Novel Insights into Evolution of Protistan Polyketide Synthases through Phylogenomic Analysis

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Polyketide synthase (PKS) enzymes are large multi-domain complexes that structurally and functionally resemble the fatty acid synthases involved in lipid metabolism. Polyketide biosynthesis of secondary metabolites and hence functional PKS genes are widespread among bacteria, fungi and streptophytes, but the Type I was formerly known only from bacteria and fungi. Recently Type I PKS genes were also uncovered in the genomes of some alveolate protists. Here we show that the newly sequenced genomes of representatives of other protist groups, specifically the chlorophytes Ostreococcus tauri, O. lucimarinus, and Chlamydomonas reinhardtii, and the haptophyte Emiliania huxleyi also contain putative modular Type I PKS genes. Based on the patchy phylogenetic distribution of this gene type among eukaryotic microorganisms, the question arises whether they originate from recent lateral gene transfer from bacteria. Our phylogenetic analyses do not indicate such an evolutionary history. Whether Type I PKS genes originated several times independently during eukaryotic evolution or were rather lost in many extant lineages cannot yet be answered. In any case, we show that environmental genome sequencing projects are likely to be a valuable resource when mining for genes resembling protistan PKS I genes.

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Key words: evolution; fatty acid synthase (FAS); ketoacyl synthase (KS); polyketide synthase (PKS); protists; phylogenomics.

Introduction

Polyketides are a structurally diverse class of natural products derived from the polymerization of acetyl and propionyl subunits in a process...
similar to fatty acid synthesis. Such compounds are of pharmaceutical and biomedical interest because many of them have potent biological effects, such as antibiotics, anti-tumor compounds, natural insecticides and immunosuppressive agents (Staunton and Weissman 2001). Numerous functions in nature have been proposed for these secondary metabolites, ranging from chemical defense to complex cell communication. The presence of polyketides in bacteria, fungi and streptophytes has been known for decades, but their occurrence in protists has only recently been confirmed. Analysis of metabolic pathways has established that certain potent polyether biotoxins found in free-living marine dinoflagellates and haptophytes are entirely or partially derived via polyketide biosynthesis (MacKinnon et al. 2006; Wright and Cembella 1998).

The polyketide synthase (PKS) enzymes responsible for the synthesis of polyketides are large multi-domain complexes that structurally and functionally resemble fatty acid synthase (FAS) enzymes involved in lipid metabolism. FAS and PKS catalyze the sequential condensation of acyl units onto a growing carbon chain and both enzymes possess a similar set of functional domains: ketoacyl synthase (KS), acyl transferase (AT), ketoacyl reductase (KR), dehydratase (DH), enoyl reductase (ER), acyl carrier protein (ACP) (or phosphopantetheine attachment site [PP]), and thioesterase (TE). Whereas FAS is dependent upon the presence of the complete set of aforementioned functional units, the minimal structure of PKS requires only the ACP, KS and AT domains for the condensation reaction. The other domains (when present) catalyze the stepwise reduction of the initial carbonyl units (Hopwood and Sherman 1990).

Polyketide synthases are generally classified into three major structural sub-groups. Type I PKSs are large, highly modular proteins, whereas Types II are aggregates of monofunctional proteins. Type I PKSs are among the largest known proteins encoded by a single open reading frame (ORF), encompassing as many as 12,000 amino acid residues. These enzymes include several modules, some of which are responsible for chain elongation while others catalyze the associated reduction steps. In most bacteria, each module directs one round of chain extension and post-condensation modification to generate non-aromatic polyketides. In fungi and some bacteria, each module/ enzyme of Type I PKS is used iteratively, yielding either aromatic or non-aromatic compounds. By comparison, Type II PKSs are multi-protein complexes whereby the individual enzymes are used iteratively for each cycle of chain extension. These Type II complexes are found exclusively in bacteria for synthesis of aromatic polyketides. The Type III PKSs, also known as chalcone synthases, are homodimeric and function iteratively as condensing enzymes. Their distribution was believed to be essentially restricted to streptophytes, within which they employ unusual starter units to act directly on acyl-CoA thioesters, independently of ACP.

Recent microbial genome sequencing, however, has revealed additional Type III PKSs in bacteria, most of which are of unknown function (Gross et al. 2006; Moore and Hopke 2001). In their study of bacterial Type I PKS evolution, Jenke-Kodama et al. (2005) concluded that FAS and PKS passed through a long joint evolutionary process with the modular PKS type arising from bacterial FAS and primary iterative PKS.

Our attention in this phylogenomic analysis focused on the origin and evolution of the biosynthetic genes for Type I PKS. These genes are well documented in bacteria and fungi, but have also recently been found in the genome of the apicomplexan parasite Cryptosporidium parvum (Zhu et al. 2002). Fragments of putative Type I PKS genes have also been identified in free-living marine protists belonging to the dinoflagellates and haptophytes, which are also known to produce polyketide-derived biotoxins (Cembella and John 2006; MacKinnon et al. 2006; Snyder et al. 2005). However, the evidence for Type I PKS genes from these marine protists has been questioned because of the difficulties in maintaining true axenic cultures (the absence of both free-living and endosymbiotic bacteria), thereby raising suspicions the PKS genes identified could be either of protistan or bacterial origin. Despite strong circumstantial evidence of their protistan origin, such as recent in situ hybridization experiments that have localized Type I PKS sequences to the nuclear genome of the toxic polyketide-producing dinoflagellate Karenia brevis (Snyder et al. 2005), large genome fragment sequencing is necessary to confirm their eukaryotic origin.

In contrast to the aforementioned distribution of putative and confirmed Type I PKS genes, no such genes have been confirmed in any other eukaryotic genome. The apparent disparate distribution of Type I PKS genes among the lineages of the eukaryotic evolutionary tree raises the question of the origin of these genes.
The increasing availability of eukaryotic genome sequences from various lineages initiated our interest in a broad survey and phylogenomic analysis of Type I PKS genes in diverse organisms. We screened most published as well as several ongoing eukaryotic genome sequence projects for candidate Type I PKS genes. Besides the included genome projects, data from several EST projects on protists are available or are in progress. We chose not to include these data sets in our analyses because the mostly short sequence fragments were not a reliable basis for sequence analysis. Furthermore, because of the potential problem of bacterial contamination, the eukaryotic origin of sequences from such projects is much more uncertain than in the case of complete or near complete genome sequences. We then established the phylogenetic relationships of the putative PKS I sequences found in our genome screen and compared their primary structure to known representatives of this gene family. This genomic analysis has provided novel insights into the distribution and evolutionary history of these key enzymes in secondary metabolism.

Results and Discussion

Distribution of Polyketide Synthases in Protists

Screening most available eukaryotic genomes from published and ongoing genome sequencing projects for candidate Type I PKS genes (Table 1 and Supplementary Table S1) showed that sequences with high similarity to Type I PKS genes were only present in a few lineages and

Table 1. List of eukaryotic genomes analyzed in this study for PKS type I genes.

<table>
<thead>
<tr>
<th>Species</th>
<th>Taxonomy</th>
<th>Source</th>
<th>Abbreviations</th>
<th>PKS I type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cryptosporidium hominis</td>
<td>Apicomplexa</td>
<td>COGENT</td>
<td></td>
<td>Yes*</td>
</tr>
<tr>
<td>Cryptosporidium parvum</td>
<td>Apicomplexa</td>
<td>COGENT</td>
<td>EAK87820_cParvum</td>
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</tr>
<tr>
<td>Eimeria tenella</td>
<td>Apicomplexa</td>
<td>SANGER</td>
<td>dev_EIMER</td>
<td>Yes</td>
</tr>
<tr>
<td>Toxoplasma gondii</td>
<td>Apicomplexa</td>
<td>SANGER</td>
<td>Toxop_KS</td>
<td>Yes</td>
</tr>
<tr>
<td>Plasmodium falciparum</td>
<td>Apicomplexa</td>
<td>COGENT</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Theileria parva</td>
<td>Apicomplexa</td>
<td>TIGR</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Tetrahymena thermophila</td>
<td>Ciliophora</td>
<td>TIGR</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Perkinsus marinus</td>
<td>Perkinsea</td>
<td>TIGR</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Emiliania huxleyi</td>
<td>Haptophyta</td>
<td>JGI#</td>
<td>Ehx_contig</td>
<td>Yes</td>
</tr>
<tr>
<td>Thalassiosira pseudonana</td>
<td>Stramenopiles</td>
<td>JGI</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Phaeodactylum tricomatum</td>
<td>Stramenopiles</td>
<td>JGI</td>
<td></td>
<td>No</td>
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<tr>
<td>Phytophthora sojae</td>
<td>Stramenopiles</td>
<td>JGI</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Phytophthora ramorum</td>
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<td>JGI</td>
<td></td>
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</tr>
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<td>Leishmania major</td>
<td>Kinetoplastida</td>
<td>SANGER</td>
<td></td>
<td>No</td>
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<tr>
<td>Trypanosoma brucei</td>
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<td></td>
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</tr>
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</tr>
<tr>
<td>Monosiga brevicollis</td>
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<td>Cyanidioschyzon merolae</td>
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<td>Galdieria sulphuraria</td>
<td>Rhodophyta</td>
<td>MSU</td>
<td></td>
<td>No</td>
</tr>
<tr>
<td>Chlamydomonas reinhardtii</td>
<td>Chlorophyta</td>
<td>JGI</td>
<td>chlamyV3</td>
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<tr>
<td>Ostreococcus lucimarinus</td>
<td>Chlorophyta</td>
<td>JGI</td>
<td>opPKS</td>
<td>Yes</td>
</tr>
<tr>
<td>Ostreococcus tauri</td>
<td>Chlorophyta</td>
<td>Genopole</td>
<td>KSotauri</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Languedoc-Roussillon</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
were dispersed throughout the eukaryotic tree (our criteria for “high similarity with PKS I genes” were: regions showing sequence similarity ($e < 10^{-5}$) to different PKS I domains in close proximity within a contiguous ORF or at least in a contiguous chromosomal region).

Among lineages of Plantae, we found candidate sequences in the genomes of unicellular green algae (Chlorophyta), namely, *Chlamydomonas reinhardtii*, and in two recently sequenced *Ostreococcus* species, *O. tauri* (Derelle et al. 2006) and *O. lucimarinus* (Palenik et al. 2007). Both *Ostreococcus* species have a compact genome of approximately 13 Mbp (Derelle et al. 2002) and both genomes contain three large PKS genes. The corresponding polyketide products and their potential function are unknown, but the fact that the three PKS ORFs account for up to 1.5% of the genome indicates that they likely have an important function in this small genome (Keeling and Slamovits 2005). In any case, the presence of Type I PKS genes among green algae was surprising because such sequences were not found in the genomes of other Plantae lineages, neither in streptophytes nor in the red algae (Rhodophyta), *Cyanydioschyzon merolae* and *Galdieria sulphuraria* (Table 1).

The alveolates, which comprise the three sublineages Apicomplexa, dinoflagellates and ciliates, also have representatives possessing Type I PKS genes. Among Apicomplexa, Type I PKS and phylogenetically closely related FAS genes have been found in the parasites *Cryptosporidium parvum* (Zhu et al. 2002, 2004), *Eimeria tenella* (http://www.sanger.ac.uk/Projects/E_tenella/), and *Toxoplasma gondii* (http://www.sanger.ac.uk/Projects/T_gondii/). Similar sequence fragments can also been found in the genome of *Cryptosporidium hominis*. Nevertheless, the existence of Type I PKS genes is not universal among alveolates because these sequences were lacking in other species, including the parasites *Perkinsus marinus*, *Plasmodium falciparum* and *Theileria parva* and the free-living ciliates, *Tetrahymena thermophila* and *Paramecium* spp. Dinoflagellates, another important group of alveolates, appear to contain Type I PKS genes as indicated by several cDNA sequencing studies (Cembella and John 2006; Jaeckisch et al. 2007; Lidie et al. 2005; Snyder et al. 2003, 2005). In these cases, only short fragments were recovered and no full genome sequences are available for this group, therefore we did not include them in our analyses.

Among the stramenopiles, genomes of the oomycetes *Phytophthora ramorum* and *P. sojae*, and the diatoms *Thalassiosira pseudonana* and *Phaeodactylum tricornutum* apparently lack Type I PKS genes. The genome of another chromalveolate, the haptophyte *Emiliania huxleyi* was recently released (Version 1, 2006, bread@csusm.edu), but it has not yet been annotated. Long contiguous ORFs containing typical Type I PKS domains (KS, AT, ACP, KR, ER) are present on several scaffolds clearly corresponding to the *E. huxleyi* genome, based on their nucleotide composition, and not to bacterial contaminants, which had been removed from the assembly. This provided strong evidence that this species has at least one and perhaps several Type I PKS genes.

Of the three available genomes of Euglenozoa (*Trypanosoma brucei*, *T. cruzi* and *Leishmania major*), which are members of the Excavata, none exhibited PKS Type I sequences.

This patchy distributional pattern of PKS genes among members of the eukaryotic crown lineages is similar to findings for prokaryotes. In a study of 138 bacterial genomes, PKS genes could be identified in only 27 (21% of total), and none of the archaean genomes possess putative PKS genes (Jenke-Kodama et al. 2005).

Our further analyses focused on the beta-ketoacyl synthase (abbreviated as KS) domain, the most conserved domain of Type I PKS genes (Kroken et al. 2003). This domain has the greatest potential for revealing divergent homologs and thus provides the most informative basis for comparative and phylogenetic analyses. In phylogenetic analysis of the conserved KS domains of these newly discovered Type I PKS genes, we compared these gene sequences to known representatives from bacteria and fungi, and to the highly similar KS domains from metazoon FAS. These phylogenetic analyses (Fig. 1) recovered the clades known from previous phylogenetic analysis of this PKS domain (Kroken et al. 2003). The bacterial clades, the metazoon FAS, and Ascomycota KS clades (both reducing and non-reducing) were supported by bootstrap (BP) and posterior probability (PP) values (Fig. 1). The putative protistan PKS I KS sequences did not resemble any of these formerly described groups, but instead, clustered distinctly from them. The KS sequences from Apicomplexa (*C. parvum*, *T. gondii*, *E. tenella*) formed a well-supported monophyletic clade. The *Ostreococcus* Type I PKS sequences were spread into two clades—Chlorophyta Clade 1 and Clade 2. Chlorophyta Clade 1, consisting of 32 *O. tauri* (KSotauri), 21 *O. lucimarinus* (opPKS), and three *C. reinhardtii* KS sequences, grouped together with the Haptophyta
clade (*E. huxleyi* KS sequences), whereas Clade 2, containing three *O. tauri* and one *O. lucimarinus* KS sequences was distinct. This grouping was surprising because chlorophytes are supposed to be a monophyletic group (Bhattacharya and Medlin 1998). Furthermore, within Chlorophyta Clade 1, sequences from *C. reinhardtii, O. tauri* and *O. lucimarinus* formed a relatively tight but mixed group, not reflecting the presumed long independent evolutionary history of these three taxa (Fig. 2). Whereas the monophyly of all described clades was well supported by BP and PP values, the relationships among them remain largely unresolved.

Considering the large divergence in the KS data set, we also tried to identify alignment positions at which substitution saturation has occurred using the program Asatura (van de Peer et al. 2002). We generated trees with different degrees of highly variable sites, but all tree topologies and bootstrap values remained basically unchanged by these manipulations (data not shown).

One of the most remarkable integrative results of our analyses was that the newly identified modular Type I PKS genes from protists differ from all hitherto known genes of this type. Whereas in bacteria the Type I PKS pathway frequently co-occurs with non-ribosomal peptide synthases (NRPS), regulating a second type of secondary metabolite pathway (Shen et al. 2001), such genes have not yet been found in protists. Furthermore, our analyses based on many currently available

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**Figure 1.** Maximum likelihood phylogenetic tree of the KS domain of Type I PKS sequences. Scale bar represents corrected evolutionary divergence. Numbers at the branches indicate bootstrap and posterior probability values (BP/PP).
Figure 2. Maximum likelihood phylogenetic trees of the protistan clades. Corresponding taxon names can be taken from Table 1. Numbers at the branches indicate bootstrap and posterior probability values (BP/PP).
protistan genome sequences indicate that the occurrence of Type I PKS genes is much wider than traditionally thought. Our survey reveals that from the four large groups of eukaryotes for which genome sequences are currently available (unikonts, Plantae, excavates, and chromalveolates), three contain PKS I gene members (up to now, no Type I PKS was found in members of the Excavata). Nevertheless, these eukaryotic PKS genes do not group with any of the previously described clades (Kroken et al. 2003; Shen et al. 2001), but rather cluster separately in well-supported monophyletic clades that do not always fit well with the proposed phylogenetic relationships of their hosts (Baldauf 2003; Keeling et al. 2006).

**Discrimination of Protistan from Bacterial and Fungal Beta-Ketoacyl Synthase Domains**

The possibilities of a sequence-based discrimination of protistan PKS fragments was approached by using hidden Markov models (HMMs) of KS domain sequences. First, HMMs for bacterial, fungal and protistan KS domains were calculated based upon those sequences included in our alignments, i.e., 68 protist, 35 fungal and 19 bacterial sequences (for graphical representation of these profile models, see Supplementary Figures S1 and S2). Each sequence included in our alignment was then scored with the three models. As expected the best scores were obtained when sequences were compared to the model of their own group (i.e., protistan sequences with the protistan HMM, etc., Supplementary Table S2). The difference between the top scoring and second best model was always large (>120). The e-values from the best model were always at least seven orders of magnitude smaller than the second best model (when ignoring two *C. reinhardtii* sequences, this difference was always larger than 30 orders of magnitude). This suggests that it is possible to discriminate protistan from bacterial and fungal PKS I KS sequences by scoring protein sequences with these HMMs. Protistan KS sequences will conform better to the protistan model (higher scores/lower e-values) than to the other models.

To illustrate the potential of this approach in finding new candidate sequences by mining sequence databases, we screened the env_nr (environmental shotgun sequences) database of NCBI for sequences showing high similarity to KS domains of PKS I by tblastn searches, using as query a KS domain from *C. parvum* translated the high scoring segments of the first 100 hits, and scored them with the three HMMs (protistan, fungal and bacterial KS). Although the differences among the scores with different models were not always as well defined as in our original data set, we identified several candidate sequences which likely encode protistan KS domains from PKS I genes (see Supplementary Table S3). This suggests that environmental sequencing projects have a high probability of identifying other novel protistan PKS genes, despite the fact that most of these projects are aimed at the prokaryotic size fractions of environmental samples.

**Evolution of PKS in Protists**

The global eukaryotic phylogeny is still unresolved and thus the phylogenomic relationships among the groups must be cautiously interpreted. Furthermore, the evidence for monophyly of each eukaryotic super-group remains inconclusive and is still the subject of debate. With these caveats in mind, our results nevertheless highlight the irregular phylogenetic distribution of Type I PKS genes in eukaryotes. Whereas representatives containing Type I PKS are known for three above-mentioned groups (unikonts, Plantae, and chromalveolates), they also contain members that do not possess this gene.

The view that Type III PKS is characteristic for Plantae is now challenged by the finding of Type I PKS in all three available chlorophyte genomes (*Chlamydomonas reinhardtii*, *Ostreococcus taurii* and *O. lucimarinus*). Taking into account the grouping of the PKS sequences of *Chlamydomonas* and *Ostreococcus*, and the fact that those taxa diverged long ago, this would put the gene divergence event near the base of chlorophyte evolution. However, no Type I PKS genes were found in another lineage of the Plantae, i.e. in the Rhodophyta *Cyanidioschyzon merolae* and *Galdieria sulphuraria*, the only two red algae thus far sequenced or in progress. A simple explanation is that these genes have been lost during evolution of the rhodophytes. Nevertheless, both rhodophyte species that have been sequenced are unicellular and live in acidic hot springs, and are thus atypical. It is therefore important to confirm whether or not PKS genes are generally absent from rhodophytes, including species living in less extreme (e.g., temperate or tropical marine) habitats.

There are several possibilities to explain the irregular phylogenetic distribution of Type I PKS genes among protists. One possibility is that a
common ancestor of the lineages Plantae, unic-onts and chromalveolates did indeed possess a
type I PKS gene, which was later lost among sev-
eral of the descendant lineages (at the least five losses, based upon the rather limited taxon
sampling available and depending upon the
accuracy of the proposed phylogenetic rela-
tionships of the taxa included). One can speculate
that these sequences were replaced by other
types of PKS genes, such as PKS III in Plantae,
and that they might have become unnecessary
and then lost in several other groups.

An alternative scenario is that PKS I genes were
obtained independently in several eukaryotic
lineages, either through duplication and diver-
genome from existing FAS genes in their genomes
lineages, either through duplication and diver-
several of the descendant lineages (at the least
three losses, based upon the rather limited taxon
sampling available and depending upon the
correlation of the taxa included). One can speculate
that these sequences were replaced by other
types of PKS genes, such as PKS III in Plantae,
and that they might have become unnecessary
and then lost in several other groups.

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gene from existing FAS genes in their genomes
lineages, either through duplication and diver-

gene from existing FAS genes in their genomes
lineages, either through duplication and diver-

The unexpected grouping of
Euryachytes
with chlorophytes (instead of alveolate) KS might
point to a common origin of E. huxleyi and green
algal PKS, as distinct from that of alveolates. Yet
we cannot exclude the possibility that the grouping
of E. huxleyi with chlorophytes (and exclusion of
alveolates) does not fully reflect the evolutionary
relationships of these genes. Sequences of some
hundreds of amino acids in length might not
contain sufficient phylogenetic information to cor-
rectly resolve relationships on time scales compar-
able to the age of eukaryotes.

Perhaps the key to understanding the origins and
evolution of the PKS genes in protists lies in the
analysis of the functional significance of these genes
and their role in secondary metabolism. The main-
tenance of PKS genes, which have no known
obligate role in primary metabolism, over long
evolutionary time suggests a persistent strong
positive selection, but which is not necessarily well
 correlated with the general evolution of the taxa. For
example, the selective pressure on secondary
metabolite production among free-living marine
flagellates of diverse phylogenetic lineages may be
more similar than that exerted upon free-living forms
of phylogenetically related obligate parasites.

Conclusion

The functions of protist polyketides are unknown, but
since they are structurally different from those of
bacterial and fungal origin it is not unreasonable to
expect they may have diverse and divergent func-
tions. In any case, the diversity of modular Type I PKS
products seems to arise from frequent recombination
events among the modules. The modular PKS
system provides an extraordinary platform for
recombination, with the evolutionary advantage that
the organisms have the ability to produce a large
chemical diversity from a limited number of different
genes.

Novel polyketides often exhibit bioactivity, with
potentially valuable functions as antibiotics
against bacterial, fungal or protozoan pathogens.
The protistan lineages have not been extensively
investigated for novel polyketides (with exception
of the known potent polyether biotoxins) and
therefore may represent a vast pool of undiscover-
ed bioactive substances, particularly critical at a
time when the severe clinical problem of multi-
drug resistant pathogens is on the rise.

Recent inferences on biosynthetic pathways
and secondary metabolite structures of bioactive
compounds based on comparative gene
sequence information in cyanobacteria (Sudek et
al. 2006) points the way to similar future dis-
covers from protists. Future work is aimed at
elucidating the structure of protistan polyketides
and determining the genetic regulation of tran-
scription of PKS genes to clarify the functional
significance of these secondary metabolites.

Methods

Data sets: Our base data set was the alignment from Kroken
et al. (2003) containing a representative subset of KS domains from
bacterial and fungal PKS, metazoan FAS and from oxoacyl-ACP
For phylogenetic analyses, multiple alignments were prepared which were performed with hmmsearch from the HMMer package. Models of regions of group-specific or overall conservation were identified genomic regions containing multiple fragments with similarity to different PKS domains as putative PKS genes. KS domain sequences longer than 300 amino acids, and showing significant similarity (e<10^-5) were used in the multiple alignments and phylogenetic analysis.

**Tentative assembly of Emiliania huxleyi PKS gene candidates:** At the time of our analyses, no assembly for the E. huxleyi genome was available. Thus, we have assembled genomic segments containing putative PKS fragments as follows. The sequence and quality files of the 3,868,954 genomic reads available (as of October 2005) from the NCBI trace database were downloaded and screened with BLASTn with the complete amino acid sequence of the C. parvum PKS (EAK87820) as query. All highly significant hits were re-blasted (BLASTn) against the traces to identify potentially overlapping reads. This selection of reads was assembled using phrap (http://www.phrap.org/) into 389 contigs, of which 17 were longer than 10,000 bases. Six of these contigs contained apparent frame-shifts. From the remaining eleven (containing combinations of all known PKS domains in large contiguous ORFs), five KS domains chosen from two contigs (showing the highest similarity to other protistan KS sequences) were further analyzed.

After the recent publication of the first draft of the E. huxleyi assembly, we checked for the presence of the KS domain sequences in the public assembly, which was cleaned of containing nucleotide sequences. The origin of these sequences from E. huxleyi was thus confirmed.

**Alignments and phylogenetic analyses:** Simultaneous HMM-based alignment and tree construction were performed with SATCHMO (Edgar and Sjölander 2003). SatchmoView was used to inspect the results and identify well-conserved regions for alignment within subgroups of the sequences. Profile hidden Markov models (HMM) for these regions were calculated and calibrated using hmmbuild and hmmpcalibrate from the HMMer package (http://hmmer.wustl.edu/). HMM logos for these models were generated from the HMM Logo web server “LogoMat-M” (http://logos.molgen.mpg.de/). HMM searches with the profile models of regions of group-specific or overall conservation were performed with hmmsearch from the HMMer package. For phylogenetic analyses, multiple alignments were prepared using kalign (Lassmann and Sonnhammer 2005). The alignment contained 130 sequences and 679 characters (provided upon request). Maximum likelihood phylogenetic trees were calculated with PhyML (Guindon and Gascuel 2003) using a Bio-NJ tree as starting tree, the WAG evolutionary model, with a gamma distribution parameter estimated from the data. Bootstrap analyses were performed with the same settings in 200 replicates. Bayesian analyses were conducted via MrBayes v.3.1.2 (Ronquist and Huelsenbeck 2003), with one million iterations, two runs with six chains each, a temperature parameter of 0.05 and with the WAG amino acid model. Burnin for construction of the consensus tree was 500,000 generations. The program Asatura (van de Peer et al. 2002) was used to explore substitution saturation in the multiple alignments.

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metabolites by heterologous expression in pseudomonads. Arch Microbiol 185: 28—38


