BOTH MODULAR AND SINGLE-DOMAIN TYPE I POLYKETIDE SYNTHASES ARE EXPRESSED IN THE BREVETOXIN-PRODUCING DINOFLAGELLATE, *KARENIA BREVIS* (DINOPHYCEAE)  

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Dinoflagellates are prolific producers of polyketide compounds, many of which are potent toxins with adverse impacts on human and marine animal health. To identify polyketide synthase (PKS) genes in the brevetoxin-producing dinoflagellate, *Karenia brevis*, we assembled a transcriptome from 595 million Illumina reads, sampled under different growth conditions. The assembly included 125,687 transcripts greater than 300 nt in length, with over half having >100× coverage. We found 121 transcripts encoding Type I ketosynthase (KS) domains, of which 99 encoded single KS domains, while 22 contained multiple KS domains arranged in 1–3 protein modules. Phylogenetic analysis placed all single domain and a majority of multidomain KSs within a monophyletic clade of protist PKSs. In contrast with the highly amplified single-domain KSs, only eight single-domain ketoreductase transcripts were found in the assembly, suggesting that they are more evolutionarily conserved. The multidomain PKSs were dominated by trans-acyltransferase architectures, which were recently shown to be prevalent in other algal protists. *Karenia brevis* also expressed several hybrid nonribosomal peptide synthetase (NRPS)/PKS sequences, including a *burA*-like sequence previously reported in a wide variety of dinoflagellates. This contrasts with a similarly deep transcriptome of *Gambierdiscus polynesiensis*, which lacked NRPS/PKS other than the *burA*-like transcript, and may reflect the presence of amide-containing polyketides in *K. brevis* and their absence from *G. polynesiensis*. In concert with other recent transcriptome analyses, this study provides evidence for both single domain and multidomain PKSs in the synthesis of polyketide compounds in dinoflagellates.

**Key index words:** algal toxin; dinoflagellate; *Karenia brevis*; polyketide synthase; toxin biosynthesis  

**Abbreviations:** ACP, acyl carrier protein; AT, acyl transferase; CEGMA, Core Eukaryotic Genes Mapping Approach; CoA, coenzyme A; DDBJ, DNA Databank of Japan; DH, dehydratase; DNAse, deoxyribonuclease; DTT, dithiothreitol; EMBL, European Molecular Biology Laboratory; ER, enoyl reductase; FAS, fatty acid synthase; FeCl₃, ferric chloride; HCS, HMG coA synthetase; HMG, hydroxymethyl glutarate; HMM, hidden Markov model; KR, ketoreductase; KS, β ketoacyl synthase; LbH, Left-handed Parallel beta-Helix; MAFFT, Multiple Alignment using Fast Fourier Transform; MT, methyl transferase; NRPS, nonribosomal peptide synthase; P450, cytochrome P450; Pfam, protein families database; PhyML, phylogenetics maximum likelihood; PKS, polyketide synthase; PP, phosphopantetheine binding site; RNA, ribonucleotide; RNAseq, RNA sequencing; RNAse, ribonuclease; SAM, S-adenosyl methionine; TE, thioesterase; TTP, thiamine pyrophosphate binding domain; VSSC, voltage sensitive sodium channel
The dinoflagellate *Karenia brevis* is endemic to the Gulf of Mexico, where it is responsible for red tides that occur almost every year on the west coast of Florida and less frequently in the western Gulf on the Yucatan peninsula and in the northern Gulf from Texas to the Florida panhandle (Steidinger 2009). *Karenia brevis* blooms have significant environmental impacts, including massive fish kills, marine mammal, bird, and turtle mortalities (Landsberg 2002, Twiner et al. 2012), adverse human health effects through both seafood consumption and respiratory exposure (Fleming et al. 2011), and economic impacts of over $26 million per bloom year (Hoagland et al. 2009, Morgan et al. 2009).

The adverse effects of *Karenia brevis* blooms are due to their elaboration of a suite of polyether ladder compounds known as brevetoxins, potent neurotoxins that both alter the voltage sensitivity and prolong the open state of voltage sensitive sodium channels (VSSC) involved in both neuronal and neuromuscular synaptic transmission (Baden et al. 2005). Many other dinoflagellate toxins with known health impacts are polyether ladder or related polyketide compounds (e.g., ciguatoxins, diarrhetic shellfish toxins, yessotoxins, amphidinols). *Karenia brevis* elaborates at least 11 different congeners of brevetoxin bearing two different backbones containing either 10 (A-type) or 11 (B-type) trans-fused ether rings, with variations in side chain constituents that alter their relative toxic potencies (Fig. 1). In addition, *K. brevis* produces an array of other polyketide compounds (Fig. 1), all with varying affinities for the brevetoxin binding site on the VSSC, including a four-ring polyether ladder hemibrevetoxin B (Prasad and Shimizu 1989), which binds to but does not activate the VSSC, five-ring brevenal, which opposes the action of brevetoxins by inhibiting the activation of VSSC (Bourdelais et al. 2004), and a six-ring “interrupted” polyether ladder, brevesin, whose action is uncharacterized (Satake et al. 2009). In addition, two polyethers contain amide bonds, the seven-ring tamulamides (Truxall et al. 2010), and a single-ring compound, brevisamide (Van Wagoner et al. 2010). The extensive elaboration of polyketide compounds by *K. brevis* and other dinoflagellates suggests they serve an important function. The total cellular concentration of brevetoxins in studied isolates ranges from 1 to 68 pg cell\(^{-1}\) (Hardison et al. 2012). Based on an estimated 500–700 pg carbon cell\(^{-1}\) (Kamykowski et al. 1998), with brevetoxins made up of 66% carbon, this equates to an investment of up to 9% of total cellular carbon in brevetoxins alone.

Stable isotope labeling studies established that brevetoxins are the products of polyketide syntheses (PKS; Lee et al. 1986, 1989, Chou and Shimizu 1987). PKSs build carbon chains in a manner analogous to fatty acid syntheses (FASs), in which a starting substrate, generally acetyl CoA, is extended through a series of sequential condensations with malonyl CoA, performed by a β-ketoacyl synthase (KS) domain, to produce a product of predetermined length. In the case of fatty acids, each acetate unit added to the growing chain then undergoes ketoreduction (KR), dehydration (DH), and enoyl reduction (ER), resulting in a fully saturated carbon chain. In contrast, PKSs may lack one or more of these catalytic domains, thereby producing carbon chains that include carbonyl groups (absence of KR domain), hydroxyl (absence of DH domain) or double bonds (absence of ER; Staunton and Weissman 2001). The growing carbon chain resides on a phosphopantetheine “arm” of an acyl carrier protein (ACP) that presents it to the catalytic sites, while an acyl transferase (AT) presents the extender units to be added to the growing chain, most often malonyl CoA. The full-length polyketide is released from the PKS complex by a thioesterase (TE), while post-PKS modifications create the final polyketide structure. Two major classifications of PKSs are found. In Type I PKS (modular) all catalytic domains are found on a single polypeptide, which are used in a progressive fashion for chain elongation, analogous to FASs in animals and fungi (Khosla et al. 1999, Jenke-Kodama et al. 2005). Type II PKSs are multiprotein complexes where each catalytic domain is found on a separate polypeptide, analogous to type II FASs in bacteria and plants. PKSs typically initiate with an acetyl CoA; however, in dinoflagellates, alternative starter units are sometimes used (Kellman et al. 2010). Furthermore, in brevetoxin, as well as other...
Polyketide synthases in *Karenia brevis*

Dinoflagellate polyketides, the carbonyl carbons of some of the acetate units added in the condensation reaction are excised from the growing polyketide by a Favorski rearrangement (Wright et al. 1996), catalysed by a P450 or flavin monooxygenase. Whereas this step is rare among bacterial and fungal polyketides, it appears to be the norm among dinoflagellate polyketides (Van Wagener et al. 2014). Another unusual event among dinoflagellate polyketides is the addition of pendant methyl groups from both methionine and acetate. These steps require the involvement of SAM methyl transferase (MT) and HMG-CoA synthetase (HCS) activity, respectively. Together, these hallmark characteristics of dinoflagellate polyketides suggest that their biosynthetic machinery may include catalytic domains that differ from the better characterized PKSs in bacteria and fungi.

The PKSs of dinoflagellates remain poorly characterized. Sequences with homology to Type I KS domains were first identified in dinoflagellates by Snyder et al. (2005) using degenerate PCR. With the advent of EST libraries, full length transcripts containing Type I-like KS and KR domains were identified by Monroe and Van Dolah (2008), which were highly unusual in that each contained a single catalytic domain, a 5′ spliced leader sequence and polyA tail, indicating their dinoflagellate origin. These single domain dinoflagellate PKS sequences group phylogenetically with a protist clade, confirming their eukaryotic origins (John et al. 2008). Since that time, single domain PKS sequences have been found in numerous dinoflagellate species (Eichholz et al. 2012, Pawlowiez et al. 2013, Kimura et al. 2015, Kohli et al. 2015) including species considered non-toxic (Salcedo et al. 2012, Meyer et al. 2015), and single domain Type I-like PKSs appear to be a characteristic of dinoflagellates. Kohli et al. (2016) mined data from 24 genera and 46 strains of dinoflagellates whose transcriptomes were sequenced under the Moore Foundation’s Marine Microbial Eukaryote Transcriptome Sequencing Project, and found that dinoflagellate single domain KSs fall into three clades. Representatives of each clade were present in most dinoflagellates sequenced, and there was no apparent relationship with the evolutionary history of the organism, and no clear relationship with the production of particular known classes of marine biotoxins (Kohli et al. 2016).

The presence of amide-containing tamulamide and brevisamide (Fig. 1) suggests that *Karenia brevis* may also encode non-ribosomal peptide synthases (NRPS) in its genome, which incorporate amino acids in a manner analogous to the incorporation of activated acetate groups by PKS. The first module in an NRPS, known as the initiation module, typically includes an adenylation domain (A) and a peptide carrier protein domain (PCP). Following this are elongation modules which contain a condensation domain (C), A, and PCP domains. Like the ACP domain of PKS, the PCP domain has a post-translationally added phosphopantetheine arm that passes the peptide-thioester intermediate from one module to the next, with a single amino acid being added at each module. Also analogous to PKS, the full length polypeptide is released by a terminating TE domain. Hybrid NRPS/PKS proteins incorporate both amino acids and acetate building blocks to create a high diversity of natural products. A hybrid NRPS/PKS has previously been reported in *K. brevis* (Lopez-Legentil et al. 2010).

With the advent of deeper sequencing afforded by second generation sequencing platforms, there have been several reports of multidomain PKSs present in dinoflagellates (Beedessee et al. 2015, Kohli et al. 2017). Here we assembled a deep transcriptome of *K. brevis* with a particular interest in mining for the diversity of single domain PKS sequences and the presence of multidomain PKSs, NRPSs, and hybrid NRPS/PKS transcripts.

**Methods**

**Culture conditions.** *Karenia brevis* (nonaxenic, Wilson strain) was grown in sterile filtered seawater at 36%, and enriched with f/2 media (Guillard et al. 1973) modified with ferric sequestrene in the place of EDTA-Na2 and FeCl3·6H2O and the addition of 0.01 µM selenious acid. All cultures were maintained in 1 L glass bottles at 25°C on a 16:8 h light:dark cycle under cool white light at 45–50 µmol photons·m−2·s−1. Heat shocked cultures were subjected to 31°C for 30 or 60 min prior to harvesting. This 6°C shock results in a significant decrease in translation, as evidenced by the loss of polysomes, a classical hallmark of stress (Fridey 2015).

**RNA extraction.** Whole cultures were harvested by centrifugation at 600g. RNA extraction was performed on the pelleted cells using TRIReagent (Life Technologies, Carlsbad, CA, USA) following the manufacturer’s protocol. Extracted RNA was DNase treated with RQ1 RNase-free DNase (Promega, Madison, WI, USA) for 10 min at room temperature and the RNA was then purified on columns using the RNeasy Mini Kit (Qiagen, Valencia, CA, USA) following the manufacturer’s protocol.

To obtain actively translated RNA pools, 50 µg·mL−1 cycloheximide was added to each L of *K. brevis* and incubated for 5 min to prevent ribosome runoff prior to harvesting. The cells were then harvested by centrifugation at 600g and pellets homogenized in polysome resuspension buffer (50 mM Tris HCl, pH 8, 100 mM NH4Cl, 20 mM sucrose, 10.5 mM Mg acetate, 0.5 mM EDTA, 1 mM DTT, 0.1% heparin, 0.01% cycloheximide, 0.1% Triton X-100, 80 U RNAsin) and centrifuged at 15,000×g for 10 min. Supernatants were layered on 12.5%–50% sucrose gradients, which were centrifuged at 180,000×g at 4°C for 2.5 h in a Beckman SW 41Ti rotor. Polysomes were then profiled by pumping the sucrose gradients through a flow cell with UV detection at 254 nm (ISC0). One milli liter sucrose fractions representing the polysome fractions (2–6 ribosomes) were pooled and RNA extracted by the addition of 1 part 8 M guanidine HCl. One part ethanol was then added and samples were incubated at −80°C overnight; then 2 µL of polyacryl carrier was added to facilitate RNA precipitation. Samples were centrifuged at 12,000g for 30 min at 4°C, washed with cold 75% ethanol, and...
resuspended in nuclease-free water with RNasin Ribonuclease Inhibitor (Promega), and DNase treated as described above. All RNA samples were quantified using a NanoDrop N-1000 Spectrophotometer (Thermo Scientific, Waltham, MA, USA) and integrity assessed using an Agilent Bioanalyzer 2100 (Agilent Technologies, Santa Clara, CA, USA).

RNAseq libraries. A total of 595 million *Karenia brevis* reads were assembled into one transcriptome. The constituent reads included several RNAseq libraries, generated under different conditions and using different library generation protocols and sequencing platforms. The first two libraries were generated from cells harvested at mid-log phase (MMETSP201, 56 M reads) or stationary phase (MMETSP202, 44 M reads) and submitted to the Moore Foundation Marine Microbial Eukaryote Transcriptome Sequencing Project for library preparation using Illumina TruSeq RNA protocol. These libraries generated ~25 million 50 nt paired-end sequencing reads on Illumina HiSeq2000. These sequences, available from NCBI Short Read Archive number SRR042159, were pooled to yield 100 million reads. Nine stranded libraries were generated using the NEBNext Ultra Directional RNA Library Prep Kit (Illumina, Hayward, CA, USA) from total RNA of log phase cultures under control, exponential growth conditions (n = 3), or cultures exposed to 30 min (n = 3) or 60 min (n = 3) of 6°C heat shock prior to harvest as described in Fridey (2015). Sequencing of these libraries was performed on an Illumina HiSeq 2500 sequencer, at a depth of ~25 million, 125 nt, single end reads per library. Pools of these libraries resulted in 225 million single end reads. Lastly, nine libraries consisted of RNA isolated from the translationally active RNA pools isolated from polyribosomes as described by Fridey (2015) from log phase *K. brevis* cells under control exponential growth conditions (n = 3), or cultures exposed to 30 min (n = 3) or 60 min (n = 3) of 6°C heat shock prior to harvest. These libraries were generated using the TruSeq RNA Sample Preparation Kit (Illumina) and on Illumina HiSeq2000 at a depth of ~14 million 50 nt paired-end reads. These sequences were pooled to yield 270 million reads. These sequences are available from NCBI Bioproject number PRJNA343279.

Transcriptome assembly and analysis. Raw reads from individual libraries were quality filtered, and then all reads were assembled together into contigs using CLC Genomics Workbench (CLC Bio, Cambridge, MA, USA) with default settings. Contigs with a length of less than 300 nt were not analyzed further. The combined transcriptome has been deposited in DDBJ/EMBL/GenBank under the accession GFLM01000000. BLASTx analysis, mapping, annotation and Kyoto Encyclopedia of Gene and Genomes analysis for both the gene catalogue and those reported from *Gambierdiscus* and *Karenia* was submitted (Parra et al. 2007). BLASTx was performed against the nr database of GenBank using an E-value cut-off of 10^-3. For mapping and annotations, the default values of Blast2GO were used. To analyze the comprehensiveness of the gene catalogues the Core Eukaryotic Genes Mapping Approach (CEGMA) tool was used (Parra et al. 2007). HMMER (Finn et al. 2011a, b) was used for the identification of contigs containing conserved PKS domains (KS, KR, ACP, AT, DH, ER, TE) using an in-house HMM database (Kohli et al. 2017). Functional prediction of sequences was further analyzed by Pfam (Punta et al. 2012) and conserved domain searches (Marchler-Bauer et al. 2014) were used for identification of conserved amino acid residues and functional prediction of PKS and PKS/NRPS transcripts. Phylogenetic analysis steps were performed in Geneious software (Kearse et al. 2012). Sequences were aligned using MAFFT v6.814b (Katoh et al. 2002) and/or ClustaW (Thompson et al. 1994). Alignments were trimmed manually to ensure they spanned the KS or KR conserved domain. Maximum likelihood phylogenetic analysis was carried out using PhyML (Guindon et al. 2010) or RAxML-7.0 (Stamatakis 2006) using LG model of rate heterogeneity with 1,000 bootstraps. Phylogenetic trees were visualised using MEGA-Version 6 (Tamura et al. 2013).

RESULTS AND DISCUSSION

A transcriptome of *Karenia brevis* assembled from 595 million Illumina reads, pooled from libraries representing different growth stages (log and stationary phase) and stress conditions (heat shock), produced 125,687 contigs greater than 300 nt in length. The assembly had an N50 of 1,829 nt and an average contig length of 1,413 nt, where the longest contig generated was 36,109 nt and over 400 contigs were >8,000 nt in length. The GC content was 52.4%, consistent with previous *K. brevis* libraries reported (Lidie et al. 2005, Ryan et al. 2014), similar to *Symbiodinium*, which varies by isolate (50.5%–56.4%; Bayer et al. 2012), and lower than those of other peridinin containing dinoflagellates (~60%; Jaeckisch et al. 2011, Kohli et al. 2017). Over half (53.6%) of the contigs had >100x coverage, supporting their assembly (Table 1). Overall, 55.1% of the contigs had BLASTx hits to the Genbank non-redundant database (E-value < 10^-3), of which 54.4% had annotated matches, while 45.6% matched uncharacterized proteins. The percentage of annotated matches observed increased with increasing contig length and coverage (Table 1).

To further assess the completeness of the transcriptome assembly, we conducted an analysis for the presence of 248 ultra-conserved core eukaryotic genes using CEGMA. We found 223 (88%) of the 248 highly conserved genes present in full length, with 90% recorded as present if partial matches are included. This is higher than previously reported transcriptome assemblies of *K. brevis* (Ryan et al. 2014) and likely reflects the greater depth of sequencing (595M reads) included in the current transcriptome. A description of sequences encoding essential enzymes for fifteen conserved metabolic pathways is presented in Table S1 in the Supporting Information.

Polyketide synthases. Using HMMER and conserved domain searches to identify ketosynthase domains, we found 121 KS containing contigs: 99 possess a single KS domain while 22 contain multiple KS domains (Table S2 in the Supporting Information). A phylogenetic analysis of KS domains from *K. brevis* and those reported from *Gambierdiscus* spp. (Kohli et al. 2017) placed all dinoflagellate single-KS domain sequences, and most dinoflagellate KS domains found in multidomain transcripts, within a clade made up of KS sequences from other protists (Fig. 2). This provides evidence that both the single domain transcripts and multidomain KS sequences are of dinoflagellate origin, and not due to bacterial or fungal contaminants known to co-occur in
K. brevis cultures. Further evidence for the dinoflagellate origin of the single domain sequences is the presence of the dinoflagellate-specific spliced leader sequence on the 5’ end of at least 12 transcripts containing the full 5' end. Within the protist clade, the single domain K. brevis KSs form a monophyletic sub-branch, while multidomain KS sequences formed several independent clades that grouped with the Apicomplexa. A small subgroup of multidomain dinoflagellate KSs fell within a bacterial clade as discussed below.

Single domain KS sequences. Within the dinoflagellate single-domain KS clade, three sub-clades exist with good bootstrap support (Fig. 2), here termed Clades 1, 2, and 3. These clades appear to correspond with three single-domain KS clades identified by Kohli et al. (2016) in a survey of dinoflagellate transcriptomes. Single domain KS sequences identified in Karenia mikimotoi similarly fall into three clades (two clades, one with two major subclades; Kimura et al. 2015) with the same K. brevis membership as described below.

Clade 1 includes previously published Karenia brevis KS proteins KB2006 and KB5361 (Monroe et al. 2008). All K. brevis sequences within this clade have the conserved active site cysteine (C) required for decarboxylative condensation, within the conserved sequence (D/E)TACS(S/T/A)(S/A/G/M; Fig. 3). All members of this clade have the conserved H within H(G/A/C)TGT sequence required for transacylation. The second conserved H required for KS activity is also present in all sequences in this clade, with a consensus of NIGH. Sequences in this clade contain a consensus of AYLG in the 5’ dinoflagellate motif, with variations in the first two positions quite frequent. This clade included the KB6736 previously described by Monroe and Van Dolah (2008).

Clade 3 included 16 Karenia brevis sequences with highly divergent active sites, including transcripts identical to previously reported KS sequences KB1008, KB4361, and KB6842. No sequences in this clade possess an active site C (in the expected sequence DTACSS) required for decarboxylative condensation. In its place is a consensus of DXEX (A/S)S (where X is too variable to obtain a consensus; Fig. 3). The conserved H in the expected HGTGT is present in only about half of the sequences in this clade, with a consensus sequence of HGXGX. The second conserved H is replaced in most sequences in this clade with N, with a consensus of NXGN. Given these differences in the active sites that are conserved in all known KSs, it is unclear what function(s) these sequences may have. When the full-length sequences are aligned, the conserved 5’ dinoflagellate motif is present as GLLG, the original sequence reported by Eichholz et al. (2012) and observed also in Gambierdiscus poly-nesiensis (Pawlowiez et al. 2014).

Clade 2 Karenia brevis KS sequences in this clade all have the conserved active site C required for decarboxylative condensation, within the sequence (D/E)TACS(S/T/A)(S/A/G/M; Fig. 3). All members of this clade have the conserved H within H(G/A/C)TGT sequence required for transacylation. The second conserved H required for KS activity is also present in all sequences in this clade, with a consensus of NIGH. Sequences in this clade contained a consensus of AYLG in the 5’ dinoflagellate motif, with variations in the first two positions quite frequent. This clade included the KB6736 previously described by Monroe and Van Dolah (2008).

Single domain PKS KS sequences lack plastid targeting peptides. Insight into the presence of organelle targeting sequences on transcripts can be informative for discerning the metabolic processes they are involved in. To this end, we analyzed the 5’ ends of all full length single domain K. brevis KS sequences for the presence of signal peptides using SignalP predictor (http://www.cbs.dtu.dk/services/SignalP/), beginning with the translation initiating.

### Table 1. Coverage and annotation statistics of Karenia brevis gene catalogue. An e-value cut-off of 10^{-3} was applied during BLASTx analysis.

| Coverage          | No. of contigs | Length (mean) | Annotated match | Non-annotated match | No match | PKS sequences |
|-------------------|----------------|---------------|------------------|---------------------|----------|---------------|
| 1×–5.00×          | 2,819          | 300–1,401     | 167              | 310                 | 2,342    | 7             |
| 5.01×–20×         | 15,033         | 300–10,974    | 2,455            | 3,046               | 9,532    | 14            |
| 20.01×–50×        | 19,906         | 300–24,286    | 5,278            | 5,425               | 9,203    | 5             |
| 50.01×–100.00×    | 20,907         | 300–36,109    | 6,804            | 5,306               | 8,707    | 10            |
| 100.01×–1,000×    | 61,676         | 300–34,047    | 20,692           | 16,265              | 24,719   | 91            |
| 1,000.00×–10,000× | 5,272          | 300–6,989     | 2,291            | 11,09               | 1,872    | –             |
| 10,000.01× and above | 74            | 311–2,254    | 58               | 5                   | 11       | –             |
| Total number of contigs (percentage) | 125,687 | 300–36,109 (1,413.3) | 37,745 (30%) | 31,556 (25.1%) | 56,386 (44.9%) | 127 |

**POLYKETIDE SYNTHASES IN KARENIA BREVIS**
methionine. Signal peptides were absent from all PKS KS domains. In contrast, signal peptides were present on the six Type II FAS KS domain transcripts (KASII and III oxoacyl ACP synthases) present in the K. brevis transcriptome (Table 2). Previous analyses show that dinoflagellate FASs reflect the phylogenetic history of their plastids and are predicted to be plastid localized based on the presence of targeting sequences (Kohli et al. 2016). Karenia brevis possesses a tertiary plastid of haptophyte origin, and its chloroplast transit peptides differ from those in peridinin dinoflagellates, in which the signal peptide cleavage site is typically followed by a conserved sequence FVAP (Patron et al. 2006). The signal cleavage sites in K. brevis KAS II sequences and several known plastid targeted proteins examined (Table 2) are preceded by G and followed by RRV(K/Q), while the sequence in KASIII is G-RRIL. A similar conserved sequence RRxQ was found in 26 plastid targets proteins in another tertiary plastid containing dinoflagellate, Karlodinium micrum (Patron et al. 2006). The absence of such plastid targeting sequences on single domain PKS KS transcripts would predict cytosolic localization. Confoundingly, the only single domain KS protein to be studied at the protein
level, KB2006 (Monroe et al. 2010) was localized to the chloroplast pyrenoid by immunolocalization and to the chloroplast by subcellular fractionation (Monroe et al. 2010, Van Dolah et al. 2013). If the PKS biosynthetic machinery is in fact located in the chloroplast, it is unclear what mechanism transports these proteins to the plastid.

**Multidomain KS containing transcripts.** Twenty-four contigs were identified as multidomain PKS (16), NRPS/PKS (6), or NRPS (2) encoding one to three modules (Table 3). Both cis- and trans-AT architectures were present. None of the multidomain sequences includes a 5′ SL sequence, suggesting that they may not be complete, or they may not be processed via SL trans-splicing.

**Trans-AT PKSs:** A majority of the *Karenia brevis* multidomain PKSs lack AT domains (Table 3). Trans-AT PKSs were first identified over a decade ago in bacteria, and since been found broadly among bacteria and protists. Trans-AT PKSs evolved independently of the better known cis-AT PKSs, both originating from FAS (Piel 2010). Whereas cis-AT PKSs have only eight possible domain architectures (KS-AT-ACP, KS-AT-KR-ACP, KS-AT-DH-KR-ACP, KS-AT-DH-ER-KR-ACP, with and without MT), there are more than 50 module types known to date in trans-AT PKSs (Piel 2010). Trans-AT PKSs are distinguished from cis-ATs not only in the lack of AT domains, but also unusual ordering of domains, inclusion of unusual domains, the presence of non-condensing KS₉ domains, and the amplification of ACP domains. Trans-AT PKSs have been found to be common among other microalgal PKSs (Shelest et al. 2015). In the phylogenetic analysis of *K. brevis* KS domains, most of the multidomain KSs fall within a clade that includes apicomplexan PKSs (Fig. 2; Dinoflagellate multidomain KS Clade 1). All *K. brevis* KS domains in this clade are from trans-AT sequences or trans-AT modules within mixed cis/trans-AT sequences from *K. brevis* and *G. polynesiensis* (Fig. 4A).

**Cis-AT PKSs:** Two PKS transcripts encode cis-AT domains. Contig 75628 falls within the protist clade but is outside of the dinoflagellate KS clades. Contig 28414 has one cis- and one trans-AT module (ACP-KS-AT-DH-ER-KR-ACP-KS-KR). The trans-AT module falls within the dinoflagellate trans-AT clade (Figs. 2 multidomain Clade 1 and Fig. 4A), while the cis-AT module falls within Dinoflagellate multidomain KS Clade 2 (Figs. 2 and 4B), along with *K. brevis* contig 10709 and several *G. polynesiensis* KS sequences, which appear to be from trans-AT PKSs. Thus, the cis-AT associated KSs from PKS sequences did not form a unique clade.

**NRPS/PKSs:** Six hybrid NRPS/PKS sequences were identified in the *Karenia brevis* assembly. Sequences with identical domain architecture to Contig 3318 (TE-A-PP-KS-AT-TE-KR-PP) have been reported previously in a wide array of dinoflagellates, from the basal dinoflagellate *Oxyrrhis* to 18

Table 2. Predicted signal peptides and cleavage sites in known plastid-localized proteins and Type II FAS sequences in *Karenia brevis*. Signal peptide prediction and score generated by signalP. The D-score (discrimination score) is used to discriminate signal peptides from non-signal peptides, using a threshold of 0.5. Cleavage site sequences were screened for the conserved FVAP sequence found in peridinin dinoflagellates. In some sequences a plausible FVAP-like sequences is present, but most possess the RRV(Q/K) also present in *Karlodinium*. No signal peptides were present in any single domain PKS KS or KR sequences.

| GenBank accession no. or contig no. | CD and blast ID | Prediction | D score | Cleavage site |
|------------------------------------|----------------|------------|---------|---------------|
| ABF73013                           | Cytochrome b₆₆ FeS subunit | No         | 0.131   | –             |
| Contig 15936                       | Cytochrome b₆₆-f | Yes        | 0.55    | G-QSPREQ     |
| ABF73015                           | Ferredoxin      | Yes        | 0.613   | R-RVVDRV     |
| Contig 15936                       | Multimeric flavodoxin | Yes    | 0.763   | A-ASLAS      |
| ABF73016                           | Flavodoxin nadph reductase | Yes    | 0.673   | G-FRVQ       |
| ABF73017                           | Oxygen enhancer | No         | 0.419   | G-RFQQK      |
| ABF73018                           | psII 12kD       | Yes        | 0.864   | A-FSPA       |
| ABF73029                           | GapDH C1        | Yes        | 0.606   | A-(-9)FEEQ   |
| kbrevis_combined_contig_386        | KASII 3 oxoacyl ACP synthase | Yes    | 0.7     | G-RVVQ or G-(-9)FKPA |
| kbrevis_combined_contig_387        | KASII 3 oxoacyl ACP synthase | Yes    | 0.681   | G-RVVQ or G-(-9)FKPA |
| kbrevis_combined_contig_46229      | KASII 3 oxoacyl ACP synthase | Yes    | 0.723   | S-DYGR or G-RRVK |
| kbrevis_combined_contig_37304      | KASII 3 oxoacyl ACP synthase | Yes    | 0.607   | G-RVVQ or A-FNPA or G-(-9)FNPA |
| kbrevis_combined_contig_49622      | KASII 3 oxoacyl ACP synthase | Yes    | 0.816   | G-RRIL       |
| kbrevis_combined_contig_275        | KASII 3 oxoacyl ACP synthase | Yes    | 0.572   | G-RRLN       |
| kbrevis_combined_contig_97889      | KR              | Yes        | 0.837   | A-OTPT or A-(-9)FPHA |
| kbrevis_combined_contig_64411      | KR              | Yes        | 0.479   | G-RPMQ       |
| kbrevis_combined_contig_86499      | KR              | Yes        | 0.798   | A-YGEEF or C-SSRE |
| kbrevis_combined_contig_28593      | ER              | Yes        | 0.646   | G-RVVQ       |
| kbrevis_combined_contig_47965      | ER              | Yes        | 0.69    | G-RRVK       |
| kbrevis_combined_contig_14484      | DH              | No         | 0.148   | –            |
| kbrevis_combined_contig_6314       | AT              | Yes        | 0.659   | G-RTLQ       |
| kbrevis_combined_contig_101182     | AT              | No         | 0.427   | G-RRLQ       |
core dinoflagellates, including *K. brevis* (Bachvaroff et al. 2014), as well as in *Gambierdiscus* spp. (Kohli et al. 2017). This domain architecture is found in the *burA* gene from the bacteria *Burkholderia*. *BurA* is involved in the synthesis of one of two polyketide precursors of burkholderic acid, an unprecedented furan derivative produced from the fusion of two polyketide chains (Franke et al. 2012). The transcript has both an N-terminal TE domain and an internal TE. The function of an N-terminal TE script has both an N-terminal TE domain and an internal TE. The function of an N-terminal TE domain at the N-terminal end. The function of an N-terminal TE domain is not yet understood. Bachvaroff et al. (2014) reported a dinoflagellate spliced leader on the 5' end of this transcript in *Amphidinium carterae*, suggesting its dinoflagellate origin, although the SL was supported by a single overlapping read. In our phylogenetic analysis, the KS domain from Contig 5155 falls in a clade with other dinoflagellate KS domains from cis-AT NRPS/PKS sequences and bacterial *burA* (Figs. 2 and 4C).

**Table 3.** Domain structure of multidomain PKS, NRPS/PKS hybrids, and NRPS contigs in the *Karenia brevis* assembly.

| Contig number | Length (nt) | Classification | Cis/Trans | No. of modules | Domain structure |
|---------------|-------------|----------------|-----------|----------------|-----------------|
| Contig 1930   | 19,025      | NRPS/PKS       | trans     | 3              | TE-A-DH-PP-C-A-PP-KS-DH-KR-PP-PP-PP-TE-(MT) |
| Contig 10563  | 18,781      | NRPS/PKS       | trans     | 3              | TE-A-DH-PP-C-A-PP-KS-DH-KR-PP-PP-PP-TE-(MT) |
| Contig 5155   | 11,047      | PKS            | trans     | 2              | PP-KS-DH-KR-PP-KS-DH-KR-PP-PP-PP-PP-TE-(MT) |
| Contig 78360  | 2,221       | PKS            | trans     | 3              | PP-KS-KR-PP-KS-KR-PP-TE-PP-LbH |
| Contig 10709  | 14,778      | PKS            | trans     | 3              | PP-KS-DH-KR-PP-KS-KR-PP-TE-PP-LbH |
| Contig 15957  | 3,031       | PKS            | trans     | 1              | KR-PP-KS-KR-PP-TE-LbH |
| Contig 81604  | 7,799       | PKS            | trans     | 3              | (KS)-KR-PP-KS-KR-PP |
| Contig 28414  | 12,319      | PKS            | cis       | 2              | PP-KS-AT-DH-KR-[ER]-KR-PP-KS-KR |
| Contig 54805  | 8,035       | PKS            | cis       | 1              | KS-AT-KR-LbH-PP-LbH-PP-LbH-PP-LbH |
| Contig 77766  | 4,288       | NRPS/PKS       | cis       | 2              | PP-C-A-PP-(C) |
| Contig 3318   | 8,238       | NRPS/PKS       | cis       | 1              | A-KS-AT-DH-ER-PP-(TE) |
| Contig 10632  | 9,386       | NRPS/PKS       | cis       | 1              | A-KS-AT-DH-ER-PP-(TE) |
| Contig 134145 | 2,544       | PKS            | ?         | 1              | PP-KS |
| Contig 34829  | 18,811      | NRPS/PKS       | trans     | 2              | tpp-3-KR-PP-KS-PP-C-TE-A-PP-C-PP-TE |
| Contig 10563  | 18,781      | NRPS/PKS       | cis       | 1              | A-KS-AT-DH-ER-PP-(TE) |
| Contig 3318   | 8,238       | NRPS/PKS       | cis       | 1              | TE-PP-KS-AT-TE-KR-PP |
| Contig 77766  | 4,288       | NRPS            | ?         | 2              | PP-C-A-PP-(C) |
| Contig 4898   | 2,638       | NRPS            | ?         | 2              | TE-A-DH-PP-C-A |

KS, ketosynthase; KR, ketoreductase; DH, dehydratase; ER, enoyl reductase; KR[ER]K, KR with embedded ER domain; AT, acyl transferase; TE, thioesterase; A, adenylation domain; C, condensation domain; PP, phosphopantetheine binding site of ACP or PCP domains; (MT), possible methyl transferase; tpp, thiamine pyrophosphate binding domain; LbH, left-handed beta helix; ( ), partial domain; [ER], embedded within KR domain.

Comparison with multidomain PKS identified in other dinoflagellates. To date few publications describe multidomain PKSs in dinoflagellates. Bachvaroff et al. (2014) provided the first report of a multidomain PKS/NRPS hybrid in a large array of dinoflagellates with architecture similar to the *burA* gene in...
FIG. 4. Details of clade membership of *Karenia brevis* KS domains in Figure 2. Multidomain PKS sequences were found primarily in three clades. Inset shows locations of clades detailed in this figure. (A) Dinoflagellate multidomain Clade 1 within the protist clade that included apicomplexans. This clade contains KS domains from *K. brevis* and *Gambierdscus polynesiensis* trans-AT PKSs. (B) Dinoflagellate multidomain Clade 2 within the protist clade includes two cis-AT and one trans-AT PKS in *K. brevis* and several trans-AT *G. polynesiensis* KSs. (C) Clade 3 includes bacterial and dinoflagellate *BurA*-like NRPS/PKS, cis-AT NRPS/PKS sequences and cis-AT bacterial Type I PKS. S. atroolivaceus, Streptococcus atroolivaceus.
Burkholderia as described above. Kohli et al. (2017) conducted a survey of PKSs in four species of Gambierdiscus, sequenced at depths between 134 million to 1 billion Illumina reads, and assembled in CLC Workbench using similar parameters as used here for K. brevis. In contrast to K. brevis, where a quarter of the sequences recovered were NRPS/PKS hybrids, only the deepest library (G. polynesiensis) revealed any NRPS/PKS sequences, a contig similar to the burA-like sequence discussed above and a short contig containing domains C-ACP-KS. Unlike K. brevis, which produces at least two amide containing compounds (talamamide and brevisamide; Fig. 1), no products of NRPS pathways are known in Gambierdiscus spp. Like K. brevis, both cis- and trans-AT PKS sequences were present in Gambierdiscus spp. The longest sequence found in G. polynesiensis was a 7-module PKS, of which the first six modules lacked AT domains while the seventh encoded a cis-AT. The predicted product of this sequence resembles the carbon backbone of a polyether ladder compound. No sequence of this length was found in K. brevis. However, the KS domains in this sequence cluster with trans-AT sequences in K. brevis that have similar domain architectures. All other contigs found in Gambierdiscus encoded only one module or partial modules, whereas in K. brevis, half of the PKS or NRPS/PKS contigs contained 2–3 modules. Also absent from the Gambierdiscus assemblies were the tandemly repeated ACP domains observed in some K. brevis NRPS/PKS and PKS sequences. Beedessee et al. (2015) reported 10 multidomain PKSs and NRPS/PKS hybrid genes in a genomic survey of Symbiodinium minutum, with transcriptome support indicating that these genes are expressed as the predicted multidomain sequences. Both cis- and trans-AT PKS sequences were present. One sequence contained a singly repeated ACP domain but none contained highly repeated ACPs as found in K. brevis. One very large 8-module NRPS/PKS hybrid sequence of 10,601 amino acids was found. Most of the PKSs reported encoded single modules (7), 2 modules (1) or 3 modules (1). In some cases, multiple gene models with transcript support were found on the same scaffold, suggesting that the smallish (1–3 module) transcripts that have been identified in K. brevis and other dinoflagellate transcriptomes may represent full transcripts and not partial assemblies of larger multimodular transcripts. Confirmation of their completeness may require genomic sequence data for K. brevis.

KR sequences. Using HMMER and conserved domain searches to identify ketoreductase domains, we found 8 transcripts encoding single KR domains, while 25 were present on the multidomain PKS sequences described above (Table 2; Table S2). All K. brevis KR domains possess the conserved active site residues YXXKN present in Type I PKS and animal FAS that distinguishes them from KR domains in Type II FAS/PKS, which possess YXXK. The number of unique single domain KR is much smaller than the number of single domain KS sequences described above (n = 95), suggesting that their selection is more highly conserved. A similar trend was observed in four Gambierdiscus spp. (expressing collectively 90 single domain KS and 7 single domain KR domains; Kohli et al. 2017). Phylogenetic analysis placed all K. brevis KR domains within a clade that included protist - chlorophyte, haptophyte, and Apicomplexa - and bacterial KR domains (Fig. 3). Within this clade, the single domain KR sequences formed a monophyletic group separate from the KR domains found in multidomain PKSs (Fig. 4). The KR domains from multidomain PKSs formed a clade with several interesting subclades (Figs. 5 and 6). Sub-clade 1 is made up entirely of KR domains from PKS modules with the architecture KS-KR-ACP. Sub-clade 2 is made up primarily of KR domains from trans-AT PKS modules with the architecture KS-DH-KR. Sub-clade 3 KR domains possess an ER domain inserted between the N-terminal and C-terminal subdomains of the KR. This architecture has been observed previously in porcine FAS (cd05275; https://www.ncbi.nlm.nih.gov/cdd/). Subclades 4 and 5 include all NRPK/KS sequences found in K. brevis. Clade 4 includes burA-like sequences in K. brevis and other dinoflagellates as well bacterial burA genes from Burkholderia spp. Clade 5 includes three highly similar contigs in K. brevis with module architecture KS-DH-KR - two NRPS/PKS (contig 1930, contig 10563) - and one PKS (contig 5155) that lacks the N-terminal NRPS module (Table 2). These sequences have in common highly amplified ACP domains (n = 6) and potential MT domains at their c-terminal ends. In general, KR domains clustered according to their module architecture rather than grouping with other KR domains in a given multimodule sequence, if their module architectures differed.

Survey of other PKS domains in Karenia brevis. Acyl transferase. Using HMMER, CD search and blast analysis we found 34 unique AT domain containing contigs (E value cut-off ≤1E-10). Of these, 9 occurred in the cis-AT PKSs described above, while 25 encode single AT domains (Table S2). Phylogenetic analysis separates the cis-AT domains from the stand-alone ATs (Fig. S1 in the Supporting Information). Twenty single-domain AT sequences have top blast hits of malonyl:acyl carrier transacylase. These sequences cluster separately from two sequences that are FabD domains of Type II FASs.

Dehydratase. By HMMER, CD search and blast analysis we found 24 unique DH domains, characterized by the presence of a “hotdog fold,” which consists of a seven-stranded antiparallel beta-sheet “bun” that wraps around a five-turn alpha-helical “sausage,” originally identified in E. coli FabA fatty acyl dehydratase (DH). The “hotdog fold” protein superfamily includes 17 subfamilies of proteins of diverse function including TEs and DHs (Dillon


and Bateman 2004). Fourteen of the DH domains found are located within multidomain PKS or PKS/NRPS sequences (Table 3) with homology to either the “hotdog fold” superfamily or specifically to the PKS DH subfamily (Table S2). Another 9 contigs contain individual, partial “hotdog fold” domains (Table S2). Of these, one contains both spliced leader and polyA tail, indicating that it is not a partial sequence, but represents a full transcript. It is uncertain if these individual “hotdog fold” containing contigs function as standalone PKS DH since their “hotdog fold” domains are truncated. These sequences differ from the single Type II fatty acid beta-hydroxyacyl-ACP DH present in the assembly (Contig 44484), which has specific homology to the FabZ subfamily of “hotdog fold” superfamily.

Phosphopantetheine attachment domains (PP): PP attachment domains are hallmarks of both ACPs in PKSs and PCP in NRPSs. HMMER analysis identified 123 contigs containing PP domains (E value cut-off ≤1E-3). Twenty of these are among the multidomain sequences described above (two are short contigs lacking PP domains). The majority of standalone PP domain contigs were <1,000 aa in length and contained 1 or 2 PP domains, with or without tpr repeat domains in the N-terminal end. Several of these sequences possessed the dinoflagellate spliced leader, indicating their dinoflagellate origin.

Thioesterase domains: Using HMMER, and CD searches we found 49 TE domain containing contigs (E value cut-off ≤1E-3). Nine were among the multidomain sequences described above. Among the 40 standalone TE domain contigs (Table S2), several included a dinoflagellate spliced leader and in some cases polyA tails, indicating full length transcripts. Discrete TE domains (termed TEII) are frequently associated with multidomain PKS and NRPSs and play roles in editing and efficiency as well as chain release. Both integrated and discrete TE domains identified are members of the alpha/beta hydrolase fold class.

Unusual domains or motifs identified in K. brevis multidomain PKS and NRPS/PKS.

**Left-handed parallel beta-helix (LbH):** Regions with high similarity to LbH domains were found at or near...
FIG. 6. Details of clade membership of *Karenia brevis* KR domains summarized in Figure 5. KR domains from multidomain PKS sequences were found in subclades that correspond more closely to module architecture than to contig membership. Sub-clade 1: KS-KR-ACP; Sub-clade 2: KS-DH-KR; Sub-clade 3: KS-DH-KR[KR]-ACP; Sub-clade 4: *burA*-like NRPS/PKS; Sub-clade 5: trans-AT modules from NRPS/PKS with architecture KS-DH-KR in sequences with highly amplified ACP domains at their C-terminal ends. *G. australis*, *Gambierdiscus australis*; *G. excentricus*, *Gambierdiscus excenctricus*; *H. triquestra*, *Heterocapse triquestra*; *A. carterae*, *Amphidinium carterae*. 

\[ \text{Fig. 6. Details of clade membership of *Karenia brevis* KR domains summarized in Figure 5. KR domains from multidomain PKS sequences were found in subclades that correspond more closely to module architecture than to contig membership. Sub-clade 1: KS-KR-ACP; Sub-clade 2: KS-DH-KR; Sub-clade 3: KS-DH-KR[KR]-ACP; Sub-clade 4: *burA*-like NRPS/PKS; Sub-clade 5: trans-AT modules from NRPS/PKS with architecture KS-DH-KR in sequences with highly amplified ACP domains at their C-terminal ends. *G. australis*, *Gambierdiscus australis*; *G. excentricus*, *Gambierdiscus excenctricus*; *H. triquestra*, *Heterocapse triquestra*; *A. carterae*, *Amphidinium carterae*.} \]
the C-terminal end of three multidomain PKSs: Contig 78360, Contig 15957, and Contig 54805 (Table 2). In contig 54805 three separate, nearly identical LbH motifs are found between pairs of ACP domains. Left handed beta helices are generated by 30 imperfect repeats of a hexapeptide motif (X-[STAV]-X-[LIV]-[GAED]-X), where three contiguous repeats specify one turn of the β-helix. Proteins containing hexapeptide repeats are often enzymes showing acyltransferase activity.

Thiamine pyrophosphate binding domain: A region with strong similarity to IlvB or other thiamine pyrophosphate binding domain (TPP) containing protein, was found at the N-terminal end of Contig 34829, a hybrid NRPS/PKS (Table 2). IlvB is a transketolase involved in isoleucine synthesis that converts pyruvate to acetoacetate. Although unusual, TPP-dependent transketolases have previously been reported to be the source of 2-carbon donors from glycolytic pathway intermediates to NRPS assemblies (Peng et al. 2012).

Methyl transferase: Three similar contigs with trans-AT architecture, Contig 1930, Contig 10563, and Contig 5155, possess a region at the C-terminal end with homology to O-methyltransferases. This family of MT utilizes S-adenosyl methionine and have been shown in PKSs to incorporate methyl branches (Piel 2010). The location of the potential MT domain in these sequences, C-terminal to a TE domain is unusual, as in Type I PKSs they usually occur in the AT-KR, DH-ER, or DH-KR linker regions, or in a trans-AT PKS between ACP domains (Ansari et al. 2008). The MT-like regions in these contigs appear to be (identically) truncated and thus may not have activity.

CONCLUSIONS

Dinoflagellates produce an enormous diversity of polyketide compounds, many with adverse effects on human and marine animal health, yet their biosynthetic machinery has been little explored in large part because the considerable size of dinoflagellate genomes has hindered genomic sequence analysis. Over the past decade, transcriptome sequencing of a variety of dinoflagellates has revealed unique, single-domain Type I PKSs likely to be involved in polyketide biosynthesis, while more typical multidomain PKS have been consistently absent. In Karenia brevis, we (Monroe and Van Dolah 2008, Sanger sequencing) and others (Ryan et al. 2014, RNAseq, 50M reads/library) reported only single domain PKS transcripts. However, with RNAseq-based access to deeper transcriptome sampling, as well as sequencing of the smallest dinoflagellate genome, Symbiodinium, there is increasing evidence for the presence of multidomain PKS in dinoflagellates, in addition to the previously identified single domain PKS proteins. In the current study, we therefore pooled 20 RNAseq libraries to yield a combined library of 595M reads, in order to obtain sufficient coverage to assemble long, multidomain PKS transcripts if present. Using this approach, in addition to 121 single domain KS contigs, we found 22 contigs containing multiple KS domains, two NRPS sequences, and five hybrid NRPS/PKSs. Trans-AT PKS architectures were prevalent, as recently reported in PKSs from other eukaryotic microalgae (Shelest et al. 2015). The longest PKS found in this study consisted of three modules, insufficient by itself to synthesize any of polyethers known to be produced by K. brevis. This suggests that multiple PKSs likely work together in the biosynthetic process. How, and if, the single domain KS sequences interact with multidomain PKSs is currently unknown, and is a question of great interest. Investigating the intracellular localization of the newly identified PKSs will be important for establishing potential interactions between single and multidomain sequences. The only dinoflagellate KS protein for which cellular localization has been investigated to date is K. brevis KB2006, a Cluster 1 single domain KS, which was found in the chloroplast (Monroe et al. 2010). This study, in concert with recent evidence in Gambierdiscus spp. and Symbiodinium suggests that dinoflagellates utilize both single domain and multidomain PKS and PKS/NRPS proteins in their toxin biosynthetic machinery.

We thank Kelly Fridley Sides for generation of the heat stressed transcriptome and translatome libraries included in this analysis, with funding from the NOAA Marine Biotoxins Program. Support for this analysis was provided by the University of Technology Sydney Distinguished Visiting Scholar program to FMVD.

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POLYKETIDE SYNTHASES IN KARENIA BREVIS

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web site:

Figure S1. Phylogenetic analysis separates the cis-AT domains from the stand-alone ATs. Phylogeny was inferred using Mafft alignment and PHyML.

Table S1. Description of sequences from Karenia brevis encoding essential enzymes for various metabolic pathways (128 out of 133 enzymes are present). Enzymes colored in red were not present in the transcriptome.

Table S2. Description of all single-domain contigs encoding KS, KR, DH, AT, ACP, and TE domains in the Karenia brevis assembly, and multidomain PKS and NRPS/PKS contigs containing incorporated modules. Contig sequences may be obtained from the NCBI Transcriptome Sequence Archive Accession number GFLM01000000.