Research article

Open Access

In silico analysis of methyltransferase domains involved in biosynthesis of secondary metabolites

Mohd Zeeshan Ansari[†], Jyoti Sharma[†], Rajesh S Gokhale and Debasisa Mohanty^{*}

Address: National Institute of Immunology, Aruna Asaf Ali Marg, New Delhi-110067, India

Email: Mohd Zeeshan Ansari - zeeshan@nii.res.in; Jyoti Sharma - jyotisharma@nii.res.in; Rajesh S Gokhale - rsg@nii.res.in; Debasisa Mohanty* - deb@nii.res.in

* Corresponding author †Equal contributors

Published: 25 October 2008

BMC Bioinformatics 2008, 9:454 doi:10.1186/1471-2105-9-454

This article is available from: http://www.biomedcentral.com/1471-2105/9/454

© 2008 Ansari et al; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<u>http://creativecommons.org/licenses/by/2.0</u>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

Background: Secondary metabolites biosynthesized by polyketide synthase (PKS) and nonribosomal peptide synthetase (NRPS) family of enzymes constitute several classes of therapeutically important natural products like erythromycin, rapamycin, cyclosporine etc. In view of their relevance for natural product based drug discovery, identification of novel secondary metabolite natural products by genome mining has been an area of active research. A number of different tailoring enzymes catalyze a variety of chemical modifications to the polyketide or nonribosomal peptide backbone of these secondary metabolites to enhance their structural diversity. Therefore, development of powerful bioinformatics methods for identification of these tailoring enzymes and assignment of their substrate specificity is crucial for deciphering novel secondary metabolites by genome mining.

Received: 21 April 2008 Accepted: 25 October 2008

Results: In this work, we have carried out a comprehensive bioinformatics analysis of methyltransferase (MT) domains present in multi functional type I PKS and NRPS proteins encoded by PKS/NRPS gene clusters having known secondary metabolite products. Based on the results of this analysis, we have developed a novel knowledge based computational approach for detecting MT domains present in PKS and NRPS megasynthases, delineating their correct boundaries and classifying them as N-MT, C-MT and O-MT using profile HMMs. Analysis of proteins in nr database of NCBI using these class specific profiles has revealed several interesting examples, namely, C-MT domains in NRPS modules, N-MT domains with significant homology to C-MT proteins, and presence of NRPS/PKS MTs in association with other catalytic domains. Our analysis of the chemical structures of the secondary metabolites and their site of methylation suggested that a possible evolutionary basis for the presence of a novel class of N-MT domains with significant homology to C-MT proteins could be the close resemblance of the chemical structures of the acceptor substrates, as in the case of pyochelin and yersiniabactin. These two classes of MTs recognize similar acceptor substrates, but transfer methyl groups to N and C positions on these substrates.

Conclusion: We have developed a novel knowledge based computational approach for identifying MT domains present in type I PKS and NRPS multifunctional enzymes and predicting their site of methylation. Analysis of nr database using this approach has revealed presence of several novel MT domains. Our analysis has also given interesting insight into the evolutionary basis of the novel substrate specificities of these MT proteins.

Background

Nonribosomal peptide synthetases (NRPSs), polyketide synthases (PKSs) and fatty acid synthases (FASs) employ a common biosynthetic strategy to synthesize their metabolic products by stepwise condensation of simple amino or carboxylic acid monomers. The core catalytic domains involved in the biosynthesis of the polyketide/nonribosomal peptide/fatty acid backbone moieties are ketosynthase (KS), acyltransferase (AT), dehydratase (DH), enoylreductase (ER), ketoreductase (KR), acyl carrier protein (ACP), condensation (C), adenylation (A) and thiolation (T) [1,2]. Apart from these core catalytic domains, a number of auxiliary functional domains, often called tailoring domains, introduce a variety of different chemical modifications to the backbone moieties of these secondary metabolites to further increase their structural diversity. Bioinformatics analysis of various catalytic domains present in NRPS and PKS proteins has been an area of active research in recent years [3-8]. These studies [3-8] have not only led to development of novel computational methods for in silico identification of secondary metabolites by genome mining [9-16], they have also guided rational reprogramming of secondary metabolite biosynthetic pathways to generate designed "natural products" [12,17-20]. However, all these studies including our earlier work have concentrated on core catalytic domains and no detailed bioinformatics analyses have been carried out for important tailoring enzymes like, methyltransferases.

Methyltransferase (MT) domains present in NRPS and PKS clusters constitute a major class of tailoring domains/ enzymes involved in biosynthesis of secondary metabolites. They catalyze the transfer of methyl group from Sadenosylmethionine (SAM or AdoMet) to the carbon, nitrogen or oxygen atoms at various positions on the backbones of polyketides, nonribosomal peptides and fatty acids and therefore have been classified as C-MT, N-MT and O-MT respectively depending upon their site of methylation. These enzymatic domains in general have a bidomain structure, where the first subdomain contains the binding site for methyl group donor, while the second subdomain harbors the binding site for acceptor substrate [21,22]. The presence of MT domains in multifunctional NRPS and PKS proteins is generally inferred from chemical structure of the secondary metabolite products. There are only few in vitro studies on enzymatic characterization of NRPS/PKS MT domains [23-27]. A recent study on MT domains from type II PKS biosynthetic pathways has revealed interesting correlation between regioselectivity of methylation and MT sequence [24]. However, no such analysis has been carried out for MT domains present in type I PKS or NRPS proteins. In contrast to type II PKS MTs which are stand alone proteins, MT domains in type I PKS and NRPS are present along with other catalytic domains on a single polypeptide chain. Therefore, it has been difficult to decipher the correct length and domain boundaries for MT domains in type I PKS or NRPS proteins. Various studies have suggested that the size of N-MT domain is typically 450 amino acids, while C-MT and O-MT are generally 300 amino acids long. A set of 3 conserved sequence motifs has been identified in most MTs [28-30]. Mutational studies of N-MTs of peptide synthetases have shown that these 3 motifs are essential for the catalysis [31]. The knowledge of these MT sequence motifs and the expected spacing between them is often used for discerning presence of MT domains in multifunctional NRPS and PKS proteins. However, because of the high degree of sequence divergence, delineating the correct boundary of these proteins is quite often a difficult task. In our earlier study, we attempted to identify MT domains in various NRPS/PKS gene clusters based on pairwise alignment with MT domain from actinomycin cluster [32]. However, this domain identification protocol failed to detect 23 out of 32 MT domains. The 23 unidentified MT domains included the three groups of MTs (C-, O- and N-MTs), for which proper templates were not available. The general purpose domain identification tools like CDD-search can identify MT domains in NRPS and PKS proteins, but can not predict the domain boundaries accurately and they also fail to classify them as C-MT, N-MT and O-MT. Such classification is crucial for prediction of chemical structures of secondary metabolites. The knowledge of substrate specificity and domain boundaries of MT domains is also important for rational design of novel secondary metabolites by introduction of heterologous MT domains.

In this manuscript, we have carried out a systematic analysis of the sequence/structural features of MT domains present in various experimentally characterized NRPS and type I PKS clusters having known metabolic products. Since crystal structures are available for many stand alone small molecule methyltransferases from several microbial organisms, we have carried out threading analysis for the experimentally characterized MT domains from NRPS and PKS biosynthetic pathways. The threading analysis has helped in elucidating the putative three dimensional structure adopted by MT domains and based on the alignment of MT containing sequences on the structural fold of MT domain it has been possible to delineate the correct boundaries for NRPS/PKS MT domains. Our threading analysis has also given novel insight into the structural features of linker sequences flanking the MT domains in NRPS and PKS proteins. Using the curated sequences of these MTs, we have carried out detailed phylogenetic analvsis to investigate whether these catalytic domains cluster as per their specificity for site of methylation i.e. C-MT, N-MT and O-MT. Based on this analysis, we have identified suitable template sequences of C-MT, N-MT and O-MT domains from representative clusters, which can be used

to identify MT domains in uncharacterized NRPS/PKS proteins. We have also developed Hidden Markov Model (HMM) profiles which can identify MT domains in a query sequence and classify them as N-MT, C-MT and O-MT. Using these HMM profiles, we have analyzed non-redundant protein sequence database of NCBI to identify other multifunctional enzymes containing C-MT, N-MT and O-MT domains.

Results

In this study, we have carried out a number of different bioinformatics analyses on MT domains present in type I PKS and NRPS proteins, to correlate the sequence of these MT domains to their substrate specificity i.e. the site of methylation. Figure 1 gives a schematic overview of the various different analyses carried out and the type of results obtained from them, while the results are discussed in detail in the following sections.

The chemical structures of the secondary metabolites produced by various PKS, NRPS and hybrid NRPS/PKS clusters cataloged in NRPS-PKS web resource were analyzed carefully to identify methyl substitutions on polyketide or nonribosomal peptide backbones. The presence of methyl substitutions on nitrogen and oxygen atoms indicated presence of N-MT or O-MT domains in the proteins encoded by these gene clusters. However, in absence of MT domains methyl substitutions on carbons in a ketide group can also result from selection of methylmalonate extender groups by the AT domains of PKS proteins.

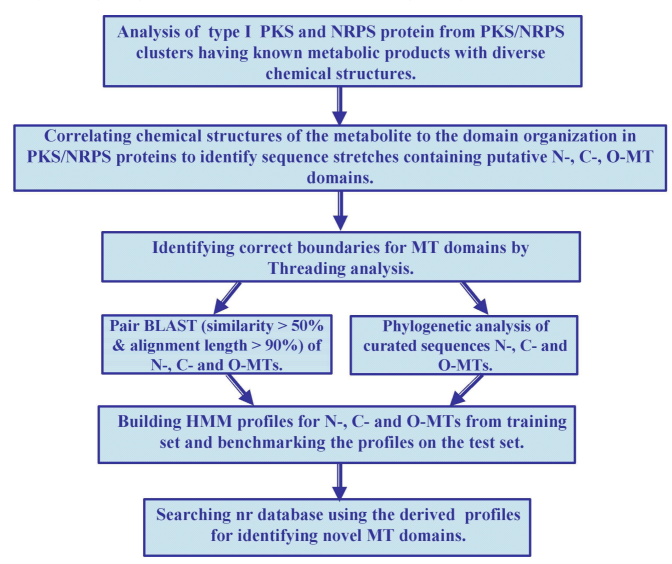


Figure I

A schematic overview of different bioinformatics analyses carried out in the current work on MT domains present in type I PKS and NRPS proteins.

Therefore, for correctly inferring presence of C-MT domains in a PKS protein, the substrate specificity of the corresponding AT domain was also checked. Table 1 lists various ORFs harboring the MT domains, their GenBank accession number and the type or substrate specificity of MT domain as deduced from the chemical structure of the metabolite. As can be seen, the data set consisted of 20 C-MT, 19 O-MT and 22 N-MT domains from 27 different NRPS/PKS clusters. Figure 2 shows the chemical structures of 5 representative secondary metabolites highlighting the methyl groups added by C-MT, N-MT and O-MT domains, while chemical structures of the remaining 22 secondary metabolites are shown in Additional file 1. Each of the ORFs listed in Table 1 were analyzed by NRPS-PKS search tool as well as CDD server of NCBI. NRPS-PKS search tool, which used a single MT domain from actinomycin cluster as template, could identify only 26 MT domains out of a total of 61. Even though the latest version of CDD server could identify 55 out of these 61 MT domains, the lengths of the MT domains detected by both these programs were notably shorter than the typical length of these domains. These programs also failed to distinguish between C-MT, N-MT and O-MT domains. Additional file 2 shows the length of each MT containing sequence stretch and other catalytic domains flanking this region. As can be seen, all N-MT domains are present in NRPS clusters only as C-A-MT-T modules and typically a 400 amino acid sequence stretch containing this domain is inserted in the adenylation domain between the conserved motifs A8 and A9 of adenylation domains and hence alignment of these N-MT containing A domains with regular A domains produces a split alignment (Figure 3). All stand alone O-MT containing sequences were typically 300 to 400 amino acids long, while the amino acid stretches containing other O-MT and C-MT domains present in type I PKS proteins were 600 to 700 amino acids long. The O-MT and C-MT domains in PKS proteins were present in four different types of modules i.e. KS-AT-MT-ACP, KS-AT-MT-KR-ACP, KS-AT-DH-MT-KR-ACP and KS-AT-DH-MT-ER-KR-ACP. Only in case of leinamycin gene cluster, which has trans-AT domains, the MT domain is present as KS-DH-KR-ACP-MT-ACP. There were only two examples where MT domain was adjacent to an aminotransferase (AMT) domain in a hybrid NRPS/PKS system. In these cases MT-AMT stretch was inserted between a PKS and a NRPS module. It appears that MT domains in type I PKS proteins are present in AT-KR, DH-ER or DH-KR linker regions which are typically more than 200 amino acids long. Hence, length of the flanking linker region could be the reason for the larger length of these MT containing sequence stretches. Therefore, we decided to carry out various structure based sequences analysis for representative MT containing stretches of each category to delineate the exact domain boundaries for MT domains.

Threading analysis of MT domains

Since domain boundaries can be identified correctly by aligning the sequence of multi domain proteins with the 3D structures of the corresponding single domain proteins, we attempted to identify other proteins in PDB which are structurally similar to these MT domains of PKS/NRPS enzymes. However, the lack of crystal structures for any MT domains from PKS/NRPS biosynthetic pathway and high degree of sequence divergence in this enzyme family prompted us to use threading or fold recognition approach. These tools can potentially reveal structural similarity in absence of high degree of similarity in sequences. GenTHREADER and PHYRE fold prediction servers were used for threading analysis. As discussed in the methods section, the MT sequence stretch identified by CDD or NRPS-PKS along with their flanking linkers was threaded on various structural folds in PDB. In cases where chemical structure of metabolite indicated presence of MT domains but no MT domain was detected by these programs, all linker stretches having unusual length were analyzed by both these fold recognition servers.

Table 2 shows the results of threading analysis for representative members of these MT containing sequences. The fold prediction hits with highest level of statistical significance corresponding to p-value lower than 0.0001 were labeled as CERTAIN by the GenTHREADER server, while hits having p-value between 0.0001 and 0.001 are labeled as HIGH. We considered only those matches which are labeled as CERTAIN or HIGH by GenTHREADER or have precision of more than 95% in case of fold prediction by PHYRE. As can be seen, all the 18 sequences matched with structure of MT proteins in PDB. This suggests that, the MT domains present in NRPS/PKS proteins would adopt a fold similar to the small molecule methyltransferase in other organisms. The N-MT domains present in our data set aligned not only with N-MT structures, but also with C-MT and O-MT structures. Similar was the case for C-MT and O-MT domains in our data set. Analysis of their alignment scores did not show any preference for structural matches from the same functional category. Therefore, the structures which aligned consistently with all the sequences by both servers and had maximum sequence identity and alignment length with the query sequences were chosen as structural templates for MT domains of NRPS/PKS proteins. The structure of histamine N-MT (PDB code 1VLM; 207 residues) and a hypothetical protein from M. tuberculosis (PDB code 1VL5; 230 residues) showed alignment with most C-MT, O-MT and N-MT by both the fold prediction servers. However, taking into consideration the alignment length and the length of the query sequence, for further analysis 1VLM was selected as template for C-MT and O-MT domains, while 1VL5 was selected as structural template for N-MT domains. These results suggest that, MT domains present in type I PKS and

Name of gene cluster	ORF	Accession no.	CDD search	Types of I	MT-domain		Tota
				C-MT	O-MT	N-MT	
				NRPS clusters			
Actinomycin	acmC	AAF42473	PF08242			2	2
Anabaenopeptilide	apdB	CAC01604	PF08242	_	_	2	2
Anabaenopeptilide	apdB	CAC01607	PF08241	-	-	2	3
Complestatin	comC	AAK81826	PF08242	-		-	J
	simA	CAA82227	PF08242	-	-	7	7
Cyclosporine Enniatin		CAA82227 CAA79245	PF08242 PF08242	-	-	/	/
	esynl			-	-		1
Pristinamycin	snbDE	T30289	PF08242	-	-		1
Pyochelin	pchF	AAD55801	PF08242	-	-		I
Thaxtomin	txtA	AAG27087	PF08242	-	-	1	-
	txtB	AAG27088	COG2226	-	-	I	2
				PKS cluste	ers		
Compactin	mlcA	BAC20564	PF08242	Ι	-	-	
	mlcB	BAC20566	PF08242	I	-	-	2
Erythromycin	eryG	CAA42929	COG2226	-	I.	-	I
Equisetin	eqiS	AAV66106	PF08242	I	-	-	i
Fumonisin	fuml	AAD43562	PF08242	i	-	_	i
Lovastatin	lovB	Q9Y8A5	PF08242	i	_	_	
Lovastatin	lovF	AAD34559	PF08242	1	-	-	2
Stiensstallin			PF08242	1	-	-	2
Stigmatellin	stiD	CAD19088		-		-	
	stiE	CAD 19089	PF08242	-	1	-	
	stiK	CAD19094	PF01209	-		-	3
				Hybrid N	RPS-PKS		
Bleomycin	blmVIII	AAG02357	PF08242	I	-	-	Ι
Barbamide	barF	AAN32980	PF08242	-	I	-	
	barG	AAN32981	PF08242	-	-	1	2
Epothilone	epoD	AAF26922	PF08242	1	-	-	Ī
amaicamide A	jam]	AAS98781	PF08242	i	-	_	-
	jamN	AAS98785	COG0500		-	_	2
Leinamycin	•	AAN85523	PF08242	-		-	1
Melithiazol	lnmJ melE	CAD89776	PF08242		-	-	
TENUIIAZUI	melF	CAD89776 CAD89777	PF08242 PF08242	-	1	-	2
Microcystic				-	I	-	2
Microcystin	mcyD	BABI2210	PF08242	1	-	-	
	mcyE	BABI2211	- DE000.40	1	-	-	
	mcyG	BAB12213	PF08242	I	-	-	
	mcyA	BAA83992	PF08242	-	-	I	4
Myxothiazol	mtaE	AAF19813	PF08242	-	I	-	
	mtaF	AAF19814	PF08242	-	I	-	2
Nodularin	ndaC	AAO64404	-	I	-	-	
	ndaD	AAO64405	PF08242	I	-	-	
	ndaF	AAO64407	-	I	-	-	
	ndaA	AAO64403	PF08242	-	-	I.	
	ndaE	AAO64406	PF08242	-	I	-	5
Onnamide	onnB	AAV97870	PF08242	-	I	-	
	onnD	AAV97872	PF08242	-	I.	-	
	onnG	AAV97875	PF08242	-	1	-	
	onnH	AAV97876	COG2226	_		_	
	onnl	AAV97877	PF08242	-	1	-	5
Pederin				-	I	-	J
euerin	pedF	AAS47564	PF08242	I	-	-	
	pedA	AAS47557	PF08241	-		-	-
	pedE	AAS47560	COG2226	-	I	-	3

Table 1: List of ORFs containing C-, O- and N-Methyltransferase domains

	•	-		,				
Tubulysin	tubF	CAF05651	PF08242	I	-	-		—
	tubB	CAF05647	PF08242	-	-	I		
	tubC	CAF05648	PF08242	-	-	I	3	
Yersiniabactin	HMWP-I	AAC69588	PF08242	2	-	-	2	
Total				20	19	22	61	

List of ORFs containing methyltransferase domains in various experimentally characterized NRPS/PKS gene clusters. Table also lists the number and type of MT domains in each of the ORFs and results from PFAM analysis using CDD search.

NRPS proteins are likely to be around 200 amino acids long.

It may be noted that the small molecule MT structures present in PDB show significant variation in their length and in case of some of these sequences alignments were found with MT structures having significant difference in length. For example, 3 of the C-MT domains and 1 N-MT domain in our data set aligned with aclacinomycin-10-hydroxylase (PDB code <u>1QZZ</u>; 340 residues) [33] and histamine N-MT (PDB code <u>1VLM</u>; 207 residues). The huge length difference between these two MT structures made the choice of correct structural template a difficult task. 1QZZ aligns with the beginning of the MT containing

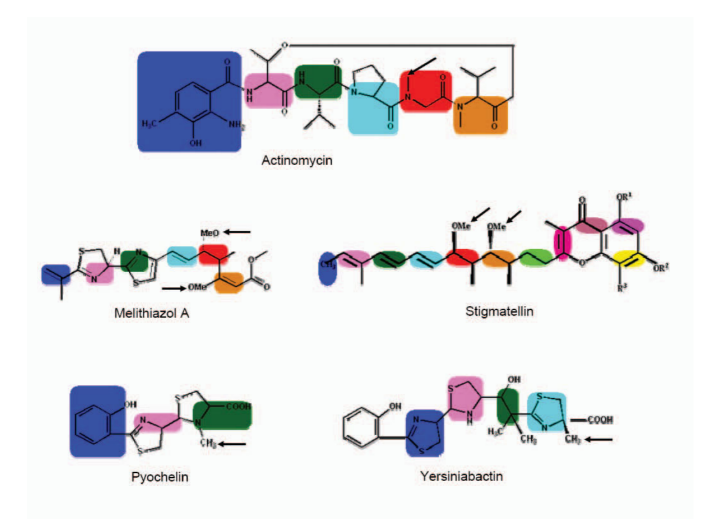
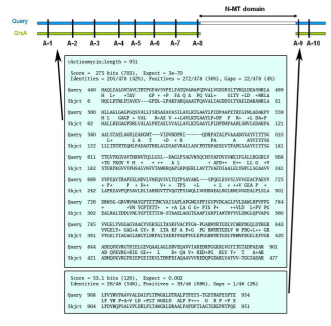


Figure 2

Chemical structures of representative secondary metabolites like nodularin, leinamycin, pyochelin, yersiniabactin and stigmatellin containing methyl groups (highlighted by arrow sign) added by C-MT, N-MT and O-MT enzymatic domains.



Alignment of the sequence stretch containing A and N-MT domains from a C-A-MT-T NRPS module with sequence of A domain from a C-A-T module. A split alignment is obtained, because N-MT domain is integrated between A-8 and A-9 signature motifs of A domain.

sequence stretch, while in the alignment of the same sequences with 1VLM, the length of the linker preceding the MT domain ranges from 50 to even 200 residues (Figure 4). However, careful manual analysis of the alignments as well as the corresponding structural templates indicated that, the crystal structure 1QZZ consists of a predominantly *a*-helical N-terminal domain which is involved in dimerization and a C-terminal SAM binding domain typical of methyltransferases. On the other hand, the crystal structure 1VLM contains the SAM dependent MT domain alone. The overlapping region of both the alignments correspond to the catalytic domain of methyltransferase, while the linker sequences preceding the MT catalytic domain showed alignment with additional dimerization domains, if present in the structural template. Earlier studies on structural analysis of small molecule methyltransferases have suggested that, these enzymes have a conserved SAM-MT fold consisting of alternating β strands and α helices [22]. Earlier bioinformatics analysis [28,30,31] of MT domains has also revealed that, despite high divergence in primary sequence, certain conserved sequence motifs are present in all SAM dependent methyltransferases. Therefore, we wanted to examine whether the conserved secondary structural elements and sequence motifs responsible for catalytic activity, substrate specificity and cofactor binding are conserved in the MT domains identified by threading alignments. Figures 5, 6 and 7 show the structure guided multiple alignment of representative N-MT, C-MT and O-MT sequences with the structural template 1VLM. As can be seen from Figure 5, the N-MT sequences have the conserved motifs I, II/Y, IV and V in N-MTs. Similarly motifs I, motif I-post, II and III are present in the C-MT and O-MT sequences (Figures 6 &7). Thus all the motifs [28,30,31] identified in earlier analysis of small molecule methyltransferase sequences were found to be conserved in the multiple sequence alignments of C-, O- and N-MT domains identified by our threading analysis. The average percent identity among the C-, O- and N-MT domains was found to be 31, 28 and 17% respectively.

The threading analysis also showed statistically significant matches with proteins other than methyltransferases. Such matches were specifically seen for MT containing sequences which were longer in length due to the presence of large flanking linker sequences. As can be seen from table 2 and figure 4, the N-terminal region of MT containing sequence from bleomycin showed an alignment (Figure 8a) with the last 60 amino acids of the KS-AT didomain structure from erythromycin PKS [34]. This stretch corresponds to the last helix of the AT domain and a segment of the AT-DH linker region (Figure 4). The Cterminal 200 amino acid stretch of the MT containing sequence from bleomycin [35], nodularin and melithiazol also showed highly significant alignments (Figure 8b) with the structural subdomain (Figure 4) of the recently elucidated structure of KR from erythromycin PKS. These results suggest that, in type I PKS proteins, the linker sequences preceding the MT domain i.e. AT-MT linkers are homologous to the AT-DH linkers and MT-KR linker regions are likely to adopt a short chain reductase (SCR) fold and would constitute the structural half of the KR domain as demonstrated in erythromycin PKS. Thus MT catalytic domain is likely to be 200 amino acids only.

Development of a computational protocol for identifying MT domains and their classification as C-MT, O-MT and N-MT

The domain boundaries were thus identified with 1VLM (for C- and O-MT) and 1VL5 (for N-MT) as templates and the regions which aligned with these templates were extracted from the 61 MT domain sequences. They represented the curated MT domains with correct boundary. A set of 42 MT domains out of these 60 sequences were used as test set to check if MT domains can be correctly classified by pairwise alignment with these 18 template sequences. Each of the 42 MT domains was queried against 18 MT templates to find the number of MT domains identified by these templates. The query MT domain was classified as C-MT, O-MT or N-MT depending on the highest scoring match from the template set. The results obtained by the pairwise comparison of 60 query

Methyltransferases	Len.	2hg4	lqzz	l vI5	lvlm	lwzn	2gpy	2aot	lg6q	l xxl	lim8	2fr0
		917	374	260	219	252	233	292	328	239	244	486
melit01_OM_001	609	С	-	C {100}	С	-	С	-	-	-	-	С
anaba01_OM_001	263	-	-	C {100}	C {100}	С	-	С	-	{100}	-	-
onnam04_OM_001	268	-	-	C {100}	С	С	-	-	-	{100}	-	-
peder01_OM_001	312	-	-	C {100}	С	-	-	-	-	{100}	-	-
stigm03_OM_001	256	-	-	C {100}	C {100}	С	-	-	-	{100}	-	-
eryth01_OM_001	306	-	-	C {100}	C	-	С	-	-	{100}	-	-
barba01_OM_001	422	-	-	C {100}	С	С	С	-	-	{100}	-	-
onnam01_OM_001	484	-	-	{100}	C {100}	-	-	С	-	{100}	{100}	-
bleom01_CM_001	640	С	-	C {100}	C {100}	С	С	-	-	{100}	{100}	С
nodul02_CM_001	720	-	С	C {100}	C {100}	-	С	С	-	{100}	-	С
compa01_CM_001	490	-	-	C {100}	C {100}	-	С	С	-	{100}	-	-
leina01_CM_001	432	-	С	C {100}	C {100}	C {100}	С	С	-	-	-	-
yersi01_CM_002	479	-	С	C	C	-	С	С	-	-	{95}	-
actin01_NM_001	422	-	-	{100}	{100}	C {100}	С	-	-	{100}	-	-
anaba01_NM_001	367	-	-	-	-	-	С	-	H {100}	-	-	-
cyclo01_NM_001	43 I	-	-	{100}	{100}	-	н	-	-	{100}	-	-
pyoch01_NM_001	390	-	н	{100}	C {100}	-	С	С	-	{100}	{100}	-
thaxt02_NM_001	362	-	-	{100}	C {100}	С	С	-	-	{100}	-	-

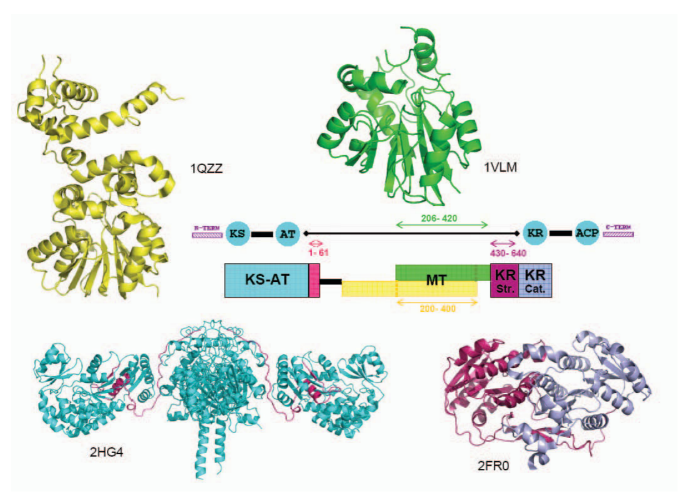
Table 2: Threading analysis of 18 representative MT containing sequences

Column I gives the unique name assigned to each MT containing sequence stretch, while the second column indicates their length. Column 3–13 lists the PDB IDs for the structures which show alignment with these sequences in fold recognition analysis using GenTHREADER and PHYRE servers. Results from GenTHREADER corresponding to confidence level CERTAIN and HIGH are labeled as C and H respectively. Similarly, PHYRE results corresponding to precision level 100% and 95% are labeled as 100 and 95 respectively in curly braces. (-) indicates the absence of that particular fold.

sequences with the 18 template sequences indicated that C-MT templates were able to identify all other C-MT sequences. However, there were a few O- and N-MTs, which were also recognized by these C-MT templates. Specifically, 2 MT sequences in onnamide gene cluster (onnB and onnI) showed very high similarity to C-MT while they are functionally annotated as O-MTs by the authors who reported experimental characterization of this gene cluster [36]. These 2 O-MTs also have motif I as ExGxG which is characteristic of C-MTs. An N-MT from pyochelin synthetase (pchF) also exhibited considerable similarity to several C-MTs and the two onnamide O-MTs. In view of this apparent anomaly in sequence similarity of these proteins, their functional assignment needs to be examined carefully. In the ORF apdB of anabaenopeptilide gene cluster, there are two MT-domains wherein the first one is entirely different from all the other MT sequences and does not show similarity with any other MT sequence. A dendrogram (Figure 9) of all the 60 MT sequences also illustrates the same pattern of results as obtained by these pairwise alignments. The two onnamide O-MTs in onnB and onnI genes and the N-MTs in pyochelin and anabaenopeptilide show clustering with C-methyltransferases. A stand alone O-MT in stigmatellin (stiK) is sequentially different from other O-methyltransferases, which was observed from the pairwise alignment results and is evident from the dendrogram. Thus it can be concluded from the above analysis that these 18 sequences can be used as templates to identify MT-domains in any given query sequence by pairwise alignment. These new MT templates were included in the current version of NRPS-PKS program for correct identification of various types of MT domains.

Profile HMMs of C-, O- and N-methyltransferases

An alternative approach to detect MT domains in a new sequence is to query that sequence against a database of HMM profiles. Individual profiles were built for C-MT, O-MT and N-MTs using the curated sequences. These profiles were then used to make a HMM profile database for MT domains. The set of 61 C-MT, O-MT and N-MT sequences from experimentally characterized NRPS/PKS clusters were queried against this HMM database and the location of the MT-domain and their class was predicted in these sequences. Table 3 lists the score and E-value for alignment of 18 representative MT sequences with the HMM profiles of N-MT, C-MT and O-MT domains. As can be seen from Table 3, most C-MT sequences show statistically significant alignment with O-MT profiles and vice versa. On the other hand, only the N-MT sequences from pyochelin aligned with the N-MT as well as C-MT profiles unlike the other N-MT sequences which aligned with N-



Schematic representation of the results of threading analysis for typical C-MT containing sequence (from bleomycin ORF blmVIII) stretches having large length. The central stretch aligns with various methyltransferase crystal structures like IVLM and IQZZ. A 200 amino acid C-terminal stretch aligns with the structural half of the KR domain in the crystal structure 2FR0, a 60 amino acid N-terminal stretch shows alignment with the terminal stretch of the KS-AT di-domain structure 2HG4. The query sequence containing the MT domain is represented as a black line, while rectangular colored boxes represent matches with various structural folds. The corresponding structures are shown in the same color.

MT profile alone. This finding is consistent with results from pairwise sequence alignment discussed in the previous section.

In order to test the predictive ability of our MT HMM profiles further, the recent version of the nr database of NCBI was also searched using these profiles to identify putative NRPS/PKS MT domains in various proteins. The sequences which matched with these profiles were grouped as C-MT, O-MT and N-MT containing sequences based on highest scoring profile match. The complete domain architecture of these MT containing proteins were also analyzed in details. The nr database search identified 4197 stand alone MT proteins and 684 multifunctional proteins containing MT domains. Out of the 4197 stand alone MT proteins, 3977 were O-MT proteins. In contrast to the very large number of stand alone O-MT proteins, there were only 155 stand alone C-MT and 55 stand alone N-MT proteins. Even though experimentally characterized NRPS/PKS biosynthetic pathways have a relatively larger number of stand alone O-MT proteins compared to stand alone C-MT or N-MT domains, it is not apparent whether all these stand alone O-MT proteins identified by our profile search would indeed be associated with secondary metabolite biosynthesis. Analysis of domain architectures in multifunctional proteins containing MT domains revealed several interesting results. These MT containing multifunctional proteins containing MT domains along with core PKS domains, core NRPS domains, PKS as well

		Motif I	Motif II/Y
1VLM_A	α_1 α_2 α_3		$\begin{array}{ccc} \alpha 4 & \beta 2 & \alpha 5 \\ 0 0 0 0 & \longrightarrow & 0 0 0 0 0 0 0 0 0 \\ \end{array}$
IVLM_A actin01_NM_001 actin01_NM_002 anaba01_NM_001 compl01_NM_001 cyclo01_NM_001 cyclo01_NM_002 cyclo01_NM_002 cyclo01_NM_004 cyclo01_NM_005 cyclo01_NM_005 cyclo01_NM_005 cyclo01_NM_007 ennia01_NM_001 micro01_NM_001 prist01_NM_001 pyoch01_NM_001	100000 000000000 000000000000000000000	22020 340 AVKCLLPEGRGVEIGU PRRVLEIGVGTGLLI KDKIVVEIGTGKDAII PQRVLEIGCGSGLLI PNSVWEVGCGTGMLI PNSVWEVGCGTGMLI PRRVLEIGVGTGTGMVI PAGRVLEVGTGTGMVI PAGRVLEVGTGTGMVI PPGHVLEIGTGTGMVI IPRVLEIGTGTGMVI IPRVLEIGCGTGLII PKKVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII PRRVLEIGCGTGLII	
	22	Motif IV	Motif V
1VLM_A	→ TT TT 80	β4 η1 200 200 20 100	$ \begin{array}{c} \alpha 6 & \beta 5 \\ 1 0 0 0 0 0 0 0 0 0 0 \\ 1 1 0 & 1 2 0 \end{array} $
<pre>IVLM A actin01 NM 001 actin01 NM 001 actin01 NM 002 anaba01 NM 002 barba01 NM 002 barba01 NM 001 cyclo01 NM 001 cyclo01 NM 003 cyclo01 NM 004 cyclo01 NM 004 cyclo01 NM 006 cyclo01 NM 007 micr001 NM 001 micr001 NM 001 prist01 NM 001 prist01 NM 001 thaxt01 NM 001 thaxt01 NM 001 tubul01 NM 001 tubul01 NM 001</pre>	AARVOLRTQPAHDFDQLPHG. ARRVTLRAGAAHBHEGLPVG. KITVIHGDATLVDIPELADVCVSEIVGAIGG. LENVRLLHKTADNFEGIPQG. YAHVTLAQKAAEBMADIADN. AAKVRLRVQAADBTDGLPEG. G.KAQVHVGTAMDTG.RLSA.L. D.RIRVEVGTATDIDRLGDD.L. G.RAEVHVGTAADVGTLQG.L. DGRSIVHVGTAADVGTLQG.L. DGRSIVHVGTAEDIN.KAGPI. G.KARVLVGTAEDIK.LVKD.F. G.SARVHVGTAEDIS.SIDG.L. G.KARVHVGTAEDIS.SIDG.L. G.KAKVHVGMATDVN.KLGE.V. LEQVRLLHSTADNFEGLBSE. WSQVQLYNQPAHNLQGFBPK. ADKVTVDARAAHBTDDLPKG ERFADQPWVRFGRFDMN. ADRVVLHQLSAHBLGSLPSG. REKVHLLHGPAHDLPGLPEG. AQRVRLMAQPADDFSGLPEA.	. FD FALMVTTICFVDDPE HFDTVVLNSVVQYFPNAG HFDTVVLNSVVQYFPNAG SEGAAVIINNARRFLKPD SEGAAVIINNARRFLKPD SPDVLLSSIVQYFPSVD SFDVULSSIVQYFPSVD SPDVLLSSIVQYFPSNAD SPDVLIVINSVAQYFPSRE HASLVVNSVAQYFPTPE QPELVVINSVAQYFPTPE QPELVVINSVAQYFPTPE SRSDLVVINSVAQYFPSRE HPDVVVINSVAQYFPSRE HPDVVVINSVAQYFPSRE HPDLVVINSVAQYFPSRE SSDLVVINSVAQYFPSRE HPDLVVINSVQYFPSID RFDTIVLNSVQYFPSDA .SDLLDQGVAPHSVDIL GFDTVVLNSVQYFPSGD YFDTVVLNSVQYFPSAD GFDTVVLNSVQYFPSAD	RALKEAYRILKKGGYLIVGIVDRESFLG YLEQVLDHALRILAPGGTVFIGDIRNERLLRTFTTA GLAQVIEQALRLLAPGGAVFIGDIRNERLLRTFTTA GLMIPERSITKMAAVTLPDEILHNFKFABIPAYYTK YLLQVLEGAMTATASQGKIFVGDVRSLPLLPYHAA YLLQVLEGAMTATASQGKIFVGDVRSLPLLPYHAA YLLQVLEGAMTATASQGKIFVGDVRSLPLLPYHAA YLLQVLEGAMTATASQGKIFVGDVRNHRLLRPPATA YLQVLEGAMTATASQGKIFVGDVRNHRLLRPPATA YLQVLEGAMTATASQCKIFPGDMRTYATHKDFLVA YLQVLEGAMTATASQCKIFPGDMRTYATHKDFLVA YLQVLEALVRIFG.VRRIFPGDMRTYATHKDFLVA YLQVLFALVRDJFS.VERIFPGDMRTYATHKDFLVA YLQVLFALVQIFS.VERIFPGDMRTNAINRDFAAA YLQVUFALVQIFS.VERIFPGDMRTNAINRDFAAA YLQVIASELIHMTS.VKTIFFGDMRTYATNKDFLVA YLLABLTANLIRLFG.VKRIFPGDMRTYATNKDFLVA YLLAVUFAVES.VKRIFFGDMRTYATNKDFLVA YLAGLUFASLIHMTS.VKTIFFGDMRTYATNKDFLVA YLLAVLEGAIKVLTPGGCIFLGDVRNIQLMEAPHAD YLVSVLEGAVEMVAPGGWIFVGDVRSLPLLPAPHAD YLVSVLEGAVEMVAPGGVFLGDVRNIQLMEAPHAD YLVSVLEGAVEMVAPGGVFLGDVRNIQLMEAPHAD YLYSVLEGAVEMVAPGGVFVGDVRSLPLLPAPHAD YLAGILNALDTPALLAGLRELSADAWLVI YLFDLLREVSRLLVFGGAVFLGDVRNIRLLRTFHAG YLVSVLEGALRVLQPGGTLFIGDVQNIRLFELPHAS YLVSVLEGALRVLQPGGTLFIGDVQNIRLFELPHAS
1VLM_A		α8 <u>β6</u> <u>60</u> <u>170</u>	$\begin{array}{ccc} \eta 2 & & & & & & \\ \underline{\rho} 2 & & & & & & \\ \underline{\rho} 2 & & & & & & \\ 18 $
1VLM A actin01_NM_001 actin01_NM_002 anaba01_NM_001 barba01_NM_001 cyclo01_NM_001 cyclo01_NM_002 cyclo01_NM_002 cyclo01_NM_003 cyclo01_NM_005 cyclo01_NM_005 cyclo01_NM_006 cyclo01_NM_006 cyclo01_NM_001 micro01_NM_001 prist01_NM_001 thaxt02_NM_001 tubul01_NM_001 tubul02_NM_001	VHT ARATDP ADHAAVRRAVERA VQT.ARABDP ADTAAVRRAVEQS KIFEEVGYPFDLRVCIKKFSQANVLSTVDVL VQL.ARABD KTVEQWQQQVHQTL VQL.ARABD KTVEQWQQQVHQTL VQL.HKAPPS LSVQQLKQGIYRL VET.ARTAD QDVFTLRAVEQN RAVHTNGSK VTRSKVQQEVARL RAVHALGDK ATKABIQREVVRM RAVHALGDK ATKABIQREVVRM RAVHALGDK ATKABIQREVVRM RAVHLGDK ATKABIQREVVRM RAVHLGDK ATKABIQREVVRM RAVRALGDN ASKEQIREKIAEL RAVRLGDN ASKAWVRQVAKL RAVHTLGSN ASKAWVRQVAKL RAVHTLGVQVAKL ATKDRVRQKIQEL VELYKANPE DAIKDFKQRVQRK VEL, RRAGPF ADPAAVRRAVEQS SQLTREHNEIS VSQSLMMENPRDL GLL, AAATHT DTPQTVCAAITORAI VRLRSATRRE ASASAVRAAVEQU VAL EQASAD LEAPALLARTRQR	LVLEKELLIDPEYPTALG LVLEKELLVDPEYPTALT EDLNFNBYISPEFFSHEVQ VAAEEELLIDPRFFIALQ MQQETELLVSPELFVALK LTIEKELLVDPEFFALG EBLEEELLVDPAFPTSLK EBSEDELLVDPAFPTSLK EBSEDELLVDPAFFTSL EBKEEELLVDPAFFTSLR EBREDELLVDPAFFTSLR EDREEELLVDPAFFTSLR EDREEELLVDPAFFTSLR EDREEFLVEPAFFTSLS EDREEEFLVEPAFFTLK LATEKELLIDPOFFTALM LATEKELLIDPOFFTALM LATEKELLUDPAFFTAV MAQEKELLVDPAFFTAV LVDEKELLUDPAFFTAV	FKHPSE. LSETEF. VKEGYGEGAFVVIRGTK HHVPD. LAGVDIRLKCGAHNELTRYRYDT HRLPD. LAGVDIRLKCGAHNELTRYRYDT LEIHKNGRMNGFLVWLNLHTIAGBEIDILEHEYCWL QRPPQ. ITWVEIQFKRGHAQNELTQFRYDV DTYPE. ITWVEIQFKRGHAQNELTQFRYDV SLSEE. IBHVBILPKNKNNELSSYRYGA QQDR. ICHVEILPKNKKANNELSSYRYAA SQLQDR. ICHVEILPKTKKATNELSSYRYAA SQLPN. IKHVEILPKTKKATNELSSYRYAA SQUPR. IKHVEILPKTKKATNELSSYRYAA SQUPR. IKHVEILPKTKKATNELSSYRYAA SQUPR. IKHVEILPKTKKATNELSYRYAA SQUPR. IKHVEILPKTKKATNELSYRYAA SQUPR. IKHVEILPKTMKATNELSYRYAA SQUPR. IKHVEILPKTMKATNELSYRYAA SQUPR. IKHVEILPKTMKATNELSYRYAA SQUPGK. IKHVEILPKTMATNELSYRYAA SQUPGK. IKHVEILPKRMATNELSYRYAA SQUPGK. IKHVEINT SQUPGK. IKHVEINT SQUPGK. IKHVEINT SQUPGK. IKHVEINT SQUPGK. IKHVEINT SQUPGK. IKHVEINT SQUPGK. IKHVEINT <td< td=""></td<>

Multiple sequence alignments of N-MT domains from experimentally characterized NRPS/PKS clusters with the structural template IVLM.

				Motif I		Motif	I-Post
1VLM_A	αι α2 00000 000000000	(13 000000000000000000000000000000000000		β1 4.0	04 0000 -	β2	- 22
IVLM A blecm01_CM 001 compa01_CM 001 epoth01_CM 001 fumon1_CM 001 famai01_CM 001 leina01_CM 001 lovas01_CM 001 lovas01_CM 001 micro02_CM 001 micro03_CM 001 modul01_CM 001 peder01_CM 001 yersi01_CM 001 yersi01_CM 002		HE PAYLS ELQAVKCLL TGRLTRLAGSLLDRLA TINKLAFGSAIHVVQDL TINKLAFGSAIHVVQDL VINIKWSRSNAQASEL GIVRGVVBSARVVA EGIGLSTANWWLAHM V YVESRTDSIDF NTILSFGPALHYAREL VDALKWSRSNAQASEL VDALKWSRSNAQASEL VQLVATTISRAIENLPS NLGISCTVELLSSYP MLGISCTVELLSSYP	ADRE PGRPLS VSQIAHRYQSII IRLCAHRYPSII IRLCAHRYPAMP PS		PÀV PLEIKI LTQALVTRA ATKRVLASP CTRLIVNAL TTAAVLPVL TGAQVILBGL TTNYILPYL SARTVLAAL SARTVLAAL SARTVLAAL TGLVVDSL ATSGLIHHL TTGQVLKAC TTGLIHL TTSTVLPQL LTASTVLPL LTAALLPEL TTAPULKAT	PGR LID WHA QLG PNS UTC QLG PNS UTC QLG PNS UTC GNTKP IDF WDP L PDRTE WHP GTS PSS UTT TNGKERLPST WAY N PNQTH WIP PAAA AAA DAGAH O'CY QLG PNS WITY GPNPP VGF 'DF P KBHIF WP PN LDHVT USP N.SRQIT WTP A.SHQIS WTP NSHIF WVP NLW.RAFIAF WAY RHHP VFHP RNTROSCHE	TDISAD TDVSAG TDVSAG TDVSAG TDISAG TDISAL TDISAG TDISAG TDISAS SEVSPF TDISSI TDISSI TDISSI TDISSI TDISSI
			Motif II	6	M	otif III	
IVIM_A	α5 <u>222222</u>	β3 TT T	β4 90 10	200 20.00	10	β5 120 +11	222
IVIM_A bleem01_CM 001 compa01_CM 001 epoth01_CM 001 equis01_CM 001 fumen01_CM 001 leina01_CM 001 lovas01_CM 001 lovas01_CM 001 micro03_CM 001 micro03_CM 001 modul02_CM 001 peder01_CM 001 tubu01_CM 001 yersi01_CM 002	FVTALGRBAARRGLDPVRAN VIGKAR, BQLSEPDGLMQFI FPBSAR, BQFADWQDVMTFI FLARAE, QR PRDYP, PLKYY FPBAAC, DK FKTYADRMVFI FPVAAQ, BRFKAYKG, LDPI F75KAQ, DK FRDYR, FLGY FLRHGE, REFGPTYPQLAFI FPBAAR, KR FAGWQNVMDFI FPBAAR, KR FAGWQNVMDFI FLRHK, ENFRNYP, FIK LLHQAR, QKFQKKY, NLNFI	EALDINKSPAEG9FK KKLDINKSPAEG9FK KVLDIT DPSEG9FK KVLDIT DPSEG9FK KVLDIT DPSEG9FK FLDIS DPK GFK FLDIS DPK GFK FLDIS DPK GFK KLDIS DPK GFK KVLDIK DPK GFC KVLDIK DPK GFC KVLDIK DFK FLDIK FFT FL FLSF KVLDIK DFK FL FLSF KLLIK FFT FL FL FL FL FL FL FL FL FL FL	ESPDFALM VTTT ERPDVVCGLDVV HSYDLIIASVI ORPDVIVACQVI ORPDVIVACQVI SYDMILASVI SYDVILASVI HQYDVIIANVI SYDVILASVI SYDVLIASVI SYDVILAQVI SYDLIIASVI SYDLIIANVI SYDLIIANVI SYDLIIANVI SYDLIIANVI SYDLIIANVI SYDLIIANVI SYDLIIANVI SYDLIIANVI	TEVDDPE.AA NATPDLR.TT HASSNPE.FK HATRCMK.RT HATRCMK.RT HATPTLN.ET HATPTLN.ET HATPDLE.TT HATPDLE.TT HATRDLE.TT HATANLQ.TT HSTRDLQGT HSTRDLQGT HATARID.TT HATRNMR.NT HATRNMR.NT HATRNMR.Q	LIEAYRILE LGRUPSLMA LAHIRSLMA LAHIRSLK LSNVRRLLS LANVRRLLS LANVRRLLS LANVRRLLS LANVRRLLA LSNVRSLLA LSNVRSLLA LTNVRRLLA LTNVRSLLA LTHIRBLLR LPHIRGLLR LQHVHSLIA LRNAKAALR LAHLESLV	KGBYLTUGIYÜR. PDGTLAIIETTAI PGGHLVIPOVTHRI. PGGHLVIPOVTHRI. PGGHLVIPOVTHRI. PGGHLVIPOTALIS PGGHLVIPOTALIS PGGHLVIPOTALIS PGGHLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLVIPOTALIS PGGKLLILS PGGKLLILS PGGKLVIPOTALIS PGGKLLILS PGGKLVIPOTALIS PGGKVIPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS PGGKVPOTALIS </td <td>SPARLA DOLDLP IPIWFD DMLRMG ARMVN. STRWVD FSEPLT SHTRLG DBLDLP ARRWVD NNSWIN DSSWID DSRWID DSRWID ARRWVD</td>	SPARLA DOLDLP IPIWFD DMLRMG ARMVN. STRWVD FSEPLT SHTRLG DBLDLP ARRWVD NNSWIN DSSWID DSRWID DSRWID ARRWVD
IVLM_A	α7 22222 ΤΤ	00000000		η2	<u>\$7</u>	β8	
lovas01 CM 001 lovas02 CM 001	LTFOLTTGWWMYEDAQCRL FIFQLFADWWAGVDDGRCT FTFOLTEGWWLCSQDLREN LTFOLTEGWWLCSQDLREN LTFOLTEGWWRFQDQQLRSI LIFOVTQGWWRFSDYHLRP LIFOVAQGWWRFSDANLRP LTFOLTEGWWLCTNDPHRN VLFOLIDGWSLAEDEHWRI MIWOLAEGWWSYEDELRTR: FVFOPLVLPLQDLDARS	3. PLLDAAGWEALLA 3. ASGSYDQWEHTLK . PSLTDLWNTMLD 0. HPLLPARTWCDVLR . PHLDLASWNSLLK . PYLDPSQWDTVLK 0. YPLLSRQQWKKVLS PHSPLLAFVQWRQSAA . PFVSPDRWDAILK . PSLSPTMWRSMLH 1. YPLISTKDWQNLLI 0. TPLLRGENWCSVLK 4. HPIITASTWESMLL 0. HPIITASTWESMLL 1. HPIITASTWESLL . PLLEVGDWEHALR 1. ELFLTAQWQQCR	TSGPNGVELEVF RVGPADAVSLPO RCGPGGIETSTE ETGPSGVDSAIY ETGPTQVVTLPE AAGLETVRTGGI RVGPSGVDSRTT TTGPNGVEVEAF ENGPEDINIIE SCGPATANSPSF KSGPSQVEPVSF ESGPAQAEPVSF 2EGPD.KVSFPA DVGPAEVEVLPA	PDGPQDAALL DREDDLIP.S DCEDDEPYMI DGSPAGILGQ VYDS.IHAVA VEGMAKALSE PGVPADELC DRDANLFPTS DCDSHEP. SQVKTRNLLN QQNIFINNGQ DQKIGTALPQ DQKIGTALPQ QAEIGSPLFQ QV.AHDLGQ AVEQRSRWD	LARQTPRPA VPS	AAA. ENTTPDTVAESEVI IAGAA RVALLK. TTTLE TVEYLDAPLASSG EQLP E. QAV.	TVKDSY

Multiple sequence alignments of C-MT and domains from experimentally characterized NRPS/PKS clusters with the structural template IVLM.

NRPS domains and other catalytic domains. Figure 10 shows the number of proteins containing N-MT, O-MT and C-MT domains for each of the four categories. As can be seen from Figure 10a, 359 PKS proteins have C-MT

domains, 14 PKS proteins have O-MT and none of them have N-MT domains. This result is consistent with the fact that polyketides have no sites for N-methylation and Omethylation in known PKS biosynthetic pathways is cata-

	Motif I
1VLM_A	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
barba01_OM_001 eryth01_OM_001 jamai01_OM_001	WHIPERFVNEYER. WFLVHRPAYLSELQAVKCLLPEGRGVEIGVCTGRPAV FDSLIFNTSTRDYYGEKBFFNVGYWHSDTQNQHEACFNLMEKLLEFTPRKQG.NILDVGCGLGATTS DYARPAPSEVVPDFSWLEVILN.PDQHP.EQTSLSLQAQAEMKAVLPRGTDPSSIKKVMDIGCGYSHDLI RARLAYALFADDHEATTEGAYINLGYWKPGCAGLEEANQELANQLAEAAGISEGDEVLDVGFGLGAQDF KSAKTYLRMTPLPEKIPGLSWLEMMLK.PEKQQ.ENIBLILKYQEEMKSVLPRGTDPSSIEKVMDIGCGYSEDLI GMSETPEHKDVLLDGQRGLRSVLPRGTDPSSIEKVMDIGCGYSEDLL GMSETPEHKDVLLDGQRGLRSVLPRGVDWSSVRKVLDPGCGYASDLL GFSWLMSVYGLSEREE.HTEVLLNGQRGLRSVLPRGVDWSSVRKVLDFGCGYASDLL RFATLREAVPGFSWISLYKAEPENMAGFQKYYDLAVQANREMSQTIFRGVDWSSVRKVLDFGCGYASDLI SSMTLVEGIYKNNLVSD.YPQVLGDVLVAFMTTRGHEEP.VRILEIGAGTGGTA HYDRFFYEQRGVERLIREETDFKNLGYWDDTTLD.LNAAAERLLAKLGEVABLSPGCVLDVGFGFABCDI BSSNLDVGUYKNNLVSD.YPQVLGDVLVAFMTTRGHEEP.VRILEIGAGTGGTA MSRSHLEBIAELYDSAEGHVGNLIFDG.QVHWGYWDETTPD.QNAAEKLQDMLLEMIPEKTG.RILDVACCMGASTR MSRSHLEBIAELYDSAEGHVGNLIFDG.QVHWGYWDETTTT.QHEASERLQDALLOFT IRKGRKFCDLGCGFGSTS HINQHYDHTFFS.EGLTSLLVDGSDYNNGYWDETTTT.QHEASERLQDALLDFI PEKSG.RILDAACCMGASTR TAIADVEKVATLYDSAEGQVGPILFGG.HMHWGYWDETTTT.QHEASERLQDALLDFI PEKSG.RILDAACCMGASTR TAIADVEKVATLYDSAEGQVGPILFGG.HMHWGYWDETTTT.QHEASERLQDALLDFI PEKSG.RILDAACCMGASTR TAIADVEKVATLYDSAEGQVGPILFGC.HINGVGCFFGESE.RMRLLLHSQRALRVVLLDSVDFSRAKSVWDFGCGFASDI LVAYGLDFSEC.RFAXSVWDFCGGFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGQKFIDMGCFFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGQKFIDMGCGFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGQKFIDMGCGFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGVKSVWDFGCGFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGVKSVWDFGCGFGESAL LVAYGLDFSEC.NFANAAERLAQTMIAKAPIKAGVLJDFGCGHGSDI SECONSSNDDEHFLTWGVFQEIVFGFSWIRTVFRSBRFE.GREFLAVAQRELRRVLFRAVDLSAIKNVMDFGCGHGSDI NWWNDDSSVSLLSTS.INPCRIAT
	$\beta 2 \alpha 5 \beta 3$ Motif II $\beta 4 \eta 1 \alpha 6$
1VLM_A	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
barba01 0M 001 eryth01 0M 001 melit01 0M 001 myxot01 0M 001 myxot02 0M 001 nodul01 0M 001 onnam01 0M 001 onnam03 0M 001 onnam03 0M 001 onnam05 0M 001 peder01 0M 001 stigm02 0M 001	PLKIKIGVEPSERMAEIARKRGVFVLKGTAENLPLKDESPEFALMVTTICFVDDPBRALKEAY HLLNYYSPADVVGINISRKQIERSIVNAPGCKFICMDAVQMEFDEDFPDNIICVBAPYFNTBKFLKBAM DLATNHVHLQLDGVNISPEQVKAGEQKIQGLGYSDRIVLYNRDSAKQPLPITVDLIFSCQVIHHIKRKBDVPLNIS FWLETRKPARIVGVDITPSHVRIASERAEENVQDRLQFKEGSATDLPFGAETPPKVTSLESALHYEPTDFFKGAF HLAEEHPHLQLDGYNISPEQVNFGKERIESLGYAQRINIYNRDSAKQPFLDEYDLGYSCQVIHHMKNQBNVLLNIS SLARRHPHLKLHGYTISAEQAAVDARRVRERGFEDRIRVFARDSAKDAFPDEYDLGYSCQVIHHMKNQBNVLLNIS SLARRHPHLKLHGYTISAEQAAVDARRVRERGFEDRIRVFARDSAKDAFPDRYDVLGYSCQVIHHMKNQBNVLLNIS SLARRHPHLKLHGYTISAEQAAVDARRVRERGFEDRIRVFARDSAKDAFPDRYDLGYSCQVIHHMKNQBNVLLNIS SLARRHPHLKLHGYTISAEQAAVDARRVRERGFEDRIRVFARDSAKDAFPDRYDLGYSCQVIHHMKNQBNVLLNIS SLARRHPHLELGCNISPDQIEVGRQRIRALGLDGRVLHYQDSSRDQFPSTYDLVIAYOVHHIRAKSDLFANIS SLGKKHPHLTLHGYTISAEQAAIDEERVRARGLQDRIRVFARDSAKDAFPDRYDLIVGQCVCAAF LWMRENNLGAIGGVTHSACGCRIRGLGLDERITLHYQDSSRDPFPSTYDLVLGPQVIHHIRNKADLFANIS LWMRENNLGAIGGVTHSQVKJAQERVRAGLEGRINLQVGSSRDPFPSTYDLVLGPQVIHHIRNKADLFANIS RLESYPPENVWAINISAKQIETTKOVKGCHAIVMNAVDMTFBDNFPDTVLSIBAAMHFETRRKFLEBSF RLAESYPENVWAINISECTESTRENAKGCHQVMSAVEMTFDNFPDTVLSIBAAMHFETRRKFLEBSF RLAELYSPENVWAINISECTESTRENAKGCHQVMSAVEMTFDNFPDTVLSIBAAMHFETRRKFPDSL
	Motif III
1VLM_A	
1VLM_A anaba01_OM_001 barba01_OM_001 jamai01_OM_001 melit02_OM_001 myxot01_OM_001 myxot01_OM_001 nodu101_OM_001 onnam01_OM_001 onnam03_OM_001 onnam04_OM_001 peder01_OM_001 stigm01_OM_001 stigm02_OM_001	RILKKGGYLIVGIVD

Figure 7

Multiple sequence alignments of O-MT domains from experimentally characterized NRPS/PKS clusters with the structural template IVLM.

lyzed by stand alone O-MTs. This indicates that our profiles are able to correctly classify the three different classes of secondary metabolite methyltransferases. Our nr database search also identified 59 NRPS proteins containing N-MT domains, 1 containing O-MT domains and 44 containing C-MT domains (Figure 10b). Since most nonri-

			-	-		10		
	CCCCCCCTT	10	20	CCCEEEEEE		40	50	60
2fr0A0				LDGTWLVAK				
211040	the second s					I J	11 I	VRELVVI
Ouerv	and the second second second second	the state of the s		LTGGCLLLG		A CONTRACTOR OF		VTTVCC
Query		430	440	450	DOD	460		0
								0
		70					00	
	CCCCUUU	70	80	90 ECCCCCCCC	recer			110
2fr0A0				SLLAVDEAE				
LIIONO	I	I	I		T FEUL	22.02.2.1		
Ouery	RPPGPER	RELVE	PATRLAV	DLWPLRDAS	HRGRAAG	AAGVETAC		THLARAT
durel	480	490		500	510	520		530
	112221	16576	25		22225	1000		
		120	130	140	1.1.2	50	160	170
			and the second se	CCCCCCCHH				and the second se
2fr0A0				PFERVRNAA				
ZIIOAO	VORLL			TTERVENAA	A CONTRACTOR OF CONTRACTOR		1	11 1
Query	GALEERHI			GDD-LAHPE			PWIACT	LDVEPGI
duct i	540		550		570		0	590
	С-НИНИИ	180	and the second se			210	220	EECCCC
2fr0A0 Query	V-AELARI DAERLADI	HHHHHHH HLAAVVS	GAARETTVT	EEECCEEEE ALRADGVYG ACRGRRRFT	EEEEECC RRWVRAA PCPVRQP	CCCCCCCC APATDDEW LPAAPERF	CCCCEEF	
	V-AELARI	HHHHHHH HLAAVVS	GAARETTVT	EEECCEEEE ALRADGVYG	EEEEECC RRWVRAA	APATDDEW	CCCCEEF	
Query	V-AELARI DAERLADI 600	HHHHHH HLAAVVS LIVRELC	CCCCCCCEE SGGAGEDQL GAARETTVT 610	EEECCEEEE ALRADGVYG ACRGRRRFT	EEEEECC RRWVRAA PCPVRQP	CCCCCCCC APATDDEW LPAAPERF	CCCCEEF	
Query	V-AELARI DAERLADI 600 gnment wit	HHHHHHH HLAAVVS LIVRELC th 2hg4	GAARETTVT 610 4FO:	EEECCEEEE ALRADGVYG ACRGRRRFT 620	EEEEECC RRWVRAA PCPVRQP 630	CCCCCCCC APATDDEW LPAAPERF	CCCCEEF	
Query	V-AELARI DAERLADI 600 gnment wit	HHHHHHH HLAAVVS LIVRELC th 2hg4 790	GAARETTVT 610 4FO: 800	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810	EEEEECC RRWVRAA PCPVRQP 630	CCCCCCCC APATDDEW LPAAPERF 64	CCCCCEEF WKPTGTVI 2A 10 830	VTGGTG
Query	V-AELARI II DAERLADI 600 gnment wit CHHHHHH	HHHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC	GAARETTVT 610 4FO: 800 CCEEEEECCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH	EEEEECC RRWVRAA PCPVRQP 630 &	CCCCCCCC APATDDEW LPAAPERF 64 320 CCCCCCCE	RECCCEEE	NTGGTGG 84 CCCHHHH
Query	V-AELARI II DAERLADI 600 gnment wit CHHHHHH	HHHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC	GAARETTVT 610 4FO: 800 CCEEEEECCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810	EEEEECC RRWVRAA PCPVRQP 630 &	CCCCCCCC APATDDEW LPAAPERF 64 320 CCCCCCCE	RECCCEEE	NTGGTGG 84 CCCHHHH
Query >>> Ali; 2hg4F0	V-AELARI II DAERLADI 600 gnment wit CHHHHHH	HHHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC	GAARETTVT 610 4FO: 800 CCEEEEECCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH	EEEEECC RRWVRAA PCPVRQP 630 &	CCCCCCCC APATDDEW LPAAPERF 64 520 CCCCCCCE PADAGAC	830 EECCCCCC VVGTLRR	84 CCCHHHH DRGGLAD
Query	V-AELARI II DAERLADI 600 gnment wit CHHHHHH	HHHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC	GAARETTVT 610 4FO: 800 CCEEEEECCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH	EEEEECC RRWVRAA PCPVRQP 630 &	CCCCCCCC APATDDEW LPAAPERF 64 520 CCCCCCCE PADAGAC	830 EECCCCC VVGTLRR	84 CCCHHH DRGGLAI PD-EAAS
Query >>> Ali; 2hg4F0	V-AELARI II DAERLADI 600 gnment wit CHHHHHH	HHHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC	GAARETTVT 610 4FO: 800 CCEEEEECCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH	EEEEECC RRWVRAA PCPVRQP 630 &	CCCCCCCC APATDDEW LPAAPERF 64 520 CCCCCCCE PADAGAC	830 EECCCCCC VVGTLRR	84 CCCHHHH DRGGLAD PD-EAAS
Query >>> Ali; 2hg4F0	V-AELARI II DAERLADI 600 gnment wit CHHHHHH RFQDATRO	HHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC QLAEAGI	SCCCCCCEE SGGAGEDQL GAARETTVT 610 4FO: 800 CCEEEECCC FDAFVEVSP	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH HPVLTVGII	EEEEEECC RRWVRAA PCPVRQP 630 8 HHHHHHH ATLDSAI	CCCCCCCC APATDDEW LPAAPERF 64 320 CCCCCCCE PADAGAC	830 EECCCCC VVGTLRR	84 CCCHHHH DRGGLAD PD-EAAS
Query >>> Ali; 2hg4F0	V-AELARI II DAERLADI 600 gnment wit CHHHHHH RFQDATRO	HHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC QLAEAGI	SCCCCCCEE SGGAGEDQL GAARETTVT 610 4FO: 800 CCEEEEECCC FDAFVEVSP 860	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH HPVLTVGIH	EEEEEECC RRWVRAA PCPVRQP 630 8 HHHHHHH ATLDSAI	CCCCCCCC APATDDEW LPAAPERF 64 320 CCCCCCCE PADAGAC PPY 880	830 EECCCCC VVGTLRR	84 CCCHHHH DRGGLAD PD-EAAS
Query >>> Ali; 2hg4FO Query	V-AELARI II DAERLADI 600 gnment wit CHHHHHH RFQDATRO	HHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC QLAEAGI 850 HHCCCCC	CCCCCCEE SGGAGEDQL GAARETTVT 610 4FO: 800 CCEEEECCC FDAFVEVSP 860 CCCHHHHCC	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH HPVLTVGII	EEEEEECC RRWVRAA 11 PCPVRQP 630 8 HHHHHHH ATLDSAI	CCCCCCCC APATDDEW 11 LPAAPERF 64 520 CCCCCCCE PADAGAC PPY 880 EECCCCC	RCCCCEEF RFTGTVI PA HO 830 EECCCCC VVGTLRR IVRCLAA 10	84 CCCHHHH DRGGLAD PD-EAAS
Query >>> Ali; 2hg4F0	V-AELARI II DAERLADI 600 gnment wit CHHHHHH RFQDATRO 	HHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC QLAEAGI 850 HHCCCCC YAQQVEV	SECCECCEE SGGAGEDQL GAARETTVT 610 4FO: 800 CCEEEECCCC FDAFVEVSP 860 CCCCHHHHCC VDWSPAFAD	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH HPVLTVGII HPVLTVGII 37 CCCCCCCC ARPVELI	EEEEECC RRWVRAA 11 PCPVRQP 630 8 HHHHHHH ATLDSAI CCCCCCCE VYPFORO	CCCCCCCC APATDDEW 11 LPAAPERF 64 520 CCCCCCCE PADAGAC PPY 880 EECCCCC RYWLPI-	RCCCCEEF RFTGTVI PA HO 830 EECCCCC VVGTLRR IVRCLAA 10	84 CCCHHHH DRGGLAD PD-EAAS
Query >>> Ali; 2hg4FO Query	V-AELARI II DAERLADI 600 gnment win CHHHHHHHH RFQDATRO 	HHHHHH HLAAVVS LIVRELC th 2hg4 790 HHHCCCC QLAEAGI 850 HHCCCCC YAQQVEV	CCCCCCEE SGGAGEDQL GAARETTVT 610 4FO: 800 CCEEEECCC FDAFVEVSP 860 CCCCHHHHCC VDWSPAFAD	EEECCEEEE ALRADGVYG ACRGRRRFT 620 810 CCCCCHHHH HPVLTVGII	EEEEEECC RRWVRAA 11 PCPVRQP 630 8 HHHHHHH ATLDSAI ATLDSAI	CCCCCCCC APATDDEW LPAAPERF 64 320 CCCCCCCE PADAGAC PPY 880 EECCCCC RYWLPI 	RCCCCEEF RFTGTVI PA RO 830 EECCCCC VVGTLRR I I IVRCLAA 10	84 CCCHHHH DRGGLAD I PD-EAAS

Threading alignment C-MT containing sequence (ORF blmVIII) stretch from bleomycin gene cluster. (a) Alignment of 60 amino acid N-terminal stretch with structure of KS-AT di-domain (2HG4) from erythromycin PKS (b) Alignment of 200 amino acid C-terminal region with structure (2FR0) of KR domain from erythromycin PKS.

------ 0.1

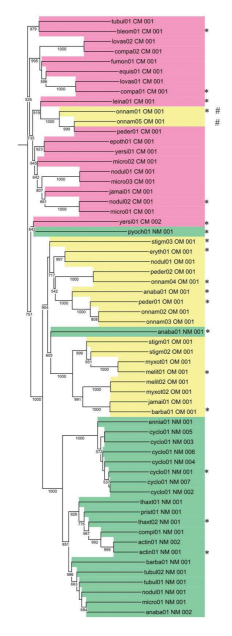


Figure 9

Dendrogram of 60 MT domains from experimentally characterized NRPS, PKS and hybrid NRPS/PKS biosynthetic clusters. The C-MT, O-MT and N-MT are colored pink, yellow and green respectively. The 18 representative MT sequences used as templates for detecting MT domains in a query are marked by "*". Two MT domains from onnamide-A which are annotated as O-MTs and cluster with C-MTs are marked with hash (#) symbol.

bosomal peptide products are N-methylated, the presence of such a large number of C-MT domains as C-A-CMT-T modules was surprising. Similarly, the hybrid NRPS/PKS set also had 121 proteins containing C-MT domains as compared to only 4 proteins containing N-MT domains (Figure 10c). In fact 17 of these C-MT domains in hybrid NRPS/PKS proteins were present next to condensation domains of NRPS as C-CMT-PCP modules, while one would a priori expect N-MT domains in such modules instead of C-MT domains. In view of this finding of anomalously large number of NRPS modules containing C-MT domains in NRPS and hybrid NRPS/PKS proteins, we decided to analyze these proteins by pairwise alignment with 18 representative MT templates from known NRPS/ PKS biosynthetic pathways. Interestingly the N-MT domain of pyochelin synthase was found to be closest match for 23 out of the 44 C-MT containing NRPS proteins identified by profile search. Most of these 23 NRPS proteins showed very high percentage identity with pyochelin synthase N-MT ranging from 27% to 69%. Thus it is very much likely that, these 23 NRPS proteins indeed contain pyochelin type N-MT domains which are different from other N-MT domains. They were annotated by profile approach as C-MTs, because N-MT domain of pyochelin synthase shows homology to C-MTs and has comparable scores with C-MT as well as N-MT profiles (Table 3). Similarly, a C-MT domain from versiniabactin synthase was found to be the closest homolog of C-MTs found in NRPS modules of hybrid NRPS/PKS proteins and the sequence similarity was also very high. This MT domain in yersiniabactin synthase catalyzes C-methylation (Figure 2), but is present as C-CMT-PCP module similar to the domain organization found in C-MT containing hybrid NRPS/PKS proteins found by our profile search. This suggests that our profile search has genuinely identified yersiniabactin type novel C-MT domains embedded in NRPS modules. In case of 20 other NRPS proteins which showed matches with C-MT profiles, the C-MT domains from versiniabactin, leinamycin and nodularin were found to be closest match. Table 4 shows the domain organization predicted for each of them by HMM profiles, and the percentage identity and similarity with closest matching MT domain in our 18 representative templates set. Even though the closest homolog approach also detects C-MT domains in these proteins in agreement with results from HMM approach, in view of the relatively low sequence identity with C-MTs from known NRPS/PKS biosynthetic pathways, it is not clear if these domains are likely to be genuine C-MT domains as in versiniabactin synthase or a different class of N-MT domains which lack homology to known N-MT domains in our data set. Thus our analysis of nr database has revealed presence of many C-MT domains in NRPS modules as in versiniabactin. It has also identified several pyochelin type N-MT domains which often show higher homology to C-MT profiles. None of these proteins are currently experimentally characterized. Experimental characterization of some of these MT domains would help in understanding how MT

	NMT		OMT		СМТ	
	Score	E-value	Score	E-value	Score	E-value
actin01_NM_001	390.1	1.10E-117	-	-	-	-
anaba01_NM_001	252.8	2.40E-076	-	-	-	-
anaba01_OM_001	-	-	375.5	2.80E-113	-	-
barba01_OM_001	-	-	385.5	2.80E-116	-	-
bleom01_CM_001	-	-	-54.3	4.70E-007	385.5	2.80E-116
compa01_CM_001	-	-	-	-	332.7	2.10E-100
cyclo01_NM_001	449.7	1.30E-135	-	-	-	-
eryth01_OM_001	-	-	369.6	1.60E-111	-	-
leina01_CM_001	-	-	-5.7	2.20E-010	395.5	2.60E-119
melit01_OM_001	-	-	178.7	4.80E-054	-	-
nodul02_CM_001	-	-			270.3	1.30E-081
onnam01_OM_001	-	-	289.7	I.80E-087	280. I	I.50E-084
onnam04_OM_001	-	-	378.1	4.60E-114	-	-
peder01_OM_001	-	-	234.7	6.90E-071	-	-
pyoch01_NM_001	187.4	1.10E-056	-		17.1	3.30E-013
stigm03_OM_001	-	-	210.1	1.70E-063	-	-
thaxt02_NM_001	318.9	3.10E-096	-	-	-	-
yersi01_CM_002	-	-	-	-	321.4	5.20E-097

Table 3: Scores and E-values for the alignment of 18 representative MT domains with the HMM profiles of N-MT, O-MT and C-MT.

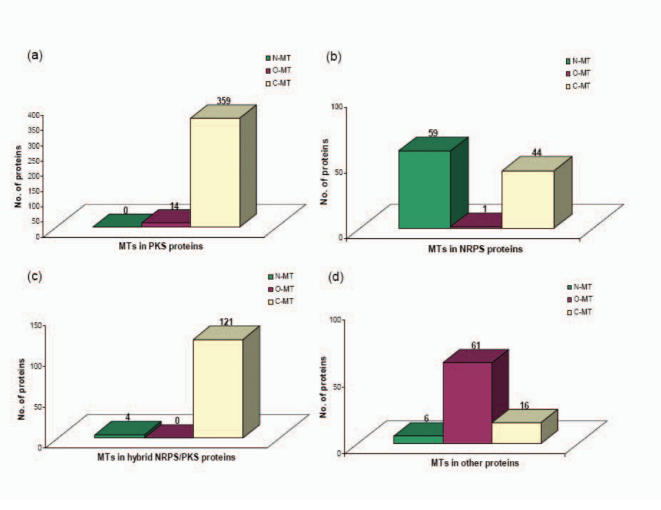
A '-'sign indicates that alignments resulted in scores with E-value higher than 1.0E-6. Several C-MTs align with O-MT profiles, while only N-MT of pyochelin synthase shows alignment with C-MT profile as well.

domains in NRPS/PKS family have evolved to acquire specificities for different substrates. However, a close examination of the chemical structures of pyochelin [37] and yersiniabactin [38] provides a rational for presence of N- MT domains with homology to C-MT sequences. As can be seen from Figure 2, in both these biosynthetic clusters MT domains transfer methyl groups to a five membered rings having very similar chemical structures. They only

Table 4: List of protein sequences from nr database which contain C-M	IT domains adjacent to the core NRPS domains
---	--

Gi. no.	Domains	ldentity (%)	Template	Organism name
108809363	ACP-C-A- CMT -ACP-C	25	yersi01_CM_002	Yersinia pestis Antiqua
108809365	C-A- CMT	33	nodul02_CM_001	Yersinia pestis Antiqua
108813376	C-A- CMT	33	nodul02_CM_001	Yersinia pestis Nepal516
116215693	CMT-ACP-C-A-ACP-TE	47	yersi01_CM_002	Vibrio cholerae RC385
4827 509	C-A- CMT -ACP	33	nodul02_CM_001	Clavibacter michiganensis subsp.michiganensis
153947007	C-A- CMT	33	nodul02_CM_001	Yersinia pseudotuberculosis IP 31758
153954130	ACP-C-A- CMT -ACP-C-ACP	29	nodul02_CM_001	Clostridium kluyveri DSM 555
16121089	C-A- CMT	33	nodul02_CM_001	Yersinia pestis CO92
16121091	ACP-C-A- CMT -ACP-C	25	yersi01_CM_002	Yersinia pestis CO92
17546530	C-A- CMT -ACP-TE	33	leina01_CM_001	Ralstonia solanacearum GMI1000
21225943	C-A- CMT -ACP	33	nodul02_CM_001	Streptomyces coelicolor A3(2)
22127285	ACP-C-A- CMT -ACP-C	25	yersi01_CM_002	Yersinia pestis KIM
26248281	ACP-C- CMT -ACP-TE	100	yersi01_CM_002	Escherichia coli CFT073
28869792	ACP-C-A- CMT -ACP-C-ACP	27	leina01_CM_001	Pseudomonas syringae pv. tomato str. DC3000
41409840	C-A- CMT	34	yersi01_CM_002	M. avium subsp. paratuberculosis K-10
45443170	С-А -СМТ	33	nodul02_CM_001	Yersinia pestis biovar Microtus str. 91001
51597596	C-A- CMT	33	nodul02_CM_001	Yersinia pseudotuberculosis IP 32953
89102588	ACP-C-A- CMT -ACP-C	25	yersi01_CM_002	Yersinia pestis biovar Orientalis str. IP275
89895567	C-A- CMT	37	yersi01_CM_002	Desulfitobacterium hafniense Y51
90424526	C-A- CMT -ACP-TE	38	yersi01_CM_002	Rhodopseudomonas palustris BisB18

List of protein sequences from nr database which contain C-MT domains adjacent to the core NRPS domains as identified by our MT HMM profile search. Gene identifier number, domain organization and organism name is listed in columns 1, 2 and 5. Columns 4 and 3 list the closest match (as identified by pair BLAST) with the representative MT domains from experimentally characterized NRPS/PKS clusters and the corresponding percentage identities.



Histograms showing the number of proteins in nr database having N-MT, O-MT and C-MT domains as identified by our HMM profile search. (a) PKS proteins, (b) NRPS proteins, (c) hybrid NRPS/PKS proteins and (d) proteins other than NRPS/PKS proteins.

differ by the site of methylation. In view of the similarity in the structure of the acceptor substrate, pyochelin type N-MT domains show homology to C-MT proteins. It may be interesting to note that, in a recent study involving MTs from type II PKS pathways, Hertweck and coworkers [24] have observed that, these type II MTs cluster according to the site of methylation not necessarily as C-MTs, O-MTs and N-MTs. Based on this observation, they have proposed that for polyketide alkylation, regioselectivity is more dominant than the type of nucleophile (C-, O- or N-) that is being alkylated. Thus the C-MT domains found in NRPS modules by our profile search could indeed be due to such correlation between sequence of type I MTs and their regioselectivity. These observations have interesting implications of prediction of chemical structures of metabolites by genome mining.

The analysis of MT containing proteins in nr database also revealed the co-occurrence of NRPS/PKS type MT domains with several other catalytic domains apart from core PKS and NRPS domains (Figure 10d). However, in most cases these proteins had O-MT domains and there were relatively few C-MT and N-MT domains. Table 5 shows domain organization for representative proteins from each category and also lists the number of such proteins identified by our HMM profile search. As can be seen, the O-MT domains are present along with a variety of catalytic domains which do not belong to core NRPS or PKS family. Interestingly 19 out of the 61 examples correspond to co-occurrence with glycosyltransferase domains. Since glycosyltransferases are a major class of tailoring enzymes involved in secondary metabolite biosynthesis, many of these proteins might indeed be associated with novel PKS/

Gi no.	Domains	No. of proteins	Organism name
113477217	Glycos_transf_1- OMT -Glycos_transf_2	19	Trichodesmium erythraeum IMS101
110599935	TPR_I-TPR_2-TPR_I-TPR_2-TPR_2-TPR_I-TPR_2- OMT	18	Geobacter sp. FRC-32
154319269	KS-KS-AT -CMT -ER-DH-KR-Carn_acyltransf	7	Botryotinia fuckeliana B05.10
126178605	OMT -Dala_Dala_lig_C	4	Methanoculleus marisnigri JR I
110634799	Abhydrolase_I- OMT	3	Mesorhizobium sp. BNCI
31794901	Radical_SAM-OMT	3	Mycobacterium bovis AF2122/97
30681189	DUF248- OMT	2	Arabidopsis thaliana
62290714	HTH_3- OMT	2	Brucella abortus biovar str. 9–941
71013608	Amidohydro_3- OMT	2	Ustilago maydis 521
107099798	FA_hydroxylase- OMT	2	Pseudomonas aeruginosa PACS2
4779 35	Amino_oxidase- OMT	2	Pseudomonas aeruginosa PACS2
156064387	E1-E2_ATPase- OMT	2	Sclerotinia sclerotiorum 1980
157382467	A-ACP-C-A- NMT -ACP-ACP-C-ApbA-ApbA_C	I	Xylaria sp. BCC 1067
157752876	OMT-NoI1_Nop2_Fmu	I	Caenorhabditis briggsae
538 375	AstE_AspA- OMT	I	Ruminococcus obeum ATCC 29174
71030506	OMT-Pox_MCEL	I	Theileria parva strain Muguga
66825109	KS-KS-AT- CMT -DH-KR-Chal_sti_synt_N-Chal_sti_synt_C	I	Dictyostelium discoideum AX4
145608084	HET-OMT	I	Magnaporthe grisea 70-15
147858936	DUF642- OMT	I	Vitis vinifera
158520366	DUF1365- OMT	I	Desulfococcus oleovorans Hxd3
115402313	OMT-MIP	I	Aspergillus terreus NIH2624
110681402	C-A- NMT -ACP-C-A-ACP-TE-PEP-utilizers	I	Chondromyces crocatus
17546523	OMT-TE-ACPS	I	Ralstonia solanacearum GMI1000
51892502	Phosphodiest- OMT	I	Symbiobacterium thermophilum IAM 14863
116207616	MFS_I-KS-KS-AT- CMT -DH-ER-KR-DH	I	Chaetomium globosum CBS 148.51
41407518	C-A-Strep_SA_rep-ACP-C-A- NMT -ACP-C-A-Strep_SA_rep-ACP- C-A-Strep_SA_rep-ACP-C-A-Strep_SA_rep-ACP-TE	2	Mycobacterium avium subsp. paratuberculosis K-10
26541536	ACP-KR-DH-KS-KS-ACP-ACP-KS-KS-KR-DH-ACP- CMT -ACP-KS- KS-DH-KR-ACP-KS-KS-ACP-ACP-Beta_elim_lyase-Abhydrolase_I	I	Streptomyces atroolivaceus
108757966	A-ACP-C-A- NMT -ACP-C-A-ACP-C-A-ACP-C-A-ACP-C-A-ACP- KS-KS-AT-ACP-C-Bac_luciferase-C-A-ACP-C-A-ACP-C-A-ACP-C- A-ACP-C-A-ACP-TE	I	Myxococcus xanthus DK 1622

Table 5: List of representative protein sequences from nr database which contain MT domains in combination with other functional domains

List of representative protein sequences from nr database which contain C-MT, N-MT or O-MT domains in combination with functional domains other than core PKS or NRPS domains. Gene identifier number, domain organization and organism names are listed in columns 1, 2 and 4 respectively, while column 3 lists total number of proteins having similar domain organization.

NRPS biosynthetic pathways [39,40]. Apart from glycosyltransferases, the other functional domains which occur with O-MT are oxidases, hydroxylases, tetratricopeptide repeat (TPR), phosphodiasterases, DNA binding helixturn helix proteins etc. It would be necessary to further analyze each of these proteins in details to understand their biochemical function. In contrast to O-MTs, the N-MT and C-MT domains are present in multifunctional enzymes which contain other catalytic domains in addition to PKS and NRPS domains. Some of the interesting examples are NRPS/PKS proteins containing chalcone synthase or carnitine acyltransferase domains along with C-MT domains. Similarly, N-MT domains are present along with domains associated with streptococcal surface antigen, bacterial luciferase, phosphoenol pyruvate (PEP) utilizing enzyme and ketopantoate reductase (ApbA) etc. Thus these results give valuable clues about organisms which can potentially make secondary metabolites with a variety of structural modifications by diverse types of tailoring enzymes. Some of these proteins would be interesting targets for detailed experimental investigation.

Discussions

We have carried out a comprehensive analysis of the sequence and structural features of the MT domains present in multi functional proteins encoded by various experimentally characterized NRPS, PKS and hybrid NRPS/PKS gene clusters with known secondary metabolite products. Even though presence of methyltransferase domains in NRPS and PKS family of megasynthases have been inferred from methylation pattern of the chemical structure of the secondary metabolite product, earlier studies have not defined the correct domain boundaries. Threading analysis of the MT containing sequence stretches suggest 1VLM and 1VL5 as possible structural templates for NRPS/PKS MT domains. Based on the alignment with these structural templates, we identify the correct boundaries and predict that, the general length of MT

domains will be in the range of 200–220 residues. Our threading analysis also reveals interesting homology between AT-MT and AT-DH linkers. Similarly, large sequence stretches C-terminus to the C-MT domains of PKS proteins are found to have homology with structural half of KR domains. These results not only explain the large variation in the length of the MT containing sequence stretch, they can also provide valuable clues for design of domain swapping experiments.

The curated sequences of these MT domains were further analyzed. From the MT sequences of correct length, a representative set of 18 MT domains covering the entire range of sequence divergence were chosen. A novel protocol for identification of MT domains by pairwise alignment and their classification as N-MT, C-MT and O-MT was developed using these 18 MTs as multiple templates. This MT domain identification protocol has been implemented in the current version of the program NRPS-PKS. Using this approach C-MT domains were annotated in the PKS proteins of *Dictyostelium discoideum* in a recent work [41].

The MT sequences with correct domain boundaries were also used to build profile HMMs for O-MT, C-MT and N-MT domains. Using these HMM profiles searches were carried out in the nr database of NCBI for identifying various other proteins containing C-MT, O-MT and N-MT domains. It is interesting to note that, apart from core PKS and NRPS domains, these secondary metabolite biosynthetic MT domains are also associated with other important catalytic domains like glycosyltransferase, oxidases, hydroxylases, phosphodiesterases and reductases. These proteins could be interesting targets for experimental characterization. Our analysis also surprisingly revealed the presence of a large number C-MT domains in NRPS modules adjacent to condensation (C) and adenylation (A) domains. These predicted C-MT domains can be classified into two groups. One group of C-MT domains showed high homology to N-MT domain of pyochelin synthase, which is different from typical N-MT domains present in NRPS modules. Unlike typical N-MT domains of NRPS proteins, this shows homology to C-MT domains but catalyzes transfer of methyl group to nitrogen. This group could indeed be pyochelin type N-MT domains, but are mis-classified as C-MT due to their homology with C-MT proteins and under representation of pyochelin type N-MT in our training data set. The closest homolog of the other group of C-MT domains present in NRPS modules is the C-MT domain of yersiniabactin synthase, which is present adjacent to a condensation domain of NRPS, but is indeed a C-MT. However, in view of the low homology between the C-MT from yersiniabactin and this second group of predicted C-MTs, it is difficult to predict whether they are versiniabactin or pyochelin type MTs. A close examination of the chemical structures of the yersiniabactin and pyochelin provides an evolutionary basis for the presence of pyochelin type N-MT domains having homology with C-MT proteins. As can be seen from Figure 2, an identical five membered ring is the acceptor moiety for these two classes of MTs, while pyochelin N-MT methylates at the N position of the ring, the yersiniabactin C-MT methylates the adjacent C position. It may be interesting to note that such correlation between regioselectivity of the site of methylation and MT sequence have also been reported earlier by Hertweck and coworkers [24] for MTs in type II PKS biosynthetic pathways. Experimental characterization of proteins identified by our analysis would help in building more specific profiles for these novel MT domains.

Conclusion

We have carried out a comprehensive bioinformatics analysis of methyltransferase (MT) domains present in PKS/ NRPS clusters having known secondary metabolite products. Based on the site of methylation of these known secondary metabolites, the MT domains have been grouped as N-MT, C-MT and O-MT proteins and sequence/structural features have been analyzed in detail for each group. Based on the results of this analysis, we have developed a novel knowledge based computational approach for detecting MT domains present in PKS and NRPS megasynthases, delineating their correct boundaries and classifying them as N-MT, C-MT and O-MT using profile HMMs. Analysis of proteins in nr database of NCBI using these class specific profiles has revealed several interesting examples of MT domains with novel substrate specificities. Our analysis has also given interesting insight into the evolutionary basis of the novel substrate specificities of these MT proteins. These results have interesting implications for identification of novel secondary metabolites by genome mining and also rational design of novel natural products by biosynthetic engineering.

Methods

For various PKS/NRPS clusters catalogued in NRPSDB [5], PKSDB [4] and ITERDB [5], the multi functional proteins containing MT domains were identified based on comparison of chemical structure of the metabolic products with the domain organization in the proteins of the corresponding biosynthetic clusters. The PKS/NRPS clusters in which the MT domains were found included actinomycin [32], metithiazol A [42], pyochelin [37], yersiniabactin [38], stigmatellin [43], anabaenopeptilide [44], enniatin [31], leinamycin [45], microcystin [46], jamaicamide A [47], complestatin [48], bleomycin [35], epothilone [49], myxothiazol [50], nodularin [51], pristinamycin [52], thaxtomin [53], tubulysin [54], onnamide [36], pederin [36], barbamide [55], cyclosporine [56], lovastatin [57], compactin [58], fumonisin [59], erythromycin [60] and equisetin [61]. The various MT containing multi func-

tional proteins found in these PKS/NRPS clusters were analyzed by NRPS-PKS web server [5] as well as CDD [62] search. Both NRPS-PKS and CDD often failed to detect full length MT domains due to lack of homology over the complete length. The sequence stretch identified by these programs along with their flanking linkers was threaded on various structural folds in PDB using GenTHREADER [63] and PHYRE fold recognition servers [64]. In cases where chemical structure of metabolites indicated presence of MT domains but no MT domain was detected by these programs, all linker stretches having unusual length were analyzed by GenTHREADER. It is known that, in many NRPS clusters, N-MT domain is inserted between A-8 and A-9 conserved signature motifs of the adenvlation domain. Therefore, if an adenylation domain produced two discrete regions of local alignments with a regular A domain, the unaligned stretch was analyzed by threading method for possible presence of MT domains. Apart from integrated MT domains present in multidomain proteins, 8 stand alone O-MT present in NRPS or PKS clusters were also analyzed by GenTHREADER.

In order to choose a minimal set of templates representing the entire range of sequence diversity, every sequence was compared with every other sequence using BLAST program [65]. Based on these alignments, sequences with similarity above 50% and alignment length greater than 90% were grouped together. A representative sequence was selected from each group and finally 18 sequences were selected as templates for identification of MT domains by pairwise alignment. These 18 templates included five C-MT, five N-MT and eight O-MT sequences.

The sequence to structure alignments obtained from Gen-THREADER were sorted according to their threading score and the top-ranking hits were selected as possible structural templates. The boundaries of the predicted domains were obtained from the aligned region between the query and the crystal structure. The sequences were then classified into three groups as N-MT, C-MT and O-MTs by correlating with the structure of the metabolic product. The curated sequences of the three classes were taken and MSA was performed using ClustalW2 [66] and ESPript [67] software. MSA was used to study the similarity of the three classes of MT sequences and also find the pattern of conservation of the motifs among these sequences. A phylogenetic tree was obtained from multiple sequence alignment using iTOL [68]. Bootstrapping was performed 1000 times to obtain support values for each node. All significant bootstrapping values (more than 500) are shown in figure 7. A local version of HMMER 2.3.2 package was used to derive profile HMMs for each class of MT domains. These profiles were used to search the nr database (non-redundant database) of NCBI for identifying NRPS/PKS family of C-MT, N-MT and O-MT domains present in various proteins. The proteins showing matches with these HMM profiles were searched locally using Pfam – A database release 22.0 [69] at E-value cut off of 10⁻⁶ for identifying other catalytic domains associated with C-MT, N-MT and O-MT.

Authors' contributions

MZA and JS performed the computations, analyzed data and wrote the manuscript; RSG and DM designed research, analyzed data and wrote the manuscript. All the authors read and approved the final manuscript.

Additional material

Additional file 1

MSWORD file containing supplementary Figure 1. Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2105-9-454-S1.doc]

Additional file 2

MSWORD file containing supplementary Table 1. Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2105-9-454-S2.doc]

Acknowledgements

Authors thank Director, NII for encouragement and support. MZA thanks CSIR, India for award of senior research fellowship. RSG is a HHMI International Research Scholar and is also recognized with Swarnajayanti Fellowship. The work has been supported by grants to National Institute of Immunology from Department of Biotechnology, Government of India, grants to DM under BTIS project of DBT, India and grants to DM and RSG from CEFIPRA.

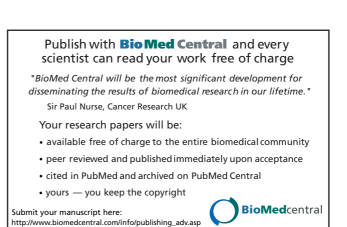
References

- Liou GF, Khosla C: Building-block selectivity of polyketide synthases. Curr Opin Chem Biol 2003, 7(2):279-284.
- Mootz HD, Schwarzer D, Marahiel MA: Ways of assembling complex natural products on modular nonribosomal peptide synthetases. Chembiochem 2002, 3(6):490-504.
- Yadav G, Gokhale RS, Mohanty D: Computational approach for prediction of domain organization and substrate specificity of modular polyketide synthases. J Mol Biol 2003, 328(2):335-363.
- Yadav G, Gokhale RS, Mohanty D: SEARCHPKS: A program for detection and analysis of polyketide synthase domains. Nucleic Acids Res 2003, 31(13):3654-3658.
- Ansari MZ, Yadav G, Gokhale RS, Mohanty D: NRPS-PKS: a knowledge-based resource for analysis of NRPS/PKS megasynthases. Nucleic Acids Res 2004:W405-413.
- Minowa Y, Araki M, Kanehisa M: Comprehensive analysis of distinctive polyketide and nonribosomal peptide structural motifs encoded in microbial genomes. J Mol Biol 2007, 368(5):1500-1517.
- Challis GL, Ravel J, Townsend CA: Predictive, structure-based model of amino acid recognition by nonribosomal peptide synthetase adenylation domains. *Chem Biol* 2000, 7(3):211-224.
- Stachelhaus T, Mootz HD, Marahiel MA: The specificity-conferring code of adenylation domains in nonribosomal peptide synthetases. Chem Biol 1999, 6(8):493-505.

- 9. Lautru S, Deeth RJ, Bailey LM, Challis GL: Discovery of a new peptide natural product by Streptomyces coelicolor genome mining. Nat Chem Biol 2005, 1(5):265-269.
- Challis GL: Mining microbial genomes for new natural products and biosynthetic pathways. *Microbiology* 2008, 154(Pt 6):1555-1569.
- Nguyen T, Ishida K, Jenke-Kodama H, Dittmann E, Gurgui C, Hochmuth T, Taudien S, Platzer M, Hertweck C, Piel J: Exploiting the mosaic structure of trans-acyltransferase polyketide synthases for natural product discovery and pathway dissection. Nat Biotechnol 2008, 26(2):225-233.
- Wilkinson B, Micklefield J: Mining and engineering natural-product biosynthetic pathways. Nat Chem Biol 2007, 3(7):379-386.
- Bergmann S, Schumann J, Scherlach K, Lange C, Brakhage AA, Hertweck C: Genomics-driven discovery of PKS-NRPS hybrid metabolites from Aspergillus nidulans. Nat Chem Biol 2007, 3(4):213-217.
- Gross H, Stockwell VO, Henkels MD, Nowak-Thompson B, Loper JE, Gerwick WH: The genomisotopic approach: a systematic method to isolate products of orphan biosynthetic gene clusters. Chem Biol 2007, 14(1):53-63.
- de Bruijn I, de Kock MJ, Yang M, de Waard P, van Beek TA, Raaijmakers JM: Genome-based discovery, structure prediction and functional analysis of cyclic lipopeptide antibiotics in Pseudomonas species. *Mol Microbiol* 2007, 63(2):417-428.
- Van Lanen SG, Shen B: Microbial genomics for the improvement of natural product discovery. Curr Opin Microbiol 2006, 9(3):252-260.
- Trivedi OA, Arora P, Vats A, Ansari MZ, Tickoo R, Sridharan V, Mohanty D, Gokhale RS: Dissecting the mechanism and assembly of a complex virulence mycobacterial lipid. *Mol Cell* 2005, 17(5):631-643.
- Menzella HG, Reeves CD: Combinatorial biosynthesis for drug development. Curr Opin Microbiol 2007, 10(3):238-245.
- Baltz RH: Molecular engineering approaches to peptide, polyketide and other antibiotics. Nat Biotechnol 2006, 24(12):1533-1540.
- Nguyen KT, Ritz D, Gu JQ, Alexander D, Chu M, Miao V, Brian P, Baltz RH: Combinatorial biosynthesis of novel antibiotics related to daptomycin. Proc Natl Acad Sci USA 2006, 103(46):17462-17467.
- Miller DJ, Ouellette N, Evdokimova E, Savchenko A, Edwards A, Anderson WF: Crystal complexes of a predicted S-adenosylmethionine-dependent methyltransferase reveal a typical AdoMet binding domain and a substrate recognition domain. Protein Sci 2003, 12(7):1432-1442.
- 22. Martin JL, McMillan FM: SAM (dependent) I AM: the S-adenosylmethionine-dependent methyltransferase fold. *Curr Opin Struct Biol* 2002, 12(6):783-793.
- 23. Hornbogen T, Riechers SP, Prinz B, Schultchen J, Lang C, Schmidt S, Mugge C, Turkanovic S, Sussmuth RD, Tauberger E, et al.: Functional characterization of the recombinant N-methyltransferase domain from the multienzyme enniatin synthetase. Chembiochem 2007, 8(9):1048-1054.
- Ishida K, Fritzsche K, Hertweck C: Geminal tandem C-methylation in the discoid resistomycin pathway. J Am Chem Soc 2007, 129(42):12648-12649.
- Schenk A, Xu Z, Pfeiffer C, Steinbeck C, Hertweck C: Geminal bismethylation prevents polyketide oxidation and dimerization in the benastatin pathway. Angew Chem Int Ed Engl 2007, 46(37):7035-7038.
- Cox RJ, Glod F, Hurley D, Lazarus CM, Nicholson TP, Rudd BA, Simpson TJ, Wilkinson B, Zhang Y: Rapid cloning and expression of a fungal polyketide synthase gene involved in squalestatin biosynthesis. Chem Commun (Camb) 2004:2260-2261.
- Song Z, Cox RJ, Lazarus CM, Simpson TT: Fusarin C biosynthesis in Fusarium moniliforme and Fusarium venenatum. Chembiochem 2004, 5(9):1196-1203.
- 28. Kagan RM, Clarke S: Widespread occurrence of three sequence motifs in diverse S-adenosylmethionine-dependent methyltransferases suggests a common structure for these enzymes. Arch Biochem Biophys 1994, 310(2):417-427.
- Gaurav K, Gupta N, Sowdhamini R: FASSM: enhanced function association in whole genome analysis using sequence and structural motifs. In Silico Biol 2005, 5(5-6):425-438.

- Katz JE, Dlakic M, Clarke S: Automated identification of putative methyltransferases from genomic open reading frames. *Mol Cell Proteomics* 2003, 2(8):525-540.
- Hacker C, Glinski M, Hornbogen T, Doller A, Zocher R: Mutational analysis of the N-methyltransferase domain of the multifunctional enzyme enniatin synthetase. J Biol Chem 2000, 275(40):30826-30832.
- Schauwecker F, Pfennig F, Schroder W, Keller U: Molecular cloning of the actinomycin synthetase gene cluster from Streptomyces chrysomallus and functional heterologous expression of the gene encoding actinomycin synthetase II. J Bacteriol 1998, 180(9):2468-2474.
- Jansson A, Niemi J, Lindqvist Y, Mantsala P, Schneider G: Crystal structure of aclacinomycin-10-hydroxylase, a S-adenosyl-Lmethionine-dependent methyltransferase homolog involved in anthracycline biosynthesis in Streptomyces purpurascens. J Mol Biol 2003, 334(2):269-280.
- 34. Gokhale RS, Sankaranarayanan R, Mohanty D: Versatility of polyketide synthases in generating metabolic diversity. *Curr* Opin Struct Biol 2007, 17(6):736-743.
- 35. Du L, Sanchez C, Chen M, Edwards DJ, Shen B: The biosynthetic gene cluster for the antitumor drug bleomycin from Streptomyces verticillus ATCC15003 supporting functional interactions between nonribosomal peptide synthetases and a polyketide synthase. *Chem Biol* 2000, **7(8)**:623-642.
- Piel J, Hui D, Wen G, Butzke D, Platzer M, Fusetani N, Matsunaga S: Antitumor polyketide biosynthesis by an uncultivated bacterial symbiont of the marine sponge Theonella swinhoei. Proc Natl Acad Sci USA 2004, 101(46):16222-16227.
- Quadri LE, Keating TA, Patel HM, Walsh CT: Assembly of the Pseudomonas aeruginosa nonribosomal peptide siderophore pyochelin: In vitro reconstitution of aryl-4, 2bisthiazoline synthetase activity from PchD, PchE, and PchF. Biochemistry 1999, 38(45):14941-14954.
- Gehring AM, DeMoll E, Fetherston JD, Mori I, Mayhew GF, Blattner FR, Walsh CT, Perry RD: Iron acquisition in plague: modular logic in enzymatic biogenesis of yersiniabactin by Yersinia pestis. Chem Biol 1998, 5(10):573-586.
- Oberthur M, Leimkuhler C, Kruger RG, Lu W, Walsh CT, Kahne D: A systematic investigation of the synthetic utility of glyco- peptide glycosyltransferases. J Am Chem Soc 2005, 127(30):10747-10752.
- Kamra P, Gokhale RS, Mohanty D: SEARCHGTr: a program for analysis of glycosyltransferases involved in glycosylation of secondary metabolites. Nucleic Acids Res 2005:W220-225.
- Ghosh R, Chhabra A, Phatale PA, Samrat SK, Sharma J, Gosain A, Mohanty D, Saran S, Gokhale RS: Dissecting functional role of polyketide synthases in dictyostelium discoideum: Biosynthesis of differentiation regulating factor MPBD. J Biol Chem 2008.
- 42. Weinig S, Hecht HJ, Mahmud T, Muller R: Melithiazol biosynthesis: further insights into myxobacterial PKS/NRPS systems and evidence for a new subclass of methyl transferases. *Chem Biol* 2003, 10(10):939-952.
- Gaitatzis N, Šilakowski B, Kunze B, Nordsiek G, Blocker H, Hofle G, Muller R: The biosynthesis of the aromatic myxobacterial electron transport inhibitor stigmatellin is directed by a novel type of modular polyketide synthase. J Biol Chem 2002, 277(15):13082-13090.
- Rouhiainen L, Paulin L, Suomalainen S, Hyytiainen H, Buikema W, Haselkorn R, Sivonen K: Genes encoding synthetases of cyclic depsipeptides, anabaenopeptilides, in Anabaena strain 90. Mol Microbiol 2000, 37(1):156-167.
- 45. Tang GL, Cheng YQ, Shen B: Leinamycin biosynthesis revealing unprecedented architectural complexity for a hybrid polyketide synthase and nonribosomal peptide synthetase. *Chem Biol* 2004, 11(1):33-45.
- Nishizawa T, Ueda A, Asayama M, Fujii K, Harada K, Ochi K, Shirai M: Polyketide synthase gene coupled to the peptide synthetase module involved in the biosynthesis of the cyclic heptapeptide microcystin. J Biochem 2000, 127(5):779-789.
- Edwards DJ, Marquez BL, Nogle LM, McPhail K, Goeger DE, Roberts MA, Gerwick WH: Structure and biosynthesis of the jamaicamides, new mixed polyketide-peptide neurotoxins from the marine cyanobacterium Lyngbya majuscula. Chem Biol 2004, 11(6):817-833.

- Chiu HT, Hubbard BK, Shah AN, Eide J, Fredenburg RA, Walsh CT, Khosla C: Molecular cloning and sequence analysis of the complestatin biosynthetic gene cluster. Proc Natl Acad Sci USA 2001, 98(15):8548-8553.
- Molnar I, Schupp T, Ono M, Zirkle R, Milnamow M, Nowak-Thompson B, Engel N, Toupet C, Stratmann A, Cyr DD, et al.: The biosynthetic gene cluster for the microtubule-stabilizing agents epothilones A and B from Sorangium cellulosum So ce90. Chem Biol 2000, 7(2):97-109.
- Silakowski B, Schairer HU, Ehret H, Kunze B, Weinig S, Nordsiek G, Brandt P, Blocker H, Hofle G, Beyer S, et al.: New lessons for combinatorial biosynthesis from myxobacteria. The myxothiazol biosynthetic gene cluster of Stigmatella aurantiaca DW4/3-I. J Biol Chem 1999, 274(52):37391-37399.
- Moffitt MC, Neilan BA: Characterization of the nodularin synthetase gene cluster and proposed theory of the evolution of cyanobacterial hepatotoxins. *Appl Environ Microbiol* 2004, 70(11):6353-6362.
- Mootz HD, Marahiel MA: Design and application of multimodular peptide synthetases. Curr Opin Biotechnol 1999, 10(4):341-348.
- 53. Healy FG, Wach M, Krasnoff SB, Gibson DM, Loria R: The txtAB genes of the plant pathogen Streptomyces acidiscables encode a peptide synthetase required for phytotoxin thaxtomin A production and pathogenicity. *Mol Microbiol* 2000, 38(4):794-804.
- Sandmann A, Sasse F, Muller R: Identification and analysis of the core biosynthetic machinery of tubulysin, a potent cytotoxin with potential anticancer activity. Chem Biol 2004, 11(8):1071-1079.
- 55. Chang Z, Flatt P, Gerwick WH, Nguyen VA, Willis CL, Sherman DH: The barbamide biosynthetic gene cluster: a novel marine cyanobacterial system of mixed polyketide synthase (PKS)non-ribosomal peptide synthetase (NRPS) origin involving an unusual trichloroleucyl starter unit. Gene 2002, 296(1-2):235-247.
- Lawen A, Traber R: Substrate specificities of cyclosporin synthetase and peptolide SDZ 214-103 synthetase. Comparison of the substrate specificities of the related multifunctional polypeptides. *J Biol Chem* 1993, 268(27):20452-20465.
- polypeptides. J Biol Chem 1993, 268(27):20452-20465.
 57. Hendrickson L, Davis CR, Roach C, Nguyen DK, Aldrich T, McAda PC, Reeves CD: Lovastatin biosynthesis in Aspergillus terreus: characterization of blocked mutants, enzyme activities and a multifunctional polyketide synthase gene. Chem Biol 1999, 6(7):429-439.
- Abe Y, Suzuki T, Ono C, Iwamoto K, Hosobuchi M, Yoshikawa H: Molecular cloning and characterization of an ML-236B (compactin) biosynthetic gene cluster in Penicillium citrinum. *Mol Genet Genomics* 2002, 267(5):636-646.
- Proctor RH, Desjardins AÈ, Plattner RD, Hohn TM: A polyketide synthase gene required for biosynthesis of fumonisin mycotoxins in Gibberella fujikuroi mating population A. Fungal Genet Biol 1999, 27(1):100-112.
- Rawlings BJ: Type I polyketide biosynthesis in bacteria (Part A – erythromycin biosynthesis). Nat Prod Rep 2001, 18(2):190-227.
- Sims JW, Fillmore JP, Warner DD, Schmidt EW: Equisetin biosynthesis in Fusarium heterosporum. Chem Commun (Camb) 2005:186-188.
- 62. Marchler-Bauer A, Bryant SH: CD-Search: protein domain annotations on the fly. Nucleic Acids Res 2004:W327-331.
- McGuffin LJ, Jones DT: Improvement of the GenTHREADER method for genomic fold recognition. Bioinformatics 2003, 19(7):874-881.
- Bennett-Lovsey RM, Herbert AD, Sternberg MJ, Kelley LA: Exploring the extremes of sequence/structure space with ensemble fold recognition in the program Phyre. Proteins 2008, 70(3):611-625.
- McGinnis S, Madden TL: BLAST: at the core of a powerful and diverse set of sequence analysis tools. Nucleic Acids Res 2004:W20-25.
- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, et al.: Clustal W and Clustal X version 2.0. Bioinformatics 2007, 23(21):2947-2948.
- Gouet P, Robert X, Courcelle E: ESPript/ENDscript: Extracting and rendering sequence and 3D information from atomic structures of proteins. Nucleic Acids Res 2003, 31(13):3320-3323.



- Letunic I, Bork P: Interactive Tree Of Life (iTOL): an online tool for phylogenetic tree display and annotation. Bioinformatics 2007, 23(1):127-128.
- Finn RD, Mistry J, Schuster-Bockler B, Griffiths-Jones S, Hollich V, Lassmann T, Moxon S, Marshall M, Khanna A, Durbin R, et al.: Pfam: clans, web tools and services. Nucleic Acids Res 2006:D247-251.