Domoic acid biosynthesis in the red alga *Chondria armata* suggests a complex evolutionary history for toxin production

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Domoic acid (DA), the causative agent of amnesic shellfish poisoning, is produced by select organisms within two distantly related algal clades: planktonic diatoms and red macroalgae. The biosynthetic pathway to isodomoic acid A was recently solved in the harmful algal bloom–forming diatom *Pseudo-nitzschia multiseries*, establishing the genetic basis for the global production of this potent neurotoxin. Herein, we sequenced the 507-Mb genome of *Chondria armata*, the red macroalgal seaweed from which DA was first isolated in the 1950s, identifying several copies of the red algal DA (rad) biosynthetic gene cluster. The *rad* genes are organized similarly to the diatom DA biosynthesis cluster in terms of gene synteny, including a cytochrome P450 (CYP450) enzyme critical to DA production that is notably absent in red algae that produce the simpler kainoid neurochemical, kainic acid. The biochemical characterization of the *N*-prenyltransferase (RadA) and kainoid synthase (RadC) enzymes support a slightly altered DA biosynthetic model in *C. armata* via the congener isodomoic acid B, with RadC behaving more like the homologous diatom enzyme despite higher amino acid similarity to red algal kainic acid synthase enzymes. A phylogenetic analysis of the *rad* genes suggests unique origins for the red macroalgal and diatom genes in their respective hosts, with native eukaryotic CYP450 neofunctionalization combining with the horizontal gene transfer of *N*-prenyltransferases and kainoid synthases to establish DA production within the algal lineages.

**Significance**

Originally isolated from the red alga *Chondria armata*, domoic acid (DA) is best known as a potent marine neurotoxin produced by oceanic harmful algal blooms of planktonic diatoms. Sequencing efforts to date of kainoid-producing red algae have focused exclusively on a closely related molecule, kainic acid, leaving a gap in the understanding of DA biosynthesis in red algae and its evolutionary linkage to diatoms. Here, we present the phylogenetic and biochemical investigation of DA biosynthesis in *C. armata*. This work demonstrates the high synteny of DA biosynthetic genes between relatively distant taxonomic groups of algae and suggests a complex evolutionary history for DA biosynthesis involving gene transfer and neofunctionalization.

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the genera *Pseudonitzschia* and *Nitzschia* contain species that have been demonstrated to produce DA, although it is possible that other understudied diatoms, such as species belonging to the genus *Amphora*, may also produce the toxin (14–16). Understanding the distribution and evolutionary history of DA biosynthesis may help identify understudied emerging sources of toxicity in the marine environment.

The genes encoding DA (Fig. 1B) biosynthesis (*dab*) and KA biosynthesis (*kab*) were recently elucidated (Fig. 1C) (17, 18). Both pathways begin with the N-prenylation of L-glutamate by either geranyl diphosphate (GPP) or dimethylallyl diphosphate (DMAPP) to establish the linear precursors to DA and KA, respectively (19). Subsequent oxidative cyclization to build the characteristic pyrrolidine ring of either molecule is catalyzed by an α-ketoglutarate (αKG)-dependent Fe^{2+} oxidase, also known as a kainoid synthase enzyme (DabC/KabC). These two unusual enzymatic transformations are diagnostic for kainoid biosynthesis in algae, forming a core biosynthetic gene cluster and serving as bioinformatic hooks to probe sequencing data for kainoid biosynthetic pathways (Fig. 1C). The biosynthesis of DA, unlike that of KA, requires further oxidation to install the carboxylic acid functionality on the monoterpene-derived alkyl side chain via a cytochrome P450-catalyzed (DabD) reaction occurring prior to oxidative cyclization of the pyrrolidine. Additionally, the biosynthesis of DA also involves an uncharacterized isomerization transformation catalyzed by a yet to be identified enzyme. The discovery of the various kainoid pathways was facilitated by transcriptomic and genomic sequencing of the DA-producing diatom *Pseudonitzschia multiseries* (17) and of the KA-producing red algae *Digenea simplex* and *Palmaria palmata* (18). Notably, the kainoid biosynthetic genes from all organisms sequenced thus far appear to cocluster within each respective genome. Follow-up sequencing efforts of red algae to date have focused exclusively on organisms producing KA, leaving a gap in our understanding of the enzymology underlying DA biosynthesis in red algae.

To address this fundamental question, we sequenced the genome of *Chondria armata* and identified unique copies of a biosynthetic gene cluster suggestive of DA biosynthesis. Subsequent in vitro characterization of the *C. armata* pyrrolidine-forming enzymes further implicates their role in DA biosynthesis, leading to the designation of these sequences as red algal domoic acid (*rad*) biosynthesis genes. Comparative enzyme substrate assays, syntenic comparisons of gene cluster organization, and phylogenetic analysis provide key insights into the evolution of DA biosynthesis in diatoms and red algae. Our findings suggest a combination of horizontal gene transfer (HGT) and neofunctionalization of native enzymology served to establish neurotoxin production in the divergent algal lineages.

**Results and Discussion**

**Genome Sequencing and Assembly.** We collected *C. armata* from Kyushu Island, Japan, and sequenced its genomic DNA with a combination of Oxford Nanopore Technologies (ONT) and Illumina platforms. Genome size was estimated to be 480 Mb using the 17-mer histogram generated by Platanus v1.2.4 (20), repeat content was estimated to be 45.49%, and kmer distributions were consistent with tetraploidy (SI Appendix, Table S3). Notably, the longest assembled contig was 3.3 Mb in length. Genome contiguity and completeness was assessed with the eukaryota benchmarking universal
single-copy orthologs (BUSCO v4.0.5) database (22). Because of a lack of rhodophyta-specific datasets, we assessed the 10 publicly available red macroalgal genomes with the eukaryota_odb10 BUSCO database (23–26). We found the *C. armata* assembly to be relatively complete, containing 69% of complete, single-copy eukaryotic gene orthologs, where current published red macroalgal genomes contain on average 65% of complete, single-copy eukaryotic gene orthologs (*SI Appendix*, Table S4 and Fig. S3). To further assess overall genome completeness, we mapped ONT reads to the assembled genome using minimap2 (27), in which 88% of ONT reads were present in the assembled genome, a clearer indicator of genome completeness. Genome sequences and assembly are deposited in the National Center for Biotechnology Information (NCBI), BioProject identification PRJNA762367.

**Kainoid Cluster Identification and Comparison.** To identify kainoid biosynthesis genes in the *C. armata* genome, we used *dab* and *kab* genes as genetic hooks (17, 18). Previous work demonstrated the remarkably low sequence similarity of *dab*A and *kab*A genes to any publicly available protein sequence (19). tBLASTn searches with the highly conserved N-prenyltransferase genes initially revealed three copies of a kainoid biosynthetic gene cluster. Notably, unlike previously characterized red algal kainoid gene clusters, two of the *C. armata* clusters included a coclustered cytochrome P450 (CYP450), which is suggestive of DA biosynthesis. We refer to these clusters as red algal domoic acid (*rad*) biosynthetic gene clusters. The first *rad* cluster (*rad1*, GenBank accession: OK169902) is contained within a 7.9-kb region, and complete coding regions for all three key DA biosynthetic genes were clearly identified (Fig. 2). The second copy of the *rad* cluster (*rad2*, GenBank accession: OK169903) is contained within a shorter 7.5-kb region, and complete coding regions of an N-prenyltransferase and kainoid synthase were identified. However, this copy of the *rad* cluster appears to contain a degraded CYP450 with premature STOP codons. The third copy of the *rad* cluster (*rad3*, GenBank accession: OK169904) is contained within a 5.8-kb region and completely lacks a CYP450. This cluster is located 271 kb into an assembled contig, 443 kb in length. Further analysis did not reveal any DA CYP450 gene homologs or general CYP450 enzymes within this assembled sequence space (*SI Appendix*, Fig. S4). Curiously, despite *C. armata* appearing to be tetraploid, only three copies of the *rad* cluster were initially identified. To determine if one copy of the *rad* clusters was a collapsed paralog, the coverage of each *rad* cluster copy was calculated using mapped ONT reads. The mean coverage of *rad1* was almost double (39×) that of *rad2* (21×) and *rad3* (21×) and is indicative of a collapsed paralog, supporting the presence of all four expected genomic copies of the *rad* cluster.

Publicly available *dab* and *kab* gene clusters were used to construct a comparative analysis with new *rad* gene clusters from *C. armata* (Fig. 2) (28). Despite core kainoid biosynthetic genes being more similar by amino acid percent identity to the KA-producing red alga *D. simplex*, a relative of *C. armata* also belonging to the family Rhodomelaceae, all-versus-all global amino acid alignments, revealed the overall structure and organization of the *rad* gene cluster to be more like *dab* clusters from diatoms. Aside from lacking the hypothetical “*DabB*” protein found in the *dab* clusters, the *rad* and *dab* clusters surprisingly display nearly identical gene organization and orientation (Fig. 2). Amino acid percent identity remains consistent across the N-prenyltransferase (*radA/dabA*) and kainoid synthase (*radC/dabC*) enzymes, at 54 and 56%, respectively. However, despite the conservation of gene location and orientation, the amino acid percent identity between the coclustered CYP450s *radD* and *dabD* decreased to 24%.

**In Vitro Validation of Biosynthetic Gene Function.** In addition to DA, several additional isomers have also been isolated from *C. armata* and other sources (29–31). Although production of isodomoic acid A, B, and C have all been described in *Pseudo-nitzschia*, these isomers are found at substantially lower levels than DA in the diatom species and isolates studied thus far (14, 32). A similar trend has also been observed with *C. armata* (33). However, high-resolution liquid chromatography–mass spectrometry (LC-MS) analysis of methanolic algal extracts suggests that isodomoic acid B is an especially abundant DA isomer in our independently collected *C. armata* isolate (Fig. S4) (*SI Appendix*, Fig. S5). We hypothesized that the differential abundance of DA isomers in *C. armata* extracts is linked to the activities of one or more kainoid synthase enzymes expressed by the seaweed, a hypothesis consistent with our previous observations that the kainoid synthase *DabC* can generate structural diversity (17).

To determine the basis for differences in isomer production, we examined the newly identified *rad1* cluster from *C. armata*, beginning with kainoid synthase. Full-length RadC1 from the intact *rad1* cluster was expressed as an N-terminal hexahistidine (N-His$_6$)–soluble construct in *Escherichia coli* (*SI Appendix*, Fig. S6). After affinity chromatography purification, RadC1 was observed to perform oxidative cyclization on the linear precursor cNGG, similar to the previously described activity for *DabC* (17). Overnight RadC1 and *DabC* reactions both proceed to near-complete substrate consumption when cNGG was added. However, RadC1 makes isodomoic acid B as its major cyclization product, whereas *DabC* produces isodomoic acid A (Fig. 3). While isodomoic acid B production by RadC1 is consistent with the relative isomer abundance observed in our *C. armata* extracts, the presence of isodomoic acid A and C in the algae is yet to be explained biochemically. Although trace quantities of isodomoic acid C are observed from the RadC1 reaction, it remains to be seen whether stereoisomer diversity is also enriched upstream of cyclization through incorporation of neryl diphosphate. Initial expression, purification, and enzymatic assay of RadC2, encoded in the *rad2* cluster, did not reveal substantial activity toward cNGG, although small quantities of isodomoic acid B are produced. As the *rad2* cluster also contains a degraded
copy of the coclustered RadD CYP450 with several premature STOP codons, we suspect that this second cluster may be undergoing pseudogenization and that the encoded enzymes may not function in vivo. The third copy of the kainoid synthase enzyme RadC3 displays 100% amino acid sequence identity to RadC2 and is assumed to be similarly inactive.

Both RadC1 and DabC generate a similar series of dainic acid molecules, a set of higher isoprene homologs of kainic acid, from the less-favored NGG substrate. RadC1 appears to approach complete substrate turnover in our preliminary, qualitative in vitro analysis, whereas DabC does not (SI Appendix, Fig. S7) (17). Because RadC1 is more closely related in amino acid sequence to the KA-forming KabC enzyme than the diatom DabC enzyme, the substrate specificity of KabC from *D. simplex* (DsKabC, GenBank accession: QCC62383.1) was also further investigated. Intriguingly, KabC does not catalyze cyclization on either the cNGG or NGG precursor (Fig. 3 and SI Appendix, Fig. S7). As demonstrated previously, KabC efficiently cyclizes the shorter prekaic acid substrate to KA, and this activity can also be seen in this study for RadC and DabC, although neither of the latter reactions proceed to completion (SI Appendix, Fig. S8) (18). This inability of KabC to catalyze cyclization on the longer cNGG and NGG substrates represents a marked difference between the DA isomer–forming RadC and DabC versus the KA-forming KabC. While no obvious single residue can be attributed to these differences in substrate specificity, structural prediction via AlphaFold2 (34, 35) reveals KabC enzyme models to have an extended loop that pushes further into the active site than in RadC or DabC models (Fig. 3B). The quantification of active site area and volume solvent–accessible surface area using CASTp (36) identified an overall decrease in KabC modeled pocket volume (115 Å²) when compared to RadC (208 Å²) and DabC (262 Å²) models (Fig. 3C and SI Appendix, Fig. S9). This follows an expected trend according to each enzyme’s observed substrate specificities (Fig. 3A).

The glutamate N-prenyltransferase enzyme RadA1 was also expressed as a soluble construct following a seven amino acid N-terminal truncation and fusion to maltose-binding protein (MBP) containing an N-His₆ tag (SI Appendix, Fig. S10). Initial overnight assays with purified MBPΔ7-RadA1 suggest both GPP and DMAPP can be accepted as substrates to make NGG and prekainic acid, respectively. This observation contrasts the selectivity previously shown by DabA and, to a lesser extent, KabA (17, 18). Ongoing exhaustive experiments on RadA kinetics and structure will provide additional information regarding potential prenyl group promiscuity to compare directly with previously described DabA and KabA activity (19). Overall, the ability of RadA1 and RadC1 to produce DA intermediates and isomers, similar to what was observed previously in diatoms, supports their role in overall biosynthesis of DA in *C. armata* (17).

**Phylogenetic Analysis.** Previous efforts have attempted to describe the evolutionary history of the glutamate N-prenyltransferase enzyme present in both DA and KA biosynthesis (19). Despite structural homology to known bacterial terpene cyclases, the enzyme does not clade well with any extant terpene cyclase because of its unique amino acid sequence (SI Appendix, Fig. S11). To further explore the evolutionary history of the remaining DA and KA biosynthetic genes, maximum likelihood (ML) phylogenetic trees were constructed for both the dKG-dependent kainoid synthases (DabC/KabC/RadC) and the CYP450s (DabD/RadD). Beginning with RadC, initial BLAST searches revealed few closely related homologs outside of known KA or DA biosynthetic genes in the NCBI protein database. The closest homologs were primarily from bacteria, including the well-studied dKG-dependent oxidase deacetoxycephalosporin C synthase (*cefE*, UniProtKB/Swiss-Prot identification: P18548.1) in *Streptomyces clavuligerus*, an enzyme that performs the penicillin to cephalosporin ring expansion. To ensure a broad understanding of kainoid synthase enzyme phylogeny in relation to other
aKG-dependent Fe$^{2+}$-containing oxidases, representative sequences of key taxonomic groups were generated from the isopenicillin N synthase–like InterPro family (IPR027443). These representative sequences, in addition to top BLAST hits and other known kainoid synthases, were used to construct a ML phylogenetic tree, with all sequence accession numbers listed in SI Appendix, Table S5. This process of using BLAST hits and representative InterPro family sequences to construct a ML tree was repeated with RadD. Unlike RadC, BLAST searches using the hypothetically assigned RadD revealed several significant alignments with other red algal CYP450s. To ensure a well-populated phylogeny, representative sequences from the cytochrome P450 InterPro family (IPR001128) were generated from the same taxonomic groups as before (SI Appendix, Table S6). These sequences, in addition to representative diatom P450 sequences from publicly available transcriptomics and top BLASTp hits for RadD and DabD, were used to construct a ML phylogenetic tree (37).

Based on our phylogenetic analysis, kainoid synthases form their own distinct branch independent of taxonomic origin (Fig. 4A), a finding that is consistent with previous phylogenetic studies on N-prenyltransferases (19). Notably, the bacterial enzyme cCeF is the only well-characterized close relative to RadC. The monophyletic clading of both the N-prenyltransferase enzymes and kainoid synthase enzymes from these distantly related taxa suggests a possible horizontal gene transfer event. This observation is further supported by high levels of overall sequence identity between the two enzymatic groups. On the other hand, the CYP450 enzymes are independently nested within their respective taxonomic groups, an observation that supports convergent evolution toward the installation of the carboxylic acid on the prenyl sidechain (Fig. 4B). Based on this taxonomic clustering, the presence of a CYP450 enzyme in DA biosynthesis may be a result of gene duplication and neofunctionalization of P450 enzymes, a phenomenon that has been previously described in plants (38, 39). This hypothesis is further strengthened by the presence of additional CYP450s closely related to RadD or DabD found within the genome of C. armata or P. multiseries, respectively (SI Appendix, Figs. S12 and S13).

The evolutionary history of CYP450 families in diatoms has been the subject of previous research, with diatoms displaying substantial P450 gene family acquisition and loss despite encoding relatively few CYP450 genes compared to land plants (40). The diatom DabD protein bears the most homology to other CYP450 enzymes found in diatoms, including Thalassiosira pseudonana and Fragilariopsis cylindrus (bootstrap > 99%), suggestive of an ancient acquisition of this CYP450 family in the diatom lineage (SI Appendix, Fig. S12). The biochemical function of these related enzymes is unknown, and the specific CYP450 family classification of DabD-like enzymes remains to be determined.

**Conclusions and Discussion**

Our discovery of the DA biosynthetic machinery within the genome of the red alga *C. armata*, the first-described DA producer, provides crucial insight into the evolutionary history of kainoid biosynthesis in diatoms and red algae. The overall sparse distribution of kainoid biosynthetic genes in these two distantly related taxa suggests a possible horizontal gene transfer event. However, direct HGT of the entire gene cluster from red algae to diatoms, or vice versa, is not likely in modern ecological contexts. Alternatively, many genes in diatoms and other microalgae were indeed acquired from red algae in distant evolutionary history through endosymbiotic gene transfer (EGT). Endosymbiosis of unicellular red algae played a major role in the evolution of the diatom clade, and several red algal genes were transferred to the host nuclear genome via EGT to contribute to the diverse origins of diatom genomes (41, 42). However, an ancient EGT event moving the DA biosynthetic gene cluster from red algae to diatoms is unlikely given the limited distribution of kainoid biosynthetic enzymes in modern diatom genera, whereas genes acquired via EGT are typically widespread in the diatom lineage. Nevertheless, the lack of clear homology of the glutamate...
N-prenyltransferase (RadA, KabA, and DabA) and kainoid synthase (RadC, KabC, and DabC) enzymes to known algal proteins suggests HGT from a potential third host as a mechanism for the acquisition of these kainoid biosynthetic enzymes. Furthermore, the observation that the enzymes RadD and DabD are most like native CYP450s within the red algal and diatom genomes, respectively, supports a history invoking two modes of evolution toward DA biosynthesis: HGT combined with gene duplication and neo-functionalization of native biochemistry.

From our biochemically supported phylogenetic analysis, it appears that the glutamate N-prenyltransferase and kainoid synthase genes form a core biosynthetic gene cluster and were acquired via HGT from an unknown source. These specific chemical transformations are unprecedented in biochemistry outside of diatom and red algal kainoid biosynthesis and do not seem to be repurposed enzymology from within the respective algal lineages as evidenced by their low similarity to terpene cyclases and dKG-dependent Fe³⁺ oxygenases from public algal genomes and transcriptomes. These enzymes may be bacterial in origin due to the structural homology of the N-prenyltransferase to bacterial terpene cyclases and the modest amino acid sequence homology of the kainoid synthase to known bacterial enzymes, although no strong candidates for bacterial kainoid biosynthetic enzymes have been found to date (19). Notably, HGT from bacteria is well described in diatom and red algae. Diatoms especially have been the subject of substantial genome sequencing efforts, revealing that ~5% of their functional genes have been acquired from bacteria in relatively recent evolutionary history (43, 44). The acquisition of HGT-derived bacterial genes, such as urea cycle enzymes and prochorhodopsins, has contributed to the unique biochemistry and subsequent ecological success of the diatom lineage (45, 46). Similarly, HGT is an important mechanism for the expansion of genes related to both primary and secondary metabolism in red algae. The extremophilic red alga *Galdieria sulphuraria* is estimated to have acquired at least 5% of protein-coding genes through HGT (47). These HGT-derived genes typically confer an adaptive advantage to the organism such as in the genus *Pyropia*, wherein an HGT-derived carbonic anhydrase gene acts as a key component of the carbon-concentrating mechanism and aids in overcoming carbon limitations (48). Red algae also seem to have acquired other terpene cyclases via HGT from bacteria, although these enzymes appear unrelated to the N-prenyltransferase enzymes in DA and KA biosynthesis (49). HGT has also been suggested to play a role in the evolution of saxitoxin biosynthesis. Saxitoxin, another notable HAB neurotoxin and the causative agent of paralytic shellfish poisoning, is produced by prokaryotic freshwater cyanobacteria and eukaryotic marine dinoflagellates (50). Genes encoding saxitoxin biosynthesis were first identified in the cyanobacterium *Cylindrospermopsis raciborskii* and later in dinoflagellates belonging to the genus *Alexandrium*, suggesting a potentially ancient HGT event from cyanobacteria to dinoflagellates based on conservation of diagnostic domain architectures between key biosynthetic enzymes (51–53).

Following acquisition of the core kainoid biosynthetic genes, both *C. armata* and *Pseudonitzschia* spp. appear to have repurposed CYP450 enzymeology to install the characteristic carbonyl scaffold has evolved independently in diatoms and red algae. An analogous independent neo-functionalization of CYP450 activities has been previously implicated in the evolution of the plant hormone diosgenin, a plant steroid saponin with a distinctive CYP450-installed spiroketal motif that is not limited to a specific taxon of plants but is instead broadly and sparsely distributed (38). An additional example of CYP450-driven convergent evolution is seen in the furanocoumarin pathway, also in plants (39). The independent selection for these enzyme activities in unrelated plants provides precedence for the neo-functionalization of CYP450s to install similar chemical features in distantly related organisms, a phenomenon that may have occurred in the evolution toward DA biosynthesis as reflected by the red algal–derived RadD1 and diatom-derived DabD.

Evolution toward DA production is further supported by the apparent substrate scope of the different kainoid synthases. Although RadC1 is more like the red algal KabC in its amino acid sequence, RadC1 displays an activity more similar to the diatom enzyme DabC in its ability to convert both NGG and cNGG to DA-like molecules. These substrates are not similarly cyclized by the KA-forming DsKabC. Despite the similar activities exhibited by the *C. armata* and *Pseudonitzschia* kainoid synthases, the difference in DA isomers generated by RadC1 (isodomoic acid B) and DabC (isodomoic acid A) raises additional biosynthetic questions. Notably, the 1,3-olefin isomerization step required to make DA itself continues to remain elusive (17, 29). Because isodomoic acid B would require both a 1,3-isomerization and a separate trans to cis isomerization of the second olefin closest to the kainoid ring, it is worth reconsidering whether the key 1,3-isomerization occurs before or after kainoid synthase cyclization to create the most abundant, canonical DA isomer in diatoms and red algae. A similar 1,3-isomerization step occurs immediately following the triple hydroxylation of a methyl group to a carbonylic acid in the putative CYP450 reaction toward installing a carboxylic acid on the DA scaffold, a process thought to be carried out by one or more CYP450 enzymes in the ergot fungus *Claviceps purpurea* (56). However, such an olefin isomerization was not revealed by previous studies on DabD expressed in *Saccharomyces cerevisiae* (17). Meanwhile, hypothetical protein DabB identified from the diatom biosynthetic cluster is missing from the respective *C. armata* gene cluster and does not appear to be encoded within the sequence *C. armata* genome. In any case, the responsible isomerase remains unidentified. Ongoing investigations of candidate genes present in these two distantly related species, including further study of the activities exhibited by the co-clustered CYP450 enzymes DabD and hypothetically assigned RadD, will shed additional light on this remaining piece of the DA biosynthetic puzzle.

The apparent evolution of kainoid synthase substrate selectivity in combination with evidence for CYP450 neo-functionalization and co-clustering suggests a potential advantage afforded by DA biosynthesis over KA biosynthesis. While DA is chemically distinct from KA and has enhanced bioactivity against ionotropic glutamate receptors, the true ecological function of DA has remained elusive (57). Previous studies have demonstrated limited trace metal chelation by DA, affording a potential selective advantage for diatoms thriving in iron-limited coastal regimes (58, 59). Other work has suggested a role for DA in grazer defense, particularly with respect to copepod grazing in the diatom system (60). However, no studies to date have provided a concrete ecological mechanism beyond reasonable doubt in both diatoms and red algae. While DA could conceivably...
serve different functions in the red alga C. armata and Pseudo-nitzschia diatoms, the evolutionary history described here provides insight to the ecological importance of this marine toxin.

Materials and Methods

**Si Appendix** includes methods for expression, purification, and mutagenesis of all enzymes used in this study. Methodologies for activity assays, sequencing, and phylogenetics are also detailed. Finally, general chemical procedures and small molecule characterization are available in **Si Appendix**. The remaining data and methods are present in **Si Appendix**. The NCBI accession numbers of sequences deposited and used in this study are listed in **Si Appendix**, Table S7.

**Data Availability.** BioProject, BioSample, Genome, Sequence Read Archive (SRA), and GenBank data have been deposited in NCBI (PRJNA762367, SAMN21392177, JAWH000000000, SR19527350, SR19527349, SR19527348).

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