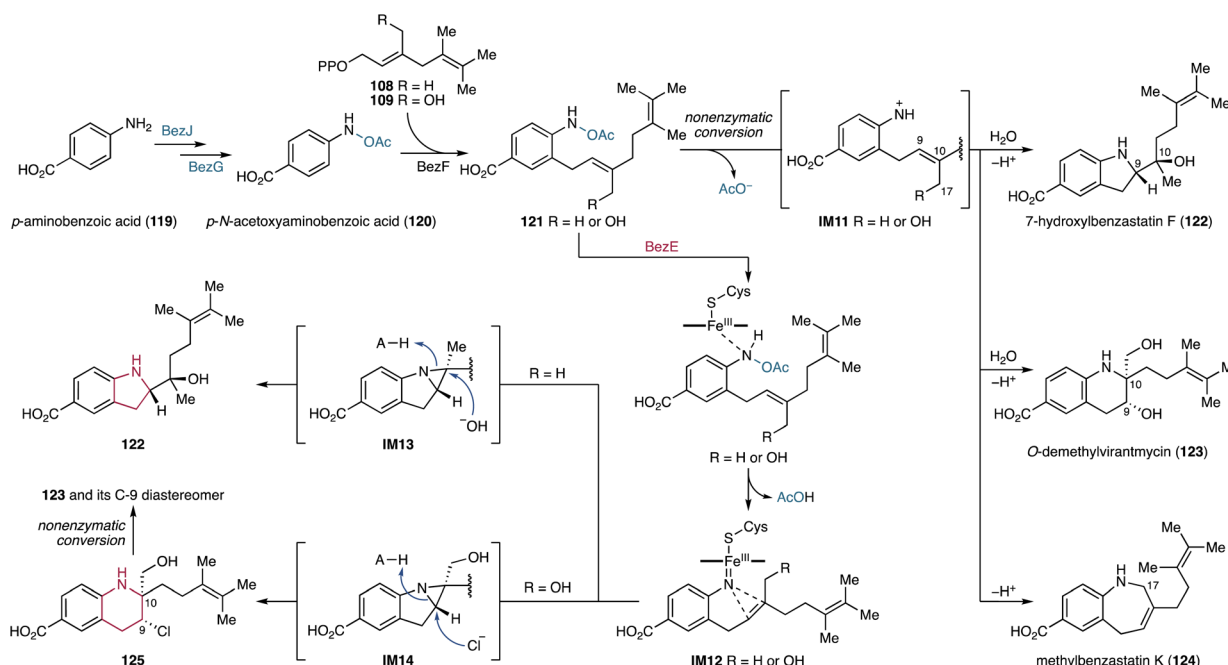


Fig. 21 Nonoxidative heterocyclization reaction catalyzed by the P450 nitrene transferase *BezF* in the benzastatin biosynthetic pathway.



4.1.5. Oxidative C–C bond formation. Some P450s favor the formation of new bonds by radical addition to π -bonds, radical coupling, and cationic rearrangements over the oxygen rebound mechanism.¹⁰⁹ In the case of (mero)terpenoids, C–C bonds formed by P450s are found in the late stage of their biosynthetic pathways as demonstrated in the fungal meroterpenoid viridicatumtoxin²⁴¹ and plant lathyrane-type diterpenoid biosynthesis.²⁴² However, there is only a single example of a characterized P450 involved in the construction of a terpene hydrocarbon scaffold by C–C bond formation in bacteria.⁹³

4.1.5.1. Aridacins. Aridacins A–C (24, 68, 69) are *cis*-eunicellane diterpenoids that feature a densely oxidized 6/7/5-tricyclic scaffold.⁹³ Heterologous expression of the *ari* BGC-encoding genes in *S. lividans* TK64 and an GGPP-overproducing *E. coli* strain and *in vitro* enzymatic experiments revealed that the P450 AriF is capable of constructing a 6/7/5 tricyclic scaffold 67 from a 6/10-bicyclic framework 126. However, an earlier report showed that heterologous expression of the *ari* BGC in *S. albus* J1074M only produced the 6/10-bicyclic eunicellane 127 decorated with a C-19 hydroxy group and a C-20 carboxylic acid.¹¹² Further feeding experiments using *S. lividans* and *S. albus* and *in vitro* assays showed that AriF oxidizes 126 to produce 6/10-bicyclic eunicellanes 127–130 as shunt products (Fig. 22). The production of the shunt metabolites 127–130 in low titers might originate from the limited activity of AriF, potentially due to the use of incompatible redox partners.⁹³

The reaction mechanism of the AriF-catalyzed C–C bond formation has been proposed to proceed through either a radical rearrangement or a carbocation-based reaction.

Density functional theory (DFT) calculations suggested that the terpenyl radical species IM15 generated by H-20 abstraction can undergo a barrierless and exothermic single electron transfer with a concomitant carbocation-based cyclization reaction (Fig. 22).⁹³ In contrast, the radical-mediated rearrangement requires a *ca.* 15 kcal mol^{−1} barrier to reach the transition state toward the tricyclic ring system in the doublet state of the heme iron species. The thermodynamically favorable electron transfer is likely due to the coupled reaction with the energetically barrierless cyclization of the bicyclic carbocation IM16 to form the tricyclic carbocation IM17. Consequently, the AriF-catalyzed cyclization is most likely a carbocation-driven reaction, although the radical rearrangement remains a possibility albeit with some energetic challenges. Further DFT calculations revealed that the canonical oxygen rebound for the formation of the C-20 hydroxy eunicellane IM18 is energetically feasible with a barrier of *ca.* 4 kcal mol^{−1} (Fig. 22).⁹³ These calculations explain the formation of 127–130, which are produced *via* the AriF-catalyzed oxygen rebound route as shunt products.

4.2. Non-canonical modification by other oxidoreductases

4.2.1. Oxidative alkyl migration. While members of the P450 family, such as PntM in pentalenolactone biosynthesis (*vide supra*), play a role in oxidative alkyl (methyl) migration, this enzymatic activity is not limited to P450s.¹⁹³ In-depth biosynthetic studies of bacterial meroterpenoids, such as aurachins, napyradiomycins (Chapter 4.2.6.1), and merochlorins (Chapter 4.2.6.2), revealed that other enzyme families also catalyze oxidative 1,2-alkyl migrations albeit utilizing distinct reaction types such as α -ketol rearrangements.^{243–245}

4.2.1.1. Aurachins. Aurachins are sesquiterpenoid quinolines that have been isolated from the myxobacterium *Stigmatella aurantiaca* Sg a15.²⁴⁶ They inhibit mitochondrial respiration and photosynthesis.^{246,247} Structurally, aurachins are grouped into A- and C-types based on the location of a farnesyl moiety at the quinolone.²⁴⁸ Feeding experiments using ¹³C- and ¹⁸O-labeled anthranilic acid, ¹³C-acetate, and ¹⁸O₂ indicate that A-type aurachins are derived from C-type aurachins through farnesyl migration.²⁴⁹ Furthermore, the oxygen atoms at C-3 in aurachins A and B (131, 132) are derived from molecular oxygen. Mining the genome of *S. aurantiaca* Sg a15 revealed that the biosynthetic genes for aurachins are split into three genetic loci.^{248,250} Based on gene inactivation experiments and *in vitro* enzymatic assays using synthetic variants of aurachin C (133), the FMO AuaG catalyzes 1,2-farnesyl migration in 133 through either of two possible reaction routes (Fig. 23, routes A and B).^{243,244,248} In route A, AuaG initially epoxidizes the quinolone double bond of 133, yielding IM19. The subsequent deprotonation of N-OH induces epoxide ring opening and 1,2-farnesyl migration through an α -ketol rearrangement to give the highly unstable ketone oxime 134. Alternatively, the 1,2-farnesyl migration in route B involves tandem sigmatropic rearrangements. Following epoxidation and base-mediated ring opening, the intermediate IM20 undergoes a concerted retro-[2,3]-Wittig rearrangement to yield the *O*-nerolidyl quinoline N-oxide IM21. A following Claisen rearrangement results in the formation of

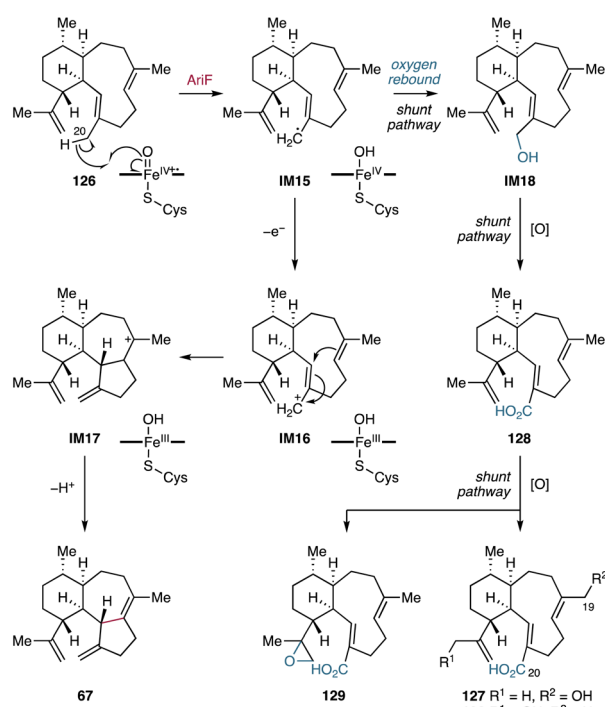


Fig. 22 AriF-mediated oxidative C–C bond formation in the aridacin biosynthetic pathway and canonical oxygen rebound-based shunt pathway.



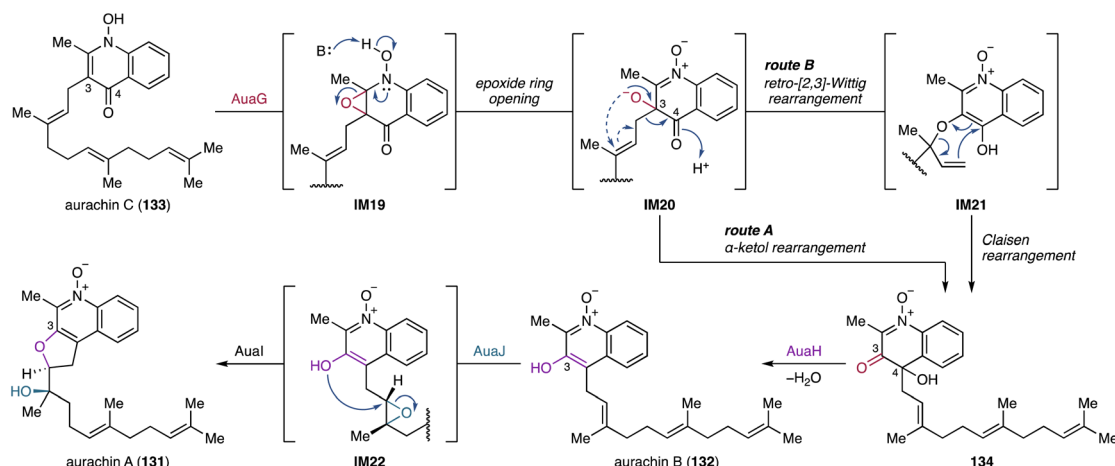


Fig. 23 FMO-catalyzed 1,2-farnesyl migration and heterocyclization in the late-stage aurachin biosynthetic pathways.

134. The NAD-dependent oxidoreductase AuaH then reduces the C-3 ketone and eliminates OH-4 to form 132. Bioinformatic analysis of *aua* genes and $^{18}\text{O}_2$ feeding experiments suggest that another FMO AuaJ likely epoxidizes an olefin proximal to the quinoline core in 132, resulting in the formation of IM22 (Fig. 23).^{248,249} This oxidation reaction follows epoxide hydrolysis and heterocyclization catalyzed by the epoxide hydrolase AuaI to afford 131.

4.2.2. Cyclization and skeletal rearrangement. Terpenoid indole alkaloids are meroterpenoids composed of an indole and a hemi-,²⁵¹ mono-,²⁵² sesqui-,²⁵³ or diterpene component.²⁵⁴ Most of these meroterpenoids exhibit remarkable and clinically important bioactivities as exemplified with the anticancer drug and drug lead, vinblastine and camptothecin, respectively.²⁵⁵ While terpenoid indole alkaloids have primarily been identified in plants and fungi,^{251–254} recent studies have demonstrated that bacteria are also capable of producing structurally complex sesquiterpenoid indole alkaloids, such as the xiamycins.^{256–260}

4.2.2.1. Xiamycins. Xiamycin A (135), sespenine (136), and their congeners have been isolated from *Streptomyces* spp. from various origins such as soil, mangrove trees, and sea sediments.^{256–260} Most of them exhibit antimicrobial and anti-viral activities.^{256–261} The *xia* BGCs have simultaneously been discovered by genome mining of *Streptomyces* sp. HKI0576 and *Streptomyces* sp. SCSIO 02999.^{262,263} Bioinformatic analyses, single gene disruption, and heterologous expression of the *xia* BGC demonstrated that two unparalleled cyclization steps construct the characteristic pentacyclic scaffolds that feature the carbazole or 2-azabicyclo[3.3.1]nonane in 135 and 136, respectively.^{262–264} The first cyclization forming a *trans*-decalin ring is catalyzed by XiaE, a homolog of the Pyr4-type trans-membrane TC.⁴¹ After a six electron oxidation catalyzed by the P450 XiaJ,²⁶⁵ the latter cyclization step forming the central ring of 135 and 136 is catalyzed by a new member of the group D FMOs, XiaF. Phylogenetic analysis of XiaF and its homologs revealed that XiaF evolved from xenobiotic degrading oxygenases and groups into the class of indigo-forming oxygenases.^{262,266} Based on feeding experiments and *in vitro*

enzymatic assays, XiaF converts indosespene (137) into the dihydrocarbazole-bearing prexiamycin (138) that spontaneously oxidizes to form 135.²⁶³ The cyclization activity of XiaF depends on flavin reduction catalyzed by a flavin reductase partner. In fact, the dedicated flavin reductase encoding gene *xiaP* is located in the flanking region of the *xia* BGC.²⁶⁶ Furthermore, XiaF also hydroxylates C-3 of the indole ring as a shunt reaction, leading to the autoxidation of the resulting hydroxyindole to form indigo. Based on the catalytic potential of XiaF, cyclization and rearrangement reactions for the formation of 135 and 136 have been proposed (Fig. 24A).^{262,263,266} Initially, the 3-hydroxy iminium species IM23 is formed from 137 *via* nucleophilic attack by the C-2/C-3 double bond to the hydroperoxide group of the activated FAD. This cation formation triggers ring closure through the addition of the *exo*-methylene (C-21 of IM23) to the iminium, yielding IM24. Subsequent dehydration and deprotonation lead to the formation of 138 that is readily converted into 135. For the construction of the 2-azabicyclo[3.3.1]nonane framework in 136, the carbocation intermediate IM24 produced through C-C bond formation between C-2 and C-21 undergoes phenyl migration. Based on the proposed rearrangement mechanism to form 136 and its stereochemical outcomes, the C-3 hydroxylation catalyzed by XiaF likely occurs from the *Si* face. Intramolecular addition of C-21 to the imine species follows to form the (2*R*,3*S*)-*cis*-fused dihydroindole moiety in IM24, which is the configuration required for the subsequent phenyl migration. However, the stereoselectivity of the XiaF-catalyzed hydroxylation has not yet been experimentally confirmed. The divergent pathways for the biosynthesis of 135 and 136 may be explained by the degree of conformational freedom in the rearrangement reaction of the carbocation species.²⁶²

X-ray crystallography analysis of XiaF's *apo* structure and docking studies with indole revealed that the regiospecific hydroxylation of indole is regulated by precisely coordinating the C-3 position close to the hydroperoxide group of the activated FAD.²⁶⁶ Broad-range xenobiotic oxygenases generally have a small active site cavity lined with bulky amino acid residues.²⁶⁷

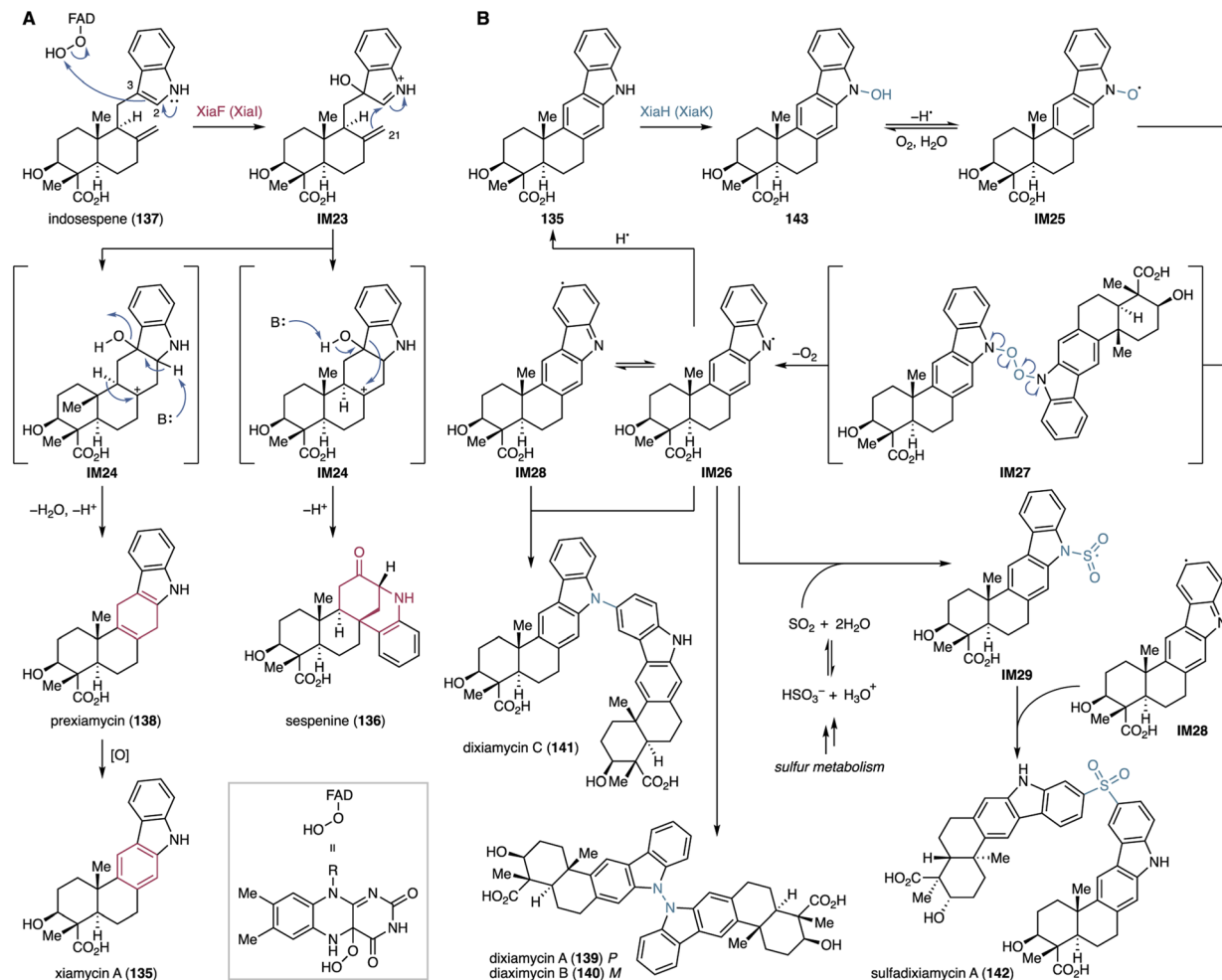


Fig. 24 Non-canonical modifications in the biosynthetic pathways of xiamycins and dixiamycins. (A) Proposed XiaF-mediated cyclization for the formation of xiamycin and sespenine. (B) Proposed dimerization reaction of the dixiamycins and sulfadixiamycins biosynthesis initiated by the XiaH-mediated *N*-hydroxylation.

In contrast, XiaF harbors a relatively large substrate binding channel to facilitate the acceptance of the bulky substrate **137**. Considering that XiaF is promiscuous enough to also hydroxylate indole but no other typical xenobiotic substrates, XiaF is a specialized tailoring enzyme that has evolved from xenobiotic detoxification enzymes to catalyze a non-canonical cyclization reaction of indolosesquiterpenoids.²⁶⁶

4.2.3. Radical-mediated dimerization. Structural diversification through dimerization is commonly observed in various types of natural products.²⁶⁸ These dimeric molecules often exhibit superior biological functions compared to their monomeric counterparts. Notable examples within the terpenoid class include the plant sesquiterpenoid dimer gossypol which has anti-fertility activity²⁶⁹ and the bisindolosesquiterpenoids dixiamycins which show antibiotic activity.²⁵⁹

4.2.3.1. Dixiamycins. Dixiamycins are dimers of xiamycin (**135**) fused via C–N, N–N, or sulfonyl linkages.^{259,262,270–272} Structurally, most C_2 or C_1 symmetric dixiamycins with aryl C–N or N–N bonds show atropoisomerism and both atropodiastereomers are found in nature.^{259,270,272}

Targeted gene deletion, heterologous expression, and feeding experiments revealed that the FMO XiaH is responsible for the production of bisindolosesquiterpenoids with aryl and sulfonyl bridges, *e.g.*, **139–142** (Fig. 24B).^{270,271} *In vitro* studies, however, show that XiaH only hydroxylates **135** to provide *N*-hydroxyxiamycin (**143**).²⁷³ NMR analysis and electron paramagnetic resonances of **143** revealed that the nitroxyl radical species **IM25** is also presented in solution. Furthermore, time-course biochemical experiments showed conversion of **143** into **135**, suggesting that the aminyl radical species **IM26** can be formed *via* homocoupling of **IM25**.²⁷³ Subsequent deoxygenation results in the formation of structurally diverse xiamycin monomer analogs.²⁷³

The formation of dimeric xiamycins may be explained by a radical-based coupling mechanism (Fig. 24B).^{270,271,273} XiaH initially *N*-hydroxylates **135** to yield **143**. The *N*-hydroxide **143** is in equilibrium with its nitroxyl radical species **IM25**. The radical dimerizes and the resulting endoperoxide **IM27** is deoxygenated to form the aminyl radical **IM26**. This intermediate readily undergoes radical migration on the π -conjugated carbazole



moiety (e.g., **IM28**). Afterwards, an intermolecular coupling reaction leads to the formation of axially chiral aryl C–N or N–N bond bearing dixiamycins **139–141**. Alternatively, the aminyl radical **IM26** reacts with sulfur dioxide to form the sulfonyl radical **IM29**, which leads to the formation of sulfadixiamycins, e.g., **142**.²⁷¹

4.2.4. Oxidative heterocyclization. As discussed in the section on non-canonical P450s (Chapter 4.1), the versatile utilization of iron species enables P450s to catalyze a wide range of unusual reactions. Similarly, Fe/αKG-dependent oxygenases employ iron in fascinating ways to catalyze transformations such as cyclizations, ring contractions, and ring expansions, particularly in the late stages of natural product biosynthesis.^{27,109} In bacterial terpenoid biosynthesis, a non-canonical Fe/αKG-dependent oxygenase has been identified to catalyze the oxidative heterocyclization of phenalinolactones.^{118,119}

4.2.4.1. Phenalinolactones. The *pla* BGC harbors a gene encoding a Fe/αKG-dependent oxygenase (*plaO1*).¹¹⁸ Inactivation of *plaO1* in the native producer resulted in the accumulation of PL CD6 (**144**) that bears an acyclic pyruvate-derived C₄ side chain at C-14, instead of a dihydroxyfuranone moiety.^{118,119} *In vitro* enzymatic assays confirmed the function of PlaO1 that converts **144** into the dihydroxyfuranone-containing **29**.¹¹⁹ A mechanism for the formation of the γ-butyrolactone moiety by PlaO1 has been proposed (Fig. 25).¹¹⁸ PlaO1 putatively initially hydroxylates C-16 of **144** to form the enol **IM30**, which is converted into the cyclopropanone **IM31** through enolization and intramolecular nucleophilic addition of C-15 to C-17. The proposed subsequent cyclopropanone ring opening results in the formation of the formyl α-ketoacid **IM32**, which undergoes ring closure to yield the hemiacetal **IM33**. Tautomerization of the C-17 ketone follows to yield **29**. Since PlaO1 is a stand-alone enzyme that produces **29** from **144**,¹¹⁹ the keto–enol tautomerization of C-17 likely occurs spontaneously or in a PlaO1-dependent fashion. Although further studies are needed to verify the putative reaction mechanism, the PlaO1-catalyzed ketolactol formation is highly unusual for Fe/αKG-dependent oxygenases.¹¹⁹

4.2.5. Non-canonical nonheme diiron oxygenase-mediated hydroxylation. Oxidative C–C bond cleavage gives rise to truncated or ring-opened terpene scaffolds, leading to the production of *nor*- and *seco*-terpenoids, respectively.^{274–276}

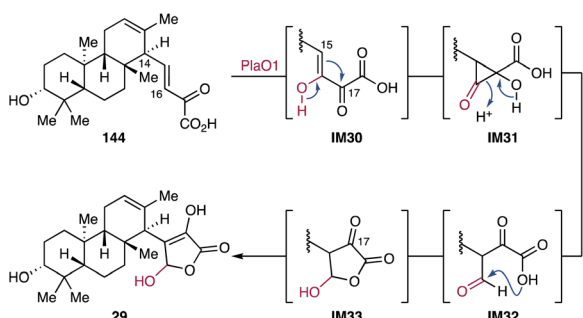


Fig. 25 Proposed ketolactol formation catalyzed by the Fe/αKG-dependent oxygenase PlaO1 during phenalinolactone biosynthesis.

4.2.5.1. Platensimycin and platencin. Platensimycin (**2**) and platencin (**44**) are examples of 4,5-*seco*-*ent*-kaurene and *seco*-*ent*-atiserene diterpenoids, respectively.^{129,277} The formation of the 4,5-*seco*-diterpene skeleton is proposed to proceed through C-5 hydroxylation with a concomitant retro-aldol reaction.^{278–280} In-depth biosynthetic studies of **2** and **44** led to the discovery of a non-canonical nonheme diiron enzyme that installs a hydroxy group at C-5 to set the stage of the intricate opening reaction.²⁸⁰

The gene encoding the C-5 hydroxylase was initially elusive as all oxygenases encoded in the *ptm* BGC had been functionally characterized to be involved in other biosynthetic steps leading up to the formation of **2** and **44**.^{130–132,167} Therefore, deletion of genes with unknown function was performed using *S. platensis* SB12029, a dual-producer of **2** and **44**.^{279,280} Comparative metabolomic studies of extracts derived from mutant strains suggest that PtmU3 is likely responsible for the C-5 hydroxylation of the C-19 CoA thioesters **145** and **146**.²⁸⁰ *In vitro* enzymatic assays imply that PtmU3 is a nonheme iron-dependent oxygenase that converts **145** and **146** into the C-5β hydroxylated CoA thioesters **147** and **148**, respectively (Fig. 26A).²⁸⁰ The generated tertiary alcohols can facilitate a retro-aldol reaction to form 4,5-*seco*-terpenyl CoAs **149** and **150**.

X-ray crystallographic studies reveal that PtmU3 forms a trioseophosphate isomerase (TIM)-like barrel structure, one of the most common tertiary structures found in nature.²⁸¹ While TIM-barrel structures are present in aldolases, glycosidases, enolases, and metal-dependent hydrolases,^{281,282} no TIM-barrel-containing metal-dependent oxygenase had been reported.²⁸⁰ Furthermore, PtmU3 harbors two iron atoms in its active site bridged by Glu241 and Asp308. Their coordination is distinct from canonical nonheme diiron enzymes that typically feature μ-oxo or μ-peroxo bridging ligands between the two metal centers.²⁸³ Mutagenesis studies of PtmU3 and QM/MM calculations of the C-5 hydroxylation reaction reveal that the two iron-bridging amino acid residues are vital for the enzymatic activity and a saturated octahedral iron plays a structural but likely not a catalytic role.^{280,284} The unprecedented diiron coordination of PtmU3 follows neither canonical diiron monooxygenases that usually use one or both iron atoms for catalysis²⁸³ nor other diiron oxygenases, e.g., *myo*-inositol oxygenase MIOX²⁸⁵ and organophosphonate oxidase PhnZ,²⁸⁶ where one iron binds to the respective substrate. In addition to the distinct structural features, QM/MM calculations suggest an unusual catalytic cycle for PtmU3 (Fig. 26B).²⁸⁴ The unsaturated iron in the active site of PtmU3 readily binds molecular oxygen to form a ferric-superoxide species. Spin density calculations for all possible states of the superperoxo-diiron species revealed electron transfers from the iron center to dioxygen and the diiron-coordinating amino acid residues, indicating that these amino acid residues play a role in the electron reorganization of the iron center.²⁸⁴ The generated Fe^{III}-superoxide species abstracts H-5β of **145** or **146** and the resulting hydroperoxide-iron(III) intermediates bind to the respective substrate radicals (**IM34**) to give **147** or **148** with the formation of an oxoiron(IV). The oxoiron(IV) species then react with the same substrates via proton abstraction and oxygen rebound to yield the

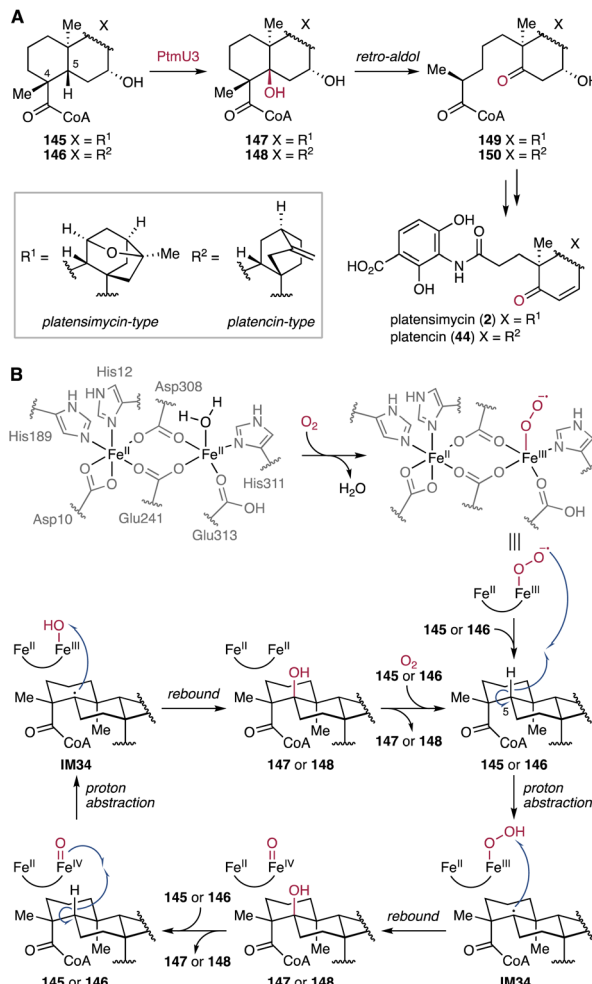


Fig. 26 PtmU3-catalyzed oxidation in the biosynthesis of platensimycin and platensin. (A) PtmU3-mediated hydroxylation to set up the retro-aldol ring cleavage reaction. (B) Proposed unusual catalytic mode of PtmU3.

hydroxylated products. Oxygen rebound in both states of ferric-superoxo and oxoiron(IV) species are feasible based on the energy barrier of the rate-limiting proton abstraction. Therefore, the proposed unusual catalytic mode of PtmU3 successively uses molecular oxygen for the efficient formation of C-5β hydroxy CoA thioesters.²⁸⁴

Sequence similarity network analysis revealed that homologs of PtmU3 are widely distributed in actinobacteria, with conserved key residues for diiron coordination and binding of CoA-activated substrates.²⁸⁰ Noteworthily, half of the genes encoding PtmU3 homologs are located in proximity to genes encoding CoA ligases. This observation suggests that these non-canonical nonheme diiron oxygenases may prefer CoA-activated substrates.²⁸⁰

4.2.6. Rearrangement and cyclization by vanadium-dependent haloperoxidases. Vanadium-dependent haloperoxidases (VHPOs) are one of the seven characterized enzyme families that catalyze halogenation reactions.^{287,288} By harnessing catalytic vanadate and hydroperoxide, VHPOs catalyze two-

electron oxidations of halides to produce short-lived and highly electrophilic hypohalous acids, which react with nucleophilic substrates.²⁸⁹ Although many VHPOs previously identified from eukaryotes and prokaryotes do not exhibit substrate- and regiospecificity,²⁸⁷ red algae-derived VHPOs have been shown to catalyze terpene cyclization reactions *via* bromonium ion formation at the terminal olefin of the acyclic sesquiterpene (+)-nerolidol.²⁹⁰ The carbocation-driven cyclization mechanism of TC-like VHPOs resembles that of type II TCs.²² Furthermore, during the past two decades, biosynthetic studies of bacterial naphthoquinone-based meroterpenoids have led to the discovery of new members of VHPOs that catalyze a wide array of reactions, such as chlorination-induced oxidative rearrangement and cyclization.^{245,291–299}

4.2.6.1. Napyradiomycins. The napyradiomycins are a family of chlorine-bearing naphthoquinone meroterpenoids that are widely found in terrestrial and marine *Streptomyces* spp.^{300–302} Based on the nucleophilicity of the 1,3,6,8-tetrahydroxynaphthalene (THN) nucleus, prenylation patterns are expected at the nucleophilic C-2 or C-4 positions. However, some napyradiomycins feature prenyl groups at the non-nucleophilic C-3 position. As a result, the napyradiomycins are classified into class I (at C-2 or C-4 positions) and class II (at C-3 position).²⁴⁵ Some napyradiomycins feature a chlorinated *gem*-dimethyl tetrahydropyran ring fused with dihydronaphthoquinone and a chlorinated *gem*-dimethyl cyclohexane ring, both of which originate from the terpene components.

Earlier biosynthetic studies using ¹³C-labeled acetic acid suggest that napyradiomycin B1 (151) is biosynthesized from a symmetric THN and two isoprene units, *i.e.*, DMAPP and FPP.³⁰³ Investigation of the genes encoding a THN-forming type III polyketide synthase (PKS) and a PT led to the identification of *nap* BGCs in the genomes of *Streptomyces* sp. CNQ-525 and *Streptomyces aculeolatus* NRRL 18422 with lengths of 36 kbp and 43 kbp, respectively.²⁹¹ Both *nap* BGCs include three genes, namely *napH1*, *napH3*, and *napH4*, encoding VHPOs.²⁹¹ Based on heterologous expression studies in *S. albus*, *in vitro* enzymatic assays, mutagenesis of VHPO genes, and biomimetic syntheses, the biosynthetic pathway of napyradiomycins was characterized (Fig. 27A).^{245,291,292,297,299} All VHPOs play key roles in the maturation of napyradiomycins. Following the biosynthesis of the THN core by NapB1, the PT NapT9 installs a geranyl group at C-4 to yield 4-geranyl THN (152). NapH1 then catalyzes dichlorination at C-2 and OH-3 of 152. The resulting dichloro THN (153) is then dearomatized through the loss of a chlorine atom at OCl-3, followed by quenching of the benzylic cation IM36 with H₂O to yield α'-hydroxyenone 153. NapH4 also converts 152 into 153, albeit with a low catalytic efficiency.²⁹⁷ With strict enzymatic stereocontrol, another PT NapT8 performs the asymmetric prenylation at C-2 of 153, resulting in the formation of the α-hydroxyketone 154.²⁴⁵ Strikingly, NapH3, a homolog of NapH1 and NapH4, is highly substrate-specific and catalyzes α-ketol rearrangement of 154 to form naphthomevalin (155) independent of Na₃VO₄ and H₂O₂.²⁴⁵ DFT calculations indicate that the 1,2-suprafacial-geranyl migration is thermodynamically favorable and requires a *gem*-substitution at C-2 to disrupt the π-conjugated system, which contributes to



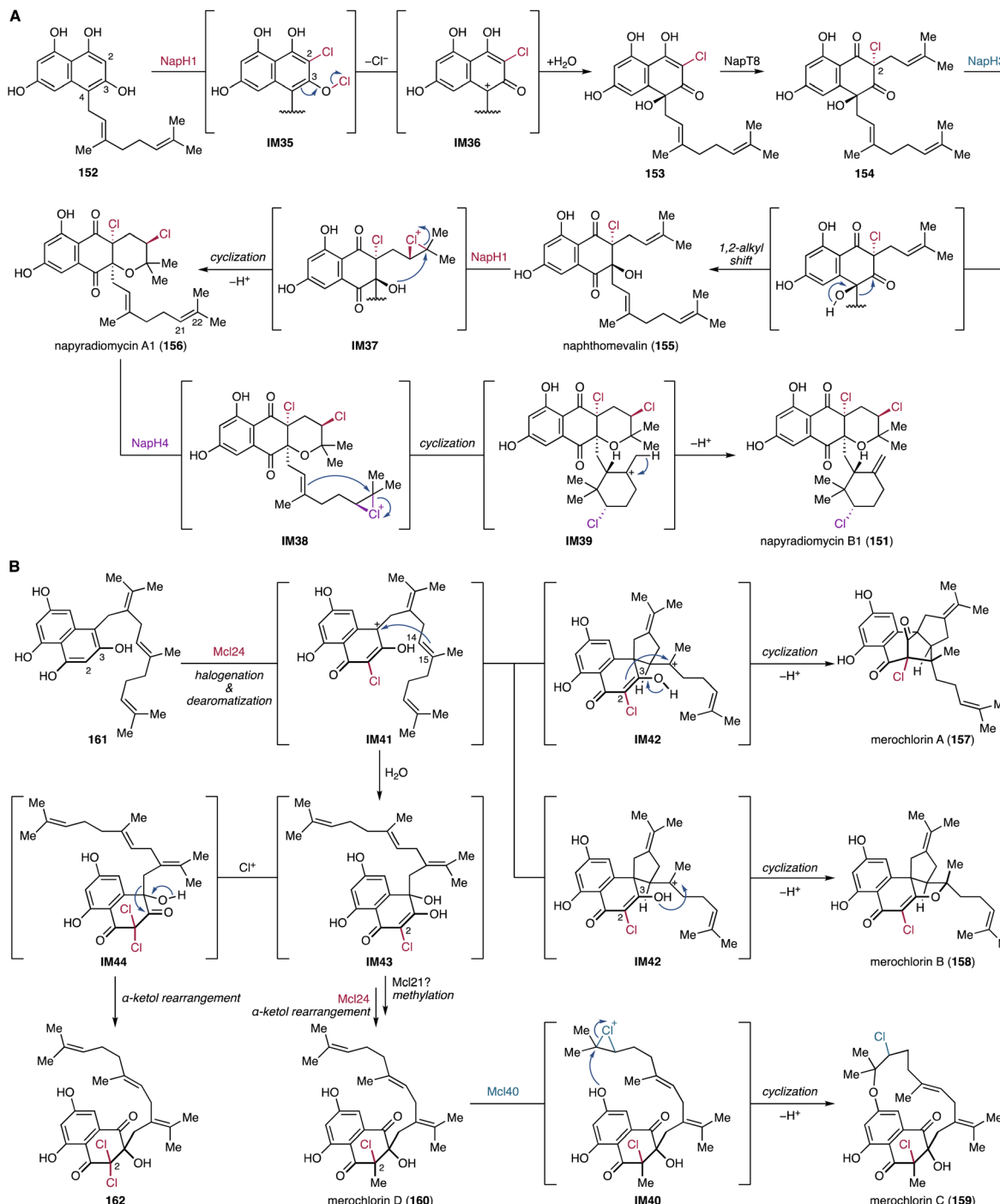


Fig. 27 VHPO-mediated rearrangement and cyclization in THN-based meroterpenoid biosynthetic pathways. (A) Biosynthetic pathway of napyradiomycin B1. (B) Biosynthetic pathway of merochlorins A–D.

the planar structure of the naphthalene-derived bicyclic compound.²⁴⁵ NapH1 is also capable of catalyzing heterocyclization to afford napyradiomycin A1 (156) *via* the enantio-specific formation of the chloronium IM37 using 155 as a substrate.²⁹² Therefore, NapH1 is a dual-functional enzyme that performs chlorination-induced dearomatization and heterocyclization in a substrate-dependent manner. In analogy to the heterocyclization reaction of NapH1, NapH4 chlorinates the

olefin C-21/C-22 of 156.²⁹⁷ The resulting intermediate IM38 undergoes chloronium-induced cyclization, forming the tertiary cyclohexyl cation IM39. Subsequent deprotonation yields 151 that features an *exo*-olefin-containing cyclohexane ring.²⁹⁷

Despite the relatively high homology between NapH1, NapH3, and NapH4, these enzymes catalyze different reactions. Phylogenetic analysis of functionally validated VHPOs demonstrated that VHPOs derived from *Streptomyces* spp., including



NapH1, NapH3, and NapH4, group separately from other microbial and eukaryotic VHPOs.²⁹⁹ Within the *Streptomyces* VHPOs clade, NapH3 is phylogenetically distant from NapH1 and NapH4, suggesting it has evolved to exclusively catalyze the α -ketol rearrangement by losing its halogenation capability.^{245,299} The *apo* structure of NapH1 and mutagenesis of its active site residues revealed the presence of two key amino acid residues that are crucial for the enantioselective chlorofunctionalization and halide oxidation, respectively.²⁹⁹ Notably, these amino acid residues are not conserved in other VHPOs that show substrate promiscuous activities.²⁹⁹ Insights from the biomimetic synthesis of merochlorins using chloramine suggest Lys324 likely forms a chloramine with hypohalous acid generated from the vanadate and regulates the stereospecific transportation of chlorine to the substrate.^{295,299} By contrast, the *apo* structure of NapH3 revealed that His445, which is corresponding to His494 of NapH1, is post-translationally phosphorylated to form a τ -phosphohistidine.²⁹⁹ Instead of the vanadate, the phosphate group non-covalently interacts with active site residues that are essential for vanadate stabilization. This post-translational modification of His445 explains the loss of the haloperoxidase activity in NapH3 by preventing vanadate binding.^{245,299} Mutagenesis studies show that His445 does not play any role in the NapH3-catalyzed α -ketol rearrangement reaction, indicating that substrate binding is independent of His445.²⁹⁹

4.2.6.2. Merochlorins. Merochlorins are another type of THN-based meroterpenoids with potent antibiotic activity.³⁰² Structurally, these compounds are derived from the TNH core and a rare isosesquilavandulyl moiety with various modifications.^{293,302} The *mcl* BGC (58 kbp) was identified by mining the genome of *Streptomyces* sp. CNH-189, which produces merochlorins A-D (**157–160**).²⁹³ Like the *nap* BGC, the *mcl* BGC harbors two genes (*mcl24*, *mcl40*) encoding VHPOs with high homology to *napH1*, *napH3*, and *napH4* respectively. Heterologous expression of the *mcl* genes in *S. coelicolor* M1152 indicated that Mcl40 is responsible for macrocyclization to produce the 15-membered cyclic ether-bearing **159** from **160** (Fig. 27B), presumably through chloronium ion formation (**IM40**) as demonstrated for NapH1.²⁹³ Further *in vitro* biochemical studies suggest that Mcl24 is a multifunctional VHPO that converts pre-merochlorin (**161**) into the structurally diverse products **157**, **158**, and **160**.^{245,295} Mcl24 precisely controls the site-selectivity of the arene and alkene halofunctionalization, as well as the timing and stereospecificity of the oxidative dearomatization and polycyclization (Fig. 27B).^{245,295} Similar to the catalytic activity of NapH1, Mcl24 initially chlorinates both C-2 and OH-3 of **161** and subsequently dearomatizes the THN core by chloride elimination. The resulting benzylic cation **IM41** is converted to the spirocyclic carbocation **IM42** via addition of the olefin C-14/C-15. Subsequent cation quenching by nucleophilic addition of the C-2/C-3 olefin or OH-3 yields the polycyclic products **157** and **158**, respectively. Alternatively, quenching of the benzylic cation by H_2O gives rise to the enone **IM43**, which is subsequently converted to the C-2 *gem*-dichloride **IM44** by Mcl24. The intermediate **IM44** undergoes a thermodynamically favored α -ketol rearrangement to produce **162**.²⁴⁵ This rearrangement reaction

is promoted under basic conditions (pH 8.0), where cation quenching by hydration is preferred over alkene addition and deprotonation.²⁴⁵ Compared to the *gem*-dichlorinated product **162** identified from the *in vitro* catalytic reaction of Mcl24,²⁴⁵ the C-2 position in both **159** and **160** is substituted with methyl and chlorine groups.²⁹³ Therefore, the pathway for the biosynthesis of **160** may proceed through C-2 methylation of the hydrated product **IM43**, presumably catalyzed by the MT Mcl21,²⁹³ followed by the Mcl24-catalyzed α -ketol rearrangement.^{295,302}

4.2.7. Cyclizations initiated by methyl transfer. MTs commonly use SAM to transfer electron-deficient methyls to nucleophilic sites on substrates.³⁰⁴ When alkenes act as methyl acceptors, SAM functions as a Lewis acid, facilitating carbocation formation.³⁰⁵ This carbocation can then undergo a cascade of C-C bond formations before being quenched by dehydrogenation.^{22,305} This catalytic mechanism is similar to that of type II TCs, where the highly conserved DXDD motif acts as a Lewis acid to protonate epoxides or olefins of oligoprenyl pyrophosphates for carbocation formation.¹⁹ The methyl transfer-initiated C-C bond formation catalyzed by non-canonical MTs is observed in biosynthetic pathways of cyclopropane ring-containing fatty acids³⁰⁶ and terpenoids.^{307–309}

4.2.7.1. Sodorifen. Sodorifen (**163**), a unique homosesquiterpene that features the polymethyl bicyclo[3.2.1]octene system, is the major component of the volatile bouquet of organic compounds produced by *Serratia plymuthica*.³¹⁰ Comparative transcriptomic analysis of **163**-producing and non-producing strains have revealed that the biosynthesis of **163** is dependent on the catalytic activity of a SAM-dependent C-MT (SodMT) and the sodorifen terpene cyclase (SodS).³¹¹ The biosynthesis of **163** has been thoroughly investigated through a combination of *in vitro* and *in vivo* experiments and complemented with DFT calculations.^{307,312,313} The complex scaffold of **163** is formed from the sequential cyclization of FPP catalyzed by SodMT and SodS (Fig. 28 and 29). SodMT methylates FPP at C-10 resulting in carbocation formation at C-11. Subsequent ring formation yields the cyclohexyl cation **IM45**. This cationic intermediate undergoes a base-induced cyclopropanation by H-9 deprotonation. In turn, cyclopropane ring opening of the resulting intermediate **IM46** by reprotonation leads to the formation of a C-8-extruded cyclopentyl cation **IM47**. The successive 1,2-hydride and 1,2-methyl shifts to **IM48**, followed by deprotonation of H-10 yield **164**.^{312,313}

SodS catalyzes the unusual cyclization reaction of **164** to produce **163** (Fig. 29).^{307,313} The reaction begins with substrate ionization by pyrophosphate abstraction in **164**, generating the allylic carbocation **IM50**. Sequential 1,4- and 1,2-hydride shifts convert **IM50** to the cyclopentyl cation **IM51**. After fragmentation of **IM51** into a cyclopentene and a pentadienyl cation (**IM52**), hydride transfer from the cyclopentene to the pentadienyl cation and rotation of the resulting pentadiene fragment take place to yield **IM53**. Subsequently, proton transfer from C-7 to C-5 results in the formation of a pentenyl cation, which rotates to align for recombination with the cyclopentadiene fragment (**IM54**). The recombination of the two generated fragments (**IM54**) proceeds through an asynchronous [4 + 3] cycloaddition to form the bicyclo[3.2.1]octene framework **IM55**.



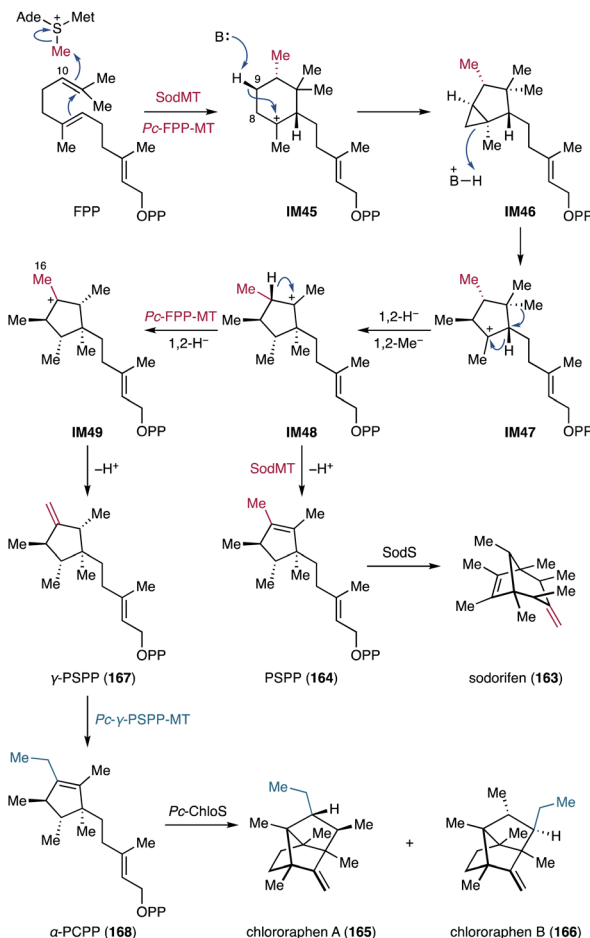


Fig. 28 Non-canonical C-MT-mediated cyclization in the biosynthesis of the homo- and bishomosesquiterpenes sodorifen and chlororaphens.

The cationic intermediate **IM55** then undergoes Wagner–Meerwein rearrangement, followed by deprotonation of H-16, to yield **163**.

Mining of the NCBI RefSeq genome database using protein sequences of SodMT and SodS as query revealed BGCs similar to the sodorifen BGC in more than 180 different bacterial species, mainly belonging to the phylum Proteobacteria.³¹⁴ The identified BGCs were classified into four types based on the presence of genes encoding MTs and TCs. Heterologous expression of selected BGCs in a yeast host led to the identification of various C₁₆ terpenes that feature a polymethylated and highly strained cage-like hydrocarbon scaffold.³¹⁴ These C₁₆ terpenes are derived from **164** as a common intermediate that undergoes TC-catalyzed cyclization. More recently, in-depth mechanistic studies of TCs that produce non-canonical homoterpenes revealed that the fragmentation–recombination mechanism is a common theme in their biosynthetic pathways,³¹⁵ as exemplified in the biosynthetic pathway of **163**.³¹³ The diverse skeletal architectures of these TC-catalyzed reaction products arise from varying modes of recombination of the two components generated by the ionization-induced fragmentation of **164**.³¹⁵

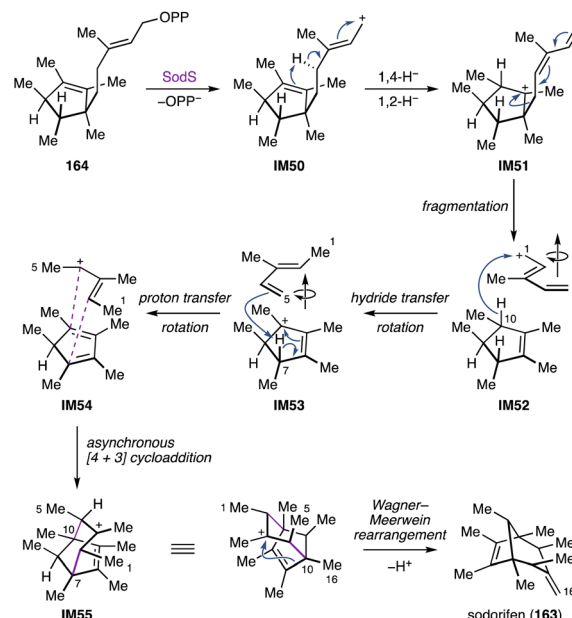


Fig. 29 Fragmentation–recombination mechanism of the SodS-catalyzed cyclization in the biosynthesis of sodorifen.

4.2.7.2. *Chlororaphens*. A similar MT-mediated cyclization has been observed in the biosynthesis of the bishomosesquiterpenes, chlororaphens A and B (**165**, **166**).^{308,316} These unusual C₁₇ terpenes have recently been identified from *Pseudomonas chlororaphis* O6, which harbors a BGC similar to the sodorifen BGCs.³⁰⁸ In addition to genes encoding the non-canonical C-MT (*Pc*-FPP-MT) and the type I TC (*Pc*-ChloS), a gene encoding another C-MT (*Pc*-γ-PSPP-MT) is contained in the chlororaphen BGC. Heterologous expression of the chlororaphen BGC and *in vitro* enzymatic assays using isotope-labeled FPP revealed that the catalytic activity of *Pc*-FPP-MT follows a similar cyclization mechanism to that of SodMT (Fig. 28).^{308,316} However, instead of cation quenching by H-10 deprotonation, an additional 1,2-hydride shift and subsequent deprotonation of H-16 follow to form γ-PSPP (**167**) with a methylidene cyclopentane moiety. Subsequently, *Pc*-γ-PSPP-MT methylates the exocyclic olefin, producing the non-canonical bishomofarnesyl pyrophosphate, α-PCPP (**168**). Finally, α-PCPP is cyclized by *Pc*-ChloS to yield **165** and **166**, as well as the premature bicyclic prechlororaphen (**169**) as a minor product.^{308,316}

Mechanistic characterization of *Pc*-ChloS revealed an unusual cyclization pathway that proceeds through the generation of the neutral compound **169** *en route* to **165** and **166** (Fig. 30).³¹⁶ The pyrophosphate of the C₁₇ intermediate **168** is abstracted by *Pc*-ChloS to generate the allylic cation **IM56**. The intermediate **IM56** undergoes C–C bond formation between C-3 and C-11, followed by 1,2-hydride shift to form the *cis*-fused bicyclo[3.3.0]octane intermediate **IM57**. Deprotonation of **IM57** yields the premature bicyclic product **169**. Afterwards, reprotonation on C-1 of **169** facilitates the second C–C bond formation between C-2 and C-7, generating the tricyclic cationic intermediate **IM58**. Subsequent 1,3-hydride shift results in the formation of **IM59**, which is further converted to **IM60** through



a Wagner–Meerwein-type 1,2-alkyl shift. Finally, cation quenching of **IM60** by deprotonation of H-14 yields **165**. Alternatively, two additional Wagner–Meerwein rearrangements (**IM61**, **IM62**), followed by deprotonation of H-12, lead to the formation of **166**. The cyclization mechanism catalyzed by *Pc*-ChloS is distinct from the fragmentation–recombination mechanism commonly observed in the biosynthesis of C_{16} terpenes like **163**.³¹⁶

4.2.7.3. Teleocidins. Teleocidins and their derivatives are terpenoid indole alkaloids that are potent protein kinase activators, produced by actinobacteria and cyanobacteria.^{317–320} Among these, teleocidin Bs feature an indole framework fused with a nine-membered lactam and a C_{11} terpene moiety. The BGC for teleocidin Bs was identified in the genome of *Streptomyces blastmyceticus* NBRC 12747 by mining for genes homologous to those responsible for the production of lyngbyatoxin A (**170**), a teleocidin analog bearing a linear C_{10} terpene moiety.^{309,321} The *tle* BGC contains genes encoding a non-ribosomal peptide synthetase (NRPS, *tleA*), a P450 (*tleB*), and a PT (*tleC*). Notably, the *tle* BGC that lacks genes encoding a TC or a C-MT is solely responsible for the production of **170** that has a linalyl group.^{309,321} Investigation of candidate MT genes located outside of the *tle* BGC led to the identification of *tleD* encoding a SAM-dependent C-MT involved in the biosynthesis of teleocidins B-1, B-4, and des-*O*-methyl-olivoretin C (**171–173**).³⁰⁹ Further *in vitro* enzymatic assays with isotope-labeled **170** indicate that TleD catalyzes methylation-induced

cyclization of the linalyl group through a cationic cascade reaction (Fig. 31).³⁰⁹ Initially, TleD installs a methyl group at C-25 to generate the tertiary cation **IM63**. After 1,2-hydride migration to **IM64**, aromatic electrophilic substitution at C-7 results in the formation of the spiro-fused cyclohexadienyl iminium intermediates (**IM65**, **IM66**). Subsequently, the intermediates undergo iminium cation-driven cascade reactions including 1,2-alkyl shift at the spiro-cyclopentane ring, followed by deprotonation to yield **171–173**. The nucleophilic addition of C-7 to the C-25 carbocation from the *Re* face is favored over the *Si* face, as indicated by the predominant production of **172** and **173**.³⁰⁹

X-ray crystallographic analysis of TleD complexed with *S*-adenosylhomocysteine and **170** revealed that the active site residues provide an ideal hydrophobic environment to accommodate **170**.³²² This hydrophobicity excludes water molecules and thus prevents the premature quenching of the cation species during methyl transfer-induced cyclization. The linalyl group of **170** appears to be flexible in the crystal structure of TleD. Molecular dynamics simulations of TleD suggest that the conformation of the linalyl group dictates the facial selectivity of the nucleophilic attack to C-25.³²² The predominant conformation of the linalyl group supports the favored formation of **172** and **173**.

4.2.8. Isomerization by hypothetical proteins. Isomerization of double bonds can significantly affect the physicochemical and physiological properties of molecules. For example, 11-*cis*-retinal, a monocyclic diterpenoid derived from all-*trans*-retinol through isomerization and oxidation, is essential for our vision.³²³ Isomerization reactions involved in natural product biosynthesis are catalyzed by various enzyme families including SDRs,³²⁴ flavoenzymes,^{325,326} glutathione *S*-transferases,³²⁷ and enzymes previously annotated as hypothetical proteins (NsrQ in the fungal tetrahydroxanthone biosynthesis, AlbU and EutC in bacterial eunicellane diterpenoid biosynthesis).^{328–330}

4.2.8.1. Albireticulones. The first *trans*-eunicellane synthase (AlbS) was recently discovered in *Streptomyces albireticuli* NRRL B-1670.³³¹ In the genome of *S. albireticuli*, *albS* is clustered with genes encoding a GGPPS (*albG*), two P450s (*albP1*, *albP2*), and a protein with unknown function (*albU*).^{112,329} Heterologous expression of the *alb* BGC in *S. albus* J1074M resulted in the production of albireticulones A and B (**174**, **175**).¹¹² Overexpression of *alb* genes in the native producer and *in vitro* enzymatic assays revealed that AlbU catalyzes the irreversible isomerization of albireticulene (**176**) to form *iso*-albireticulene (**177**).³²⁹ Since AlbU exhibits its enzymatic activity in the absence of cofactors, an acid–base catalytic mechanism, involving the generation of **IM67**, has been proposed for the isomerization reaction (Fig. 32A).³²⁹ Although the suggested mechanism is similar to that of type 1 isopentenyl diphosphate isomerases,³³² AlbU has neither a detectable Pfam domain nor an InterPro designation, indicating that it is the first member of a novel family of isomerases. Moreover, AlbU may isomerize the allyl alcohol **IM68** derived from **177** through the AlbP1-catalyzed hydroxylation at C-6.³²⁹ Specifically, protonation at C-19 of **IM68** generates the tertiary cation **IM69**, which undergoes deprotonation to form the enol intermediate **IM70**. Subsequent

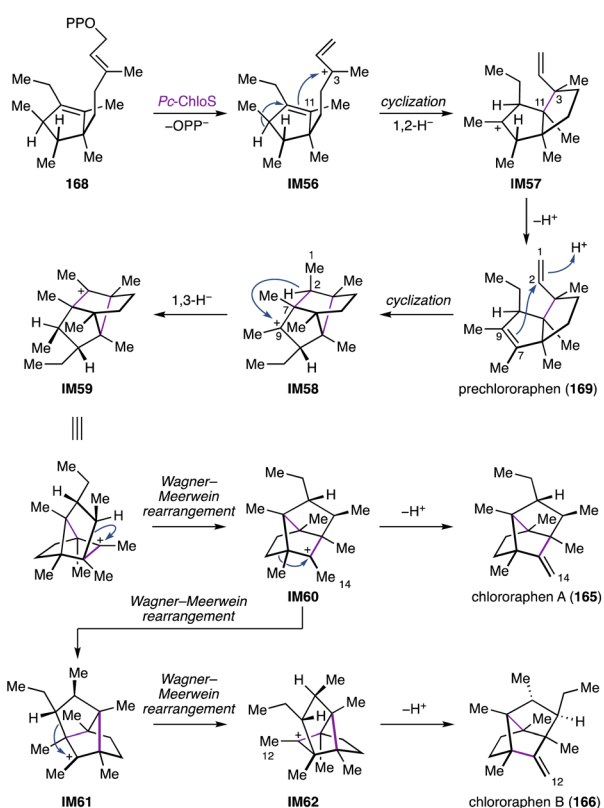


Fig. 30 *Pc*-ChloS-catalyzed cyclization mechanism in the biosynthesis of chlorophaphens.



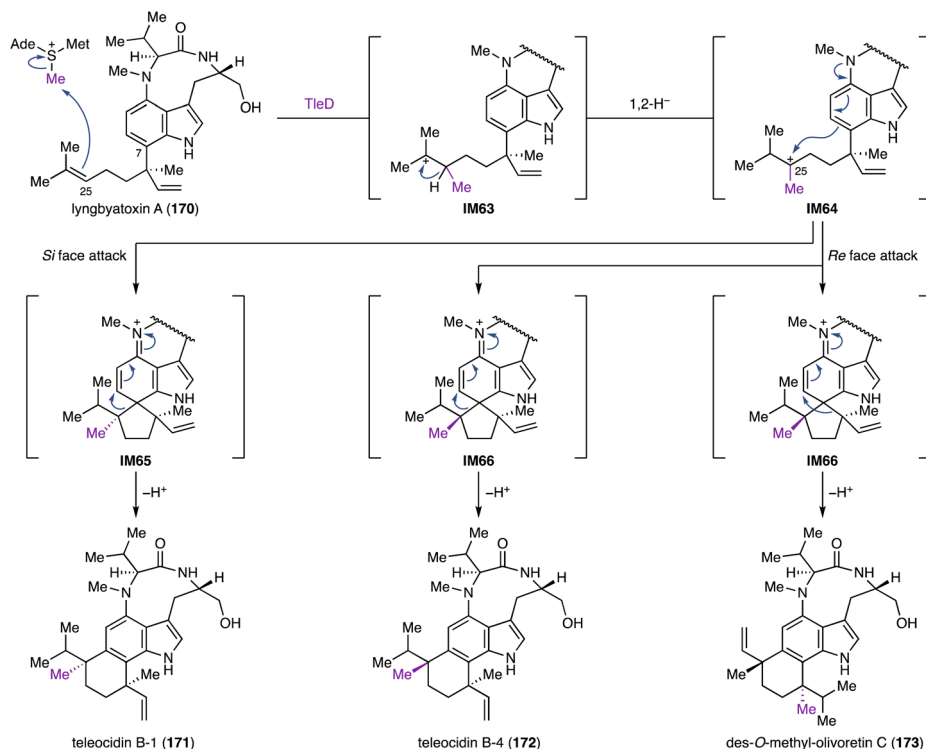


Fig. 31 Non-canonical C-MT-mediated cyclization in the biosynthesis of teleocidins.

tautomerization of **170** results in the formation of **174**.³²⁹ Further mechanistic studies of AlbU are needed to ascertain the proposed dual functions in isomerizing both **176** and the C-6 hydroxylated intermediate of **177**.

4.2.8.2. *Euthailols*. More recently, the *eut* BGC from *Streptomyces euthainensis* N2458 was identified to be responsible for

the biosynthesis of *trans*-fused eunicellane diterpenoids.³³⁰ Although the *eut* BGC shows high similarity to the *alb* BGC, it harbors unique genes encoding a TC fused with a truncated (*E*)-4-hydroxy-3-methyl-but-2-enyl pyrophosphate reductase (*eutB*) and a P450 that is split into three fragments (*eutEFG*).³³⁰ Heterologous expression of the *eut* BGC in *S. avermitilis* SUKA22

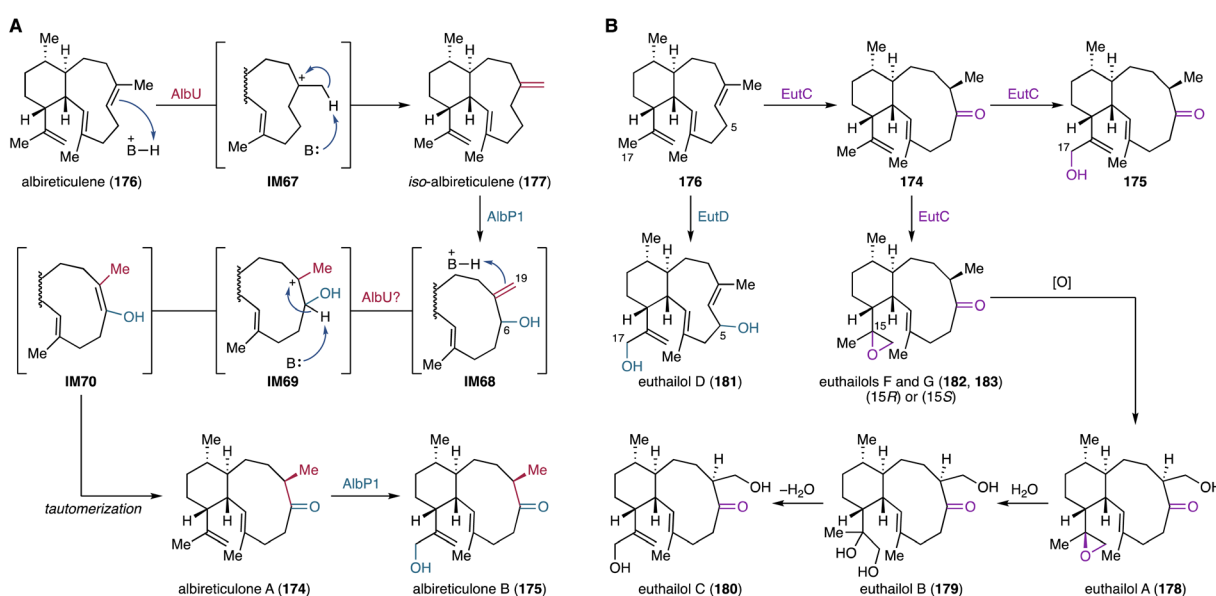


Fig. 32 Scaffold functionalization of *trans*-eunicellane diterpenoids by enzymes previously annotated as hypothetical proteins. (A) Proposed isomerization mechanism of AlbU and biosynthetic pathway of albireticulones. (B) Proposed isomerization and oxidation activity of EutD and biosynthetic pathway of euthailols.



led to the identification of eutailols A–C (**178–180**).³³⁰ *In vivo* functional characterization of the *eut* genes revealed partial functional redundancy between the P450 EutD and the hypothetical protein EutC, both of which hydroxylate C-17 of **176** (Fig. 32B).³³⁰ While EutD hydroxylates C-5 and C-17 of **176** to produce euthailol D (**181**), EutC oxidates **176** to yield **174** and **175**, as well as the epoxide diastereomers, euthailols F and G (**182, 183**). Additionally, EutC is also capable of isomerizing **176** to generate **177**.³³⁰ This observation suggests that EutC is a bifunctional enzyme, acting as an isomerase and an oxygenase. Despite the high sequence similarity of EutC to AlbU, which solely acts as an isomerase,³²⁹ the dual functionality of EutC suggests a functional divergence between two enzymes.³³⁰ The requirement of cofactors for the oxygenation activity of EutC remains unknown,³³⁰ leaving its catalytic mechanism to be elucidated in future studies.

5. Potential and limitations of genome mining for the targeted discovery of terpenoids

Natural product biosynthesis, much like textbook terpene biosynthesis, is a two-step process consisting of scaffold formation and decoration. The importance of the decoration phase varies among different natural product classes, with terpenoids relying more heavily on this phase to achieve structural diversity compared to other classes. Natural product biosynthesis can be classified into two fundamentally different strategies:³³³

(1) Complex polyketides and non-ribosomal peptides (NRPs) are biosynthesized by large multi-enzyme complexes in an assembly line-like fashion. Each assembly line is composed of individual modules, which can be further subdivided into multiple enzymatic domains that work together to add or modify a building block to the nascent natural product backbone.^{334–336} Diversity in these systems is achieved through the incorporation of a wide array of building blocks recognized by specific domains, as well as facultative enzymatic domains integrated into the modules of the assembly line.^{335,337} Consequently, assembly line-like systems generate structural diversity using distinct architectures composed of vectorial arrangements of modules with varying domain compositions. Post-assembly line tailoring reactions play a subordinate role, as a large number of functional groups are already present in the building blocks or are generated during the assembly process. The biosynthesis of these systems is well understood, and seemingly universal biosynthetic principles have been deciphered to predict the core structure of the assembly line product from genome sequence information.^{335,338} The biosynthetic logic of assembly line-like pathways, along with a set of conserved enzymatic domains that exhibit a high degree of sequence homology, facilitates the mapping of the full biosynthetic diversity of these pathways. It is assumed that the majority of assembly line-like pathways can be identified and annotated by current genome mining tools. Assembly line biosynthetic gene clusters are identified using profile Hidden

Markov Models (pHMMs) trained to recognize each commonly occurring enzymatic domain in assembly line modules.³³⁹ Module composition, number, and order, as well as the substrate specificity of individual enzymatic domains within each module, are used to predict the core structure of the assembly line product.^{338,340,341}

(2) In contrast, complex terpenes are produced by a linear arrangement of discrete, typically monofunctional enzymes.³³³ Terpene hydrocarbon scaffolds are composed of multiple isoprene units condensed into a small subset of achiral oligoprenyl precursors. Terpene BGCs do not encode large proteins composed of highly conserved enzymatic domains. Instead, TC genes are typically the only genes consistently present in all canonical terpene BGCs.³³⁹ In many cases, oligoprenyl synthases are encoded within these clusters, indicating the specific subclass of terpenes produced by a pathway.³⁴² The structural diversity of terpenes arises partly from cyclization and rearrangement processes catalyzed by TCs. Unlike the domains in assembly line-like pathways, the primary sequence of TCs is not highly conserved, with conservation typically limited to a set of catalytically important motifs.³⁷ Nevertheless, their tertiary structures are relatively conserved. State-of-the-art genome mining algorithms are capable of identifying the most common types of TCs.³⁴⁰ There have been multiple reports, however, of TCs lacking characteristic motifs³⁴³ or entirely different enzyme families acting as TCs.^{21,22,39,344} This non-canonical TC subpopulation currently evades detection due to the limited number of characterized examples available to train pHMMs to map the full biosynthetic diversity of TCs. Consequently, it is difficult to estimate the proportion of TCs and their associated BGCs that remain unrecognized. TCs merely act as chaperones that guide cationic intermediates through a cascade of cyclization and rearrangement reactions.¹¹ During the cyclization process, the achiral precursors are transformed into stereochemically intricate polycyclic structures. This process, however, currently cannot be predicted. The challenge is further compounded by the fact that a single TC can yield multiple scaffolds.^{10,345} Moreover, some TCs are only a few mutations away from producing an almost non-overlapping spectrum of products.²⁶⁸ Due to the limited number of oligoprenyl precursors, the diversity achievable through TC-mediated cyclization reactions is inherently constrained. As a result, the structural diversity of complex terpenoids is further enhanced through a wide range of tailoring reactions that decorate the terpene scaffolds. Genes encoding these biosynthetic enzymes can be annotated by genome mining algorithms in the vicinity of TC-encoding genes using pHMMs. Predicting the regio- and stereochemistry of the catalyzed reactions, let alone the type of reaction catalyzed, however, is currently impossible.

Several dozen different enzymatic transformations involved in terpene maturation have been described, yet the number of characterized enzymes per enzyme family and type of reaction catalyzed is too small to serve as training datasets for either hard-coded or artificial intelligence-based prediction tools. In summary, the principles of terpene biosynthesis, along with the small number of characterized BGCs, hinder the development of genome mining algorithms that are truly capable of charting



the full biosynthetic diversity of terpene BGCs beyond the canonical terpene BGCs identified by state-of-the-art tools. Recently, a protein 3D structure-model-based genome mining approach has been developed to identify non-canonical TCs that lack conserved motifs typically used for the identification of canonical TCs.³⁴³ BGCs were selected by focusing on the presence of genes encoding oligoprenyl synthases and hypothetical proteins upstream and downstream of oligoprenyl synthase encoding genes. The authors speculated that these hypothetical proteins serve as non-canonical TCs.³⁴³ AlphaFold predictions suggested that the identified hypothetical proteins exhibit high structural similarity to canonical type I TCs indicating their potential role as TCs.³⁴³ Sequence homology searches of the identified non-canonical TCs revealed such TSS to be encoded either within a BGCs or as standalone genes in a wide range of bacterial genera and in eukaryotes.³⁴³ The outlined strategy is one of many approaches to expanding terpene biosynthetic space, and to bring us closer to the goal of charting this space as comprehensively as we have for assembly line-like pathways.

The development of genome mining tools to comprehensively chart terpene biosynthetic space and predict terpenoid structures associated with a BGC of interest require a larger number of characterized terpene BGCs. Artificial intelligence-based approaches could help us predict both the type of scaffold formed by any given TC and the regio- and stereospecificity of tailoring enzymes, along with the types of reactions they catalyze.

6. Conclusion

Tailoring reactions of terpene hydrocarbon scaffolds are crucial biosynthetic steps *en route* to complex terpenoids. For the longest time, bacteria had not been considered as a rich source for the discovery of heavily modified terpenoids. However, as outlined in this review, bacteria undoubtedly produce complex bioactive terpenoids using dedicated sets of tailoring enzymes, such as P450s, FMOs, and MTs. These versatile modifications range from the regio- and stereospecific functionalization of unactivated C–H bonds that are chemically largely undistinguishable to unprecedented oxidative or nonoxidative structural rearrangements, executed in either promiscuous or enantiospecific fashion. This remarkable enzymatic repertoire is the bases for the extraordinary structural diversity of bacterial terpenoids. Particularly, P450s play a central role in the oxidative functionalization of terpene scaffolds, with most bacterial terpenoid BGCs including multiple P450 genes alongside TC encoding genes. The functional characterization of the full biosynthetic repertoire of P450s and other oxidoreductases will fundamentally change our understanding of how terpenoids strategically gain their unique chemical and biological features.

Ongoing efforts to characterize bacterial terpenoid biosynthetic pathways have uncovered unusual functions of tailoring enzymes. Unlike the canonical enzymes that follow well-established mechanistic routes, such as the oxygen rebound mechanism of iron-dependent oxygenases, the characterized non-canonical tailoring enzymes catalyze fascinating reactions

mostly involved in skeletal transformations through unusual enzymatic mechanisms. Notable examples discussed in this review include oxidative alkyl migration (PntM, AuaG, NapH3), ring contraction (CYP114), and cyclization (BezE, XiaE, PlaO1, TleD) reactions. These non-canonical tailoring enzyme-catalyzed transformations frequently occur in the later stage of biosynthetic pathways, contributing to the structural complexity of terpenoids. Intriguingly, some enzymes act earlier to forge unusual terpene scaffolds rather than merely decorating hydrocarbon frameworks, as shown in the biosynthesis of aridacins (AriF) and sodorifen (SodMT). The catalytic diversity of non-canonical tailoring enzymes underscores the immense potential for the characterization of novel enzymes with unique and distinct activities from bacterial terpenoid biosynthetic pathways.

Motivated by the sophisticated catalytic function of bacterial terpenoid tailoring enzymes, both native and engineered variants have been applied for the efficient synthesis of biologically important terpenoids.⁹ Hence, the discovery of new enzymology opens up opportunities to develop elaborate biocatalysts for (chemo)enzymatic late-stage functionalization and inspires biomimetic syntheses of naturally occurring complex terpenoids and even new-to-nature molecules.

Although almost all bacterial terpenoid BGCs are silent or expressed at low levels under standard laboratory culture conditions, recent advancements in molecular biology, such as the efficient heterologous expression of terpene BGCs in heterologous hosts, optimized for terpene production,^{112,346} have paved the way for the discovery of novel terpenoids and the identification of new enzymatic functions. Coupled with the ever-increasing availability of genome data, state-of-the-art genome mining techniques readily identify canonical terpenoid BGCs. To date, the antiSMASH database, an online repository of BGCs predicted by antiSMASH, lists more than 28 800 BGCs from publicly available genomes of bacteria.³⁴⁷ However, the targeted identification of non-canonical bacterial terpenoid BGCs still remains challenging. Many terpenoid BGCs evade detection by current genome mining platforms due to the relatively low sequence homology between TCs and the frequent absence of detectable Pfam domains in non-canonical TCs responsible for terpene scaffold formation.^{22,38} Moreover, unlike thiotemplate assembly line-based BGCs, structure predictions of terpenoids from BGC information remains challenging. The difficulty of structural predictions hinders the prioritization of orphan BGCs for the discovery of novel terpenoids. A deeper understanding of the principles that govern bacterial terpenoid biosynthesis may enable the construction of training datasets for the development of artificial intelligence-based structure prediction tools specific to modifications introduced by tailoring enzymes. These tools can be integrated into the genome mining workflow to thoroughly analyze BGCs and facilitate the targeted discovery of novel terpenoid biosynthetic pathways.

For the majority of the past two decades, research on bacterial terpenoid biosynthesis has predominantly concentrated on the cyclization phase mediated by TCs. During this period, various tailoring enzymes encoded in the BGCs studied



for their TC have been discovered. As the mechanistic characterization of terpenoid tailoring enzymes rapidly advances, it is evident that multidisciplinary approaches—combining bioinformatics, synthetic biology, organic chemistry, enzymology, and computational chemistry—are crucial for deepening our understanding of bacterial terpenoid biosynthesis. Emerging evidence now suggests that more and more characterized tailoring enzymes are involved in the formation of terpene backbones, rather than merely decorating pre-formed hydrocarbon scaffolds. These insights might necessitate a re-evaluation of the traditional ‘two-phase’ model of terpenoid biosynthesis, which strictly separates scaffold formation from decoration processes.

7. Conflicts of interest

There are no conflicts to declare.

8. Acknowledgements

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