

ORIGINAL ARTICLE

Deep sequencing of non-ribosomal peptide synthetases and polyketide synthases from the microbiomes of Australian marine sponges

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The biosynthesis of non-ribosomal peptide and polyketide natural products is facilitated by multimodular enzymes that contain domains responsible for the sequential condensation of amino and carboxylic subunits. These conserved domains provide molecular targets for the discovery of natural products from microbial metagenomes. This study demonstrates the application of tag-encoded FLX amplicon pyrosequencing (TEFAP) targeting non-ribosomal peptide synthetase (NRPS) and polyketide synthase (PKS) genes as a method for determining the identity and diversity of natural product biosynthesis genes. To validate this approach, we assessed the diversity of NRPS and PKS genes within the microbiomes of six Australian marine sponge species using both TEFAP and metagenomic whole-genome shotgun sequencing approaches. The TEFAP approach identified 100 novel ketosynthase (KS) domain sequences and 400 novel condensation domain sequences within the microbiomes of the six sponges. The diversity of KS domains within the microbiome of a single sponge species *Scopalina* sp. exceeded that of any previously surveyed marine sponge. Furthermore, this study represented the first to target the condensation domain from NRPS biosynthesis and resulted in the identification of a novel condensation domain lineage. This study highlights the untapped potential of Australian marine sponges for the isolation of novel bioactive natural products. Furthermore, this study demonstrates that TEFAP approaches can be applied to functional genes, involved in natural product biosynthesis, as a tool to aid natural product discovery. It is envisaged that this approach will be used across multiple environments, offering an insight into the biological processes that influence the production of secondary metabolites.

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Introduction

Non-ribosomal peptides (NRPs) and polyketides (PKs) are the cornerstones of many modern pharmaceuticals, and provide a molecular and chemical source for the isolation of novel bioactive compounds. The last 30 years have seen the majority of these compounds derived from microorganisms that were isolated from the environment and cultured within the laboratory (Newman and Cragg, 2012). Although advances in culturing methodology are enabling the isolation of previously recalcitrant microbes (Ferrari *et al.*, 2008), culturable organisms still represent only a small proportion of the total microbial diversity in many environments

(Keller and Zengler, 2004; Donachie *et al.*, 2007). The uncertainty regarding the depth of metabolic diversity that exists within a given sample remains.

Marine sponges represent a significant resource for the isolation of novel bioactive compounds (Fusetani and Matsunaga, 1993; Blunt *et al.*, 2006; Thomas *et al.*, 2010). The majority of these compounds exhibit structural features indicative of bacterial biosynthetic routes and microorganisms have been isolated that are capable of producing sponge-associated metabolites (Boot *et al.*, 2006). Previous molecular studies, utilising vector-dependent approaches, identified a lack of functional diversity among some sponge species due to the dominance of sponge-specific polyketide synthases (PKSs) (Piel *et al.*, 2004b; Schirmer *et al.*, 2005; Fieseler *et al.*, 2007; Hochmuth *et al.*, 2010). This has since led to the development of assays that amplify specific ketosynthase (KS) groups involved in the biosynthesis of complex secondary metabolites (Piel *et al.*, 2004a; Fisch *et al.*, 2009). However,

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these highly specific approaches are only applicable for the identification of biosynthetic pathways when the product is known. The feasibility of using high-throughput sequencing methods to determine the true metabolic diversity of KS domains from a single marine sponge has been previously reported (Trindade-Silva *et al.*, 2012). In contrast, those studies targeting non-ribosomal peptide synthetases (NRPS) from marine sponge microbiomes, limited to the use of vector-dependent approaches targeting the adenylation (A) domain, revealed minimal biosynthetic diversity (Kennedy *et al.*, 2008; Pimentel-Elardo *et al.*, 2012).

The biosynthesis of NRPs and PKs is facilitated by the action of a series of multimodular enzymes, with each module containing a core set of three domains required for the activation and condensation of a single amino or carboxylic subunit. The KS and condensation (C) domains, that catalyse the condensation of two activated subunits, contain motifs that are conserved at an amino-acid level (Stachelhaus *et al.*, 1998; Walsh and Fischbach, 2010). Between these motifs, variation is observed, typically within catalytic centres, giving rise to domains with functional specificity, reflecting the nature of the subunits being condensed (Moffitt and Neilan, 2003; Rausch *et al.*, 2007), the arrangement of other domains within the module and the expected product (Moffitt and Neilan, 2003; Piel *et al.*, 2004a; Schmitt *et al.*, 2008). The conserved nature for part of these domains ensures their suitability as targets for amplification using the PCR. Combining PCR with tag-encoded FLX amplicon pyrosequencing (TEFAP) (Sun *et al.*, 2011) provides a possible means for evaluating the composition and diversity of NRPS and PKS genes from microbiomes. TEFAP generates thousands of reads per sample with sufficient read length (~400 bp) and allows for phylogenetic discrimination amongst both taxonomic markers and functional genes, for example, *nifH* and *dmdA*. By multiplexing large numbers of environmental samples, the cost and time involved in evaluating the composition and diversity of a single target within an environment is drastically reduced.

The strength of targeting these domains is that they describe the diversity of gene clusters encoding bioactive molecules from organisms that are either difficult to isolate or produced by 'rare biosphere' representatives, as defined by their occurrence at very low abundances (Piel *et al.*, 2004a; Sogin *et al.*, 2006; Fisch *et al.*, 2009). Traditional methods, involving exhaustive extractions of sponge, do not reveal the chemical diversity produced by these rare biosphere representatives, or pathways that are inactive within the native host (Chiang *et al.*, 2010). Harnessing this potentially novel chemistry is, therefore, dependent on the ability of molecular approaches to identify where this diversity exists (Fieseler *et al.*, 2007; Hochmuth *et al.*, 2010; Trindade-Silva *et al.*, 2012), to provide access to

the genetic basis of biosynthesis (Piel *et al.*, 2004a; Fisch *et al.*, 2009; Banik and Brady, 2010; Brady *et al.*, 2010) and ultimately to facilitate production of these molecules through heterologous expression within a suitable host (Fu *et al.*, 2008; Craig *et al.*, 2010).

In this study, we aim to highlight the untapped potential of Australian marine sponges by evaluating the diversity of NRP and PK biosynthesis genes using a deep-sequencing TEFAP approach. The limitations and advantages of the TEFAP method, in comparison to a non-targeted metagenomic whole-genome shotgun sequencing (mWGS) approach are determined by comparing the two independent outcomes. In addition, we provide insight into how these data sets could be applied for identifying and exploiting environments containing rich biosynthetic potential. Hence, this study demonstrates the suitability of a TEFAP-based approach for evaluating and improving access to the genetic basis for natural product biosynthesis within marine sponges and other environments with untapped genomic diversity.

Materials and methods

Degenerate PCR and TEFAP

DNA derived from microbial enrichments of six Australian marine sponges was obtained in triplicate as previously described (Fan *et al.*, 2012). Extracted DNA from three replicates for each marine sponge species were pooled in equimolar amounts and screened using two sets of degenerate primers. KS sequences were amplified using the DKF/DKR (see Supplementary Information Table S1) primer pair as previously described (Moffitt and Neilan, 2003). C domain sequences from NRPS modules were amplified using the primers CnDmF and DCCR (see Supplementary Information Table S1). To facilitate subsequent incorporation of sample-specific barcodes and sequencing primer sites, amplification of each positive sample was repeated using the corresponding forward primer containing a 5' T7 promoter sequence and the corresponding reverse primer containing a 5' M13R sequence (see Supplementary Information Table S1). Amplification was performed for 35 rounds of thermal cycling at an annealing temperature of 50 °C for KS domains and 45 °C for C domains. Following PCR amplification products were gel extracted using the Zymo-Clean Gel Extraction Kit (Zymo Research, Irvine, CA, USA). For each positive sample 10 ng of purified PCR product was used as a template in a second PCR using the same cycling conditions. In this PCR a single universal reverse primer (454R-M13R) and sample-specific forward primer (454F-T7Prom) were used to incorporate priming sites of the sequencing primer and sample-specific barcodes (see Supplementary Tables S1 and S2). Samples were combined into two separate pools

corresponding to each gene target. Library preparation and unidirectional amplicon sequencing was performed at the Clive and Vera Ramaciotti Centre for Gene Function Analysis (Sydney, Australia) using the 454 FLX Titanium platform.

Identification of NRPS and PKS from mWGS data

Metagenomic DNA for the microbial communities was isolated for each replicate of the six sponges and sequenced separately on a 454 FLX pyrosequencer (Roche, Branford, CT, USA) (Fan *et al.*, 2012). Samples received between 360 000 and 1300 000 shotgun reads which were subsequently separately assembled using the Newbler software (Roche). Amino-acid sequences, translated from metagenomic contigs, for six sponges were obtained as previously described (Fan *et al.*, 2012). Sequences containing KS and C domains were recovered using the HMMER algorithm (Finn *et al.*, 2011) by searching the PFAM profiles (Finn *et al.*, 2010) PF00109 and PF00668, respectively. Hits that obtained a bit score >25 were retained. For each protein sequence recovered, the corresponding nucleotide sequence was retained for further analysis.

Sequence processing

TEFAP reads were subjected to initial pre-processing that included noise reduction using the shhh.flows algorithm (Schloss *et al.*, 2009), removal of sequences containing ambiguous bases and long (>8) homopolymers, multiplex-barcode dependent binning of sequences and removal of primer sequences.

Reference nucleotide and amino-acid alignments were generated using MUSCLE (Edgar, 2004), using reference sequences obtained from the Genbank database (Accessed: 23 January 2012) (Benson *et al.*, 1997). Reference sequences were selected on the basis of either belonging to known, characterised biosynthetic pathways or due to their previously established phylogenetic distribution (Moffitt and Neilan, 2003; Roongsawang *et al.*, 2005; Fieseler *et al.*, 2007; Rausch *et al.*, 2007). As a result, the C domain alignment contained 162 reference sequences while the KS domain alignment had 190 reference sequences.

Pre-processed TEFAP reads were screened against the respective nucleotide alignment to remove non-target sequences that were amplified as a result of the degenerate PCR conditions. Nucleotide sequences, derived from PFAM-dependent screening of the mWGS data sets, were also screened using this same approach. Following screening, the TEFAP reads and mWGS (open reading frames) were combined. Previous phylogenetic analyses of KS and C domains have indicated that these domains typically exhibit at least 5% amino-acid dissimilarity (Moffitt and Neilan, 2003; Roongsawang *et al.*, 2005; Rausch *et al.*, 2007). Previous TEFAP methods have reflected

this amino-acid dissimilarity by clustering nucleotides at a 90% nucleotide similarity cutoff (Varaljay *et al.*, 2010). Therefore, in this study, clustering of sequences into operational taxonomic units (OTUs) (Schloss and Westcott, 2011), rarefaction analyses, and generation of diversity indices was performed using the average neighbour method (Schloss *et al.*, 2009) as implemented in Mothur version 1.23.1 at a 0.10 distance threshold.

Taxonomic and functional classification

To determine whether the two methods accessed comparable genetic diversities, a qualitative assessment of the approximate taxonomic identity of sequences from both methods was made. TEFAP OTUs and mWGS open reading frames were assigned to taxa using the last common ancestor algorithm incorporated in the MEGAN 4.70.4 software package (Huson *et al.*, 2011). Individual sequences were assigned to a taxon where at least 10% of hits with a bit score >35 were in agreement. A direct comparison was also made by matching TEFAP OTUs with unprocessed mWGS sequences using the BLASTn algorithm (Altschul *et al.*, 1997). A successful match was identified where at least 90% nucleotide identity was observed across at least 100 bp. The GC content of TEFAP reads and unprocessed mWGS was determined using the GEECEE algorithm in the EMBOSS package (Rice *et al.*, 2000).

TEFAP and mWGS OTU representative sequences were individually scrutinised for correct translation into amino-acid sequences. Derived amino-acid sequences were aligned against the amino-acid reference alignment by MUSCLE (Edgar, 2004). Phylogenetic inference of amino-acid sequences was made using PhyML v3.0 (Guindon *et al.*, 2005).

Results

Identification of novel NRPS and PKS biosynthetic genes by TEFAP

Using degenerate primers, PKSs (KS domains) were detected in *Cymbastela concentrica*, *Cymbastela coralliophila*, *Sylissa* sp., *Scopalina* sp. and *Rhopaloeides odorabile*, while NRPSs (C domains) were detected in *Scopalina* sp., *C. concentrica* and *C. coralliophila*. NRP and PK biosynthetic pathways were not detected in the microbiome of the sponge *Tedania anhelens*. Following processing through the pipeline described, approximately two-thirds of the sequences generated were discarded due to length and specificity requirements (see Supplementary Figure S1). In total, 1097 KS and 4469 C domain sequences of at least 240 bp in length were retained.

A search of PFAM sequence profiles recovered 2012 KS and 133 C domain sequences from the six mWGS data sets. The corresponding nucleic acid sequences were obtained with the length of KS

domain sequences ranging between 87 and 1885 bp and the length of C domain sequences ranging between 79 and 792 bp. These nucleic acid sequences were subjected to the same pipeline that was applied to the TEFAP reads and 847 KS and 14 C domain sequences were retained (see Supplementary Figure S1). The low retention rate reflected the fragmented nature and short read length of the mWGS reads, resulting in the removal of partial KS domain and C domain sequences that did not contain the targeted domain region. Manual inspection, using the BLASTx algorithm, revealed many of the 847 KS sequences corresponded to ketoacylsynthase (I/II) sequences that, while belonging to the KS superfamily (Moffitt and Neilan, 2003), relate to aspects of primary metabolism. Following removal of these sequences, 114 KS sequences remained (Table 1). Processed TEFAP and mWGS sequences were combined and clustered into OTUs at a 0.10 distance threshold resulting in the generation of 133 KS domain OTUs and 396 C domain OTUs (Table 1). To assess whether any reads corresponded to previously characterised biosynthetic pathways, dereplication of TEFAP and mWGS OTUs was performed by comparison against the Genbank database (Benson *et al.*, 1997) using the BLASTx algorithm (Altschul *et al.*, 1997). Only one C domain and 10 KS TEFAP OTUs exhibited significant (>90%) translated amino-acid identity to previously identified NRPS and PKS protein sequences, respectively.

Table 1 Summary of sampling and gene discovery among both amplicon and shotgun-derived data sets

Target	Sponge	Method	No. of sequences	OTUs ^a	Coverage
Ketosynthase	<i>Cymbastela concentrica</i>	TEFAP	324	7	0.99
		mWGS	1 (17)	1	0
	<i>Scopalina</i> sp.	TEFAP	369	41	0.95
		mWGS	1 (8)	1	0
	<i>Cymbastela coralliophila</i>	TEFAP	15	11	0.53
		mWGS	5 (23)	5	0
	<i>Rhopaloeides odorabile</i>	TEFAP	350	17	0.97
		mWGS	107 (346)	49	0.71
	<i>Stylissa</i> sp.	TEFAP	39	16	0.74
		mWGS	0 (3)	0	N/A
Condensation	<i>Cymbastela concentrica</i>	TEFAP	14	9	0.64
		mWGS	4 (18)	4	0
	<i>Scopalina</i> sp.	TEFAP	3336	325	0.94
		mWGS	10 (51)	4	0.70
	<i>Cymbastela coralliophila</i>	TEFAP	1119	54	0.97
		mWGS	0 (18)	0	N/A

Abbreviations: mWGS, metagenomic whole-genome shotgun sequencing; OUT, operational taxonomic units; TEFAP, tag-encoded FLX amplicon pyrosequencing.

^aOTUs were defined at 0.10 distance threshold. Numbers in brackets represent the total number of KS and C domain sequences obtained using PFAM identifiers.

Comparison of TEFAP and mWGS approaches for the recovery of KS and C domain diversity

TEFAP reads were in excess of mWGS sequences in each sample for both gene targets. As a consequence, the number of OTUs observed by the TEFAP approach exceeded that observed by the mWGS approach (Table 1; Supplementary Figure S2), indicating a more comprehensive evaluation of the diversity of these secondary metabolite biosynthesis genes. The only exception was the sponge *R. odorabile*, in which more OTUs were observed by the mWGS approach (Table 1; Supplementary Figure S2).

BLASTn searches were used to match OTUs derived from the TEFAP approach to unprocessed mWGS reads. 57.14% of *C. concentrica* KS domain OTUs and 29.41% of *R. odorabile* KS domain OTUs matched to the mWGS sequences. Only 0.62% of *C. coralliophila* and 2.15% of *Scopalina* sp. C domain OTUs were matched to mWGS sequences. No overlap was observed for C domain sequences from *C. concentrica* or KS domain sequences from *Scopalina* sp., *C. coralliophila* and *Stylissa* sp. In order to scrutinise this lack of overlap, the taxonomic identity of reads from both the TEFAP and mWGS data sets were compared. Distinct differences were observed between the two methods in regard to the confidence at which sequences were assigned to more resolved taxonomic groups (see Supplementary Figure S3). Furthermore, the TEFAP approach enriched for sequences with GC contents ranging between 45 and 50%, and 60 and 65%, while the mWGS data showed a dominance of sequences with GC contents ranging between 25 and 35% for C domains and 65 and 75% for KS domains (see Supplementary Figure S4).

Reads derived from the TEFAP methods were used to calculate diversity and richness indices for each sponge where at least 324 reads were obtained. As a result of low sequence coverage, only three sponges could be analysed for their KS domains and two for their C domain sequences. Overall, the C domains presented greater observed richness (ACE), whereas the KS domains had higher observed diversity (invSimpson) (Table 2). KS and C domain

Table 2 Alpha diversity indices for select amplicon-derived samples following normalisation

Target	Sponge	Coverage	OTUs	ACE	invSimpson
Ketosynthase	<i>Cymbastela concentrica</i>	0.99	6	14.89	1.82
	<i>Scopalina</i> sp.	0.95	37	61.28	7.91
	<i>Rhopaloeides odorabile</i>	0.97	16	38.24	3.06
Condensation	<i>Scopalina</i> sp.	0.86	70	379.78	5.90
	<i>Cymbastela coralliophila</i>	0.95	28	121.98	2.99

Abbreviation: OTU, operational taxonomic units. Both condensation and ketosynthase diversity indices were determined by examining 324 sequences from each sample.

richness and diversity were highest within the sponge *Scopalina* sp. Despite being dominated by a single functional group of KS domains (Figure 1), *R. odorabile* contained a more diverse population of KS domains compared with *C. concentrica*.

The TEFAP approach and mWGS revealed different phylogenetic compositions of C and KS domains with the marine sponges (Figure 1 and Figure 2).

The two sponges *C. concentrica* and *Scopalina* sp. were shown, by both methods, to contain C domains responsible for the condensation of two L-amino subunits, annotated as symbiont LCL Clade 1 and Symbiont LCL Clade 2, that contained only sequences obtained from this study (Figure 2). The specific taxonomic assignment of sequences to these clades varied between the two methods. Both

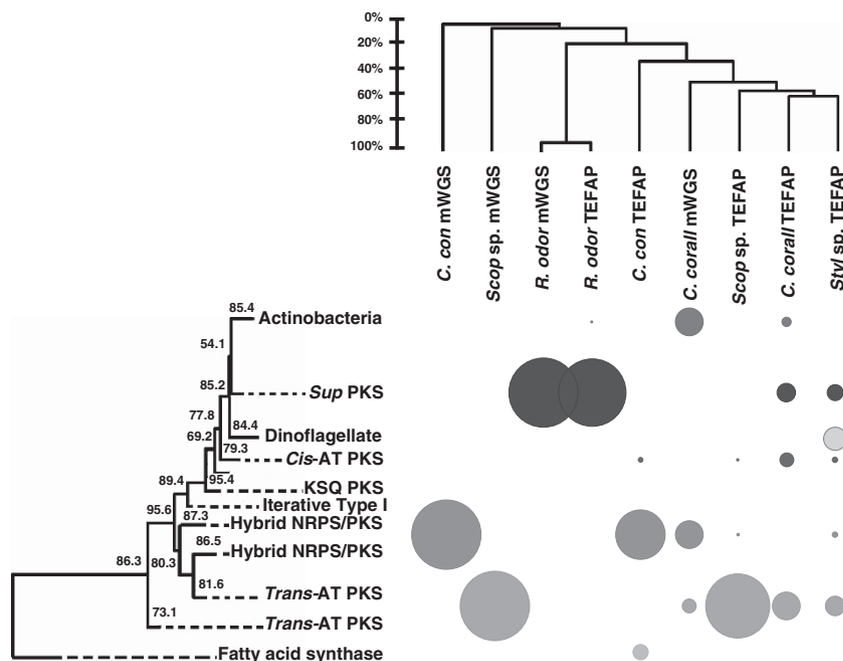


Figure 1 Taxonomic composition of KS domain sequences derived from amplicon pyrosequencing and shotgun sequencing of individual sponges. The size of a dot reflects the relative abundance of that taxon in a sample. Support values for phylogenetic groups are provided.

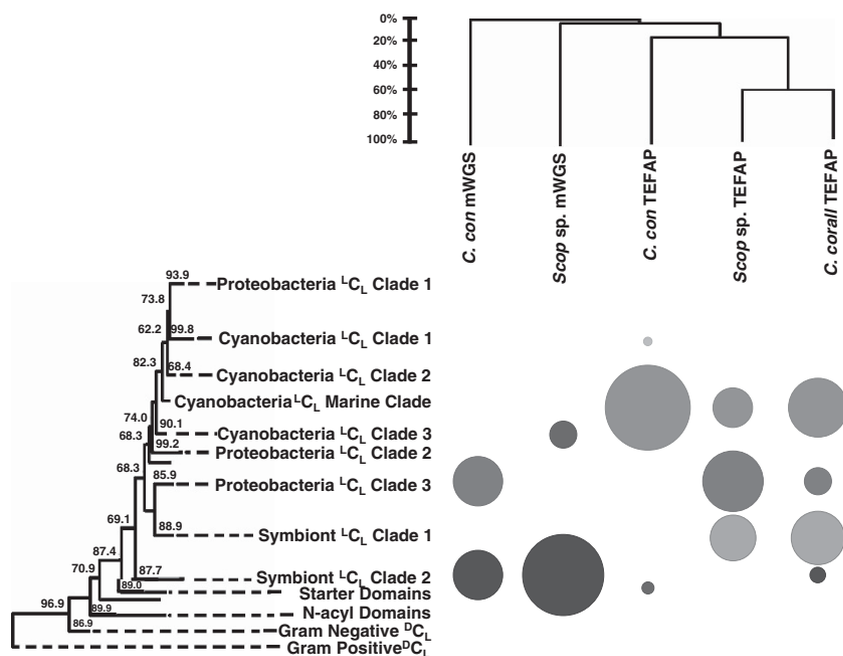


Figure 2 Taxonomic composition of C domain sequences derived from amplicon pyrosequencing and shotgun sequencing of individual sponges. The size of a dot reflects the relative abundance of that taxon in a sample. Support values for phylogenetic clades are provided.

methods were also in agreement as to the presence of a clade of proteobacteria-like LCL domain sequences in the sponge *C. concentrica* and the presence of cyanobacteria-like LCL domain sequences in the sponge *Scopalina* sp. The TEFAP method alone revealed the presence of Proteobacteria-like LCL sequences in *Scopalina*, and cyanobacteria-like LCL in *C. concentrica* and *C. coralliophila*.

Among KS domains a similar pattern was observed (Figure 1). Both methods were in agreement as to the dominance of hybrid NRPS/PKS sequences in *C. concentrica*, *trans*-AT sequences in *Scopalina* sp. and sponge ubiquitous PKS (*sup*) sequences in *R. odorabile*. In each of these three instances, additional phylogenetic groups were observed when the sponge was surveyed by the TEFAP approach. The mWGS approach was able to identify hybrid NRPS/PKS sequences within the sponge *C. coralliophila*, though these sequences were not detected using the TEFAP approach.

Discussion

Amplicon pyrosequencing enables unprecedented discovery of natural product biosynthesis

In this study, a TEFAP approach targeting natural product biosynthetic genes was applied to six marine sponge species. This study reports the use of a TEFAP approach for simultaneously evaluating the diversity of multiple domains involved in natural product biosynthesis from a number of environments. Both NRPS and PKS biosynthetic pathways were detected within the microbiomes of *Scopalina* sp., *C. coralliophila* and *C. concentrica*, whereas only PKS biosynthetic pathways were identified within the microbiomes of *Stylissa* sp. and *R. odorabile*. The TEFAP approach resulted in the identification of ~100 novel KS domain and 400 novel C domain sequences. While there was an apparent lack of coverage for these domains in some of the sponge samples, overall the diversity among the Australian sponge species as revealed by the TEFAP approach exceeded that revealed by mWGS. In addition the diversity of KS and C domains within the microbiomes of individual sponges, particular *Scopalina* sp., was greater than that of any other marine sponge (Fieseler *et al.*, 2007; Kennedy *et al.*, 2008; Pimentel-Elardo *et al.*, 2012). This is despite a relaxed clustering approach, whereby a 0.10 distance threshold was adopted (Varaljay *et al.*, 2010; Howard *et al.*, 2011) for the identification of novel sequences, in contrast to the 97% nucleotide similarity cutoff reported in comparable studies (Fieseler *et al.*, 2007; Pimentel-Elardo *et al.*, 2012). The relaxed cutoff would also account for any inflation of the discovery rate due to random sequencing errors. Within individual samples, the TEFAP and mWGS methods applied were in some disagreement in regards to the average GC

content (see Supplementary Figure S4), as well as the functional (Figures 1 and 2) and taxonomic (see Supplementary Figure S3) composition. This would indicate a potential technical biases of amplicon-dependent approaches leading to the artificial enrichment of certain sequences as observed by others (Piel, 2002; Piel *et al.*, 2004a, b; Fisch *et al.*, 2009; Pimentel-Elardo *et al.*, 2012).

The occurrence of technical biases associated with the TEFAP method implies that the reported relative composition of KS and C domain sequences within each sample is skewed. The most notable implication is the failure to detect KS and C domains corresponding to the high GC Gram-positive actinobacteria and firmicutes. This is directly reflected within GC plots (see Supplementary Figure S4), with the TEFAP method selecting for two GC ranges, likely corresponding to cyanobacterial and proteobacterial groups (Figures 1 and 2). Ultimately, this suggests that the diversity of both KS and C domain sequences within a sponge is possibly higher than that observed.

An additional consideration is that of modular redundancy, whereby multiple C and KS domains cooperate within a single biosynthetic pathway. Efforts to avoid excessive redundancy were made by utilising the C domain in favour of the commonly applied A domain, an approach, which has already been adopted by others (Ziemert *et al.*, 2012). In addition, we adopted a low distance threshold to account for gene duplications that may give rise to multimodular systems. However, the assembly-line logic of these biosynthetic pathways stipulates that multiple genetically distinct C and KS modules will be present in a single pathway leading to over-estimations of diversity. Although one could assume that typical pathways feature 3–7 condensing domains per biosynthetic pathway, targeted recovery of large metagenomic fragments is ultimately required to determine the extent of this redundancy (Piel *et al.*, 2004a; Fisch *et al.*, 2009). Regardless of these considerations, ensuring that multiple environments are sampled using a standardised approach, the TEFAP method allows for comparative analysis of composition and diversity between multiple environments.

A novel TEFAP approach highlights the high diversity of PKS genes in the microbiomes of Australian marine sponges

This study represents the first analysis of PKS diversity from Australian marine sponges. A previous study, utilising vector-based approaches, reported the identification of 150 unique KS OTUs (97% nucleotide identity) that were derived by undertaking Sanger sequencing of nearly 500 amplicons from the three marine sponges *Theonella swinhoei*, *Cacospongia mycofijiensis* and *Aplysina aerophoba* (Fieseler *et al.*, 2007). In contrast, TEFAP enabled the discovery of nearly 100 KS domain

sequences from the five Australian marine sponges, albeit at a lower distance threshold of 0.10. At a comparable distance threshold of 0.03, the TEFAP method reported the discovery of 296 OTUs (data not shown), far exceeding previous vector-based studies (Fieseler *et al.*, 2007) but comparable with other amplicon sequencing approaches (Trindade-Silva *et al.*, 2012).

Schirmer *et al.* (2005) and Fieseler *et al.* (2007) both reported the dominance of *sup* type KS domains within amplicon clone libraries from *Discoderma dissoluta*, *Theonella swinhoei*, *Cacospongia mycofijiensis* and *Aplysina aerophoba*, as well as the detection of *sup* type KS in *Verongula gigantea*, *Aiolochoira crassa*, *Xestospongia muta* and *Siphonodictyon coralliphagum*. In this study, *sup* KS domains were detected in abundance within the sponge *R. odorabile*, along with a small number of *cis*-KS domains phylogenetically affiliated with actinobacteria. This co-occurrence of these two domain types was also observed from the sponge *Cacospongia mycofijiensis* (Fieseler *et al.*, 2007; Hochmuth *et al.*, 2010), which belongs to the same order as *R. odorabile*. The occurrence of these *sup* type KS domains in *R. odorabile* coincided with the detection of poribacteria (Fan *et al.*, 2012), which are known to harbour the *sup* biosynthetic gene cluster (Siegl and Hentschel, 2010; Siegl *et al.*, 2011). The calculated richness index (Table 2), indicated a moderate number of *sup* KS domains within the microbiome of this sponge. That this moderate richness corresponded with a low invSimpson value, indicating that a few *sup* domains were dominant, reflects the shallow phylogenetic branching nature of this group (Fieseler *et al.*, 2007; Hochmuth and Piel, 2009). In contrast to *R. odorabile*, poribacteria were not previously detected in the sponges *C. coralliophila* and *Stylissa* sp. (Fan *et al.*, 2012) using the mWGS technique. However, the TEFAP approach detected *sup* type KS domains, which are thought to be limited to poribacterial species (Hochmuth and Piel, 2009; Siegl and Hentschel, 2010; Siegl *et al.*, 2011). The detection of *sup* domains at very low abundance has also been reported within the marine sponge *Arenosclera brasiliensis* (Trindade-Silva *et al.*, 2012). This discrepancy may be explained by poribacteria constituting only a small proportion of the sponge microbiome in these samples, reflective of levels observed in seawater (Taylor *et al.*, 2012). As such the level of sequencing achieved in the mWGS approach (Fan *et al.*, 2012), which targets the most abundant taxa, may not have been sufficient to retrieve rare sequences. However, it is possible that the *sup* type KS domain is not exclusive to poribacteria, but is present in other lineages.

While Fieseler *et al.* (2007) reported that *sup* type KS domains dominated the majority of sponges screened, the presence of *trans*-acyltransferase (*trans*-AT) type KS domains are more relevant for the discovery of novel bioactive small molecules.

Previously identified pathways containing *trans*-AT domains are responsible for the biosynthesis of complex polyketides with bioactivities relevant to the pharmaceutical industry (Piel, 2002; Piel *et al.*, 2004b; Fisch *et al.*, 2009). The *trans*-AT KS domains, detected in this study within the sponges *C. coralliophila*, *Scopalina* sp. and *Stylissa* sp. (Figure 1), are distinct from typical Type I or *cis*-AT KS in that one of the core catalytic domains is absent. In these systems, the AT domain, responsible for activation of the carboxylic acid precursor, is found as a mono-functional enzyme proximal to the biosynthetic gene cluster. To date, a number of *trans*-AT biosynthetic pathways have been reported primarily from organisms in symbiotic associations (Piel, 2002; Piel *et al.*, 2004b; Fisch *et al.*, 2009). *Scopalina* sp. contained 41 KS domains, the majority of which were *trans*-AT type. The *trans*-AT domains from *Scopalina* sp., *C. coralliophila* and *Stylissa*, did not exhibit >90% amino-acid identity to any known biosynthetic pathways, indicating the presence of novel biosynthetic pathways within these sponges. Surprisingly, a survey of the literature revealed a lack of bioactive compounds isolated from *Scopalina* spp., which in the context of this study may represent a large untapped resource for natural product discovery.

Metagenomic mining of condensation domains reveals an unprecedented diversity of NRP biosynthesis

Despite the large chemical diversity of cyclic peptides isolated from sponges and other marine invertebrates (Fusetani and Matsunaga, 1993; De Rosa *et al.*, 2003; Thomas *et al.*, 2010), C domains were only detected within the microbial communities of half the surveyed sponges. A high richness was observed among C domains within the sponges *Scopalina* sp. and *C. coralliophila*, particularly in contrast to that observed among KS domains (Table 2). The higher invSimpson values, among the C domain sequences suggested a more even distribution with far fewer rare sequences. This is likely to reflect either the presence of a single super-producing organism, or indicate that many of these C domains are present in a small number of clusters. The phylogenetic inference of the C domains is enabled by the presence of two binding pockets adjacent to the catalytic centre (Stachelhaus *et al.*, 1998). The C-terminus binding pocket, considered the acceptor site, exhibits specific selectivity for the activated substrate, which is conferred within the amino-acid sequence. This allows for the discrimination between activated L-amino, D-amino or N-acyl substrates (Roongsawang *et al.*, 2005; Rausch *et al.*, 2007). Within this study, both the TEFAP and mWGS approaches were in agreement as to the presence of only LCL domains within *Scopalina* sp., *C. concentrica* and *C. coralliophila*. These domain types are defined by the presence of an L-amino acid in both the acceptor and donor binding pocket

(Roongsawang *et al.*, 2005; Rausch *et al.*, 2007). Further phylogenetic analysis revealed both methods were in agreement that these LCL domains were limited to Gram-negative organisms, particularly members of the γ -proteobacteria, δ -proteobacteria and cyanobacteria (Figure 2).

Despite not detecting cyanobacteria within *Scopalina* sp. and *C. concentrica* (Fan *et al.*, 2012), the TEFAP approach amplified reads that form a phylogenetic clade with C domains from the marine cyanobacteria *Moorea producens* (Jones *et al.*, 2011) and *Acaryochloris marina* (Swingley *et al.*, 2008). Within *Scopalina* sp. and *C. concentrica*, a number of sequences also formed a phylogenetic clade with C domains from proteobacteria (Figure 2). Proteobacteria LCL Clade 3 contained exclusively reference sequences from members of the γ -proteobacteria, a group that was not identified from mWGS of *Scopalina* sp. and *C. concentrica*. For *Scopalina* sp., this may indicate the presence of γ -proteobacteria in the 'rare biosphere' (Sogin *et al.*, 2006). However, in the instance of *C. concentrica*, the mWGS method also identified sequences that grouped within proteobacteria LCL Clade 3, suggesting that in this case the organism harbouring this pathway is most likely abundant.

A large number of C domain sequences from the two marine sponges, *C. concentrica* and *Scopalina* sp., as surveyed by the TEFAP method, formed a phylogenetic clade independent of any reference sequences (Figure 2). A second phylogenetic clade containing exclusively mWGS derived sequences was also observed. These clades were annotated as Symbiont LCL Clade 1 and 2, respectively. The presence of these two phylogenetic clades, comprised entirely of C domains from sponge symbionts, is particularly remarkable considering that previous analyses of KS from marine sponges have revealed distinct taxonomic groups (Piel *et al.*, 2004b; Trindade-Silva *et al.*, 2012), leading to the definition of, in one instance, the *sup* type KS group (Fieseler *et al.*, 2007).

Conclusions

Bioactive natural products of NRPS and PKS origin are highly valued by the pharmaceutical industry as lead compounds against existing and emerging diseases. Researchers within the natural product discipline are under increasing pressures to find new resources for these compounds. During the last 10 years, significant focus has been placed on screening environments in an effort to identify natural products produced by organisms intractable to traditional methods. The approaches applied to date have been successful, however, these technologies have always been applied to environments where there is sufficient prior knowledge regarding natural product diversity. In this study, we have highlighted the untapped potential of Australian

marine sponges, in particular that of the genus *Scopalina* sp., for the discovery of natural products. Furthermore, we have done so using a novel TEFAP approach that allows for the screening of multiple environments simultaneously. Although, it is clear from comparisons with mWGS data that the TEFAP approach has some bias regarding the relative composition of particular sequences. The TEFAP method is superior to other amplicon-dependent methods, in that the generation of substantially larger numbers of sequences accounts for this intrinsic bias, while providing sufficient information for the targeted recovery of large metagenomic regions from cloned genomic libraries. In this study, we have highlighted how such an approach can reveal, and provide access to, the biosynthetic potential of previously uncharacterised environments. Following from this, it is envisaged that this approach can be applied to multiple different ecosystems in order to present a picture of how these environments select for the presence of organisms producing natural products.

Conflict of Interest

The authors declare no conflict of interest.

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Author Contributions

JNW, MVB and BAN designed research; JNW and LF performed research; JNW, MVB, LF, TT and BAN analysed data; and JNW, MVB, TT and BAN wrote the manuscript.

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