Cytochrome P450-Mediated Cyclization in Eunicellane-Derived Diterpenoid Biosynthesis

Zengyuan Wang,1,5 Qian Yang,1,5* Jingyi He,1 Haixin Li,1 Xingming Pan,1 Zining Li,3 Hui-Min Xu,4 Jeffrey D. Rudolf,3 Dean J. Tantillo,2* and Liao-Bin Dong1*

1State Key Laboratory of Natural Medicines, School of Traditional Chinese Pharmacy, China Pharmaceutical University, Nanjing 211198, Jiangsu, China
2Department of Chemistry, University of California, Davis, California 95616, United States
3Department of Chemistry, University of Florida, Gainesville, Florida 32611-7011, United States
4The Public Laboratory Platform, China Pharmaceutical University, Nanjing 211198, China
5These authors contributed to this work equally

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Corresponding to: yangqian117@cpu.edu.cn (Q.Y.), djtantillo@ucdavis.edu (D.J.T.), or ldong@cpu.edu.cn (L.-B.D.)
Abstract

Terpene cyclization, one of the most complex chemical reactions in nature, is generally catalyzed by two classes of terpene cyclases (TCs). Cytochrome P450s that act as unexpected TC-like enzymes are known but are very rare. Here, we genome-mined a cryptic bacterial terpenoid gene cluster, named *ari*, from the thermophilic actinomycete strain *Amycolatopsis arida*. By employing a heterologous production system, we isolated and characterized three highly oxidized eunicellane-derived diterpenoids, aridacins A–C (1–3), that possess a rare 6/7/5-fused tricyclic scaffold. *In vivo* and *in vitro* experiments systematically established a non-canonical two-step biosynthetic pathway for diterpene skeleton formation. First, a class I TC (AriE) cyclizes geranylgeranyl diphosphate (GGPP) into a 6/10-fused bicyclic cis-eunicellane skeleton. Next, a cytochrome P450 (AriF) catalyzes cyclization of the eunicellane skeleton into the 6/7/5-fused tricyclic scaffold via C2-C6 bond formation. Quantum chemical computations support a hydrogen abstraction and subsequent oxidation mechanism for AriF catalyzed carbocation cyclization. The biosynthetic logic of skeleton construction in the aridacin diterpenoids is unprecedented, expanding the catalytic capacity and diversity of P450s and setting the stage to investigate the inherent principles of carbocation generation by P450s in the biosynthesis of terpenoids.
Introduction

The eunicellanes, a unique subfamily of diterpenoids that contain more than 360 examples, are mainly found in Octocorallia soft corals with only a few known members isolated from plants and bacteria.\(^1\)–\(^8\) Structurally, all eunicellanes share a 6/10-bicyclic skeleton with most representatives, particularly those from corals, exhibiting diverse oxidation patterns. These structures imbue eunicellanes with promising bioactivities\(^1,2\) with examples including the anti-inflammatory klysimplexin R,\(^9\) the antimetastatic polyanthellin A,\(^10\) and the potent tubulin inhibitor eleutherobin.\(^11\) Therefore, they are fascinating targets for both chemists and biologists (Figure 1A). The biosynthetic logic of eunicellanes has remained poorly understood; the diterpene cyclases responsible for construction of the cis and trans 6/10-bicyclic skeletons were only recently disclosed from bacteria and corals.\(^3,4,8,12,13\)

Figure 1. (A) Representative bioactive eunicellane diterpenoids; (B) P450s serve as unexpected TCs in terpenoid biosynthesis.

Terpene cyclizations are one of the most complex chemical reactions in nature. These intriguing reactions are catalyzed by a superfamily of enzymes named terpene cyclases (TCs), which utilize
nature’s library of acyclic C$_{5n}$ diphosphate precursors to generate complex mono- or polycyclic terpene skeletons via carbocation intermediates. TCs are grouped into two canonical classes according to their biochemical strategies to generate the initial carbocation: class I TCs generate carbocations by abstracting the diphosphate group and class II TCs protonate an alkene or epoxide of the prenyl diphosphate; the products of class II TCs often serve as substrates for class I TCs providing a two-step sequential route to structural diversity in terpenoid biosynthesis (Figure S1).$^{14}$

Cytochrome P450 enzymes (P450s) are heme-containing monooxygenases that serve as ubiquitous tailoring enzymes that are capable of catalyzing diverse oxidation reactions including carbon-carbon bond formation in natural products biosynthesis.$^{15,16}$ P450s utilize a single-electron process to produce radical intermediates via Compound I (a high-valent oxoiron cationic radical) which readily abstracts a hydrogen from the substrate to yield reactive radical species. Typically, the radical species rapidly rebounds with the hydroxyl radical species, which is derived from molecular oxygen, to generate the hydroxylation product.$^{15,16}$ Alternatively, though rarely observed, when the rate of oxygen rebound is reduced sufficiently, an electron from the substrate can be transferred to the radical center resulting in a reactive carbocation, which can stimulate a series of cyclization or rearrangement reactions as seen in TC chemistry.$^{14,17}$ P450s that serve as TC-like enzymes are known but rare in the biosynthesis of terpenoids. The three known examples are (i) PenM/PntM catalyzing the final step of oxidative rearrangement via a neopentyl cation intermediate in the biosynthesis of the bacterial sesquiterpenoid pentalenolactone;$^{18,19}$ (ii) $T.w$CYP71BE86, a plant P450, mediating a methyl shift of the abietane-type diterpene scaffold in triptonide biosynthesis;$^{20}$ and (iii) VrtK, which resembles class II TCs and initiates cyclization in the biosynthesis of the fungal meroterpenoid viridicatumtoxin (Figure 1B).$^{21}$

Here we first genome-mined for novel eunicellane diterpenoids by targeting the bacterial biosynthetic gene cluster ari from the thermophilic actinomycete Amycolatopis arida. We then
activated this cryptic biosynthetic gene cluster (BGC) by heterologous expression in model *Streptomyces* hosts and isolated three highly oxidized and eunicellane-derived 6/7/5-tricyclic diterpenoids, aridacins A−C (1−3). Their structures were unambiguously established by extensive spectroscopic analyses, including phenylglycine methyl ester (PGME) transformation and single crystal X-ray diffraction. Using a combination of heterologous production, *in vivo* inactivation, *in vitro* biochemical experiments, and quantum chemical calculations, we established that a P450, AriF, mediates the formation of the 6/7/5-tricyclic skeleton from the 6/10-bicyclic eunicellane substrate benditerpe-2,6,15-triene (4) via a class II-like TC mechanism.

**Results and Discussion**

**Genome Mining Reveals a Novel Eunicellane Biosynthetic Gene Cluster.** Terpenoids are the largest family of natural products with over 95,000 known compounds (http://dnp.chemnetbase.com),22 the vast majority of which have been isolated from plants and fungi; less than 1.5% are from bacteria.23 However, the genomes of bacteria showcase extensive terpenoid biosynthetic potential that has yet to be revealed.23 As genes from bacterial secondary metabolism tend to be clustered together, we envisioned that targeting the gene clusters with multiple P450s neighboring TCs might be a practical approach for the discovery of highly oxidized terpenoids.24 We targeted a putative 16-kb *ari* BGC from a thermophilic actinomycete *Amycolatopsis arida* CGMCC 4.5579 (formerly known as *Yuhushiella deserti*, Figures 2A and S2).25,26 The *ari* BGC encodes a class I TC, a UbiA family prenyltransferase, four P450s, a ferredoxin, a geranylgeranyl diphosphate (GGPP) synthase, two genes, 4-hydroxy-3-methylbut-2-enyl diphosphate reductase (IspH) and 1-deoxy-d-xylulose 5-phosphate synthase (DXS), found in the non-mevalonate pathway, and three unknown genes (Figures 2A and Table S1). The sole TC in the *ari* BGC (named AriE) shares 71.6% protein sequence identity with Bnd4 from the benditerpenoic acid biosynthetic pathway,8,12 suggesting that this BGC might encode eunicellane-derived diterpenoids.
Figure 2. Biosynthesis of aridacins. (A) The *ari* biosynthetic gene cluster. (B) The proposed biosynthetic pathway of aridacins. Dashed arrows depict biosynthetic steps that were unable to be established by *in vivo* or *in vitro* experiments. (C) Δδ values (δ(S) – δ(R)) of the phenylglycine methyl ester (PGME) amides of 1a and 1b, and the X-ray crystal structure of 1a. (D) EIC (333–349 m/z; negative mode) analysis of metabolites from engineered *Streptomyces* strains. *S. lividans* TK64 with empty pSET152 was used as a control. (E) HPLC analysis of the metabolites of ΔariD and ΔariF constructs in *S. lividans*. (F) HPLC analysis of the metabolites of engineered *E. coli* strains. DL10025 is a GGPP-overproducing *E. coli* strain and is used as a control. DL10026–DL10028 are DL10025 harboring *ariE*, *ariE + ariD*, and *ariE + ariF*, respectively.

**Heterologous Expression of the *ari* BGC Produces Three Highly Oxidized Eunicellane-Derived Diterpenoids.** Initial attempts to isolate *ari* BGC-related terpenoids from *A. arida* were unsuccessful, presumably due to the BGC being transcriptionally silent under laboratory culture conditions. To activate the *ari* BGC, we cloned and heterologously expressed it in three model *Streptomyces* hosts. *Streptomyces lividans* DL10011 (*Tables S2–S4*), which harbors the entire *ari* BGC (*orf1–orf6*), showed three new peaks (1–3) by HPLC-MS with m/z [M–H]⁻ at 333, 349, and 349, respectively, when cultured in several terpenoid production media (*Figures 2D and S3–S6*). PTMM medium provided higher titers of 2 and 3 while XTM medium was more
conducive to produce 1. Therefore, DL10011 was individually fermented on a 15-L scale in both PTMM and XTM media to isolate aridacins A–C (1–3, Figure 2B).

Aridacin A (1) had the molecular formula C_{20}H_{30}O_{4} as deduced by the (−)-HRESIMS ion at m/z 333.2063 (calcd 333.2071), indicating six degrees of unsaturation (Figure S7). The $^{1}$H NMR (Figure S8 and Table S5) exhibited characteristic signals for two olefinic protons at δ_{H} (4.90 and 4.81) and two methyl groups at δ_{H} (1.61 and 0.90). Its $^{13}$C NMR and DEPT spectra showed 20 carbon resonances including two methyls, eight methylenes, five methines, and five quaternary carbons (Figures S9 and S10). The planar structure of 1, a 6/7/5-fused tricyclic diene, was elucidated by the spin systems in the $^{1}$H-$^{1}$H COSY spectrum of H-2/H-4/H-5 and H-2/H-9/H-10/H-11/H-12/H-13/H-14/H-1/H-2, and the key HMBC correlations of H-1 with C-15, H-2/H-17 with C-15 and C-16, and H-20 with C-2, C-3, and C-4 (Figures S11–S13). The key ROESY correlations of H-17a/H-1, H-17a/H-10, along with H-2/H-11 suggested that H-10 and H-11 were β- and α-orientated, respectively (Figure S14). Analyses of the NMR data of 2 and 3 concluded they were analogs of 1 (Figures S15–S28, and Tables S6 and S7). In comparison with 1, aridacins B (2) and C (3) have additional hydroxyl groups at C-19 and C-12, respectively (Figure 2B). To determine the stereochemical configuration of C-15 of the glyceric acid moiety in 1, two derivatives, 1a and 1b, were synthesized using (R)- and (S)-PGME (Figures S29–S35 and Table S8); C-15 was determined as S configuration based on the Δδ-values ($\delta_{(S)} - \delta_{(R)}$) shown in Figure 2C. The absolute configuration of 1a was unambiguously determined by X-ray crystal structure (CCDC 2216506, Figure 2C).

The aridacins A–C (1–3) are the first bona fide bacterial diterpenoids possessing a 6/7/5-fused tricyclic scaffold, despite the scaffold being previously seen in TC studies (Figure S36). Odyverdiene B was produced by heterologously expressing nd90_0354 from Streptomyces sp. ND90 in a Streptomyces host; isocatenula-2, 14-diene was produced by incubating GGPP with CaCS from Catenulispora acidiphila. The highly oxidized glyceric acid moiety, which may
originate from a six-electron oxidation of the C-17 methyl group and sequential epoxidation and hydrolysis of the C-15 alkene, is also uncommon in terpenoids.\textsuperscript{34} The aridacins A–C were screened for cytotoxicity in five cancer cells and antibacterial activity in a panel of Gram-positive and Gram-negative bacteria. However, 1–3 showed no significant biological activities (Tables S9 and S10). The continued screening of other biological models is currently underway.

**In vivo Inactivation Partially Elucidates the Biosynthetic Pathway.** To probe the biosynthetic pathway of 1–3, we first identified the boundary of *ari* BGC (Figures S37–S38). We heterologously expressed *ariA–G* in *S. lividans* TK64 to yield *S. lividans* DL10016, which, based on the HPLC-MS analysis, produced equal amounts of 1–3 with that of *S. lividans* DL10011 (Figure 2D, panels iii and iv). This result supported that (i) *ariA–G* can be assigned putative roles in the biosynthesis of 1–3; (ii) the UbiA-like gene (*orfI*), of which several members of this family are known as TCs, is not involved in cyclization or any other step, and (iii) three P450s genes, *ariD, ariF*, and *ariG*, are sufficient to decorate the diterpene skeleton. When *ariA–C* were removed (i.e., *ariD–G*), the engineered *S. lividans* DL10017 strain produced significantly less (<10%) of 2–3 than DL10011, suggestive of their important roles in precursor flux but not essential roles in the biosynthesis of the aridacins (Figure 2D, panel v). To further determine the function of AriD–G, we constructed a series of heterologous expression strains, each lacking one of the *ariA–G* genes in *S. lividans* DL10016 (Δ*ariD, ΔariE, ΔariF*, and Δ*ariG*; Figures 2D, panels vi–ix, and S39). In the *ariE* knockout strain (*S. lividans* DL10019), 1–3 was expectedly abolished. The Δ*ariG* mutant (*S. lividans* DL10021) only accumulated 1 and 2, indicating AriG is responsible for installing the C-12 hydroxyl group of 3. Both Δ*ariD* (*S. lividans* DL10018) and Δ*ariF* mutants (*S. lividans* DL10020) completely abolished the production of 1–3, but intriguingly produced the highly hydrophobic products 4 and 5, respectively (Figures 2E and S40). Due to low titers of 4 and 5 from these heterologous hosts, however, we were unable to accumulate enough material for structural elucidation.
**AriE Constructs the 6/10-Fused Bicyclic cis-Eunicellane Skeleton.** We first cloned *ariE* into our GGPP-overproduction *E. coli* system\textsuperscript{35,36} to yield *E. coli* DL10026, and produced 20 mg of 4. \textsuperscript{1}H and \textsuperscript{13}C NMR analyses confirmed its structure as the cis-eunicellane benditerpe-2,6,15-triene (Figures 2F and S41–S43).\textsuperscript{8} We also cloned and heterologously produced *ariE* in *E. coli* (Figure S44). AriE, incubated with GGPP and Mg\textsuperscript{2+}, resulted in the production of 4 as well as the known 14-membered (\(R\)-(−)-cembrene A (6, Figures 3B and S45–S48)).\textsuperscript{37,38} Compound 4 was easily transformed into a new cis, trans-6/6/6-fused tricyclic product gersemiene C (7) under mild acidic conditions (Figures 3A and S49–S56, and Table S11);\textsuperscript{39} a similar transformation was seen for the trans-eunicellane albireticulene resulting in two trans, trans-6/6/6 isomers.\textsuperscript{13} This non-enzymatic cyclization occurs via protonation at C6 of 4 followed by 2,7-annulation and deprotonation at C-20 to access 7 (Figure S49). Taken together, AriE merely produces the 6/10-fused bicyclic eunicellane diterpene skeleton but not the 6/7/5-fused tricyclic scaffold seen in 1–3.

![Diagram of biochemical reactions](image)

**Figure 3.** Biochemical characterizations of AriE and AriF. (A) Scheme of the reactions of AriE and AriF. (B) HPLC profiles of the *in vitro* reactions of AriE with GGPP. (C) HPLC profiles of the *in vitro* reactions of AriF with benditerpe-2,6,15-triene (4).
AriF, a P450, Mediates Cyclization of the Eunicellane Scaffold into the 6/7/5-Tricyclic Aridacins. Since 4 accumulated in the ΔariF mutant, we proposed that the P450 AriF catalyzes the third ring formation immediately after terpene cyclization. To test this hypothesis, we cloned a codon-optimized version of *ariF* into *E. coli* DL10026, the producer of 4, with the redox partner RhfRed to yield *E. coli* DL10028. Besides 4, DL10028 produced a new compound with an identical retention time with that of 5 seen in the ΔariF mutant (Figure 2F). A large-scale (30-L) fermentation was performed to collect 3 mg of 5. The GC-MS of 5 showed a molecular ion peak at m/z 270.17 (Figure S57), corresponding to a diterpene hydrocarbon of molecular formula C_{20}H_{30} with six degrees of unsaturation. Its 1D and 2D NMR spectra suggested that 5 was an analogue of aridacin A (1) with an isopropenyl group in place of the glyceric acid side chain in 1 (Figures S58–S63 and Table S12). This structural difference was supported by ^1^H NMR data (δ_H 4.85 and 4.81, CH_2_16; 1.65, CH_3_17) and ^1^C NMR data (δ_C 147.8, C_15; 110.7, CH_2_16; and 21.7, CH_3_17). Thus, 5 was assigned as a 6/7/5-fused tricyclic diterpene and named arida-3,6,15-triene.

To obtain direct evidence that AriF catalyzes the unusual cyclization of 4 into 5, we sought to confirm enzymatic activity via *in vitro* experiments. Unfortunately, due to unsuccessful expression of either wild-type or codon-optimized *ariF* in *E. coli*, we produced AriF in *S. albus* J1074 (Figure S64). AriF was assayed in the presence of 4, NADPH, and potential redox partners (CamA/CamB, RhfRed, or FdR/Fdx).^40^ AriF produced a single enzymatic product 5 when paired with either CamA/CamB or RhfRed, with much higher production of 5 observed with CamA/CamB (Figures 3C and S65). Overall, the data conclusively supports that the nascent 6/7/5-fused tricyclic skeleton of 1–3 is created by a step-wise combination of a class I TC (AriE) and P450 (AriF), an unprecedented event in the biosynthesis of core terpene skeletons.

Quantum Chemical Calculations Support the Carbocation Cyclization Mechanism of AriF. The P450-catalyzed conversion of 4 to 5 resembles the terpene cyclization of VrtK and rearrangements of PenM/PntM and *T. wCYP71BE86*, where oxidation of a radical provides a
We initially proposed two possible pathways (Figure 4). In pathway (i), AriF abstracts a hydrogen from CH$_3$-20 of 4 via Compound I to yield radical intermediate a, which undergoes electron transfer rather than oxygen rebound to form allylic carbocation b. A subsequent 2,6-ring closure yields the tricyclic skeleton with deprotonation at C-6 providing 5. In pathway (ii), hydrogen abstraction and cation formation occur at C-4 followed by a 1,3-hydride shift to form intermediate b; cyclization and deprotonation then follow that in pathway (i).

Figure 4. Putative catalytic mechanism of AriF.

Quantum chemical computations were executed to examine the possible cyclization mechanisms (Figure 4). Geometry optimizations and energies were obtained using the mPW1PW91/6-31+G(d,p) density functional theory method, which has been used extensively for modeling terpene-forming carbocation reactions. Structures for radicals a and d were readily located. A transition state structure for the 1,3-H shift that interconverts radicals a and d was also located, but the barrier for that shift is predicted to be very large (>50 kcal/mol). While a structure for carbocation c was readily located, attempts to locate b and e with the relative stereochemical configuration corresponding to 5 led directly to structures with 6/7/5-fused tricyclic frameworks,
i.e., c was formed when attempting to locate b and its analog when attempting to locate e. These results suggest that the cations formed from radicals a and d have their $\pi$-bonds and cation centers so close in space that there is no barrier for them to combine. In addition, since product 5 is observed, rather than a product of direct cyclization of carbocation e (i.e., 6/7/5 tricycle with an alkene between C-3 and C-4), it seems plausible that initial hydrogen abstraction occurs at C-20. Altogether, pathway (i) is supported as the most likely mechanistic route for AriF cyclization of the cis-eunicellane skeleton of 4 into arida-3,6,15-tiene (5).

### Conclusion

Most of the skeletal diversity of terpenes in nature are generated by TCs, enzymes that generate structurally and stereochemically complex polycyclic hydrocarbons from acyclic precursors. However, the enzymatic repertoire capable of performing terpene cyclization is far more diverse than a single enzyme family. In this study, we used genome mining and discovered a new class of highly oxidized eunicellane-derived 6/7/5-fused tricyclic diterpenoids aridacins A–C (1−3) from the thermophilic actinomycete *A. arida*. Heterologous production, *in vivo* inactivation, and *in vitro* enzyme characterization systematically established that a TC and P450, AriE and AriF, respectively, successively generate the 6/10-fused bicyclic cis-eunicellane skeleton and the 6/7/5-fused tricyclic scaffold. Quantum chemical computations supported the proposal that AriF catalyzes a carbocation mechanism for the third ring formation. Although three P450s have been previously reported to mediate terpene rearrangements or cyclization in (mero)terpenoid biosynthesis, they all function toward the end of their biosynthetic pathways. In the biosynthesis of the aridacins, a TC and P450 work in tandem to initially create the backbone of the aridacins, an unprecedented strategy in terpene skeletal construction. Our discovery expands the catalytic capacity and diversity of P450s and sets the stage for future efforts to investigate the inherent principles of carbocation generation by P450s in the biosynthesis of terpenoids. During submission of this manuscript, *ariD–G* was heterologously expressed in *S. albus* J1074M producing a benditerpenoid acid-like eunicellane diterpenoid. The production of our 6/7/5-tricyclic
diterpenoids and the 6/10-bicyclic product from the same set of genes raises an intriguing question of how the *ari* BGC is being controlled in different heterologous hosts and what are the genuine natural products from the original strain *A. arida*.

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**Author contributions**

L.-B.D. conceived the project; Z.W., Q.Y., D.J.T., and L.-B.D. designed the experiments; Z.W., Q.Y., J.H., H.L., X.P., and H.-M.X. performed the experiments; Z.W., Q.Y., Z.L., J.D.R., D.J.T., and L.-B.D. analyzed the results; Z.W., Q.Y., J.D.R., D.J.T., and L.-B.D. wrote the paper with inputs from all co-authors.

**Competing financial interests**

The authors declare no competing financial interest.

**Correspondence and requests for materials** should be addressed to Q.Y., D.J.T., or L.-B.D.
References

(18) Zhu, D.; Seo, M.-J.; Ikeda, H.; Cane, D. E. Genome mining in Streptomyces. Discovery of an unprecedented


(30) Deposition Number 2216506 (for 1a) contains the supplementary crystallographic data for this paper. This data is provided free of charge by the joint Cambridge Crystallographic Data Centre and FIZ Karlsruhe Access Structures service.


