

ORIGINAL ARTICLE

Coral-associated marine fungi form novel lineages and heterogeneous assemblages

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Coral stress tolerance is intricately tied to the animal's association with microbial symbionts. The most well-known of these symbioses is that between corals and their dinoflagellate photobionts (*Symbiodinium* spp.), whose genotype indirectly affects whether a coral can survive cyclical and anthropogenic warming events. Fungi comprise a lesser-known coral symbiotic community whose taxonomy, stability and function is largely un-examined. To assess how fungal communities inside a coral host correlate with water temperature and the genotype of co-occurring *Symbiodinium*, we sampled *Acropora hyacinthus* coral colonies from adjacent natural pools with different water temperatures and *Symbiodinium* identities. Phylogenetic analysis of coral-associated fungal ribosomal DNA amplicons showed a high diversity of Basidiomycetes and Ascomycetes, including several clades separated from known fungal taxa by long and well-supported branches. Community similarity did not correlate with any measured variables, and total fungal community composition was highly variable among *A. hyacinthus* coral colonies. Colonies in the warmer pool contained more phylogenetically diverse fungal communities than the colder pool and contained statistically significant 'indicator' species. Four taxa were present in all coral colonies sampled, and may represent obligate associates. Messenger RNA sequenced from a subset of these same colonies contained an abundance of transcripts involved in metabolism of complex biological molecules. Coincidence between the taxonomic diversity found in the DNA and RNA analysis indicates a metabolically active and diverse resident marine fungal community.

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Introduction

Microbial impacts on a host's physiology, metabolism and fitness are a hallmark of symbiotic microbe–host interactions. A clear example of one such obligate symbiosis is the interaction between corals and their co-evolved photo-symbionts: dinoflagellates in the genus *Symbiodinium*. Although commonly presented as a simple model mutualism, technological advances such as scanning electron microscopy and environmental DNA sequencing have demonstrated additional complexity to this symbiosis involving a wide array of microbial symbionts associated with corals. A high diversity of bacteria, archaea, viruses, algae, protozoa and fungi comprise a complex assemblage that, including the coral animal, has been termed the coral holobiont (Rohwer *et al.*, 2002; Knowlton and Rohwer, 2003). These organisms have been found

to differ from those in the adjacent water column and represent potentially co-evolved symbionts (Rosenberg *et al.*, 2007).

Although the association between corals and fungi (henceforth 'coral fungi') has been reported from a wide host, geographic and climatic range (Freiwald *et al.*, 1997), very little is known about their identity or the nature of their interaction with the holobiont. Coral fungi were long believed to be parasitic to the coral itself (Kendrick *et al.*, 1982) or to endolithic algae within the coral skeleton (Priess *et al.*, 2000). More recent hypotheses portray coral fungi as potentially mutualistic; either protecting the holobiont from infection and disease (Rohwer *et al.*, 2002; Reshef *et al.*, 2006; Shnit Orland and Kushmaro, 2009), or else by cycling recalcitrant nitrogen molecules for uptake by the *Symbiodinium* (Wegley *et al.*, 2007). Alternatively, coral fungi may span a continuum from mutualist to commensalist to parasite depending on environmental context and overall coral health (Le Campion-Alsumard *et al.*, 1995; Bentis *et al.*, 2000; Golubic *et al.*, 2005; Wegley *et al.*, 2007; Lesser *et al.*, 2007a; Thurber *et al.*, 2009).

In contrast to the relationship between coral and fungus, comparatively much is understood about

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interactions between the coral animal and *Symbiodinium*, which passes photosynthetically fixed carbon to its host and greatly accelerates the process of calcification (Muscatine *et al.*, 2005). The majority of *Symbiodinium* are obligately symbiotic, and genetically distinct *Symbiodinium* can confer differing levels of heat tolerance to the coral holobiont (Rowan *et al.*, 1997). For example, in both field observations and laboratory experiments, corals hosting specific members of clade D *Symbiodinium* have been shown to be more resistant to warmer waters and less prone to bleaching than corals hosting common members of clade C (Rowan, 2004; Berkelmans and Van Oppen, 2006). *Acroporid* corals are known to associate with multiple genetically distinct *Symbiodinium* types, including heat-resistant members of clade D (Baker, 2003; Oliver and Palumbi, 2011).

Our objectives here are to investigate the relationship between *Symbiodinium*, the environment and fungi associated with *Acropora hyacinthus* a reef building, tropical coral. First, we assess the phylogenetic diversity of fungi associated with *A. hyacinthus* to detect whether these comprise lineages distinct from terrestrial fungi. While a growing body of literature has found novel lineages of marine fungi in pelagic, sedimentary and vent environments, very few have examined the phylogenetic placement of marine biotrophic fungi, which presumably share many functional and nutritive traits with terrestrial biotrophic relatives.

Second, we assess whether *Symbiodinium* genotype or pool environment correlate with *A. hyacinthus* fungal community composition. Both environment and host identity have been shown to determine community composition in biotrophic fungi on land (Schechter and Bruns, 2008; Smith *et al.*, 2009), even over relatively small spatial and phylogenetic scales (Sthultz *et al.*, 2009). As *Symbiodinium* produces the majority of the holobiont's carbon, and because of presumed algal–fungal interactions (Wegley *et al.*, 2007), we hypothesize that *Symbiodinium* genotype would significantly impact *A. hyacinthus* fungal community composition. We sample replicate coral colonies containing either *Symbiodinium* C or D from two natural pools with different water temperatures in order to assess how these variables impact coral fungi. Since environmental conditions covary with *Symbiodinium* genotype, and because global change is predicted to increase both water temperature and the prevalence of corals with heat tolerant D-clade *Symbiodinium*, we also hypothesize that the water temperature–genotype interaction significantly partitions fungal community variance.

Finally, because there are relatively few macroscopic marine fungi, it is unclear whether the taxonomic diversity of fungal communities detected via culture-independent DNA methods reflects active marine residents or terrestrial debris. To infer whether metabolically active community

membership overlaps that detected via DNA-based techniques, we examine the fungal mRNA transcripts of a subset of colonies under both ambient and warmed conditions. This method has the additional benefit of providing insight into which metabolic pathways are used in symbiosis, enabling us to generate new hypothesis and experiments to determine coral fungal functional roles.

Materials and methods

Sample site and collections

A. hyacinthus coral colonies were sampled from each of two back-reef pools on Ofu Island in American Samoa (14°11'S, 169°36'W). These pools are approximately 500 m apart and are characterized by varying degrees of daily fluctuations in a suite of environmental variables (for example, temperature, flow, salinity) described in detail elsewhere (Smith *et al.*, 2007, 2008; Barshis *et al.*, 2010; Oliver and Palumbi, 2011). The magnitude of the environmental fluctuations correlate with the amount of open ocean admixture, providing variable environmental conditions over short distances, while presumably maintaining comparable access to pools of microbial immigrants.

In the low mixture pool (warm pool) sampled in this study, water temperature fluctuates up to 6 °C daily, and summer water temperature maxima are as much as 1.5 °C warmer than in the adjacent high-mixture pool (cold pool). Owing to differences in thermal tolerance, only *Symbiodinium* D (ITS1, *sensu* Van Oppen *et al.*, 2001) are found in the warm pool, whereas both *Symbiodinium* C and D associate with corals in the cold pool, occasionally forming mixed assemblages within a colony.

For DNA analysis, individual coral fragments (~2 cm³ per fragment) were sampled from 18 colonies of *A. hyacinthus* in each pool (36 total colonies) and stored in 70% ethanol until subsequent nucleic acid extraction.

For mRNA analysis, six individuals were removed from the cold pool and divided into duplicate samples. One duplicate was maintained in a laboratory tank at approximately ambient temperature (26.8–34.5 °C), whereas the other was maintained in a warmer-temperature tank (27–37.6 °C). Colonies were retrieved after 72 h and stored in a high-concentration trisodium citrate buffer until subsequent nucleic acid extraction.

Nucleic acid extractions

Genomic DNA was extracted from colonies preserved in 70% ethanol using a Nucleospin Tissue kit (Clontech, Mountain View, CA, USA) as per the manufacturer's instructions.

Total RNA was extracted using a modified TRIzol (GibcoBRL/Invitrogen, Carlsbad, CA, USA) protocol. Approximately, 150–200 mg of coral tissue and

skeleton was placed in 1 ml of TRIzol and homogenized for 2 min by vortexing with ~100 µl of 0.5 mm Zirconia/Silica Beads (BioSpec Products, Inc., Bartlesville, OK, USA). Resulting tissue/TRIzol slurry was removed by centrifugation and standard TRIzol extraction was performed according to the manufacturer's specifications with the replacement of 250 µl of 100% isopropanol with 250 µl of high salt buffer (0.8 M Na citrate, 1.2 M NaCl) during the final precipitation step. Resulting RNA pellet was resuspended in 12 µl of diethyl pyrocarbonate treated H₂O.

Symbiodinium genotyping

Symbiodinium genotyping was performed by assessing PCR amplicon size polymorphism of chloroplast gene cp23s on an agarose gel using methods as described by Oliver and Palumbi (2011). Briefly, because of indel variants of chloroplast genes among *Symbiodinium* clades C and D, the size of the PCR product can be used to detect the presence of one or both genotypes. Although most colonies in the cold pool presumably contain both *Symbiodinium* genotypes to some degree, quantitative PCR (data not shown) demonstrates that this method reliably detects the genotype if present at >10% proportional abundance.

PCR and 454 sequencing of genomic DNA

Approximately 10 ng DNA was used to amplify ~950 bp of the gene containing the D1 and D2 variable regions of the large-subunit (28S) ribosomal DNA, using oligos containing the LRORf and LR5f primers, the 454 'A' and 'B' adaptors and a 10-bp multiplex tag as in Amend *et al.* (2010). Negative controls were run on PCR reactions. PCR products were cleaned using the QIAquick PCR Purification Kit (Qiagen, Valencia, CA, USA) following the manufacturer's instructions, quantified using a Qubit Fluorometer (Invitrogen), pooled into equimolar concentrations and pyrosequenced at the Duke University ISGP Sequencing Facility on one of eight regions of a plate using the 454 Life Sciences FLX Titanium platform. Raw sequences were deposited in the Short Read Archives of NCBI under accession SRP005168.

Bioinformatic processing of 454 DNA amplicon sequences

Base calls and quality scores derived from the 454 sequencer were 'de-multiplexed', quality-control processed and clustered into operation taxonomic units (OTUs) using the MOTHR v.1.14.0 analytic pipeline (Schloss *et al.*, 2009). Reads were culled if they were shorter than 250 bp, imperfectly matched the priming site or barcode sequences, contained a mean Q-score <30, or contained a homopolymer run longer than 12 bases.

The remaining sequences were queried against the NCBI nr/nt database using BLASTN (Altschul *et al.*, 1997) on the Biportal server at the University of

Oslo. The resulting BLAST output was imported into MEGAN 3.9 and taxonomy was determined by 'last common ancestor' analysis (with parameters as described in Amend *et al.* (2010)). Non-fungal sequences (mostly from corals) were removed. Fungal sequences were aligned in MOTHR, using a published 28S ribosomal DNA alignment (James *et al.*, 2006) as a template. Putative chimeras were screened and removed using the 'Chimera-Slayer' algorithm in MOTHR with default settings.

Phylogenetic analysis of DNA amplicons

Non-singleton OTUs were aligned with a non-redundant set of their top BLAST matches ($N=497$) to the NCBI nr/nt database and the 28S sequences from the phylogeny of James *et al.* (2006) using the local pairwise alignment setting of MAFFT. The resulting alignment contained 529 characters. A maximum likelihood tree was calculated using a web-server implementation of RAXML (v. 7.2.8; Stamatakis *et al.*, 2008) with Gamma rates of nucleotide heterogeneity and the six-gene Fungal Bayesian tree from James *et al.* (2006) as a backbone to constrain the resulting topology (Figure 1). The tree was visualized using the Interactive Tree of Life Program (Letunic and Bork, 2006).

Computing and comparing community similarity

For phylogenetic diversity analyses, all remaining sequences were included in a minimum evolution tree computed using the FastTree program (Price *et al.*, 2009), using the six-gene Bayesian tree as a constraint as above.

OTUs were calculated using MOTHR's pre-cluster option followed by average-linkage clustering. An OTU sequence identity threshold of 97% was used for subsequent analyses. We are unaware of any estimates of the rate of evolution for this locus in fungi, but presume that this cutoff resolves taxa at approximately the genus or family level, rather than at approximately the species level inferred using the same cut-off level with the ITS cistron (Nilsson *et al.*, 2008).

Differences in community composition were tested using the non-parametric ANOSIM analysis (available in the VEGAN package of R; Dixon, 2003) on matrices of the unweighted Unifrac (Phylogenetic Diversity; Lozupone and Knight, 2005) and Jaccard (OTU diversity) distances. For each diversity index calculated, sample sizes were normalized by randomly selecting the number of sequences in the smallest sample from all other samples, computing the distance matrix, and then calculating the mean matrix values from 1000 such random samplings. Distance matrices calculated to include relative abundance of sequencing reads (weighted Unifrac and Bray Curtis index) did not significantly change statistical results and are not discussed further. *Symbiodinium* genotype effects were tested by comparing groups of individuals from the cold

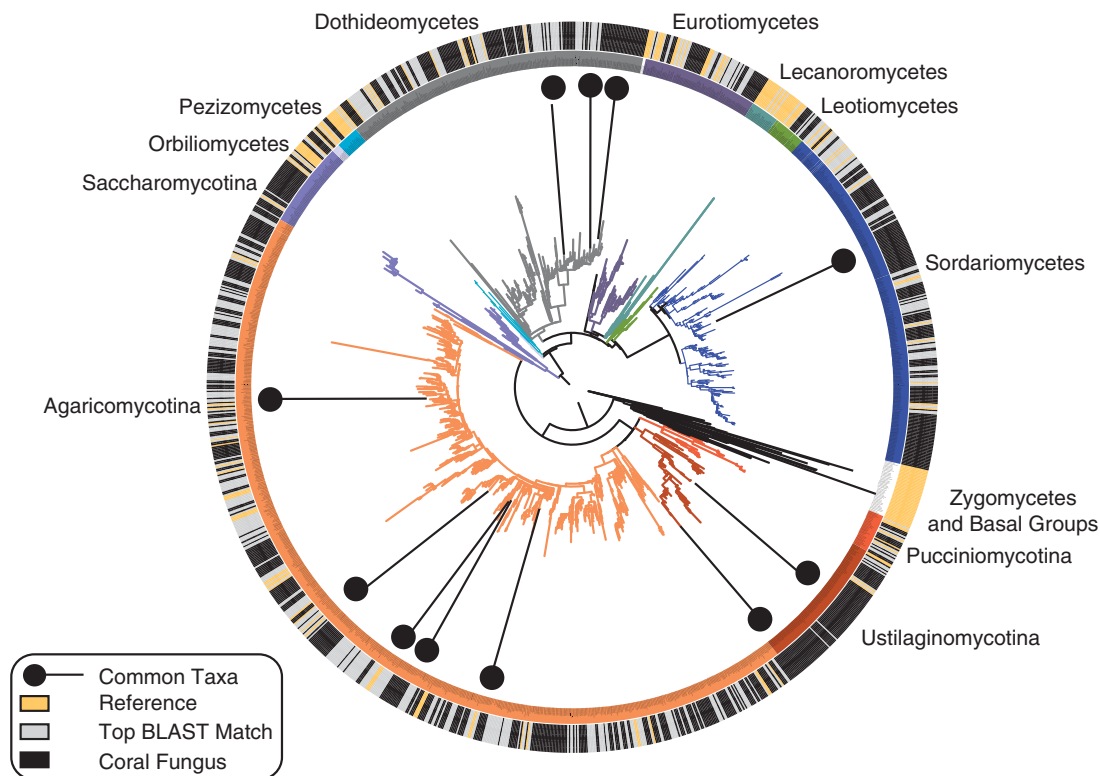


Figure 1 Best of 100 maximum likelihood trees of non-singleton coral-fungi OTUs. The tree used a previously published phylogeny as a constraining reference, and was augmented with 497 sequences, which were the top BLAST matches of OTUs downloaded from NCBI nr/nt database. Superimposed on the phylogeny are color-coded cladistic information (inner ring) and sequence source (outer ring). Basidiomycetes are designated with warm colors, Ascomycetes with cool. Uninterrupted black bars on the outer ring indicate clusters of *A. hyacinthus* fungi, which are more closely related to other coral fungi than to previously sequenced species. The 11 taxa that were found in >90% of the corals are indicated with black pushpins. A larger image with legible labels is provided as supplementary information. Sequences lacking a taxonomic color-code belong to lineages absent in this study.

pool only, as well as by comparing all clade D individuals with all clade C individuals while accounting for variance contributed by pool of origin. Environmental effects were tested by comparing only clade D individuals grouped by pool, as well as by comparing all individuals grouped by pool while accounting for variance contributed by *Symbiodinium* genotype. Genotype–environment interaction effects were tested by grouping individuals by both genotype and pool (clade D warm, clade D cold, clade C cold, mixed genotype cold).

Evaluating fungal OTU distribution patterns

Non-random positive co-occurrence patterns can indicate symbiotic relationships among organisms. To test whether the number of OTUs found in every coral sample differed from what would be expected given random occurrence, we used a randomization procedure in which we maintained sequence totals per sample and sequence totals per OTU (row and column sums of the OTU matrix) while randomizing OTU locations. Observed co-occurrence patterns were compared with those of 100 such randomizations to calculate whether deviations from a randomized null distribution were statistically significant.

To test whether individual OTUs were statistically associated with either pool of origin or *Symbiodinium* genotype, we conducted a species indicator analysis. As low abundance OTUs are likelier to be unique to a grouping variable (such as pool or *Symbiodinium* type) by random chance, this procedure uses randomizations to calculate the probability that an association between an OTU and an environmental variable indicates a statistical deviation from null expectations given that OTU's observed frequency. Indicator values were calculated with the multipatt algorithm in the Indicspecies package (Cáceres and Legendre, 2009) in R, computed with 999 permutations. The procedure was parameterized to consider each grouping variable independently in addition to interactions among grouping variables, while assigning significance to OTUs among all possible permutations of variable combinations.

Illumina sequencing of mRNA

Messenger RNA was isolated from total RNA using the micro fast-track mRNA isolation kit (Invitrogen) and an overnight precipitation at -80°C . Between 40 ng and 1 μg of mRNA was used in Illumina library

construction as in Beck *et al.* (2010). Briefly, mRNA was converted to double-stranded complementary DNA in a PCR reaction containing random hexamer primers, Superscript III Reverse Transcriptase (Invitrogen) and supplied buffer. Reactions were cleaned with the MinElute Reaction Cleanup Kit (Qiagen). Double-stranded, paired-end oligonucleotide adapters were ligated onto the ds-complementary DNA using T4 DNA Ligase (Invitrogen) at 16 °C for 4 h. Resulting libraries were size selected for 200–300-bp fragments using agarose gel electrophoresis and purified using the MinElute Gel Extraction Kit (Qiagen). The final library was generated by PCR amplification of the Linker-ligated complementary DNA using P5 and P7 primers and Phusion PCR Master Mix (New England Biolabs, Ipswich, MA, USA) using the following cycle program: initial denature at 98 °C for 30 s, 15 cycles of 98 °C for 10 s, 65 °C for 30 s, 72 °C for 30 s and a final extension at 72 °C for 5 min.

The sample of one individual's ambient-temperature treatment was not sequenced, thus 11 total libraries were constructed from 6 individuals: 5 from ambient and 6 from heat-stressed duplicates. Seven of these libraries were sequenced by Illumina Corp. with a 76-bp paired-end sequencing length (152 bp per sequence total) and the remaining four libraries were sequenced using single-end sequencing and a length of 36 bp in the lab of Dr Arend Sidow (Stanford University).

Fungal transcriptome assembly and annotation

All 11 lanes of Illumina sequence data were assembled, *de novo*, using CLC Genomics Workbench (v.4, CLC Bio, Mülhltal, Germany) with a mismatch cost of 1, insertion cost of 2, deletion cost of 2, length fraction of 0.27, similarity of 0.8, paired-end minimum distance of 1 and maximum distance of 250, single-end limit of 7, voting conflict resolution, random assignment of nonspecific matches and minimum contig length of 150. All assembled contigs were compared with the NCBI NR protein database using BLASTX. The resulting output was imported into MEGAN 3.9 for taxonomic assignment. BLAST output of putatively fungal transcripts (only transcripts that were assigned by MEGAN/LCA to kingdom Fungi) was imported into Blast2Go (Gotz *et al.*, 2008) for gene mapping and annotation. Matches that mapped at >40% similarity and with E-values < 10⁻⁵ to putative proteins were retained and annotated with general GO-slim categories.

Results

DNA sequencing

Following initial DNA sequence processing, 57 415 reads were retained. In all, 28 083 sequences, nearly half of all reads, derived from the coral animals themselves and were subsequently removed from the data set, leaving 26 022 high-quality fungal sequences.

Phylogenetic placement of A. hyacinthus-associated fungi

A large number of well-supported clades separated from terrestrial taxa by long branch-lengths were recovered within, or sister to, many Orders (Supplementary Figures S1–S3). The average identity of the top alignment of each OTU to a Genbank fungal sequence (including unidentified environmental sequences) was low: 97.39% ± 1.83 (s.d.) aligned over 325.16 ± 36.68 bp, reflecting the novelty of diversity detected here.

A. hyacinthus contained a high diversity of fungi, almost entirely dominated by Ascomycetes and Basidiomycetes (Figure 1). Sordariomycetes and Dothideomycetes (Ascomycota), and Agaricomycetes and Ustilaginomycetes (Basidiomycota) were the most species-rich. Notably absent was the causative agent of coral aspergillosis: *Aspergillus sydowii*, as well as most genera of the polyphyletic 'Halosphaeriales' commonly found on algae and woody debris in marine environments. Although several of the species detected here are closely related to species found in culture-based studies of marine fungi, including *Dendryphiella* sp., *Debaryomyces* sp. and *Lulworthia* sp., the majority of *A. hyacinthus* fungi were considerably divergent from known species, marine or otherwise.

Community similarity patterns

The high number of coral sequences in the libraries resulted in unequal numbers of fungal sequences derived from each sample, thus for comparisons of beta-diversity and fungal OTU distributions, only samples containing >464 fungal sequences were included in the analyses. Of these 19 remaining samples, 7 were from the warm pool (all *Symbiodinium* D) and 12 were from the cold pool (7 *Symbiodinium* C, 3 *Symbiodinium* D and 2 mixed genotype). These remaining samples contained a mean of 1319 sequences ± 1290 (s.d.).

Fungal communities were not statistically differentiated when grouped by *Symbiodinium* genotype, pool of origin (warm or cold), *Symbiodinium* genotype within the cold pool, or pool × genotype interaction. Results of the ANOSIM analysis showed no significant differences in community composition as inferred from either OTU-based or phylogenetic dissimilarity metrics (no *P*-value < 0.2).

The majority (4570 of 5410) of OTUs detected were singletons, suggesting either that *A. hyacinthus*-associated fungal communities are highly diverse and undersampled, or else that a large component of the fungi we detected are transient and not obligately associated with these coral hosts. Whereas singletons comprised a large proportion of OTUs, they made up only 15% of all fungal sequences. Even with singletons removed, the majority of OTUs appear to be restricted to either a single pool and/or *Symbiodinium* genotype, with less than half shared between any two groups (Table 1).

Fungal OTU distribution patterns

A core set of 11 OTUs were detected in >90% of all colonies assayed, 4 of which were present in all corals examined. None of these sequences matched perfectly to known isolates. Phylogenetic analysis demonstrates that five of these taxa are members of the Agaricomycetes, two are contained within the Ustilaginomycetes, three are Dothideomycetes and one is in the Sordariomycetes (Figure 1).

The observed number of core OTUs differed significantly from that expected given the null randomized distribution. Given random co-occurrence patterns between coral samples and fungi, the null distribution predicted more core OTUs: a median expectation of 39 OTUs distributed among 90% of the corals vs 11 observed ($P \leq 0.01$), and an expectation of 34 occurring in all vs 4 observed ($P \leq 0.01$).

Of these core OTUs, those within the Ustilaginomycetes are closely allied to and nested within the Malasseziales (Supplementary Figure S3), a group of fungi generally associated with sebaceous glands of mammalian hosts (Guého *et al.*, 1996). Although other species of *Malassezia* are common on terrestrial animals, this is the first instance of a Cnidarian host reported. Several *Malassezia sensu lato* were also detected in a recent study of marine sponges

Table 1 Shared non-singleton OTUs among grouping variables shows relatively little overlap

Group	#OTUs	#Shared	%Shared
Variable pool	528	348	43.23
Stable pool	625		
<i>Symbiodinium</i> D	625	256 (C and D)	24.38 (C and D)
<i>Symbiodinium</i> C	425	89 (All)	11.80 (All)
Mixed Genotype	219		
Warm D	528	189	30.24
Cold D	286		
Cold D	286	172	31.91
Cold C	425		

Abbreviation: OTU, operation taxonomic unit. Parentheses denote which groups are compared in the adjacent value.

Table 2 Description of statistically significant indicator species

Sequence	Statistic	P-value	Taxonomic group	Best BLAST match	Match ID	% ID
<i>Warm pool clade D</i>						
GLSBCHB02BVQZG	0.884	0.005	Pleosporales	<i>Stagnospora/</i> <i>Alternaria/Edenia</i>	HM216201.1/ HM216200.1/FJ839654.1	98
GLSBCHB02CE29K	0.784	0.044	Sordariomycetes	<i>Verticillium</i>	AY312607.1	99
GLSBCHB02BU5TQ	0.781	0.044	Agaricomycetes	<i>Phlebia</i>	AF141624.1	98
GLSBCHB02BW3T6	0.756	0.009	Pleosporales	<i>Dendriphiella</i>	EU848587.1	99
GLSBCHB02CK398	0.756	0.011	Ustilaginomycotina	<i>Malassezia globosa</i>	AY743604.1	98
GLSBCHB02B3JQH	0.655	0.031	Pleosporales	<i>Westerdykella</i>	GQ203754.1	94
GLSBCHB02CENAB	0.655	0.04	Pleosporales	<i>Bipolaris/Cochliobolus</i>	GU183125.1/GQ328852.1	98
GLSBCHB02B8O23	0.655	0.038	Agaricomycetes	<i>Cerrena</i>	FJ821522.1	99
GLSBCHB02CGFVF	0.655	0.035	Pleosporales	<i>Phaeosphaeriopsis musae</i>	DQ885894.1	99
<i>Clade D and mixed genotype</i>						
GLSBCHB02B1NDB	0.707	0.047	Agaricomycetes	<i>Exidia/Exidiopsis</i>	AY509555.1/AY885162.1	97

(Gao *et al.*, 2008). However, because different molecular markers were used here and in the sponge study, the sequences cannot be compared directly between these two studies. The sequences of the Dothideomycetes fall within a group of fungi related to *Hortaea werneckii*, consisting of so-called 'black yeasts' that thrive in high salt environments and occasionally form rashes in human hosts (Petrovic *et al.*, 2002). The sequence in the Sordariomycetes appears closely related to *Lindra* (species undetermined), a genus in which species have previously been described on marine algae (Roth *et al.*, 1964).

Finally, every coral contained four distinct fungi in the Agaricomycetes. One of these falls within the Marasmiaceae (from which marine fungi have been reported previously (Kis-Papo, 2005)), and the remainder appear nested within the Polyporales, an order of Basidiomycota that is typically saprotrophic and forms multi-cellular fruiting bodies in terrestrial environments.

Although community similarity was not correlated with *Symbiodinium* genotype or pool, the distributions of some individual OTUs were. Indicator species analysis showed several species statistically associated with corals in the warm pool containing *Symbiodinium* clade D, and a single species that was associated with corals containing clade D and mixed genotype *Symbiodinium* populations (Table 2).

Alpha-diversity of A. hyacinthus-associated fungi

Somewhat different patterns emerged from phylogenetic and OTU-based diversity analyses of fungal associates (Figure 2). Rarefied phylogenetic diversity was significantly higher in the warm pool compared with the cold pool (one-tailed *t*-test, $t = 3.87$, $df = 10$, $P = 0.003$). No significant differences in fungal diversity were found among corals grouped by the *Symbiodinium* genotypes. No significant differences were found among means of rarefied OTU richness for any grouping.

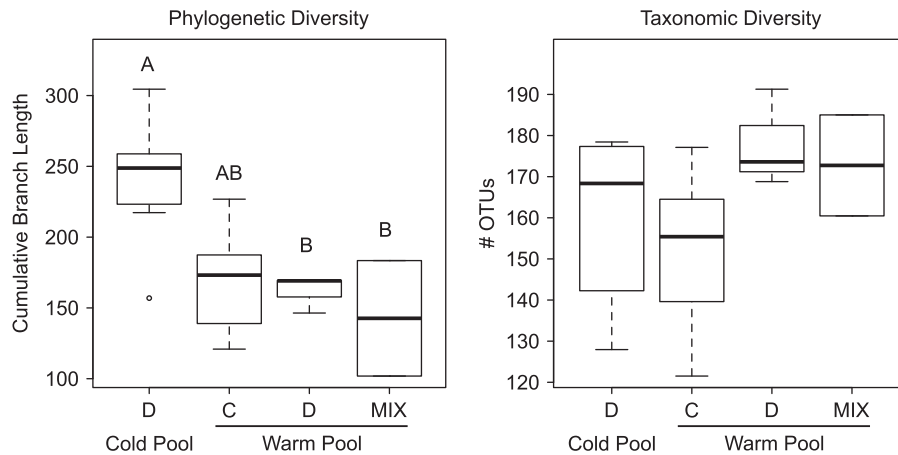


Figure 2 Box and whisker plots of alpha diversity of *A. hyacinthus* fungi communities. Boxes contain the upper and lower quartiles, and the median is displayed as the band within. The outlier (open circle) is > 1.5 times the interquartile range, and whiskers display the extent of non-outlier values. Phylogenetic diversity was significantly higher in the warm pools (*t*-test, $P = 0.003$). Box plots sharing a letter (above) do not significantly differ in a Tukey *post hoc* test of an analysis of variance analysis, although phylogenetic diversity of the fungal communities associated with D clade from the warm pool were marginally higher than those associated with C clade from the cold pool ($P = 0.06$). There were no significant differences in taxonomic diversity among variables.

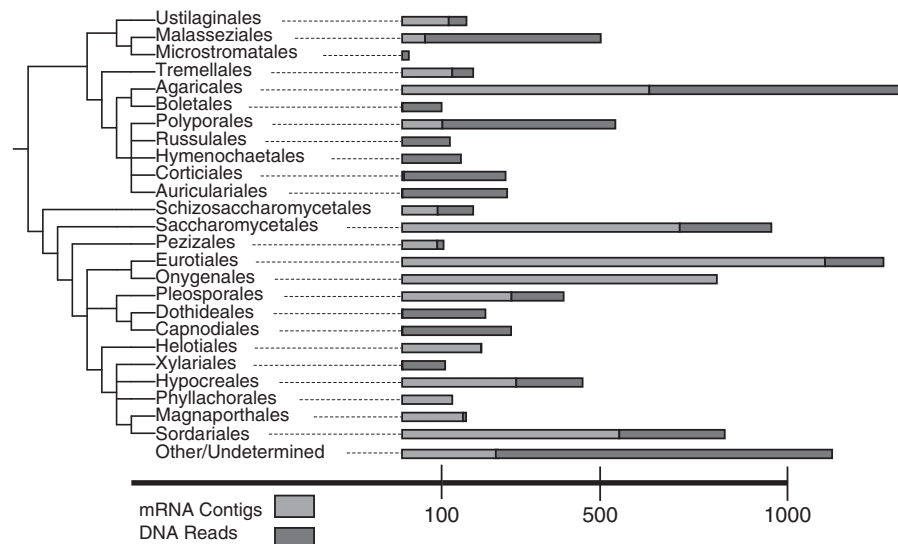


Figure 3 Order-level taxonomic diversity of mRNA contigs is broadly congruent with that of DNA amplicon OTUs for most orders, although several orders were exclusive, or nearly so, to one method over the other. All orders shown here, except the Hymenochaetales, had at least a single draft genome deposited in NCBI Genbank at the time of writing. The abundance shown here is a measure of diversity, neither transcript nor DNA copy numbers were measured directly.

Fungal community transcriptomics

Illumina sequencing resulted in 294 624 883 sequences, which were assembled into 371 363 contigs averaging 283.5 ± 203.4 bp (s.d.) in length, and $136 \times$ coverage. The vast majority were identified as coral, *Symbiodinium* or other non-fungal microbial transcripts. In all, 5736 contigs matched fungal sequences with BLAST matches above our cutoff thresholds ($> 40\%$ similarity and with E -values $< 10^{-5}$). Out of these matches, 4515 contigs were mapped to hypothetical fungal proteins in Genbank. Of these, 530 were annotated within the general GO-slim gene ontology framework (Figure 4).

Taxonomic diversity of mRNA mirrored that of DNA amplicons within the Agaricales, Sordariomycetales and Ustilaginomycetales: the orders—aside from the Dothideales—containing the core fungal associates (Figure 3). Although DNA amplicons revealed a higher diversity within the Agaricomycetes, each method revealed several different orders of Ascomycota.

The majority of GO-slim annotated fungal transcripts related to metabolic processes: notably enzymes involved in metabolism of complex molecules such as proteins, polysaccharides, carbohydrates and lipids. Nitrogen metabolism was

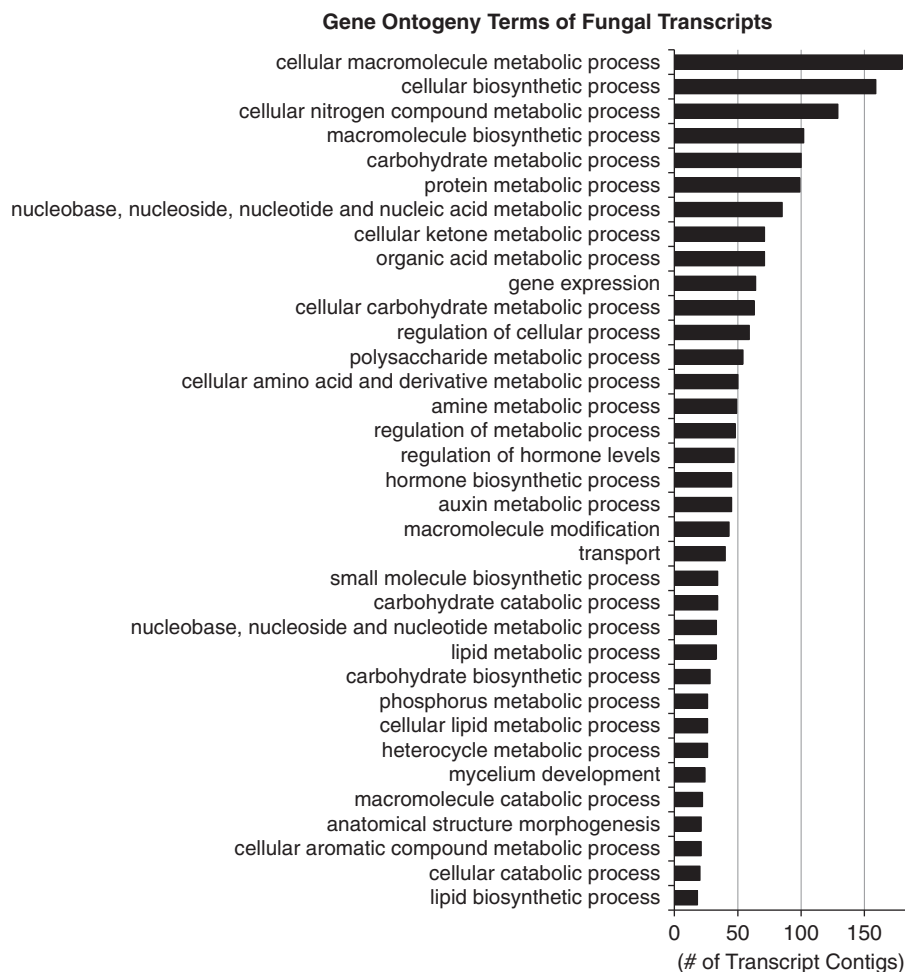


Figure 4 Diversity of generic Gene Ontology (GO) terms to which fungal transcripts were assigned. The top 20 percentile of 173 terms are shown here.

particularly well-represented, and included enzymes involved in nucleic acid, amine and cellular nitrogen compound metabolism. Specific enzymes included fungal ureases, enzymes involved in glutamate and glutamine pathways, nitrate and nitrite reductase, as well as multiple chitinases, peptidases and a relative of β -N-acetylglucosaminidase.

Discussion

Phylogenetic diversity in A. hyacinthus-associated fungal communities

We found several groups in the Agaricomycetes, Ustilaginomycetes, Eurotiomycetes and Dothideomycetes that form separate, well-supported clades with branch lengths suggesting novel lineages (Supplementary Figures S1–S3). Considering the relative paucity of culture-independent fungal studies in marine environments, such novelty reflects a presumed vast amount of yet undescribed fungal diversity. The fact that our average BLAST match

was only 97% similar to the best reference sequence of the commonly sequenced and slowly evolving 28S gene indicates a large degree of divergence from known species.

Phylogenetic analyses also demonstrate marine fungi nested within or allied to lineages observed in terrestrial environments, raising important questions about multiple transitions to or from marine environments and the frequency of amphibious lifestyles. Obtaining axenic cultures of some of these individuals will enable greater phylogenetic resolution and hypothesis testing about these evolutionary trajectories.

In a broad sense, the taxonomy of fungi we encountered in *A. hyacinthus* overlap with a recent culture-independent marine sponge study that detected fungi from seven orders of Ascomycota, including eight well-supported clades that group with sequences derived from other marine environments (Gao *et al.*, 2008). As with our own study, Gao *et al.* (2008) found a high diversity of Basidiomycete sequences within the Malasseziales, Corticiales, Agaricales and Polyporales. In another study of planktonic marine

fungi (Gao *et al.*, 2009), clone libraries were dominated by sequences within the Pucciniomycetes, a group relatively underrepresented here.

Effect of Symbiodinium genotype and water temperature on fungal diversity

Only modest differences in fungal diversity were detected between the warm and cold pools. The warm pool contained a higher phylogenetic, although not taxonomic, diversity. Indicator analysis also demonstrated the exclusive association of some OTUs with the warm pool, which exceeding expectations given a random distribution of taxa. Neither pool of origin nor *Symbiodinium* genotype differentiated overall fungal community structure, and less than half of all OTUs were shared between the pools and *Symbiodinium* genotypes. Similar to patterns described for communities of coral symbiotic bacteria (Rohwer *et al.*, 2001, 2002; Sunagawa *et al.*, 2010), archaea (Wegley *et al.*, 2004) and fungal endophyte communities on tropical leaves (Arnold and Lutzoni, 2007), there appears to be overlap of a few select 'core' species among hosts, whereas the majority of 'satellite' species are either transient or endemic to a specific host and locale.

Several factors may account for the high level of fungal community dissimilarity found within the pools and the *Symbiodinium* genotypes. Prior work has shown that physiology of conspecific corals can vary considerably over short distances (Brown *et al.*, 2002a, b; Smith *et al.*, 2007; Bay *et al.*, 2009; Barshis *et al.*, 2010), which could obscure fungal community patterns associated with our measured variables. Furthermore, stochastic recruitment of large numbers of planktonic, benthic or even terrestrial fungi could mask signatures of specialization or obligacy of a small number of taxa. Finally, because many OTUs were detected in low abundances, it is possible that increased sequencing effort would demonstrate their presence in a greater number of coral samples than observed here. However, given the typical 'long tailed' frequency distribution of the fungal communities, it is unlikely that increased sequencing effort would elucidate additional structure of the total fungal community within this experiment.

Core fungal associates

Eleven fungal OTUs, spanning two phyla and four classes were found in 90% of coral holobionts, and four of these were found in all samples. These fungi are the likeliest candidates for an ecologically significant association and/or coevolution with the *A. hyacinthus* holobiont. Detecting the limits of the geographical distribution and the host phylogenetic breadth of these associates in other natural coral systems will aid in understanding the drivers of coral fungal community composition.

In this study, we detect significantly fewer 'core' OTUs than is implicitly assumed by our null model.

This under-representation of core OTUs may suggest competitive exclusion in fungal interactions, or else that fewer fungi are adapted to extremes in host physiological heterogeneity discussed above.

Activity and function of A. hyacinthus-associated fungi

Data from fungal meta-transcriptomics provide insight into the diversity of the active community members. Concordance between the order-level taxonomic diversity of the mRNA and DNA libraries suggest that a large portion of the fungal diversity detected in DNA-based analyses are metabolically active marine residents, and not just terrestrial debris. The resolution of this comparison is directly proportional with the number of sequenced fungal genomes available to the public, and at this point we identify transcripts from >62% of fungal orders from which we detect DNA.

The role of coral-associated fungi has been described at nearly all points along the parasite-mutualist continuum and may be context dependent. Phylogenetic placement of environmental sequences provides relatively little insight into these roles because trophic status, functionality and even morphology have been found to vary widely among closely related species. For this reason, the fact that many of the fungi detected in our study belong to clades of parasitic fungi, including plant pathogenic fungi such as Pleosporaceae and dermatophytes such as *Malassezia*, may not necessarily indicate parasitism in this particular environmental context. Lichenic symbioses, for example, show evidence of repeated shifts from symbiosis to pathogen (Arnold *et al.*, 2009), and mutation in a single gene has been shown to convert a plant endosymbiotic fungus from a mutualist to a parasite (Takemoto *et al.*, 2006).

Meta-transcriptomic and meta-genomic analyses offer alternative methods for determining the role of coral-fungi, because function may be ascribed independent of presumptions based on relatedness to terrestrial relatives. In a meta-genomic DNA analysis of a *Porites astreoides* coral holobiont (Wegley *et al.*, 2007), for example, the investigators characterized a large proportion of fungal DNA as similar to genes coding for enzymes involved in nitrite and nitrate ammonification, by which the investigators hypothesize fungi convert nitrates and nitrites to NH₄ for uptake by the *Symbiodinium*. Although we did not detect signatures of this metabolic pathway in fungal mRNA, we did detect active transcription of genes related to metabolism of N-containing bio-polymers such as proteins, peptides and chitin. Linking these fungal transcripts to functions within the holobiont may be possible using food web tracer techniques or comparative genomic methods coupled with experimental manipulations of environments and the coral holobiont community. The utility of fungal meta-transcriptomic methods will undoubtedly improve as the number of sequenced fungal genomes increases.

Conclusion

Our analyses revealed several novel clades of fungi that are divergent from known and described isolates. We show that within our study, *Symbiodinium* genotype and water temperature had no discernable impact on the fungal community as a whole, but did impact the distribution patterns of some individual OTUs. We have shown the persistent association of a small, phylogenetically diverse core assemblage of fungi within *A. hyacinthus* hosts from distinct environments and with differing *Symbiodinium* partners. The finding of this core subset in all or most of our colonies suggests a locally stable, though functionally undetermined relationship between specific fungal taxa and the *A. hyacinthus* coral holobiont. Taxonomic and functional annotation of the transcriptome indicates that coral-associated fungi represent a diverse and metabolically active community. There is much to learn about fungi in corals, and we hope that with future work we can help unravel the functional and nutritive relationship between fungi and coral hosts.

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