



Terpene biosynthesis in marine sponge animals

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Sea sponges are the largest marine source of small-molecule natural products described to date. Sponge-derived molecules, such as the chemotherapeutic eribulin, the calcium-channel blocker manoalide, and antimalarial compound kalihinol A, are renowned for their impressive medicinal, chemical, and biological properties. Sponges contain microbiomes that control the production of many natural products isolated from these marine invertebrates. In fact, all genomic studies to date investigating the metabolic origins of sponge-derived small molecules concluded that microbes—not the sponge animal host—are the biosynthetic producers. However, early cell-sorting studies suggested the sponge animal host may play a role particularly in the production of terpenoid molecules. To investigate the genetic underpinnings of sponge terpenoid biosynthesis, we sequenced the metagenome and transcriptome of an isonitrile sesquiterpenoid-containing sponge of the order Bubarida. Using bioinformatic searches and biochemical validation, we identified a group of type I terpene synthases (TSs) from this sponge and multiple other species, the first of this enzyme class characterized from the sponge holobiome. The Bubarida TS-associated contigs consist of intron-containing genes homologous to sponge genes and feature GC percentage and coverage consistent with other eukaryotic sequences. We identified and characterized TS homologs from five different sponge species isolated from geographically distant locations, thereby suggesting a broad distribution amongst sponges. This work sheds light on the role of sponges in secondary metabolite production and speaks to the possibility that other sponge-specific molecules originate from the animal host.

natural products | terpene synthase | marine sponge | biosynthesis

Marine sponges are the richest source of bioactive small molecules from the ocean, accounting for nearly 30% of all known marine natural products (1). Over 9,000 sponge-derived molecules have been characterized to date and represent nearly all biosynthetic molecular classes, such as polyketides, peptides, alkaloids, and terpenoids (1, 2). Not only do sponge-derived compounds serve important ecological roles as feeding deterrents, antioxidants, and antifouling agents, but they also feature impressive medicinal activity, such as the antimalarial activity of kalihinol A, the antitumor activity of avarol, and the anti-inflammatory, antimicrobial, and analgesic activities of manoalide. (Fig. 1A) (3–11). In recent times, the vast majority of investigations into the biosynthetic origins of sponge-derived molecules have focused on the sponge microbiome—the consortium of bacteria, fungi, archaea, and viruses that live in the sponge host—and not on the sponge animal host itself (*SI Appendix, Table S1*) (12–21). To date, all biosynthetic genes responsible for sponge-derived natural product production have been found in microbial symbionts, which has led to a paradigm that the sponge microbiome is the general source of these molecules.

Pregenomic era cell sorting studies determined that certain sponge metabolites are enriched in sponge animal cells, while others are concentrated in microbial symbiont cells. These experiments suggested that both the sponge animal and its microbiome control the production of sponge-derived metabolites (23–26); however, very few recent studies have followed up on the sponge animal's role in specialized metabolite production. Notably, most of the sponge cell-associated metabolites were identified as terpenoids, making this class of sponge-derived molecules an appealing target for the study of sponge animal biosynthesis.

Terpenoids are the largest known class of secondary metabolites (12, 27), and sponges are an especially rich source, with approximately 4,100 of the 9,400 reported sponge natural products belonging to this compound class (2). Biosynthetically, terpenoids are derived from linear oligoprenyl diphosphate precursors, which are assembled from C₅ isoprenoid building blocks by isopentyl diphosphate synthases (IDSs). These linear precursors are cyclized by terpene synthases (TSs) through cationic cascade reactions initiated by either diphosphate elimination (type I) or protonation (type II) (28). Across various lifeforms, type II TSs are generally more common and conserved than type I TSs, since

Significance

Over 9,000 unique molecules have been isolated from marine sponges, including several bioactive molecules such as the calcium-channel blocker manoalide, antimalarial compounds kalihinol A and axisonitrile-3, and anticancer drug eribulin. Until now, all recent work on the origins of bioactive sponge-derived molecules has indicated that they are made by the symbiotic microorganisms that live within the sponge animal host. Herein, we describe type I terpene synthases characterized from the sponge holobiome and report that these enzymes originate from the sponge animal itself and not a microbial symbiont.

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type II TSs function in the production of key cellular molecules such as sterol triterpenoids, which are involved in membrane stabilization, hormonal signaling, and other essential processes (29). Accordingly, the majority of (>3,000) sponge terpenoids likely originate from type II terpene synthase chemistry, most of which are steroids, carotenoids, and meroterpenoids. We estimate 6% (~550) of all sponge natural products originate from type I TSs. Most type I TS-derived terpenoids from plants, fungi, and bacteria are usually oxidatively tailored by cytochrome p450s and short-chain dehydrogenases (28, 30, 31). In sponges, notably, roughly 2/3 of probable type I TS-type terpene products belong to the sponge-exclusive class known as nitrogenous terpenes, which feature nitrogen-containing functional groups like isonitriles, isothiocyanates, and formamides that originate from inorganic cyanide (32, 33).

Type I TSs are found in a wide variety of organisms but are scarce in the animal kingdom, where they have, so far, only been described in insects (34–37) and most recently octocorals (38, 39). While all type I TSs feature a conserved IDS-type α -helical fold and are thought to share an ancient evolutionary origin with IDSs (40), insect type I TSs appear to have more recently evolved independently from duplicated IDSs and, therefore, do not bear much sequence similarity to canonical type I TSs (34–37). On the other hand, coral TSs may be the result of ancient horizontal gene transfer from bacteria, given that they share a similar overall structure with key active site residues but still form their own distinct type I TS clade (34). While rare in animals, the presence of type I TSs in insects and corals does suggest that the sponge host animal could potentially house the genes necessary to produce terpenoids. Early radiolabeled feeding work investigated the biosynthesis of sponge nitrogenous terpenoids and, though key information about the origin of the nitrogen-containing functional groups was obtained, the origin of the terpenoid backbone could

not be established (33, 41–48). As far as we know, no studies have successfully characterized a TS from a sponge or a sponge-associated microbe, leaving a gap in our understanding of the origin and enzymology underlying sponge terpene biosynthesis.

To address this fundamental gap, we investigated terpeneoid biosynthesis in an undescribed San Diego sponge in the order Bubarida [formal species description in process (49), previously identified as *Axinella* sp. (50)], which is known to contain isonitrile sesquiterpenoids (Fig. 1 *B* and *C*) (50). By performing metagenomic and transcriptomic sequencing of the sponge and its microbiota, we identified several candidate TS genes resident on sponge genomic contigs. Heterologous expression of these putative TSs and homologs from other *Agelas*, *Stylissa*, and *Phakellia* sponges demonstrated the activity of seven type I sesquiterpene synthases, which are amongst the first biochemically characterized small molecule biosynthetic enzymes of sponge animal origin.

Results

Metagenome Sequencing and Assembly. We first queried publicly available sponge metagenomes for canonical microbial and plant-type terpene synthases. We also queried all four sponge genomes available on National Center for Biotechnology Information (NCBI)—including the *Amphimedon queenslandica* reference genome (51)—as well as genomes from the widespread sponge symbiont genus *Entotheonella* (52). While we readily identified multiple type II TSs, we detected only one putative type I TS gene, from a *Lamellodysidea herbacea* sponge-derived “*Candidatus* Paraprochloron terpiosi SP5CPC1” Metagenome Associated Genome (accession number GCA_014323965.1) (53, 54). Given the broad distribution of characterized sponge terpenoids, we were surprised we did not observe more candidate type I TSs in sponge

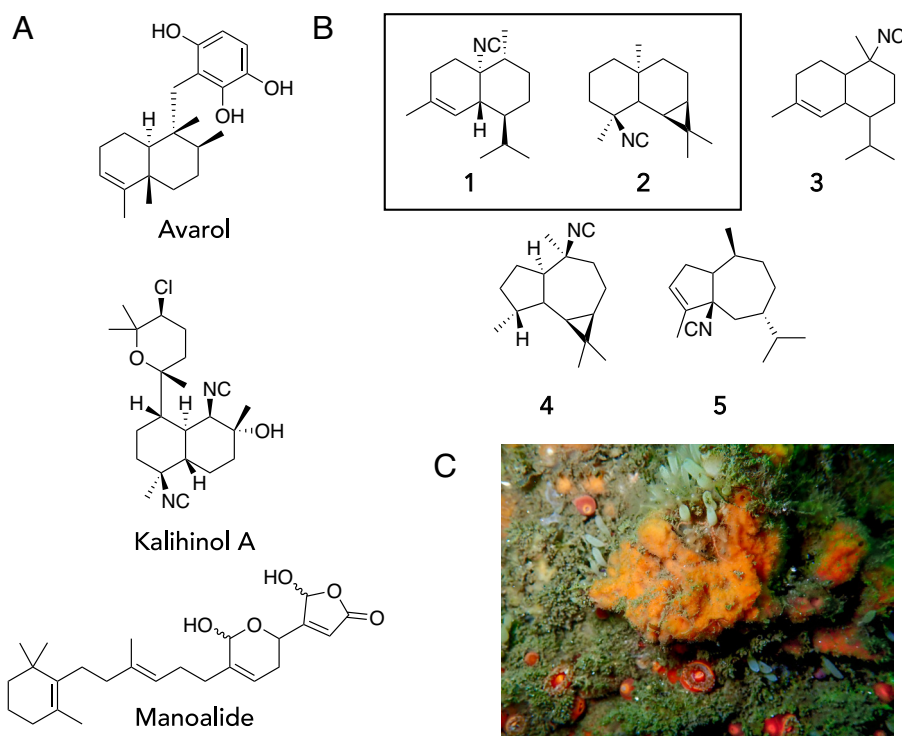


Fig. 1. Terpenoids from marine sponges. (A) Bioactive sponge-derived terpenoids: avarol from *Dysidea avara*, kalihinol A from *Acanthella* sp., and manoalide from *Luffariella variabilis*. (B) Reported isonitrile sesquiterpenes **1** to **5** from Bubarida sponges from San Diego. Boxed compounds are molecules validated from the uncharacterized Bubarida sample that was sequenced in this study and represent relative stereochemistry only (22). (C) in situ uncharacterized Bubarida sponge from San Diego.

metagenomes, which we hypothesized was because of the small number of public sponge sequences and a lack of sequences with paired metabolomics data indicative of terpenoids. To address these challenges, we sequenced the metagenome and metatranscriptome of a sponge known to produce isonitrile sesquiterpenoids **1** to **5** (50), belonging to an undescribed species of the order Bubarida.

We collected the San Diego Bubarida sponge by SCUBA and performed gas chromatography mass spectrometry (GCMS) analysis of crude sponge extract, which indicated the presence of several previously reported isonitrile sesquiterpenoids **1** to **5** (*SI Appendix, Figs. S1–S3*) (50). We further independently established the structures of **1** and **2** using an isonitrile chemoselective chlorooxime probe (22). Next, we generated a hybrid metagenomic assembly from a combination of Illumina paired-end (PE250) and Oxford Nanopore reads, as well as a metatranscriptomic assembly from Illumina data (*SI Appendix, Methods and Table S2*). The hybrid assembly features 5.5k contigs larger than 5k bp, totaling 180M bp. The N50 value was 61 kb, and the maximum contig size was 960 kb (*SI Appendix, Table S2 and Fig. S4*).

Terpene Synthase Genome Mining and Identification. To guide our bioinformatic analysis, we hypothesized that the TSs responsible for sesquiterpene production belonged to one of three previously described groups: 1. Canonical microbial type I TSs from a microbial symbiont; 2. IDS-like TSs akin to those found in insects; or 3. Coral-like type I TSs in the sponge animal. We first searched for canonical microbial type I TSs in our metagenome using BLAST and HMM searches with published TS HMMs, which were solely built from either plant or bacterial TSs (55, 56); however, these initial searches did not uncover any promising candidate TSs from the sponge microbiome. We next investigated the IDSs within the metagenome to search for evidence of neofunctionalization, like that seen in insect TSs. We identified three intron-containing IDS homologs in our metagenomic assembly that we heterologously expressed in *Escherichia coli* and found to perform typical IDS biochemistry. Two functioned as canonical farnesyl diphosphate (FPP) synthases, while the third functioned as a geranylgeranyl diphosphate (GGPP) synthase (*SI Appendix, Figs. S5–S11*). None had terpene cyclase-like activities.

Next, we queried the sponge metatranscriptome with our custom type I TS HMM (TSHMM1), which we created from a diverse set of characterized fungal, bacterial, plant, and recently discovered coral TSs (*SI Appendix, Table S3*) (38). We hypothesized this more expansive TS HMM would cast a broader net than established TS HMMs to help uncover unusual TSs that were not revealed in earlier searches. Using TSHMM1, we identified 20 transcripts (representing eight unique ORFs) with relatively weak E-values ($>10E-7$) that contained key terpene synthase amino acid sequence motifs (i.e., DDXXD, NSD/DTE, and RY) and resembled other type I TSs based on homology modeling (*SI Appendix, Tables S3 and S4 and Fig. S12*). Notably, we detected transcripts derived from five unique putative type I TS genes in the polyadenylation-enriched transcriptome, one of which had a poly-A tail (uBuTS-10), suggestive of eukaryotic origin (*SI Appendix, Fig. S13*). We mapped four of the putative TSs from the transcriptome to our assembled metagenome, and we additionally identified six unique putative type I TSs only found in the metagenome (*SI Appendix, Table S4*). After identifying several single-exon candidate gene sequences in the metagenomic dataset, we examined the gene neighborhoods of these putative TSs on assembled metagenomic contigs. We determined that several of the surrounding genes contained introns and had homology to genes from *A. queenslandica* and other publicly

available sponge animal genomes (Fig. 2A and *SI Appendix, Table S5*). The GC percentages and coverages of these TS-containing contigs were consistent with contigs from our assemblies predicted to belong to the sponge animal host (Fig. 2B). Taking all these features into consideration, we conclude that these genes are part of the sponge animal host genome.

We repeated our TSHMM1 search on several other public and in-house sponge metagenomic datasets. We identified two additional putative TSs from two in-house metagenomes of Caribbean *Agelas clathrodes* and *Agelas tubulata*, sponges which are not known to produce sesquiterpenoids. By searching all publicly available Sequence Read Archive sponge entries, we also identified a complete putative TS from a Norwegian *Phakellia ventilabrum* and two overlapping partial TS sequences from a Guamanian *Stylissa massa* (*SI Appendix, Fig. S14*) (57, 58). Neither of these sponges had paired metabolomics data available to verify the presence of terpenoids; however, both *S. massa* and several different sponge species from the *Phakellia* and *Stylissa* genera have previously been reported to contain isonitrile terpenoids (59–61). Interestingly, other Norwegian *P. ventilabrum* specimens from the same area as the sequenced TS-containing *P. ventilabrum* specimen have been shown to contain isonitrile sesquiterpenoids (62). *P. ventilabrum* has also recently been reported to form a subclade with the San Diego uncharacterized Bubarida sponge and *Axinella cannabina*, another nitrogenous sesquiterpene-producing sponge (49).

In Vitro Validation of Sponge Terpene Synthase Function.

To investigate the activity of these candidate sponge TSs, we heterologously expressed 13 codon-optimized TS genes from five sponge species in *E. coli* BL21 (DE3). We purified the recombinant proteins and performed in vitro assays with geranyl diphosphate (GPP), FPP, and GGPP. Seven of the 13 TSs showed sesquiterpene synthase activity with selectivity for FPP, five showed low-level monoterpene synthase activity with GPP, and one showed diterpene synthase activity with GGPP (Fig. 3 and *SI Appendix, Table S4 and Figs. S15–S35*). Five of the active sesquiterpene synthases, which are named according to the taxonomy of their sponge of origin, (uBuTS-1, uBuTS-2, and uBuTS-3 from the undescribed Bubarida, AgTS-1 from *A. clathrodes*, and PhTS-1 from *P. ventilabrum*) displayed high product specificity, while the other two (uBuTS-4 and uBuTS-5) produced a mixture of sesquiterpene products (*SI Appendix, Figs. S15–S21*). The products of sponge sesquiterpene synthases uBuTS-1, uBuTS-2, and AgTS-1 were isolated from scaled-up *in vitro* assays and purified, and their structures were characterized by NMR spectroscopy and polarimetry to be (+)-bicyclogermacrene (**6**), (–)-germacrene D (**8**), and (+)-alloaromadendrene (**7**), respectively (Fig. 3 and *SI Appendix, Supplementary Note*). The products of uBuTS-3 and PhTS-1 were determined via GCMS using purified **6** and **8** from uBuTS-1 and uBuTS-2, respectively, as standards. uBuTS-4 and uBuTS-5 produced several sesquiterpenes distinct from **6** to **8** which were observed by GCMS but not fully characterized (*SI Appendix, Figs. S18 and S19*). Four of the TSs with low-level monoterpene synthase activity produced only linear myrcene and ocimene type products (uBuTS-2, uBuTS-3, uBuTS-4, and AgTS-1) and one produced both cyclized and linear monoterpenes (PhTS-1) (*SI Appendix, Figs. S24–S26, S32, and S34*). StTS-1 from *Stylissa* produced a mixture of diterpenes when incubated with GGPP (*SI Appendix, Fig. S35*). The remaining five enzymes were not active with the tested substrates or had low levels of expression (*SI Appendix, Figs. S22–S35*).

To better understand the sponge TSs in the context of other known type I TSs, we collected sequences of type I TSs with characterized activity from several domains of life, including

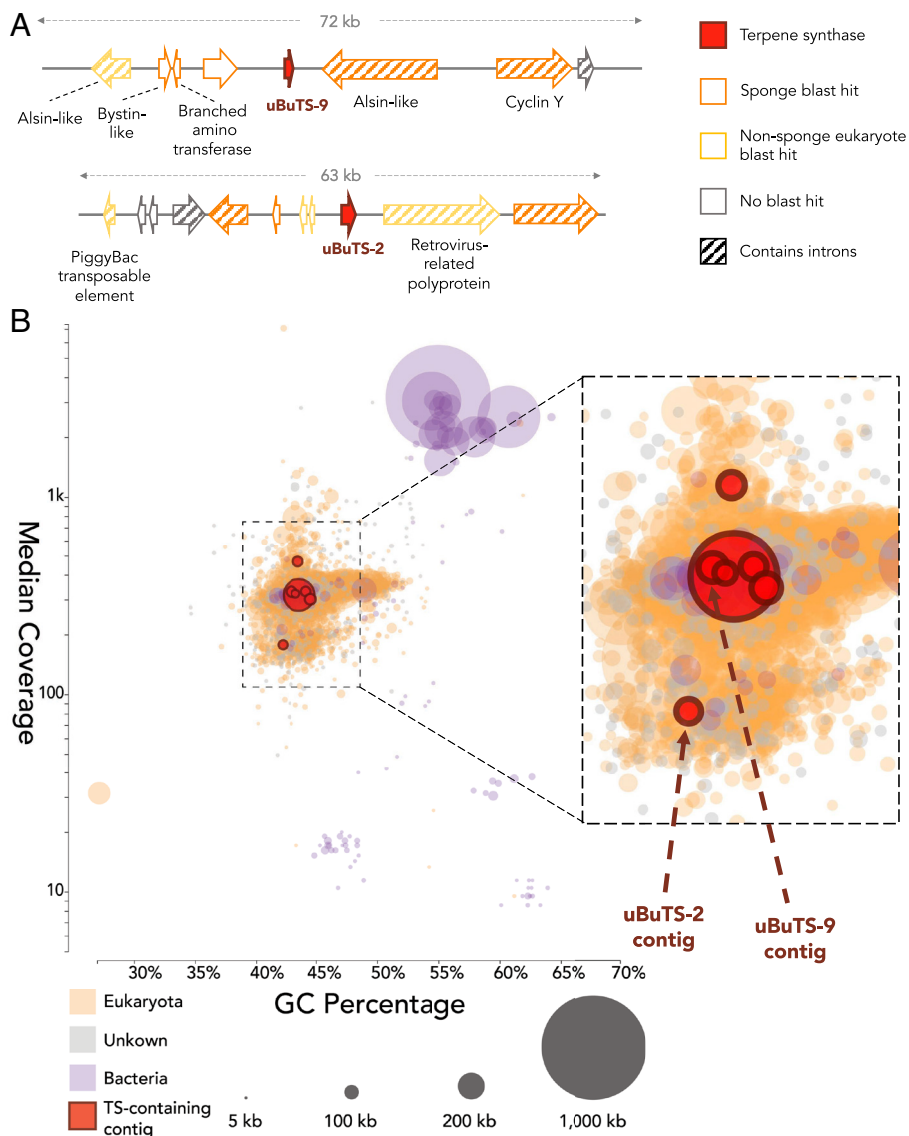


Fig. 2. Eukaryotic features of terpene synthase-containing contigs. (A) TS-containing contigs from the uncharacterized Bubarida sponge. Annotations based on closest blastx hit from the NCBI GenBank nonredundant database (*SI Appendix, Table S5*). (B) GC percentage vs. median coverage plot for all uncharacterized Bubarida metagenomic contigs greater than 5 kb in length. Each circle represents an individual contig. The size of the circle indicates the length of the contig, and for contigs not harboring a TS, the circle's color indicates taxonomic assignment as inferred by homology. Total contig count, sum, contig length, and n50 are as follows: Eukaryota: 2,700, 140 Mb, 74 kb; Unknown: 433, 7.1 Mb, 21 kb; Bacteria: 122, 8.8 Mb, 180 kb. TS-containing contigs: 7, 650 kb, 74 kb. Maximum contig size is 960,860 kb.

sequences of several IDSs as an outgroup, and conducted a phylogenetic analysis. The sponge TSs distinctly clade apart from all known terpene synthases and form a monophyletic group (Fig. 3). Despite clading separately from other TSs, they still share many of the key residues that define canonical TSs (Fig. 4). The sponge TSs have the standard aspartate-rich DDXXD motif, which is characteristic of TSs and IDSs and critical for metal binding. Canonical TSs also contain an additional metal binding region exemplified by (N/D)DXX(S/T)XXX(D/E), commonly known as the NSD/DTE motif, which has been hypothesized to arise from the second aspartate-rich DDXXD motif found in IDSs (40, 63). Twelve of the 13 characterized sponge TSs share an unusual NDXXGXXXD motif which aligns with the traditional NSD/DTE domain. While rare in microbial TSs, this unusual Gly residue in the “middle” Ser/Thr position is seen in plant TSs (64). All of the active sponge TSs we characterized feature an RY dimer near the C terminus that's also present in microbial and octocoral terpene synthases but is not present in known plant terpene synthases (27).

The Role of Sponge TSs in Isonitrile Sesquiterpenoid Biosynthesis.

Given the predominant terpenoid natural products in the Bubarida sponge are isonitrile sesquiterpenoids, including **1** and **2** that resemble uBuTS products **9** and **7**, respectively, we performed exploratory experiments to investigate the role of the newly discovered sponge TSs in the biosynthesis of isonitrile sesquiterpenoids. Garson and colleagues have proposed that inorganic cyanide is the source of the isonitrile and other nitrogenous functional groups unique to sponge terpenoids, by means of radiolabel feeding experiments (33, 41–48, 65). They proposed that a rearranged terpene carbocation intermediate generated by a TS during cyclization might react with the cyanide ion. To test this hypothesis, we performed a series of *in vitro* TS assays and controls with FPP and potassium cyanide. We tested the 13 sponge TS enzymes, including the five TS enzyme homologs that were not functional in the terpene cyclase assay. In all cases, we did not observe the production of nitrogenous sesquiterpenoids by GCMS in the enzymatic assays or noenzyme controls (*SI Appendix, Figs. S36–S47*). We also combined uBuTS1–9 and ran an *in vitro* assay with FPP and

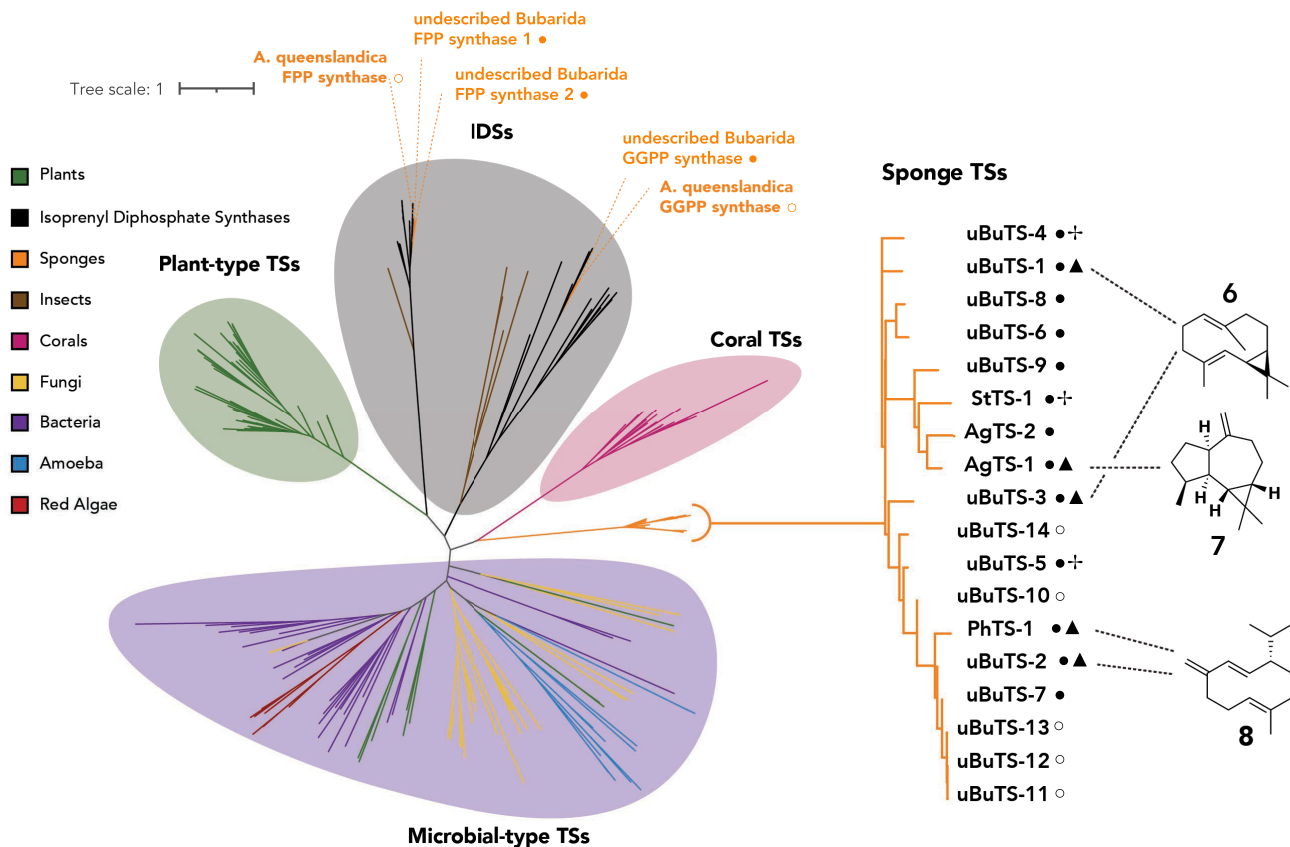


Fig. 3. Phylogenetic tree of selected type I TSs from several domains of life. Isoprenyl diphosphate synthases (IDSs) are included as an outgroup. The first two or three letters of each TS name represent the taxonomy of sponge from which it originated. Enzymes with ● have been biochemically tested, while those with ○ have not been tested. The major sesquiterpene products of enzymes marked with ▲ were fully characterized (*SI Appendix, Supplementary Note*), whereas enzymes with + produce a mixture of terpenes identified only by GC-MS (*SI Appendix, Figs. S18, S19, and S35*). Entries marked with only ● did not produce terpenes under our experimental conditions.

cyanide to test whether there was cross-reactivity between any of the nine uBuTS enzymes, but again we observed only the unmodified sesquiterpenes (*SI Appendix, Fig. S48*). These results suggest that a separate enzyme is required for the regio- and stereoselective installation of the isonitrile. We inspected the gene neighborhoods of the uBuTS genes for clustered genes that may

be involved in cyanide generation or installation; however, no candidates were identified.

To further interrogate the role of cyanide in this system, we tested the sponge tissue for the presence of free cyanide using 4-(4,6-dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride, a selective reagent that allows for cyanide detection by

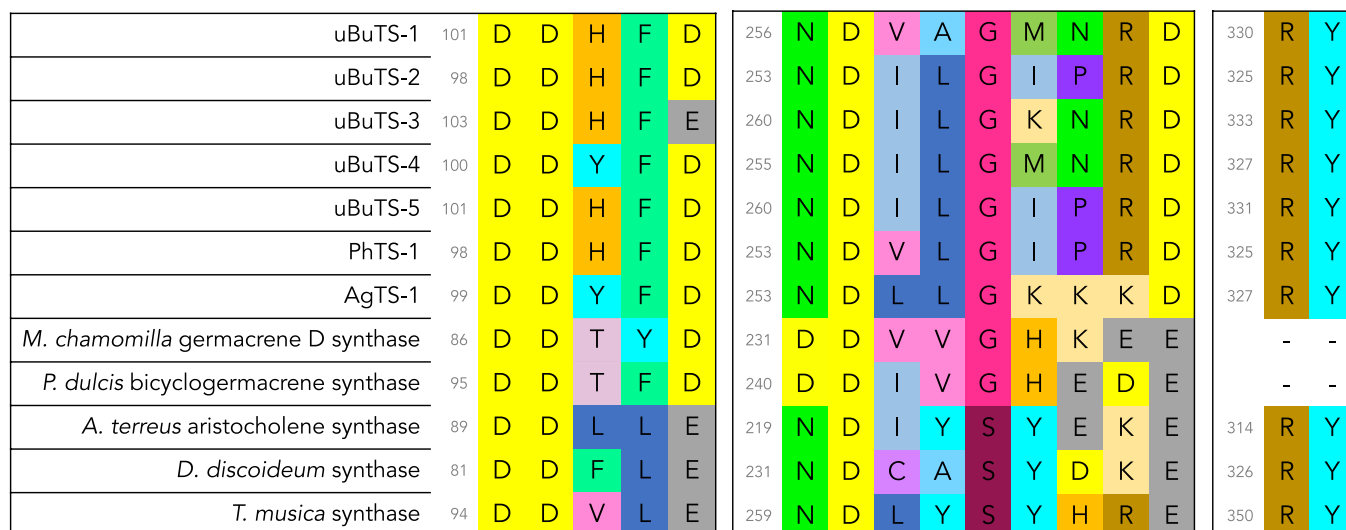


Fig. 4. Multiple sequence alignment of key active site residues for sponge TSs and selected plant, microbial, and coral TSs.

GCMS (66). Freshly collected Bubarida sponge tissue did not contain a detectable amount of cyanide (*SI Appendix, Fig. S49*). Additional work to understand the biosynthesis of isonitrile sesquiterpenoids is ongoing.

Discussion

We have discovered a new class of type I TSs from marine sponges. The Bubarida TS genes are surrounded by intron-containing genes that closely resemble genes from publicly available sponge genomes and are located on contigs with nucleotide compositions (percent GC) and coverages characteristic of other contigs matching eukaryotic, rather than bacterial sequences. Additionally, the polyadenylation of a TS transcript in our RNA-seq data and the fact we identified type I TSs in multiple sponge species from geographically distant locations strongly support the origin of these TS sequences in the sponge animal host and not a microbial symbiont. The discovery of sponge animal TSs contradicts the current paradigm that microbes are the only source of natural products isolated from sponges. Besides octocorals and insects, sponges are now the third class of animals recognized to biosynthesize specialized terpenes other than steroids and carotenoids.

Given that marine sponges have been studied as sources of bioactive natural products—including hundreds of terpenoids—over 70 y, it is notable that TSs were identified in only a few sponges. We believe that this is due to a scarcity of sponge-specific genomic and transcriptomic data, as relatively few species of sponges have any sequencing data associated with them, and even fewer projects focus on the sponge animal as opposed to the associated microbiota. Additionally, nitrogenous terpene production is largely concentrated in a relatively small number of sponges belonging to the order Bubarida (67), for which sequencing data are especially sparse. Furthermore, the order Bubarida is a recently erected and highly debated order that arose because of genetic evidence conflicting with earlier morphology-based classifications of *Axinella*, *Acanthella*, *Phakellia*, *Aximyssa*, and other nitrogenous terpene-producing genera (49, 68). It has even been suggested that nitrogenous terpenes are a chemotaxonomic marker of Bubarida (67), and the monophyletic nature of the newly identified TS sequences points to one common ancestor sequence that was inherited with evolutionary radiation of sponge species. Therefore, our unexpected discovery of functional TSs encoded by sponges outside of Bubarida and not previously reported to produce terpenoids suggests either a need for

further expansion of the Bubarida order or perhaps that this terpene producing capacity may be more widespread than currently appreciated. Unlike coral TSs, which are ubiquitous across all octocorals and are descended from an ancestral TS in the last common octocoral ancestor, sponge TSs are confined to a smaller group of sponge species. Our observation suggests that these TSs may have evolved more recently to serve a specific evolutionary need; however, more research is needed to access the greater distribution and evolutionary history of these genes. As such, further enzyme discovery efforts could greatly benefit from the increased pairing of public sequencing and metabolomics datasets (69).

Many terpenoids found in sponges are uniquely functionalized with isonitrile and related nitrogen-containing functional groups like isothiocyanates, formamides, isocyanates, and amines, suggesting the presence of enzymes that stereoselectively install these functional groups onto terpene hydrocarbons. We hypothesize that the sesquiterpenes produced by the Bubarida TS enzymes are intermediates in the biosynthesis of the isonitrile natural products (22, 50) given their uncanny structural resemblance and the fact *Axinella cannabina*, another nitrogenous terpene-producing sponge closely related to the San Diego Bubarida, is reported to produce bicyclogermacrene **6** (70). The genes and enzymes responsible for these distinctive sponge functionalizations, however, remain unknown, though work from Garson, Scheuer, and others has suggested the origin of these functional groups is inorganic cyanide (44–47). Our observation that the sponge TSs do not install the isonitrile functionality upon the addition of cyanide to TS enzyme assays suggests that there is at least one additional enzyme involved in this biotransformation. We posit that the mechanism to generate **1** from **8** and **2** from **6** proceeds via protonation of an alkene, ring formation, and carbocation quenching by free cyanide to form the isonitrile (Fig. 5), a reaction that may be catalyzed by an enzyme akin to a type II TS, which are known to perform cyclizations on isoprenoid substrates (28). Currently, there is no known precedent for such an “isonitrile synthase” which utilizes cyanide as a substrate; however, the study of enzymes such as FlvF, a repurposed terpene cyclase involved in C-N bond formation, could provide insight into how an isonitrile synthase might evolve from isoprenoid biosynthesis machinery (71). While plant, bacteria, and fungi TS genes often have genes coding for tailoring enzymes like cytochrome P450s clustered near them on the genome (72), the Bubarida sponge TSs do not cluster with obvious candidate tailoring enzyme genes. There is still much to

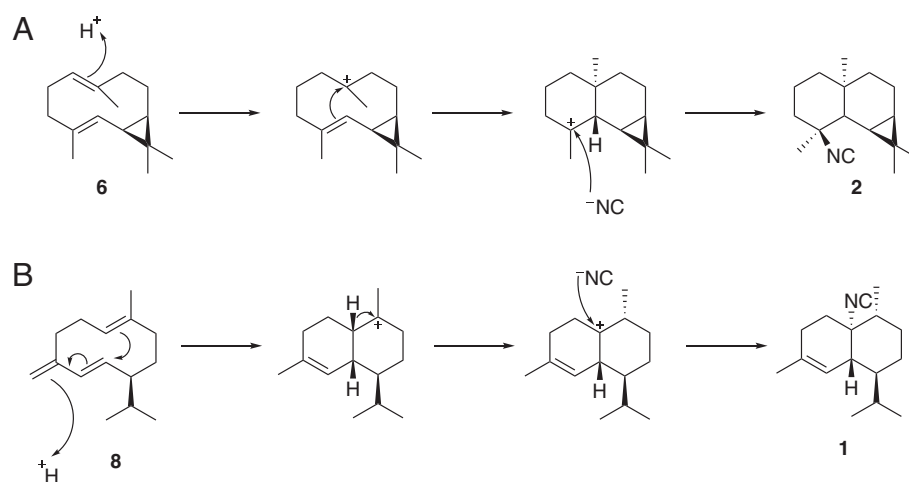


Fig. 5. Two proposed mechanistic routes to nitrogenous sesquiterpenoids from sponge TS products. (A) Proposed mechanism to yield undescribed Bubarida natural product **2** from uBuTS-1 product **6**. (B) Proposed mechanism to yield undescribed Bubarida natural product **1** from uBuTS-2 product **8**.

be learned about this fascinating pathway, of which the late D. John Faulkner once claimed that there is “no more interesting biosynthetic study among the marine natural products” (73).

The discovery of sponge animal biosynthesis genes sheds new light on other classes of sponge natural products that have long been speculated to be sponge animal-derived, such as the pyrrole-imidazole alkaloids, bromotyrosine alkaloids, and other amino acid-derived sponge alkaloids. These sponge-specific alkaloids are also enriched in sponge cell cultures and sponge tissues (74, 75), and recent work has shown that bromotyrosine alkaloids are shared between sponges with vastly different microbiomes (76). Accordingly, the sponge terpene synthase work presented herein not only lends credence to the possibility of sponge animal biosynthesis in other natural product classes but also provides a framework for identifying novel biosynthetic genes from sponge animal hosts.

Methods and Materials

Methods for sponge collection, sequencing, enzyme discovery, and chemical analysis are included in the *SI Appendix, Methods*. The *SI Appendix, Methods* also includes methodologies for

enzyme expression, purification, and activity assays. The structural characterization of enzyme products is detailed in the *SI Appendix, Supplementary Note*. Remaining data and methods are included in the *SI Appendix*. The accession numbers of sequences deposited and used in this study are listed in *SI Appendix, Table S6*.

Data, Materials, and Software Availability. DNA data have been deposited in NCBI Bioproject (PRJNA907134 and PRJNA824609) (77, 78).

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