

Biosynthesis of coral settlement cue tetrabromopyrrole in marine bacteria by a uniquely adapted brominase–thioesterase enzyme pair

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Halogenated pyrroles (halopyrroles) are common chemical moieties found in bioactive bacterial natural products. The halopyrrole moieties of mono- and dihalopyrrole-containing compounds arise from a conserved mechanism in which a proline-derived pyrrolyl group bound to a carrier protein is first halogenated and then elaborated by peptidic or polyketide extensions. This paradigm is broken during the marine pseudalteromonad bacterial biosynthesis of the coral larval settlement cue tetrabromopyrrole (**1**), which arises from the substitution of the proline-derived carboxylate by a bromine atom. To understand the molecular basis for decarboxylative bromination in the biosynthesis of **1**, we sequenced two *Pseudoalteromonas* genomes and identified a conserved four-gene locus encoding the enzymes involved in its complete biosynthesis. Through total in vitro reconstitution of the biosynthesis of **1** using purified enzymes and biochemical interrogation of individual biochemical steps, we show that all four bromine atoms in **1** are installed by the action of a single flavin-dependent halogenase: Bmp2. Tetrabromination of the pyrrole induces a thioesterase-mediated offloading reaction from the carrier protein and activates the biosynthetic intermediate for decarboxylation. Insights into the tetrabrominating activity of Bmp2 were obtained from the high-resolution crystal structure of the halogenase contrasted against structurally homologous halogenase Mpy16 that forms only a dihalogenated pyrrole in marinopyrrole biosynthesis. Structure-guided mutagenesis of the proposed substrate-binding pocket of Bmp2 led to a reduction in the degree of halogenation catalyzed. Our study provides a biogenetic basis for the biosynthesis of **1** and sets a firm foundation for querying the biosynthetic potential for the production of **1** in marine (meta)genomes.

natural products | biosynthesis | halogenation | enzymology

Marine bacteria of the genus *Pseudoalteromonas* produce numerous small molecule natural products with varied roles in marine chemical ecology (1–4). Recently, tetrabromopyrrole (**1**, Fig. 1) was established as a chemical cue produced by *Pseudoalteromonas* that induces larval settlement and metamorphosis in reef-building Caribbean coral species (3). Additionally, the chemical structure of **1** is notable for its degree of halogenation (one-to-one carbon-to-halogen ratio), unique among naturally occurring aromatic organohalogenes (5).

Halopyrrole-containing microbial natural products are biosynthesized starting from L-proline. A highly conserved halopyrrole biosynthetic route involves oxidation of an acyl carrier protein (ACP)-loaded prolyl side chain and subsequent halogenation by a flavin-dependent halogenase (Fig. 1) (6). At this stage, two biosynthetic routes are possible. In the first route, the ACP-loaded halopyrrole is extended using modular assembly lines to yield nonribosomally synthesized peptide, or polyketide natural products (e.g., **2–4**, Fig. 1) (7–10). During the biosynthesis of **2–4**, and other related natural products, the halopyrrole carboxylic acid is

transthiosterified to downstream ACPs during modular elongation reactions. Molecules arising from these modular biosynthetic pipelines are characterized by the preservation of the prolyl alpha-carboxyl carbon atom as a carbonyl, or embedded in the final natural product skeleton in varying oxidation states (in red, Fig. 1). In a recently discovered alternate route, the prolyl alpha-carboxyl carbon atom is lost during the biosynthesis of **1** and the cytotoxic marine bacterial natural product pentabromopseudilin (**5**, Fig. 1) (4, 11). Uniquely, **5** is biosynthesized via the coupling of 2,3,4-tribromopyrrole (**6**, Fig. 1) to 2,4-dibromophenol rather than via a modular assembly line. The molecular mechanism for the pyrrole offloading from the ACP and elimination of prolyl alpha-carboxyl carbon en route to the biosyntheses of **1** and **5–6** has not been determined. The structure of **1** poses an additional biosynthetic challenge due to the presence of a bromine atom in place of an acyl side chain. Although all previously described pyrrole halogenases

Significance

The majority of pharmaceuticals are inspired by natural product scaffolds that are functionalized by tailoring enzymes, such as halogenases. The degree of halogenation is an important determinant of natural product bioactivity, yet little is known regarding the molecular basis for the exquisite control exhibited by tailoring halogenases. Known pyrrole halogenases commonly perform up to two halogenations on the pyrrole. Our study of tetrabromopyrrole biosynthesis revealed a uniquely adapted halogenase–thioesterase enzyme pair that catalyzes an unprecedented series of halogenations on a pyrrole. Structural comparison of the pyrrole tetrahalogenase to a pyrrole dihalogenase revealed key residues involved in controlling the degree of halogenation. Our findings provide fundamental insights that might be applied in the rational design of biocatalysts toward directed biosynthesis of new chemicals.

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Data deposition: The sequences reported in this paper have been deposited in the GenBank database [accession nos. **KR011923** (*Pseudoalteromonas* sp. P55-derived bmp gene clusters) and **KT808878** (*Pseudoalteromonas* sp. A757-derived bmp gene clusters)]. The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.rcsb.org (PDB ID codes **5BUK** (Mpy16), **5BVA** (Bmp2), and **5BUL** (Bmp2-TM)).

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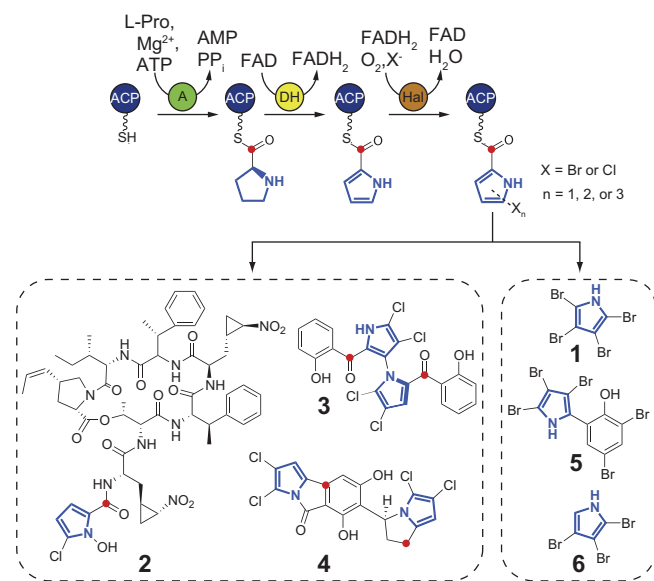


Fig. 1. Structures and biosynthesis of halopyrrole natural products. Halopyrrole biosynthesis starts from the loading of L-proline onto an ACP phosphopantetheine thiol by an adenylation (A) enzyme, followed by a $4e^-$ oxidation of the pyrrole ring to a pyrrole (in blue) catalyzed by a dehydrogenase (DH) enzyme. Subsequent halogenation by a flavin-dependent halogenase (Hal) then installs one, two, or three halogens on the pyrrole ring. At this stage, the thiotemplated halopyrrole can proceed down assembly line biosynthetic routes to afford natural products, such as hormaomycin (2), marinopyrrole A (3), and chlorizidine A (4) (8–10). However, the biosyntheses of 1 and 5–6 do not entail modular assembly line extension of the L-proline-derived halopyrrole, but rather involve the loss of the prolyl α -carboxylate carbon atom (in red) via an unexplained mechanism.

catalyze one, two, or three halogen additions upon the ACP-loaded pyrrole ring, a halogenase capable of four halogenations on the pyrrole ring, as implied by the structure of 1, has not been characterized. These open biosynthetic questions, together with the ecological significance of 1 in coral larval settlement, motivated us to explore the genetic and molecular logic for its biosynthesis in marine bacteria.

Herein, we establish the biosynthesis of 1 by the total in vitro enzymatic reconstitution of its biosynthetic machinery. We show that a single flavin-dependent brominase installs an unprecedented four halogens on an ACP-bound pyrrole required for the progression of thioesterase (TE)-mediated offloading and decarboxylation reactions. We also exploit an opportunity to structurally characterize and investigate via site-directed mutagenesis the molecular basis for a differential number of halogen additions catalyzed by flavin-dependent halogenases on aromatic substrates. Our structural comparison of a highly homologous pyrrole dihalogenase to the pyrrole tetrahalogenase reveals subtle variations in biocatalyst design that lead to divergent molecular outcomes.

Results

Genetic Basis for the Biosynthesis of 1. We recently reported that 1 produced by the marine bacterium *Pseudoalteromonas* sp. PS5 induces settlement of larvae associated with several Caribbean coral species (3). To investigate the genetic basis for the biosynthesis of 1, we sequenced and assembled a 5.08-Mbp draft genome for *P. sp.* PS5. Querying the draft genome for the presence of the halopyrrole biosynthetic genes, we identified a gene locus in *P. sp.* PS5 with high homology to a subset of bromopyrrole biosynthetic genes present in the previously reported *bmp* gene locus from marine bacteria of the genera *Pseudoalteromonas* and *Marinomonas* (11) (Fig. 2A and SI Appendix, Table S1). The organization of the *bmp* homologs identified in the genome of *P.*

sp. PS5 (*PS5_bmp*) is identical to that of the *bmp* gene cluster found in *Marinomonas mediterranea* MMB-1 *bmp* (*Mm_bmp*) and others (11). Notably, the *PS5_bmp* gene locus maintains the bromopyrrole biosynthetic module *Mm_bmp1–4* and the bromopyrrole/phenol coupling cytochrome P450 (CYP450) accessory genes *Mm_bmp9–10*, but lacks the genes associated with bromophenol biosynthesis (*Mm_bmp5–6*) and the CYP450-*Mm_bmp7* (Fig. 2A). In addition, we also sequenced and assembled a 5.13-Mbp draft genome for *Pseudoalteromonas* sp. A757 that also produces 1 (12). Querying the genome of *P. sp.* PS5 as before revealed a gene cluster with high homology to *PS5_bmp* (*A757_bmp*), providing a second example of a stand-alone bromopyrrole biosynthetic pathway from the genus *Pseudoalteromonas* (Fig. 2A and SI Appendix, Table S1).

To evaluate the role of *PS5_bmp1–4* genes in the production of 1, we cloned and heterologously expressed *PS5_bmp1–4* in *Escherichia coli*. Only in the presence of bromide in the culture media, we observed robust heterologous production of 1 along with a minor amount of 6, consistent with the product profile observed in organic extracts of cultures of *P. sp.* PS5 and *P. sp.* A757 (Fig. 2B). Although these results demonstrate that *bmp1–4* genes are necessary for the production of 1, the molecular detail underlying the offloading of the L-proline-derived pyrrole moiety from the ACP, its timing relative to the bromination events, and the chemical logic for the loss of L-proline derived α -carboxyl was not discernible at this stage. Therefore, we next examined the biosynthetic pathway by total in vitro enzymatic reconstitution of the production of 1 using purified enzyme catalysts.

Decarboxylative Bromination of Thiotemplated Pyrrole. To examine the individual roles of *Bmp1–4*, we performed the total in vitro reconstitution of the biosynthesis of 1 (Fig. 3A). Due to the high sequence similarity of *PS5_Bmp1–4* to *Mm_Bmp1–4* (SI Appendix, Table S1), and production of 1 by both *P. sp.* PS5 and *M. mediterranea* MMB-1 (11), we used recombinant *Mm_Bmp1–4* proteins in our in vitro investigations. *Bmp2* and *Bmp4* were individually purified as N-His₆-tagged proteins whereas N-His₆-tagged apo-*Bmp1* was purified in complex with untagged *Bmp3* and converted to its pantetheine-loaded holo-form by the promiscuous *Bacillus subtilis* phosphopantetheinyl transferase Sfp (11, 13) (Fig. 3A). Incubation of L-proline and bromide with purified *Bmp1–4* enzymes, along with flavin and nicotinamide cofactors (FAD and NADP⁺) and cofactor regeneration enzymes (PtdH and SsuE), led to the formation of 1 as the major product (Fig. 3 and SI Appendix,

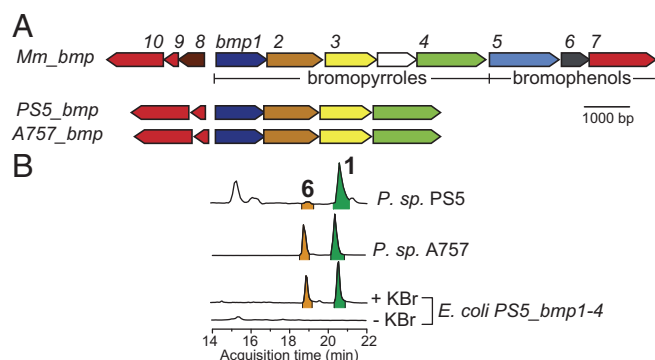


Fig. 2. A genetic basis for production of 1. (A) *Mm_bmp*, *PS5_bmp*, and *A757_bmp* gene clusters with bromopyrrole and bromophenol biosynthetic genes indicated below the *Mm_bmp* gene cluster; a putative permease (uncolored) is inserted between *Mm_bmp3* and *Mm_bmp4*. *Mm_Bmp1–4* are colored per their catalytic roles shown in Fig. 1. Note that *bmp1* encodes a di-domain protein with an ACP domain at the N terminus followed by a TE domain (11). (B) LC/MS extracted ion chromatograms (EICs) for $[M-H]^+$ ions corresponding to 1 and 6 in organic extracts of *P. sp.* PS5, *P. sp.* A757, and *E. coli* expressing *PS5_bmp1–4* grown in media with (+) or without (–) bromide.

reconstitution reactions (*SI Appendix, Fig. S6*). Together with our previous results, we propose that **7** and **6** are both off-pathway products arising, respectively, at stages preceding and after formation of **1** (Fig. 5).

In sum, our data support the Bmp2-catalyzed bromination of **8** leading to transient intermediate **i** as shown in Fig. 5, which would undergo transesterification to generate a Bmp1(TE) S202 side chain-bound oxoester **ii**, thus offloading the pyrrole from Bmp1(ACP). Hydrolytic offloading from Bmp1(TE) would generate the labile alpha-bromo acid **iii**. Driven by the rearomatization of the pyrrole ring, **iii** would spontaneously decarboxylate to **1**. Direct mass spectrometric evidence for **i** proved elusive, suggesting that **i** rapidly converts back to **8** via spontaneous debromination in the absence of Bmp1(TE). Therefore, the role of the Bmp1(TE) could be to trap the tetrabrominated pyrrole species by means of an irreversible transesterification step whereas the fourth bromination by Bmp2 activates the substrate for decarboxylation subsequent to hydrolytic offloading from Bmp1(TE).

Nonenzymatic hydrolysis of **i** may also lead to release of **iii** to afford **1**, consistent with our observation of low levels of **1** from reactions excluding Bmp1(TE) and with inactive Bmp1(TE)S202A. The observation of similar levels of **7** in all reactions further supports direct hydrolytic bypass of TE-mediated offloading. Furthermore, the fourth bromination of **8** by Bmp2, followed by transesterification and decarboxylation of **i**, neatly differentiates the biosynthesis of **1** from nonribosomal peptide synthetase- and polyketide synthase-derived pathways involving halopyrrole units in which the L-proline-derived alpha-carboxylate carbon atom is preserved in the mature natural product (Fig. 1).

Structural Basis for Unique Tetrahalogenating Activity of Bmp2. To realize the scheme shown in Fig. 5, the flavin-dependent brominase Bmp2 would need to halogenate all four positions of the pyrrole ring. This mechanistic requirement for the production of **1** is in stark contrast to all previously characterized pyrrole halogenases that halogenate the pyrrole ring only once or twice. To understand the structural basis for the tetrahalogenating activity of Bmp2, we determined the high-resolution crystal structure of FAD-bound holo-Bmp2 at a limiting resolution of 1.87 Å (Fig. 6A) and compared it with the FAD-bound holo-structure of dihalogenase Mpy16 participating in the biosynthesis of **3** that we determined at 1.95 Å resolution (Fig. 6B and *SI Appendix, Table S2*) (8). Bmp2 and Mpy16 structures are highly homologous with all secondary structural elements in the vicinity of the active site strictly conserved between the two enzymes (*SI Appendix, Fig. S7*). Furthermore, the FAD cofactor isoalloxazine rings and the postulated active site lysine residues (15) (K74 for Bmp2, K72 for Mpy16) (Fig. 6A and B) are superimposable. Although we could not determine a substrate pyrrolyl-S-ACP cocrystal structure for either Bmp2 or Mpy16, the position of the pyrrole binding sites for Bmp2 and Mpy16 was inferred by a structural alignment of Bmp2 and Mpy16 with the crystal structures of substrate-bound forms of the flavin-dependent tryptophan-7-chlorinases PrnA (16) and RebH

(17). Fittingly, the postulated pyrrole-binding site in Bmp2 and Mpy16 (Fig. 6A and B) is in close proximity to the side chain of the catalytic lysine residue. An examination of the amino acids lining this putative pyrrole binding site in Bmp2 reveals a lack of conservation of three key sites (Y302, F306, and A345) that are otherwise strictly conserved among all pyrrolyl-S-ACP halogenases that selectively catalyze one [HrmQ (10)] or two [PltA (7), Mpy16 (8), Clz5 (9), and Pyr29 (18)] halogenations on the pyrrole ring (Fig. 6D). We thus mutated these three residues in Bmp2 to the corresponding residues in Mpy16 (Fig. 6B) to generate Bmp2-Y302S/F306V/A345W triple mutant enzyme (henceforth referred to as Bmp2-TM). We next investigated the effect of the triple mutation on the in vivo heterologous production of **1** in *E. coli*. Coexpression of *Mm_bmp1*, *Mm_bmp3*, and *Mm_bmp4*, with *Mm_bmp2-TM* in *E. coli*, led to a complete loss in the in vivo production of **1** in contrast to robust production with wild-type (WT) *Mm_bmp2* (*SI Appendix, Fig. S9*).

To explore the molecular basis for the in vivo loss in production of **1** by the Bmp2-TM enzyme, we compared the in vitro enzymatic activities of Bmp2, Mpy16, and Bmp2-TM enzymes. We chemoenzymatically synthesized pyrrolyl-S-Bmp1(ACP) and pyrrolyl-S-Mpy15, the requisite substrates for Bmp2 and Mpy16, respectively (8, 11, 14). Using a previously reported mass spectrometry-based proteomic assay that relies on detection of acyl-(cyclo)pantetheine MS2 product ions (14), we verified that WT Bmp2 enzyme catalyzed three brominations on the pyrrole ring (Fig. 6E) and that Mpy16 catalyzed no more than two chlorinations on the pyrrole ring acylated to Mpy15 (Fig. 6F) (14). Note that, although Bmp2 tetrabrominates the pyrrole ring as established by our previous findings, the assay used here detects only up to tribromination of pyrrolyl-S-Bmp1(ACP). In contrast to the WT Bmp2, Bmp2-TM could catalyze only a single bromination on the pyrrole ring (Fig. 6G and *SI Appendix, Fig. S8*), thereby precluding the synthesis of **1**.

Lastly, to eliminate the possibility that the reduced degree of halogenation catalyzed by Bmp2-TM was due to disruption of the enzyme active site by the three mutations, we determined the crystal structure of holo-Bmp2-TM at a resolution of 1.98 Å (Fig. 6C). Structural comparison of Bmp2-TM to Bmp2 and Mpy16 confirmed that the mutation of the three Bmp2 active site residues did not alter the positioning of the FAD isoalloxazine ring or the catalytic lysine side chain.

Discussion

In addition to characterizing the biosynthesis of a microbially produced coral settlement cue, our study reveals previously unidentified enzymology stemming from a biosynthetic motif ubiquitous among halopyrrole-containing natural products. Although our previous work had established tribromination of pyrrolyl-S-Bmp1(ACP) by Bmp2, the enzymatic activities responsible for the addition of the fourth bromine atom in **1** and offloading and decarboxylation of the pyrrolyl moiety from the ACP were unknown (11). Our current work demonstrates that a single halogenase, Bmp2, catalyzes the unprecedented tetrabromination of the ACP-bound pyrrole, which subsequently undergoes Bmp1(TE)-mediated offloading from the ACP and spontaneous decarboxylation. In light of the elucidation of the molecular and genetic details for the construction of **1**, we posit that **1** is an intermediate en route to the production of **5**, reconciling a previous report implicating an L-proline-derived symmetrical pyrrolic intermediate in the biosynthetic scheme for the production of **5** (19). Previously, based on primary sequence homology, we postulated that the enzyme Bmp8 (Fig. 24) participates in the decarboxylation of **7** to produce **6** (11), a hypothetical route that is likely not operative in light of the biochemical data presented above. Therefore, the physiological transformations for the production of **6** en route to **5** remain to be elucidated. Furthermore, the previously reported production of **1** during the CYP450-Bmp7-mediated biradical homodimerization of **6** is likely an off-pathway route, with Bmp1–4 being the primary players in the production of **1**. Successive

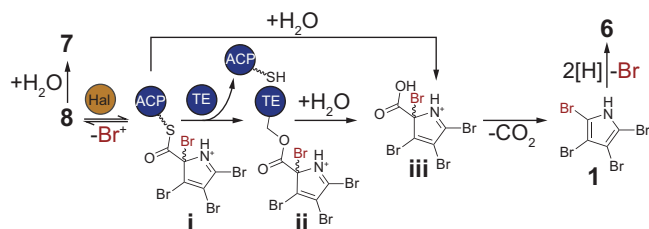


Fig. 5. Scheme for bromination-dependent pyrrole offloading and decarboxylation. Proposed reaction mechanisms for terminal bromination, offloading, and decarboxylation reactions with **8**; inferred intermediates are indicated by bold Roman numerals.

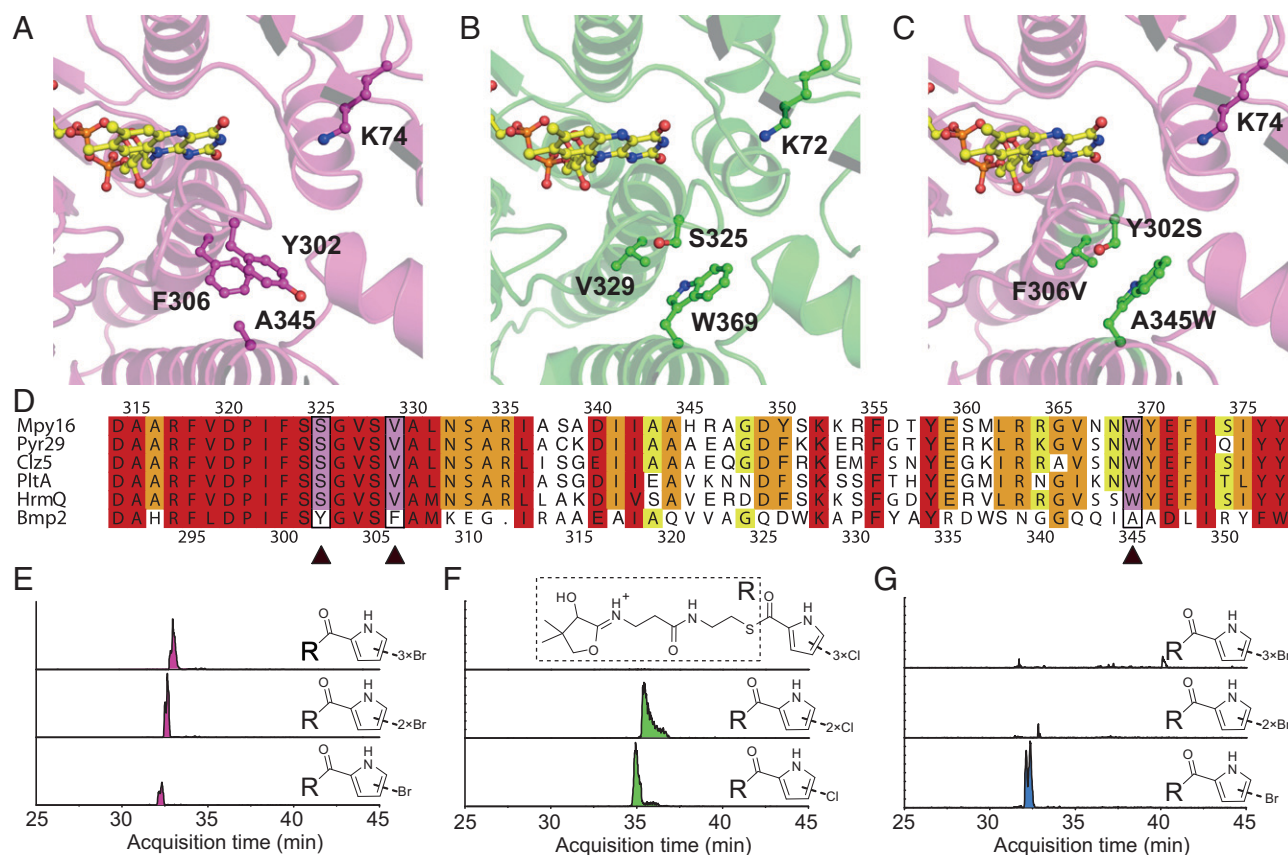


Fig. 6. Structural basis for tetrahalogenation activity of Bmp2. Comparison of the active sites of (A) holo-Bmp2, (B) holo-Mpy16, and (C) holo-Bmp2-TM. Note that side chains of Bmp2-TM residues S302, V306, and W345 in C structurally map to the corresponding side chains of Mpy16 residues S325, V329, and W369 shown in B. (D) Primary sequence alignments of tetrahalogenase Bmp2 with mono- and dihalogenating pyrrolyl-S-ACP halogenases demonstrate that the amino acids that were mutated on the basis of structural comparison with form Bmp2-TM (indicated by ▲) are conserved in all pyrrolyl-S-ACP halogenases but Bmp2 (see *SI Appendix, Fig. S20* for alignment of Bmp2 homologs from different bacterial species). EICs for acyl-(cyclo)pentathione MS2 product ions demonstrate that (E) WT Bmp2 generates mono-, di- and tribromopyrrolyl-S-Bmp1(ACP) products whereas (F) Mpy16 generates only mono- and dichloropyrrolyl-S-Mpy15, and (G) Bmp2-TM generates only monobromopyrrolyl-S-Bmp1(ACP) product.

biochemical studies promise to address these open questions to characterize all steps in the biosynthesis of **5** and assign physiological roles for each of the Bmp enzymes.

The complete in vitro reconstitution of the production of **1** using purified enzyme catalysts suggests a mechanism for the acylphosphopantetheine thioester to be sequestered and thereby inaccessible to Bmp1(TE) until the terminal fourth bromination. Sequestration of a pyrrole tethered to a type II peptidyl carrier protein (PCP) was recently demonstrated for a highly homologous PCP-PltL [57% amino acid similarity to Mm_Bmp1(ACP)] participating in the biosynthesis of the dichloropyrrole-containing natural product pyoluteorin (7, 20). By analogy, in the biosynthesis of **1**, it is possible that the terminal halogenation on the pyrrole partially liberates the sequestered substrate from the ACP, allowing access to the thioesterase. Therefore, the terminal fourth bromination by Bmp2 serves the dual role of triggering the release of the sequestered pyrrole moiety by presumably making the thioester susceptible to hydrolysis, in addition to activating the substrate for elimination of the L-proline-derived alpha-carboxyl. Activating halogenation reactions in natural product biosynthetic pathways has been demonstrated in the biosynthesis of NRPS-PKS hybrid curacin A, in which a cryptic halogenation catalyzed by α -ketoglutarate-dependent halogenase-CurA promotes cyclopropane ring formation (21), and in the biosynthesis of meroterpenoid merochlorins A and B, in which a dearomatization/terpene cyclization reaction cascade is initiated by a chlorination catalyzed by vanadium-dependent haloperoxidase-

Mcl24 (22). Most analogously, paralleling the decarboxylative bromination strategy used by Bmp2, flavin-dependent brominase-Bmp5 catalyzes a bromination reaction that drives decarboxylation in the conversion of free *p*-hydroxybenzoic acid to 2,4-dibromophenol, the bromophenol building block of **5** (11).

The tetrabromination activity of Bmp2 is unprecedented among the flavin-dependent halogenases described to date that selectively add a specific number of halogen atoms to an aromatic substrate. In this study, we embraced the opportunity to interrogate the structural basis for this halogenation control in flavin-dependent pyrrolyl-S-ACP halogenases. Although the previously reported 2.1-Å crystal structure of putative tyrosyl-S-ACP flavin-dependent halogenase CndH highlighted the general differences in the architectures of flavin-dependent halogenases acting on free versus ACP-bound substrates (23), our study sheds mechanistic insight into the tuning of the enzyme active site in the context of confirmed biochemistry. Comparison of the putative substrate-binding cavities of the highly conserved structures of Bmp2 and Mpy16 resulted in a catalyst, Bmp2-TM, exhibiting an altered halogenation profile. Notably, the active site of Bmp2-TM exhibited no perturbation of the bound FAD cofactor with respect to WT Bmp2, nor did it lead to any apparent change in halogen binding properties of the enzyme, as demonstrated by conservation of its specificity for bromide. Already, for flavin-dependent tryptophan halogenases, it has been demonstrated that amino acid side chains that constitute the substrate binding site control the regiochemical outcomes for halogen additions (17, 24, 25). Our findings extend

this observation to flavin-dependent halogenases that catalyze halogenation of aromatic substrates acylated to ACPs in demonstrating that side chains of residues lining the putative halogenase active site play a role in controlling substrate access, and potentially in specifying the positions on the pyrrole ring that are accessible to halogenation. Unfortunately, in the case of Mpy16, efforts to alter the putative substrate-binding cavity to resemble that of Bmp2 resulted in insolubility of the mutant Mpy16 enzymes. Nonetheless, our results demonstrate that the active sites of otherwise highly homologous flavin-dependent halogenases are uniquely evolved to afford distinct product profiles. Biological activities of natural products are influenced by the number of halogens decorating their organic scaffolds (26, 27), and it would seem that Nature has taken note through the evolution of highly specialized halogenation catalysts.

In light of the existing literature and the findings of this study, several open questions remain regarding enzymatic halogenation (28). Of note is the question of the binding mode of both the ACP and the aromatic substrate by flavin-dependent halogenases that require acyl-S-ACP substrates. In addition, the primary question across all classes of halogenating enzymes is the structural determinant for halide specificity among these catalysts. Although Bmp2 and Mpy16 demonstrate different halide specificities, the amino acid side chains and the positioning of the FAD isoalloxazine ring relative to these side chains are remarkably conserved. Furthermore, the conservation of halide specificity in Bmp2-TM with respect to the WT enzyme raises the possibility that halide specificity in flavin-dependent halogenases is dictated not only by steric factors, such as halide ion radii, but also by the magnitude of the enthalpic penalty associated with desolvation of the halide ion in the halogenase active site before $2e^-$ oxidation to generate the electrophilic halonium. Furthermore, the contribution of the redox potential of the flavin isoalloxazine ring necessary to oxidize the halide, and the stability of the activated lysine-amine

intermediate generated en route to the transfer of the halonium to the aromatic substrate should be considered (15). As such, the currently proposed halide binding site for flavin-dependent halogenases is itself debatable (29), thus underscoring the challenges associated with teasing apart the role of the enzymatic halogenase catalyst in each of the requisite mechanistic steps. Discovery and characterization of additional flavin-dependent halogenases promises to provide opportunities to answer these mechanistic questions, ultimately leading to engineerable toolkits to tailor the biosynthesis of halogenated natural products (30).

Materials and Methods

Detailed materials and methods are provided in *SI Appendix, Materials and Methods*.

Database Deposition Information. The sequences for *P. sp.* P55- and *P. sp.* A757-derived bmp gene clusters have been deposited in GenBank under accession numbers KR011923 and KT808878, respectively. Structures for Mpy16, Bmp2, and Bmp2-TM are deposited in the Protein Data Bank (www.rcsb.org) under accession numbers 5BUK, 5BVA, and 5BUL, respectively.

Note. During the review and publication of this manuscript, the crystal structure of the flavin-dependent halogenase PItA was reported in the literature (31).

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