

## ORIGINAL ARTICLE

# Assessing the complex sponge microbiota: core, variable and species-specific bacterial communities in marine sponges

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**Marine sponges are well known for their associations with highly diverse, yet very specific and often highly similar microbiota. The aim of this study was to identify potential bacterial sub-populations in relation to sponge phylogeny and sampling sites and to define the core bacterial community. 16S ribosomal RNA gene amplicon pyrosequencing was applied to 32 sponge species from eight locations around the world's oceans, thereby generating 2567 operational taxonomic units (OTUs at the 97% sequence similarity level) in total and up to 364 different OTUs per sponge species. The taxonomic richness detected in this study comprised 25 bacterial phyla with *Proteobacteria*, *Chloroflexi* and *Poribacteria* being most diverse in sponges. Among these phyla were nine candidate phyla, six of them found for the first time in sponges. Similarity comparison of bacterial communities revealed no correlation with host phylogeny but a tropical sub-population in that tropical sponges have more similar bacterial communities to each other than to subtropical sponges. A minimal core bacterial community consisting of very few OTUs (97%, 95% and 90%) was found. These microbes have a global distribution and are probably acquired via environmental transmission. In contrast, a large species-specific bacterial community was detected, which is represented by OTUs present in only a single sponge species. The species-specific bacterial community is probably mainly vertically transmitted. It is proposed that different sponges contain different bacterial species, however, these bacteria are still closely related to each other explaining the observed similarity of bacterial communities in sponges in this and previous studies. This global analysis represents the most comprehensive study of bacterial symbionts in sponges to date and provides novel insights into the complex structure of these unique associations.**

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## Introduction

The recent advent of massively parallel sequencing technologies has revolutionized microbial diversity and ecology studies. Deep sequencing of 16S ribosomal RNA (rRNA) gene amplicon libraries showed that microbial communities in many environments including marine habitats, soil, plants and humans, are much more diverse than previously thought (Huber *et al.*, 2007; Costello *et al.*, 2009; Turnbaugh *et al.*, 2009; Andersson *et al.*, 2010; Hollister *et al.*, 2010; Teixeira *et al.*, 2010). Many of these newly detected microbes represent the rare biosphere (Sogin *et al.*, 2006). They are not abundant but collectively represent most of the diversity within a sample. A recent 454 pyrosequencing study showed that marine sponges host bacterial communities with a diversity that is unparalleled in any invertebrate host (Webster *et al.*, 2010).

Sponges are the oldest and most primitive of the metazoan phyla with a global distribution in essentially all aquatic habitats. They are ecologically important and a rich source of novel, biotechnologically relevant natural products. The association of some species with dense and complex microbial consortia has long been recognized (Vacelet 1975; Vacelet and Donadey 1977; Wilkinson *et al.*, 1981) and these sponges were termed bacteriosponges or 'high microbial abundance' sponges to distinguish them from co-occurring sponges in the same habitat that lack dense microbial communities (which were thus termed 'low microbial abundance' sponges) (Reiswig 1981; Hentschel *et al.*, 2003). In high microbial abundance sponges, microbes can comprise as much as 40% of the sponge biomass and represent a variety of different morphotypes (Vacelet 1975). Phylogenetic studies identified members of 26 different bacterial phyla (Taylor *et al.*, 2007; Webster *et al.*, 2008, 2010; Sipkema *et al.*, 2009) including the candidate phylum '*Poribacteria*' that is almost exclusively found within sponges (Fieseler *et al.*, 2004; Lafi *et al.*, 2009). Sponge symbionts are capable of diverse metabolic processes such as nitrification, nitrogen fixation, sulfate reduction and photosynthesis (Wilkinson 1979; Hoffmann *et al.*, 2006, 2009; Bayer *et al.*, 2008; Mohamed *et al.*, 2010) and probably contribute to sponge nutrition (Weisz *et al.*, 2007). Additionally, certain sponge symbionts produce secondary metabolites that might be involved in the chemical defense of their hosts (Kennedy *et al.*, 2007; Siegl and Hentschel 2010; Thomas *et al.*, 2010a). Genomic information is currently being used to further characterize features of symbiosis and physiological properties and the lifestyle of sponge symbionts (Hochmuth *et al.*, 2010; Thomas *et al.*, 2010b; Siegl *et al.*, 2011).

A recurring phenomenon in sponge microbiology is the high similarity among sponge-associated bacterial communities. This was first described by Hentschel *et al.*, (2002) who showed that sponge-derived 16S rRNA gene sequences cluster together regardless of

their origin (host sponge and/or sampling location). The term 'sponge-specific' was introduced for these clusters to describe a phylogenetically complex community that is repeatedly detected in sponges around the world but that is different from microbial seawater communities (Hentschel *et al.*, 2002; Taylor *et al.*, 2007). Interestingly, sponge-specific microbes are mainly found in high microbial abundance sponges whereas low microbial abundance sponges usually have a much lower microbial diversity and a different microbial profile (Hentschel *et al.*, 2006; Kamke *et al.*, 2010). Given the observed uniformity of sponge-specific communities, the aim of this study was to identify geographical or host-dependent bacterial subpopulations in sponges and to define the bacterial core community that is shared among most sponges as well as the species-specific community that is unique to a certain sponge species.

## Materials and methods

### *Sponge collection, sample preparation and 454 pyrosequencing*

Tissue samples from three individuals of each of 31 sponge species and from a single individual of an unidentified sponge S1 (Table 1) were collected at eight different locations in the Indian and Pacific Ocean, the Caribbean, Mediterranean and Red Sea at depths between 1.5 and 18 m while SCUBA diving (for sampling details see Supplementary Table S1). All samples were frozen at  $-80^{\circ}\text{C}$  for at least 24 h, then freeze-dried for another 24 h and kept at  $-20^{\circ}\text{C}$  until further processing. DNA was extracted using a hexadecyltrimethylammonium bromide (CTAB)-based protocol (Taylor *et al.*, 2004) from each individual of all sponges except sponge S1 where three different tissue samples of one individual were used. Briefly, cells were disrupted by bead-beating in an ammonium acetate extraction buffer containing chloroform:isoamyl alcohol (24:1). DNA was precipitated with 3 M sodium acetate and isopropanol, then washed in 70% ethanol, dried and redissolved in  $\text{H}_2\text{O}_{\text{dd}}$ . Purity and quantity of DNA was determined with a Nanodrop 1000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA) and only high quality DNA was used for subsequent PCR reactions. PCRs were performed using the FastStart high fidelity PCR system (Roche Diagnostics N.Z. Ltd, Auckland, New Zealand) and the modified primer pair 338f and 533r (338f<sub>deg</sub>: 5'-ACW CCT ACG GGW GGC WGC AG-3', 533r<sub>deg</sub>: 5'-TKA CCG CRG CTG CTG GCA C-3') to amplify a ca 145-bp fragment of the 16S rRNA gene including the hypervariable V3 region. The degenerate primers 338f<sub>deg</sub> and 533r<sub>deg</sub> were complemented with adaptors A and B, respectively, as recommended by Roche, and the forward primer contained one of the three multiplex identifier barcodes CATG, CGAT or CTGA. For the downstream pyrosequencing step, two samples were combined for sequencing per plate region (that is, 32 samples on 16 regions) hence two different multiplex identifiers were used per plate

**Table 1** Sponge samples and sequence data summary

Sampling location	Sponge sample	Sponge order	Seqs after RDP <sup>a</sup>	Seqs after LUCY <sup>b</sup>	No. of unique seqs	No. of 97% OTUs
ANZ	<i>Ancorina alata</i>	Astrophorida	3730	3327	736	352
	<i>Polymastia</i> sp.	Hadromerida	1294	1276	127	45
	sponge S1	unidentified	1063	990	293	163
	<i>Stelletta maori</i>	Astrophorida	1850	1533	459	227
CAR	<i>Aplysina archeri</i>	Verongida	790	746	245	159
	<i>Ircinia felix</i>	Dictyoceratida	1359	1269	323	179
	<i>Pseudoceratina crassa</i>	Verongida	820	743	280	176
	<i>Xestospongia muta</i>	Haplosclerida	1509	1395	440	243
GBR	<i>Cymbastela coralliophila</i>	Halichondrida	282	259	71	38
	<i>Hyrtios</i> sp.	Dictyoceratida	1429	1347	304	162
	<i>Ircinia gigantea</i>	Dictyoceratida	377	365	108	86
	<i>Xestospongia testudinaria</i>	Haplosclerida	3166	2751	679	364
GUAM	<i>Hyrtios altum</i>	Dictyoceratida	2164	2056	295	157
	<i>Pseudoceratina</i> sp.	Verongida	618	575	209	141
	<i>Rhabdastrella globostellata</i>	Astrophorida	1925	1779	490	286
	<i>Stylissa massa</i>	Halichondrida	86	78	37	25
	<i>Xestospongia</i> aff. <i>carbonaria</i>	Haplosclerida	201	162	78	36
IND	<i>Chondrilla australiensis</i>	Chondrosida	106	96	60	42
	<i>Hippospongia</i> sp.	Dictyoceratida	1753	1635	377	188
	<i>Ircinia</i> sp.	Dictyoceratida	1211	1179	319	159
MED	<i>Aplysina aerophoba</i>	Verongida	868	831	229	133
	<i>Aplysina cavernicola</i>	Verongida	1006	908	188	119
	<i>Ircinia variabilis</i>	Dictyoceratida	741	709	178	111
	<i>Petrosia ficiformis</i>	Haplosclerida	660	595	225	142
	<i>Pseudocortidium jarrei</i>	Homosclerophorida	269	255	60	44
RS	<i>Biemna ehrenbergi</i>	Poecilosclerida	704	648	176	111
	<i>Hyrtios erectus</i>	Dictyoceratida	806	756	198	124
	<i>Theonella swinhoei</i>	Lithistida	429	373	189	131
WNZ	<i>Ancorina</i> sp.	Astrophorida	787	748	308	190
	<i>Plakina trilopha</i>	Homosclerophorida	856	823	224	109
	<i>Stelletta aremaria</i>	Astrophorida	1916	1852	440	250
	<i>Xestospongia</i> sp.	Haplosclerida	166	149	64	55

Abbreviations: ANZ, Auckland, New Zealand; CAR, Caribbean Sea; GBR, Great Barrier Reef; GUAM, Guam, Pacific Ocean; IND, Indian Ocean; MED, Mediterranean Sea; OTU, operational taxonomic unit; RDP, Ribosomal Database Project; RS, Red Sea; WNZ, Wellington, New Zealand.

<sup>a</sup>No. of tag sequences after initial quality control using the pyrosequencing pipeline in the Ribosomal Database Project (RDP).

<sup>b</sup>No. of tag sequences after end-trimming using the software LUCY.

region. PCR conditions were as follows: initial denaturing step at 94 °C for 3 min, 30 cycles of denaturing at 94 °C for 30 s, primer annealing at 57 °C for 45 s and extension at 72 °C for 20 s, followed by a final extension step at 72 °C for 5 min. Product quantity was assessed using a Nanodrop 1000 spectrophotometer and product purity was determined with Agilent 1200 Bioanalyzer DNA 1000 chips (Agilent technologies, Santa Clara, CA, USA). Finally, equal amounts of PCR products were pooled from all three individuals per species and the three tissue parts of sponge S1. Amplicon libraries were sequenced with a 454 Life Sciences FLX pyrosequencer (University of Otago, Dunedin, New Zealand). Pyrotag sequence data were deposited in the National Center for Biotechnology Information Sequence Read Archive under accession number SRP003545.

#### Tag sequence analyses

The Ribosomal Database Project pyrosequencing pipeline (<http://pyro.cme.msu.edu/>) was used to

sort tag sequences according to the multiplex identifiers and to remove low quality sequences (all sequences with ambiguous nucleotides and/or without an identical forward primer and/or a length shorter than 125 bp). To avoid overestimating the true diversity because of erroneous tag reads as a result of sequencing errors and formation of homopolymers and chimeras (Reeder and Knight 2009), we followed the approach of Kunin *et al.* (2010) who used quality score-based filtering and a low operational taxonomic unit (OTU) threshold clustering. Our tag sequences were end-trimmed based on quality scores at a stringency of 0.2% per base error probability using LUCY (Chou and Holmes 2001). Unique sequences were identified with Mothur 1.9.0 (Schloss *et al.*, 2009), aligned against a SILVA alignment (available at [http://www.mothur.org/wiki/Alignment\\_database](http://www.mothur.org/wiki/Alignment_database)) using a kmer search and a Needleman algorithm, and then grouped into 90%, 95% and 97% OTUs based on uncorrected pairwise distance matrices (furthest neighbor algorithm). The last steps were performed separately for all tag

sequences from each sponge species, for all tag sequences from each of the eight locations, and for all tag sequences combined. A representative sequence (defined as implemented in Mothur) of each OTU was used for the taxonomic assignment using customized perl scripts similar to the approach used by Sogin *et al.*, (2006) and Webster *et al.*, (2010). For each tag sequence, a BLAST search (Altschul *et al.*, 1990) was performed against a manually modified SILVA version 98 database. Pairwise global alignments were performed between each of the 10 best hits against the tag sequence using a Smith–Waterman algorithm. The most similar sequence to the tag sequence (or multiple sequences if within a range of 0.1% sequence divergence) was then used for assignment according to the Ribosomal Database Project taxonomy implemented in the SILVA database. For assignment at phylum, class, order, family and genus level, sequence similarity thresholds of 75%, 80%, 85%, 90% and 95% were applied, following Webster *et al.*, (2010). In cases where the taxonomy of the most similar sequences was inconsistent, a majority rule was applied and the tag was only assigned if at least 60% of all reference sequences shared the same taxonomic annotation at the respective taxonomic level. All previously published, sponge-derived sequences in the SILVA reference database were labeled as such and it was noted when a tag sequence was assigned to a sponge-derived sequence. On the basis of the taxonomic assignment, Bray–Curtis similarities were calculated using the program PRIMER 6 (Primer-E Ltd, Plymouth, UK) and visualized as heatmaps using JColorGrid (Joachimiak *et al.*, 2006). Bray–Curtis similarities were also used for unweighted pair-group average cluster analyses with Primer-6.

## Results and Discussion

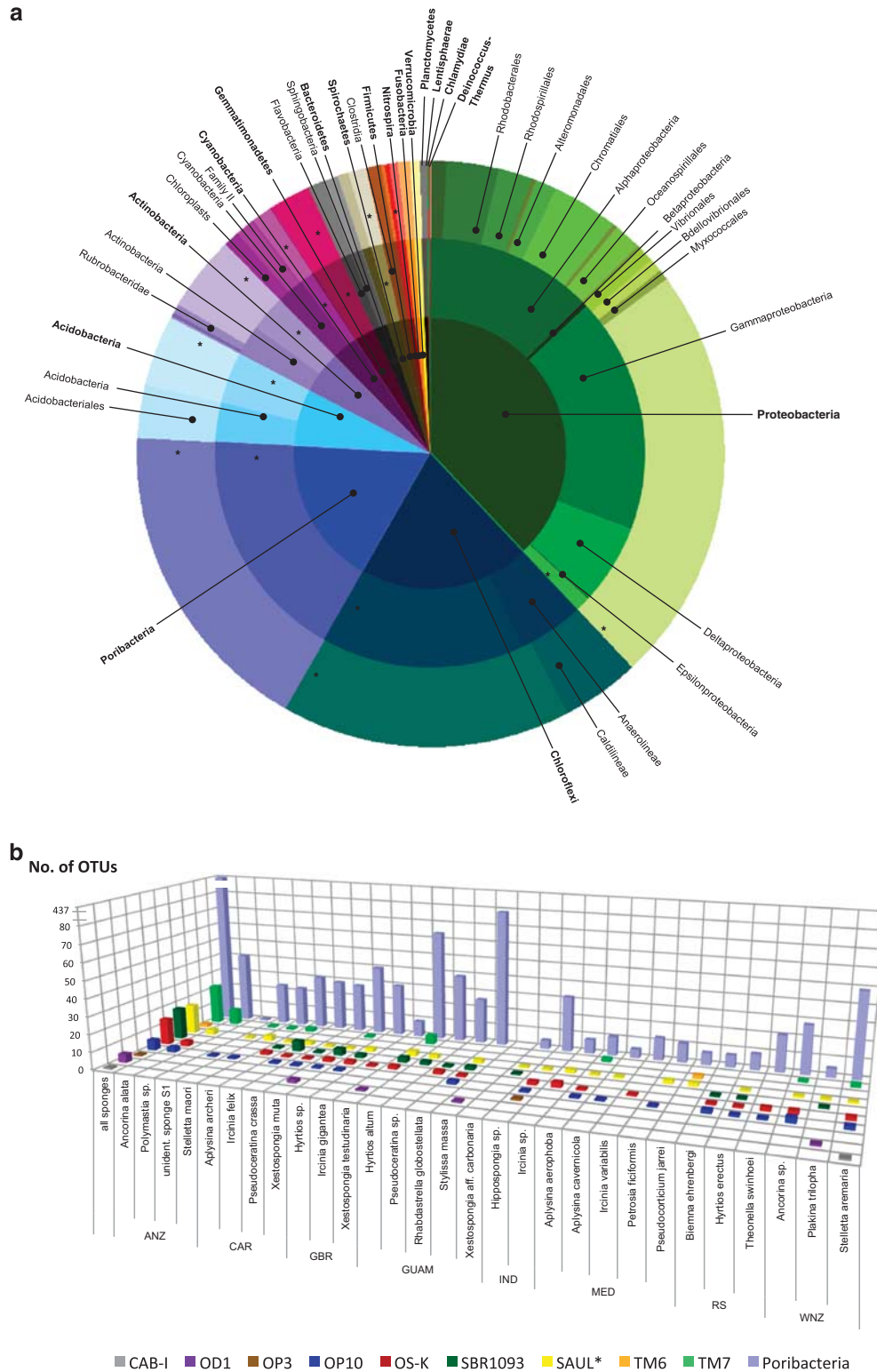
### *Taxonomic richness of the sponge microbiota*

Overall, 2567 different 97% OTUs were obtained from the 32 sponges and up to 364 different 97% OTUs from a single sponge species (Table 1). At phylum level, all 97% OTUs could be classified and belonged to 16 formally described bacterial phyla, 9 candidate phyla and 1 unclassified lineage (sponge-associated unclassified lineage (SAUL)) (Figure 1). Hence, the overall known diversity in sponges—from this and previous studies—increases to 32 different bacterial phyla and candidate phyla, which is similar to the reported number of 35 phyla found in other marine habitats such as the Arctic Ocean and the Western English Channel (Gilbert *et al.*, 2009; Galand *et al.*, 2009b). Given the sequencing depth of next-generation sequencing technologies and the large number of different sponges used in this study, we may now be approaching the total bacterial diversity at phylum level. However, it should be noted that detection of

novel phyla also depends on the tag assignment methods and the taxonomy implemented in the reference database. This is particularly relevant for candidate phyla, which are often inadequately labeled in those databases (see below).

The most diverse phyla among those detected in this study are *Proteobacteria* with 942 different 97% OTUs distributed across all five classes (*Alpha*, *Beta*, *Gamma*, *Delta* and *Epsilon*), and *Chloroflexi* with 502 different 97% OTUs (Figure 1a). This is consistent with data from sponge-derived 16S rRNA gene libraries where both of these phyla are often represented by many different phylotypes (Hentschel *et al.*, 2002; Taylor *et al.*, 2007; Lee *et al.*, 2009). The third most diverse group in this study is the *Poribacteria*, a candidate phylum that was so far barely detected outside of sponges (but see Mohamed *et al.*, 2010; Pham *et al.*, 2008). *Poribacteria* are known to be abundant in and widespread among sponges (Fieseler *et al.*, 2004; Lafi *et al.*, 2009). A total of 437 different 97% OTUs in the whole data set and up to 79 different 97% OTUs in one host species shows that *Poribacteria* are also extremely diverse. Other detected phyla that are well known from sponges include *Acidobacteria*, *Actinobacteria*, *Cyanobacteria*, *Gemmatimonadetes* and *Bacteroidetes*, each represented by 42–169 different 97% OTUs. Also noteworthy is the phylum *Fusobacteria* as it represents the only bacterial phylum that was found exclusively in one species, the New Zealand sponge *Stelletta maori*. Members of *Fusobacteria* were recently found to be associated with diseased corals (Thurber *et al.*, 2009) and it is conceivable that one or all of the sampled *S. maori* individuals was unhealthy although visually this did not appear to be the case. Alternatively, members of *Fusobacteria* may be specifically associated with *S. maori* alone.

In addition to *Poribacteria*, eight more candidate phyla were detected in the sponge data set (Figure 1b). The term ‘candidate division’ was previously created for phyla that lack cultivated members and are typically known only from 16S rRNA gene sequence data (Hugenholtz *et al.*, 1998). Before our tag sequence assignment, the taxonomic annotation of various candidate phyla in the reference SILVA database was manually corrected and sequence similarities to tag sequences were in many cases above 90%. Where sequence similarity was lower, for example, 80% in the case of the OP3-related 97% OTU, the taxonomic status was further verified by additional sequence analysis (for example, further BLAST searches). Thus, the detection of these phyla is not due to ambiguous taxonomic assignment of tag sequences, which sometimes can be problematic because of their short length (Webster *et al.*, 2010). Members of all of these eight candidate phyla were found before in other marine habitats such as seawater, marine sediments or hydrothermal vents (Stott *et al.*, 2008; Harrison *et al.*, 2009; Portillo and Gonzalez 2009) but with the



**Figure 1** Bacterial richness in sponges. **(a)** Phylogenetic distribution of 97% OTUs on phylum (inner circle), class (middle circle) and order level (outer circle). Only taxonomically described phyla and the candidate phylum *Poribacteria* are shown. Phylum names are given in bold. Selected class and order groups are labeled. \* represents unclassified groups. SAUL\* indicates a sponge-associated unclassified lineage. **(b)** Taxonomic richness of 97% OTUs that were assigned to candidate phyla, for all sponges and for each sponge species. \* indicates a sponge-associated unclassified lineage (SAUL). ANZ, Auckland, New Zealand; CAR, Caribbean Sea; GBR, Great Barrier Reef; GUAM, Guam, Pacific Ocean; IND, Indian Ocean; MED, Mediterranean Sea; RS, Red Sea; WNZ, Wellington, New Zealand.

exception of TM6 and TM7, were not previously known to inhabit sponges. Among the additionally detected candidate phyla, TM7 was most diverse with 23 recorded 97% OTUs and was previously found to be present and vertically transmitted in the sponge *Xestospongia muta* (Schmitt *et al.*, 2008). The phyla OS-K and OP10 were most widely distributed (in 19 and 17 sponge species, respectively, from all eight sampling locations) whereas SBR1093 was most diverse within a single host species, with six 97% OTUs found in *Ircinia felix*. However, most candidate phyla were represented by only one or two 97% OTUs per sponge species. It seems that these candidate phyla are generally not diverse but widespread among sponges. The SAUL lineage was found before in sponges (Taylor *et al.*, 2007; Kamke *et al.*, 2010) and, according to previous phylogenetic analyses, belongs to the *Planctomycetes–Verrucomicrobia–Chlamydiae* (PVC) superphylum (Wagner and Horn 2006). However, it remains unclear whether the SAUL lineage falls within one of the phyla in this group or whether it represents a novel candidate phylum alongside *Poribacteria* and WS3 in the *Planctomycetes–Verrucomicrobia–Chlamydiae* superphylum.

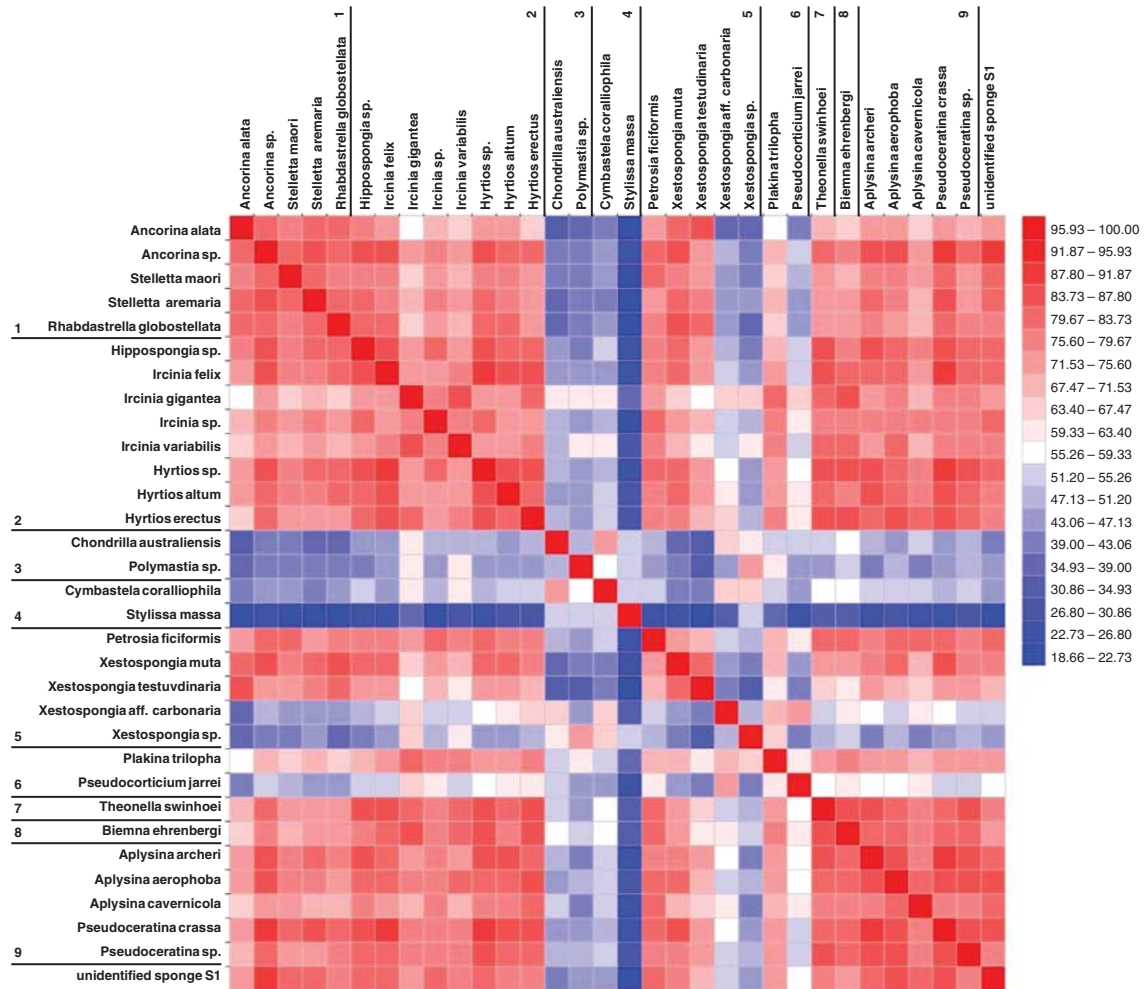
Analysis of lower taxonomic levels revealed several interesting bacterial groups in the sponge data set (Figure 1a). For example, the genera *Nitrosospora*, *Nitrospina* and *Nitrospira* all represent nitrifying bacteria. Both ammonia-oxidizing and nitrite-oxidizing bacteria were found in sponges before using specific primers (Bayer *et al.*, 2008; Hoffmann *et al.*, 2009), however, our data suggest that only the genus *Nitrospira* (which contains nitrite oxidizers) is widely distributed in these hosts. Potential sulfate reducers were also detected, for example, 97% OTUs in the genus *Desulfobrevium* and the family *Desulfobulbaceae*, confirming previous data on sulfate reduction in other sponges (Hoffmann *et al.*, 2005). However, with few exceptions (for example, nitrifying microbes), 16S rRNA gene data can generally not be used to make inferences about metabolic capabilities of microbes. For sulfate reducers, additional physiological experiments would be needed. Included among the detected bacterial groups of this study are also well-known producers of bioactive substances. Examples are *Streptosporangineae* within the *Actinobacteria* and *Rhodobacteraceae*, *Pseudoalteromonas* and *Sphingomonadaceae* within the *Proteobacteria*, which highlights the potential of sponges as sources of novel natural products.

#### *Are there bacterial sub-populations defined by host-dependency or geography?*

To determine the distribution and biogeography of sponge symbionts and to test whether host- or geography-dependent sub-populations exist among the sponge symbionts, the 454 data were analyzed in relation to sponge phylogeny and sampling loca-

tions. Figure 2 shows the similarity of microbial communities among the 32 sponge species of this study. With some exceptions, for example, the sponges *Polymastia* sp. and *Cymbastela coralliophila* that both have low microbial abundances (Kamke *et al.*, 2010, J Vacelet, personal observation), the similarity is generally above 60% and often much higher (Figure 2). However, sponges within each one of the nine sponge orders represented in this study do not contain more similar microbial communities to each other than to species from the other orders (Figure 2). In addition, a clear correlation between microbial community similarity and host phylogeny was also not evident in a more detailed cluster analysis using three species of each of the genera *Aplysina*, *Hyrtios* and *Ircinia*, although some clustering according to sponge phylogeny was found at order level (Figure 3). Such a co-speciation pattern, where closely related sponges contain more similar microbial communities compared with distantly related sponges, would be expected if sponge symbionts were strictly transmitted via reproductive stages to the next generation. Vertical transmission of single lineages and of complex microbial communities has been shown for several sponges including some of the species investigated in this study (Usher *et al.*, 2001; Oren *et al.*, 2005; Enticknap *et al.*, 2006; Schmitt *et al.*, 2007; Sharp *et al.*, 2007). However, it was previously proposed that vertical transmission is not the only mechanism of symbiont acquisition and that sponge symbionts may additionally be acquired from the seawater (Taylor *et al.*, 2007; Schmitt *et al.*, 2008; Webster *et al.*, 2010). Such a combination of vertical and horizontal/environmental symbiont transmission would explain the lack of correlation between microbial community similarity and sponge phylogeny seen in this and previous studies.

Biogeographic distribution patterns of sponge symbionts were investigated by combining all tag sequences from each location and then defining and comparing 97% OTUs. Microbial community similarities among different locations were generally high (>75%, Figure 4). Assuming that sponge symbionts occur outside of sponges (as environmental transmission would suggest) and considering the world's ocean currents, it is conceivable that sponges from some locations that are connected by ocean currents share more similar microbial communities than sponges from more isolated collection sites, as has been shown for marine microbes (Galand *et al.*, 2009a, 2010). For example, while the Mediterranean Sea and the Red Sea each represent almost separate bodies of water, the three sampling sites in the South Pacific (Great Barrier Reef and Auckland and Wellington in New Zealand) are connected by the East Australian Current that moves water along the east coast of Australia and causes the mixture of the Tasman Sea between Australia and New Zealand. However, analysis of Bray–Curtis similarity values does not show such a



**Figure 2** Microbial community similarity among 32 sponge species. Heat map displaying Bray–Curtis similarities based on abundances of assigned 97% OTUs at phylum level (75% sequence similarity) is shown. Sponges are listed according to their taxonomy and orders are labeled as follows: 1, *Astrophorida*; 2, *Dictyoceratida*; 3, *Hadromerida*; 4, *Halichondrida*; 5, *Haplosclerida*; 6, *Homosclerophorida*; 7, *Lithistida*; 8, *Poecilosclerida*; 9, *Verongida*.

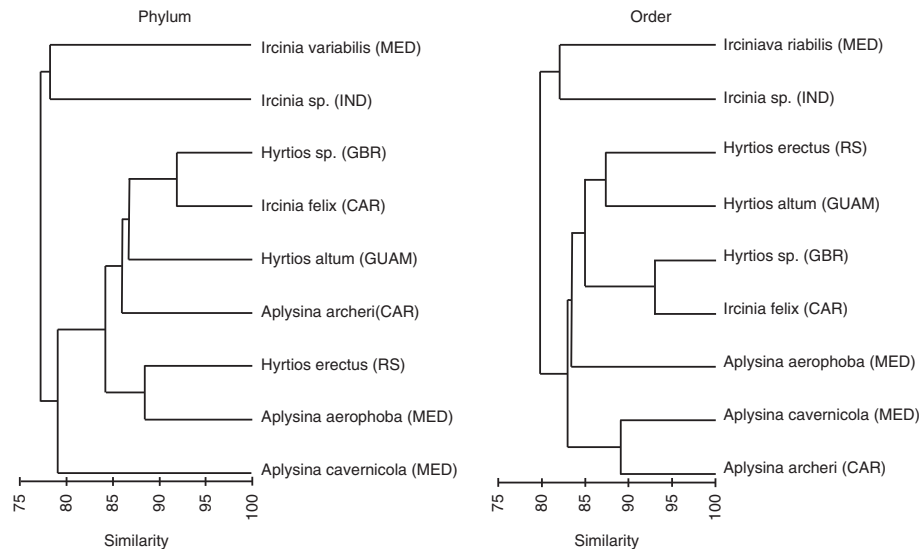
correlation between microbial community similarity and ocean currents. Instead, the Great Barrier Reef samples cluster with those from Guam (Northern Pacific Ocean) and the Caribbean Sea with a high similarity of >90% (Figure 4). Interestingly, these three locations are the only tropical sites whereas all remaining locations are in the subtropics. The finding of such a tropical clade might indicate the existence of bacterial sub-populations among sponge symbionts defined by temperature or water salinity rather than ocean currents.

#### *Core, variable and species-specific bacterial communities in marine sponges*

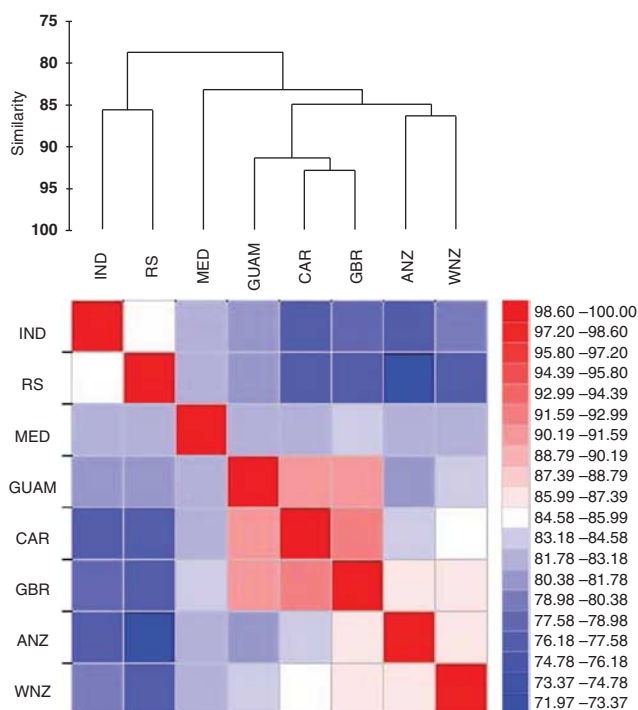
The distribution of OTUs within the sponge samples was investigated by combining all tag sequences, defining 97%, 95% and 90% OTUs, and determining their presence in different sponges. A surprising negative exponential distribution pattern was observed. More than half of all OTUs were present in only a single sponge whereas only very few OTUs

were present in many sponges (Figure 5). On the basis of this result, the microbiota in sponges was divided into the following groups: (1) core community (defined as presence of OTUs in at least 70% ( $N=22$ ) of the analyzed sponges); (2) variable community (defined as presence in <70% of sponges but in at least two species); and (3) species-specific community (defined as presence in only a single sponge species).

All OTUs were also divided into Plus- and Minus-OTUs depending on whether they were assigned—during the taxonomic assignment—to a previously sponge-derived sequence in the database (Plus-OTU) or to a non-sponge-derived sequence (Minus-OTU). This definition was used because a proper phylogenetic tree analysis and determination of sponge-specific clusters according to Hentschel *et al.* (2002) was not feasible with the short tag sequences. Nevertheless, the distinction into Plus- and Minus-OTUs allowed a comparison with previously published, sponge-derived sequences. Figure 6 shows the percentage of Plus-OTUs within



**Figure 3** Microbial community similarity among *Aplysina*, *Hyrtios* and *Ircinia* sponges. Cluster analysis based on Bray–Curtis similarities of assigned 97% OTUs on phylum (75% sequence similarity) and order (80% sequence similarity) level is shown. CAR, Caribbean Sea; GBR, Great Barrier Reef; GUAM, Guam, Pacific Ocean; IND, Indian Ocean; MED, Mediterranean Sea; RS, Red Sea.



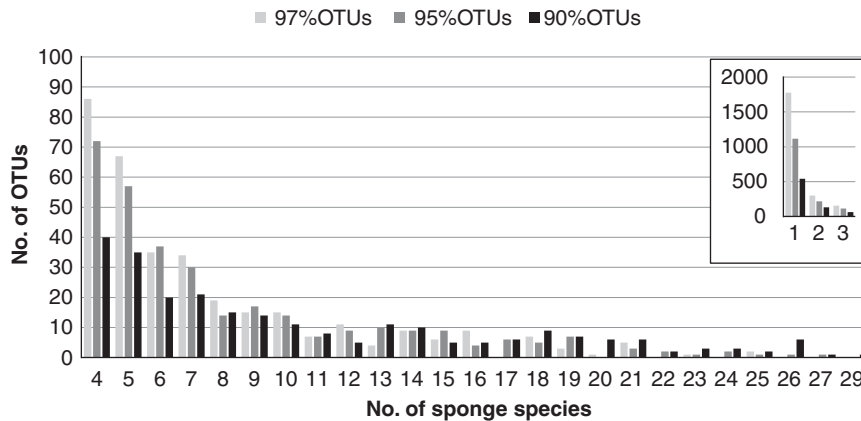
**Figure 4** Microbial community similarity among eight sampling locations. Cluster analysis and heat map displaying Bray–Curtis similarities based on abundances of assigned 97% OTUs at phylum level (75% sequence similarity) is shown. Ninety-seven percent OTUs were defined after combining tag sequences from each location. ANZ, Auckland, New Zealand; CAR, Caribbean Sea; GBR, Great Barrier Reef; GUAM, Guam, Pacific Ocean; IND, Indian Ocean; MED, Mediterranean Sea; RS, Red Sea; WNZ, Wellington, New Zealand.

each of the 32 sponges. An average of 64%, and as much as 80% of all OTUs in a single sponge, were designated as Plus-OTUs showing a large overlap with previously published sponge-derived

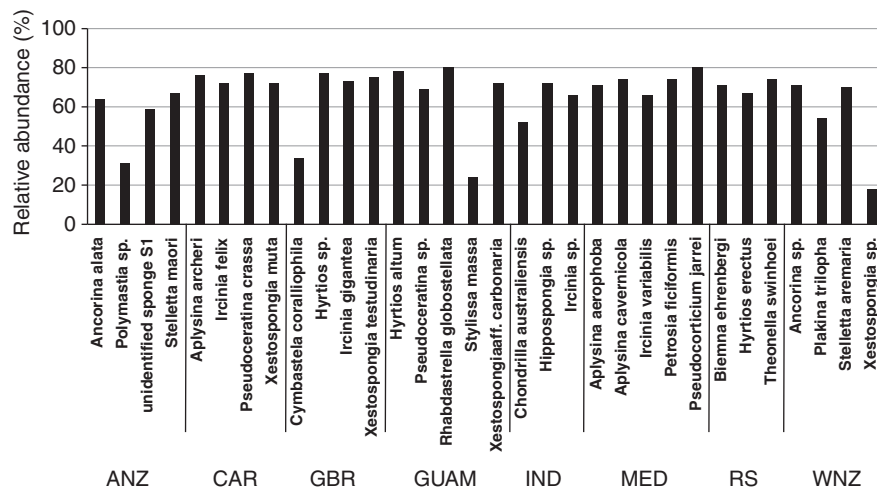
sequences. There are four notable exceptions with much fewer Plus-OTUs: *Polymastia* sp., *Cymbastela coralliophila*, *Stylissa massa* and *Xestospongia* sp. A possible explanation, at least for the first two mentioned sponges, is that they are low microbial abundance sponges that contain less diverse microbiota with a different compositional profile to high microbial abundance sponges (Hentschel *et al.*, 2003; Kamke *et al.*, 2010).

The core bacterial community consists of only three 97%, eight 95% and 18 different 90% OTUs (Table 2). Not a single OTU was found in all 32 species. Therefore, the core bacterial community in sponges is rather small. All three 97% core OTUs and many of the 95% and 90% core OTUs were affiliated with either *Proteobacteria* or *Chloroflexi* but none belonged to *Poribacteria* (Table 2). The fact that all core OTUs were detected in sponges from all eight sampling locations suggests a global distribution of the respective bacteria. Similarly, some free-living bacteria such as the SAR11 clade (*Alphaproteobacteria*), *Alteromonas macleodii* (*Gammaproteobacteria*) or *Prochlorococcus/Synechococcus* (*Cyanobacteria*) are believed to be ubiquitous marine bacteria with a global distribution (Morris *et al.*, 2002; Bouman *et al.*, 2006; Ivars-Martinez *et al.*, 2008a). Often, several ecotypes of these bacteria exist that are adapted to different environmental conditions (Rocap *et al.*, 2003; Ahlgren and Rocap 2006; Ivars-Martinez *et al.*, 2008b). It is conceivable that the core OTUs represent bacterial sponge ecotypes that are adapted to the niche sponge and are probably environmentally transmitted. This is also supported by the finding that most core OTUs represent Plus-OTUs (Table 2). Such a scenario would explain the presence of the same 97% OTU (closely related bacteria) in geographically distinct sponges.





**Figure 5** Distribution of 97 (light grey bar), 95 (dark grey bar) and 90% (black bar) OTUs within 32 sponge species. Note that the inset (presence in 1, 2 or 3 sponge species) has a different bar.



**Figure 6** Relative abundance of Plus-OTUs within each sponge species. Distribution of 97% OTUs that were assigned to a previously published sponge-derived 16S rRNA gene sequence (Plus-OTU). ANZ, Auckland, New Zealand; CAR, Caribbean Sea; GBR, Great Barrier Reef; GUAM, Guam, Pacific Ocean; IND, Indian Ocean; MED, Mediterranean Sea; RS, Red Sea; WNZ, Wellington, New Zealand.

In stark contrast to the core bacterial community, the species-specific community is very large and consists of 70% ( $N=1774$ ) of 97% OTUs and two-thirds ( $N=1116$ ) and half ( $N=542$ ) of 95% and 90% OTUs, respectively (Figure 7). The species-specific bacterial community contains members of all bacterial phyla and candidate phyla detected in this study with the exception of *Deinococcus-Thermus* (which is represented by a single OTU that was found in six species). Generally, the species-specific bacterial community consists of Plus- and Minus-OTUs, however, the proportions of both OTU types within single phyla differ (Figure 7). *Poribacteria* are represented by only Plus-OTUs, which is in agreement with the fact that they were so far almost exclusively found within sponges (but see Mohamed *et al.*, 2010; Pham *et al.*, 2008). *Chloroflexi*, *Acidobacteria* and *Gemmatimonadetes* contain many more Plus- than Minus-OTUs whereas the opposite is true for e.g. *Bacteroidetes* and *Planctomycetes*. The host species-specific bacterial com-

munity in sponges is probably mainly vertically transmitted. This would result in a separation of the symbionts within their hosts and would explain the observation that many 97% OTUs are only present in one sponge but are still most similar to other sponge-derived 16S rRNA gene sequences. Given that some of these bacteria show sequence divergences of  $>10\%$ , such separation must have happened over a long time, pointing to an ancient association with their host sponges.

Figure 8 summarizes the different community types defined for the sponge microbiota and their proportions when applying different OTU definitions. A very similar distribution pattern of core, variable, and species-specific Plus- and Minus-OTUs was observed when only one location (Mediterranean Sea) was investigated (Schmitt *et al.*, 2011). We did not analyze our data quantitatively and therefore cannot make conclusions about relative abundances of certain bacteria in different hosts. If the core microbiota consists also of rare

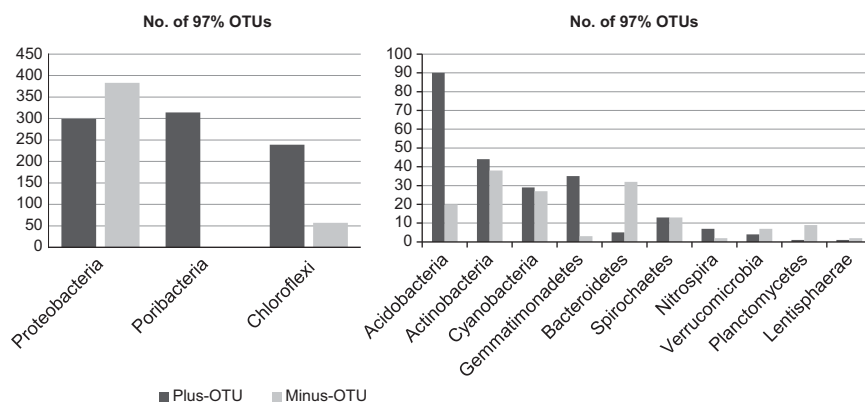
**Table 2** Core bacterial community in sponges (List of 97%, 95% and 90% OTUs present in 22 or more species. All OTUs were found at all eight locations)

Tag ID	N <sub>species</sub>	Score <sup>a</sup>	Taxonomic affiliation	Plus-OTU <sup>b</sup>
<b>97% OTUs</b>				
INDIRC F7F67TR11G0FCT	25	0.963	Proteobacteria (alpha)	NA
CARAPL F7F67TR15I524M	25	0.992	Chloroflexi	Yes
GBRXES F3FI3IU02HOXZ	23	0.985	Chloroflexi	Yes
<b>95% OTUs</b>				
AUKSTE F7F67TR01ALMDZ	27	0.985	Proteobacteria (alpha)	Yes
INDIRC F7F67TR11G0FCT	26	0.992	Proteobacteria (alpha)	NA
MEDAERO F7F67TR12HEAY1	25	0.970	Chloroflexi	Yes
RSTHEO F7F67TR04CBKLL	24	0.992	Actinobacteria	Yes
INDIRC F7F67TR11GR1PV	24	1	Proteobacteria (gamma)	Yes
GBRXES F3FI3IU02HOXZ	23	0.985	Chloroflexi	Yes
INDIRC F7F67TR11GXKZ2	22	0.992	Proteobacteria (alpha)	Yes
GBRXES F7F67TR03BTEAA	22	0.992	Chloroflexi	Yes
<b>90% OTUs</b>				
INDIRC F7F67TR11GYCL4	29	0.993	Nitrospira	Yes
AUKSTE F7F67TR01ALMDZ	27	0.985	Proteobacteria (alpha)	Yes
AUKANC F3FI3IU02JBCBF	26	0.994	Proteobacteria (gamma)	Yes
AUKS1 F7F67TR02A4WJ7	26	0.992	Proteobacteria (alpha)	Yes
AUKANC F7F67TR01ARLEH	26	0.992	Proteobacteria (alpha)	No
RSTHEO F7F67TR04CBKLL	26	0.992	Actinobacteria	Yes
AUKSTE F7F67TR01APO5O	26	0.993	Cyanobacteria	Yes
INDHIP F7F67TR12HE9V4	26	0.993	Chloroflexi	Yes
MEDAERO F7F67TR12HEAY1	25	0.970	Chloroflexi	Yes
AUKANC F3FI3IU02JPC91	25	0.992	Chloroflexi	Yes
GBRXES F3FI3IU02HOXZ	24	0.985	Chloroflexi	Yes
GBRXES F3FI3IU02GYNJB	24	0.977	Chloroflexi	Yes
CARXES F7F67TR16JQN2I	24	0.992	Chloroflexi	Yes
CARIRC F7F67TR15I91FO	23	0.987	Acidobacteria	Yes
GUAMPSEUDO F7F67TR09FO9DJ	23	0.927	Firmicutes	No
AUKANC F7F67TR01AIB3U	23	0.962	Proteobacteria (gamma)	No
INDIRC F7F67TR11GXKZ2	22	0.992	Proteobacteria (alpha)	Yes
MEDPET F7F67TR08ERRYN	22	0.992	Proteobacteria (alpha)	No

Abbreviations: NA, not assigned; OTU, operational taxonomic unit.

<sup>a</sup>Score indicates similarity to reference sequence used for taxonomic assignment.

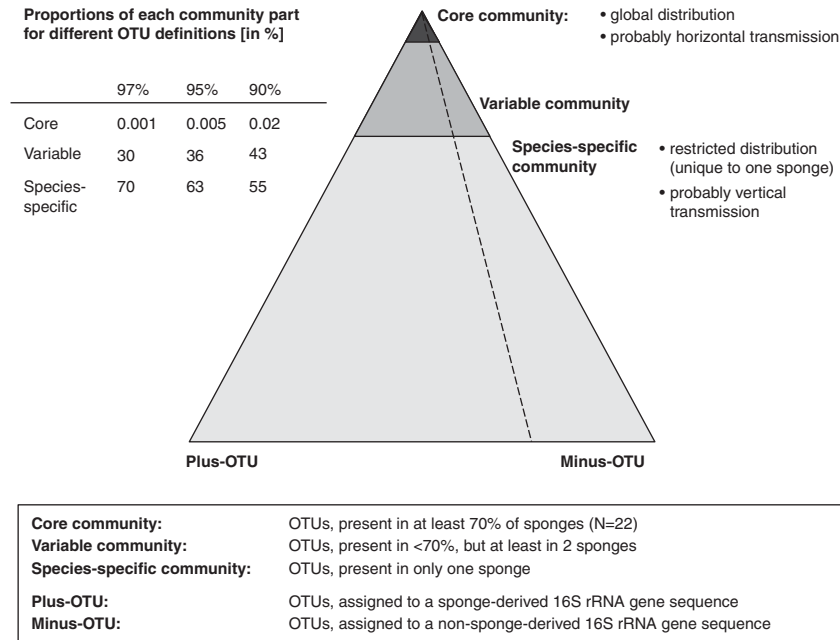
<sup>b</sup>As defined by having a previously sponge-derived 16S rRNA gene sequence as most similar reference sequence.



**Figure 7** The species-specific bacterial community in sponges. Distribution of 97% Plus- and Minus-OTUs among different bacterial phyla. Note that the species-specific community contains members of all phyla and candidate phyla detected in this study, with the exception of *Deinococcus-Thermus*, but only those phyla that contain Plus-OTUs are shown.

members then it is possible that the core is in fact larger than described here because very rare sequences might have been missed with our approach. However, it can safely be said that sponges do not share many abundant microbes. Interestingly, our findings are very similar to the results of the human gut microbiota that also

consists of a small core of shared abundant bacterial species (Hamady and Knight 2009; Tschop *et al.*, 2009; Turnbaugh *et al.*, 2009). However, despite high variability at the species level, there is a core of shared gene families in the human gut microbiome, which suggests that different combinations of species can fulfill the same functional roles



**Figure 8** Schematic representation of the sponge microbiota that was divided into core, variable and species-specific bacterial communities.

(Turnbaugh *et al.*, 2009). It is tempting to speculate that, similar to the human gut microbiota, the microbiota in different sponges vary extensively in the composition of bacterial species but are similar in their functional profiles. A recent metagenomic study provided one of the first detailed functional profiles of the microbial community within a sponge (Thomas *et al.*, 2010b). Similar studies in the future using different sponges will enable testing of this hypothesis.

## Conclusions

Analysis of 16S rRNA gene amplicon pyrosequencing data from 32 sponges around the world's oceans allowed novel insights into the complex composition of sponge-microbe associations. Comparison of bacterial communities at broader taxonomic levels (phylum, order) revealed a high overall similarity but no correlation with sponge phylogeny. However, a tropical, biogeographical clade was identified. Comparison of bacterial communities using different OTU definitions revealed a minimal core and a large host species-specific bacterial community. Finally, comparison of bacterial communities to previously published data from sponges using Plus- and Minus-OTU definitions indicated an overall large overlap with other, previously investigated sponge microbiota. On the basis of these results, we propose the following hypothesis: different sponges contain bacterial communities consisting of mainly different bacterial species (species-specific community) and share very few bacterial species (core community). However, the bacterial species in different sponges are still more closely related to each other than to, for example, seawater bacteria (indicated by Plus-OTUs

and sponge-specific clusters), consistent with previous studies suggesting at least partially overlapping communities among different sponges. Sponges therefore contain a uniform, sponge-specific bacterial community although each sponge species contains different bacterial species.

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