

ORIGINAL ARTICLE

Microbiome shifts and the inhibition of quorum sensing by Black Band Disease cyanobacteria

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Disruption of the microbiome often correlates with the appearance of disease symptoms in metaorganisms such as corals. In Black Band Disease (BBD), a polymicrobial disease consortium dominated by the filamentous cyanobacterium *Roseofilum reptotaenium* displaces members of the epibiotic microbiome. We examined both normal surface microbiomes and BBD consortia on Caribbean corals and found that the microbiomes of healthy corals were dominated by Gammaproteobacteria, in particular *Halomonas* spp., and were remarkably stable across spatial and temporal scales. In contrast, the microbial community structure in black band consortia was more variable and more diverse. Nevertheless, deep sequencing revealed that members of the disease consortium were present in every sampled surface microbiome of *Montastraea*, *Orbicella* and *Pseudodiploria* corals, regardless of the health status. Within the BBD consortium, we identified lyngbic acid, a cyanobacterial secondary metabolite. It strongly inhibited quorum sensing (QS) in the *Vibrio harveyi* QS reporters. The effects of lyngbic acid on the QS reporters depended on the presence of the CAI-1 receptor CqsS. Lyngbic acid inhibited luminescence in native coral *Vibrio* spp. that also possess the CAI-1-mediated QS. The effects of this naturally occurring QS inhibitor on bacterial regulatory networks potentially contribute to the structuring of the interactions within BBD consortia.

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Introduction

Corals are holobionts, complex metaorganisms that consist of the invertebrate animal host, dinoflagellate endosymbionts and microbial communities associated with them. Coral microbiota have important roles in the health of the holobiont and contribute importantly to nutrient acquisition (Bourne *et al.*, 2009; Bosch and McFall-Ngai, 2011). Displacement of primary commensals with other members of the microbiome or shifts in the abundance of microbiome members have been linked with the appearance of disease signs and polymicrobial diseases (Cardenas *et al.*, 2012; Vezzulli *et al.*, 2013; Roder *et al.*, 2014b).

Black Band Disease (BBD) is the first reported coral disease and the most widely distributed polymicrobial disease of corals, affecting at least 40 species worldwide (Edmunds, 1991; Littler and Littler, 1996; Sussman *et al.*, 2006; Montano *et al.*, 2013). The disease is recognized by the appearance

of a dense, very dark-purple (black) mat that is the visible accumulation of filamentous cyanobacteria. BBD migrates across the surface of the coral at a rate of up to 1 cm per day (Richardson, 1998), leaving behind bare coral skeleton. Microscopic examination has shown that both thick cyanobacterial filaments and smaller unidentified filaments are present in the leading edge of the disease front (Miller *et al.*, 2011). Similar to other cyanobacterial mats, the black band consortium is stratified, with increasing levels of anoxia and sulfide with depth (Glas *et al.*, 2012), effectively smothering and poisoning the coral tissue beneath.

Meta-analysis of published clone library studies of BBD microbial communities revealed that, with few exceptions, microbial species composition of BBD communities did not correlate with the species of the host coral (Miller and Richardson, 2011). Nevertheless, the same cyanobacterial Operational Taxonomic Unit (OTU) (now classified as *Roseofilum reptotaenium*; Frias-Lopez *et al.*, 2003; Casamatta *et al.*, 2012) was present in over 70% of the samples (Miller and Richardson, 2011), indicating that this organism is critical for the appearance or progression of BBD. Similar to other filamentous cyanobacteria, *Roseofilum* can be cultured in unialgal, but not axenic cultures; therefore, it can only be confirmed as a pathogen of BBD with

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modified Koch's postulates, that is, it is part of a polymicrobial consortium and cannot be confirmed as the primary pathogen (Casamatta *et al.*, 2012). In addition to *R. reptotaenium*, three OTUs (Bacteroidetes and Alphaproteobacteria) were present in 13% of the samples, with other OTUs found in <7% of the samples (Miller and Richardson, 2011). Even though sulfate reduction was shown to be critical to the progression of BBD (Glas *et al.*, 2012), *Desulfovibrio profundus* (which is thought to be responsible for the H₂S production) was detected in only ~5% of the clone libraries analyzed (Miller and Richardson, 2011). These discrepancies could be due to technical issues (e.g., amplification biases, low coverage of the microbiome in clonal libraries), seasonal and/or regional differences in the BBD composition or function-based (rather than taxonomy-based) assembly of the BBD community. Therefore, there is a need for a comprehensive characterization of BBD microbiota across multiple scales. Furthermore, to better understand initiation of BBD, its progression over a coral colony and ecosystem-wide spread, it is important to characterize the composition of the epimicrobiota of asymptomatic corals across temporal and spatial scales within ecosystems where BBD is known (or not known) to occur. These were some of the objectives of this study.

How BBD is initiated and transmitted remains uncertain. It spikes in warmer, sunnier months, and once a BBD outbreak takes place within a given ecosystem, it tends to reoccur each year (Zvuloni *et al.*, 2009; Sato *et al.*, 2009; Kuehl *et al.*, 2011; Muller and van Woesik, 2011; Sato *et al.*, 2011). Individual coral colonies in which BBD has halted are 3.5 times more likely to develop signs of the disease in subsequent years (Sato *et al.*, 2009). Analyses of the within-ecosystem, spatiotemporal patterns of the appearance of BBD point to various models, including transmission by water currents, stochastic appearance of lesions and direct colony-to-colony transmission (Bruckner *et al.*, 1997; Sato *et al.*, 2009; Zvuloni *et al.*, 2009). These observations also do not rule out a hypothesis that the spatiotemporal distribution of BBD cases on the reef correlates with water quality parameters (Page and Willis, 2006), which may serve as environmental triggers for community shifts mediated by BBD members. Alternate hypotheses, including vector-borne transmission from non-coral reservoirs or between diseased and healthy corals have also been suggested (Aeby and Santavy, 2006; Casey *et al.*, 2014). Cyanobacteria similar to those associated with BBD, including the genera *Leptolyngbya*, *Geitleriinema*, *Oscillatoria* and *Phormidium*, were found in microbiomes associated with turf, crustose coralline and macroalgae, potentially implicating these organisms as alternate hosts or reservoirs of the pathogens (Barott *et al.*, 2011).

Microbe–microbe interactions within BBD remain only partially understood. It is clear that complex transitions within coral microbiota are a prelude to

the development of BBD (Sato *et al.*, 2010). Changes in the biogeochemical conditions (gradients of H₂S, pH, oxygen and light availability) within the BBD mat contribute to the stratification of microorganisms (Sato *et al.*, 2011; Glas *et al.*, 2012), although mechanisms of cell-to-cell interactions driving the structuring of BBD consortia remain unknown. Recent discovery of *N*-acyl-homoserine lactones (AHLs) and AI-2 quorum sensing (QS) signals within BBD (Zimmer *et al.*, 2014) indicates that QS is likely involved in the structuring or functionality of BBD. QS is a mechanism of bacterial population density-dependent gene regulation, which has been shown to contribute importantly to structuring of host-associated microbiota (Miller and Bassler, 2001) and to interactions within microbial communities (Ng and Bassler, 2009). Bacteria use QS to express virulence genes, spread over surfaces, form and disperse biofilms and produce 'public goods' that can be used by the entire microbial community. However, it is not known how QS and signal interference contribute to coral diseases in general and to BBD in particular.

Here, we documented differences in the composition of microbiomes of healthy corals and those afflicted with BBD in distinct Caribbean ecosystems. To better understand interactions within BBD, secondary metabolites were extracted from the mat. We tested the hypothesis that an abundant cyanobacterial compound, lyngbic acid, isolated from BBD, interfered with QS, a signal perception and regulatory mechanism involved in structuring of microbial communities and their functions.

Materials and methods

Sample collection

Samples of BBD consortia and surface mucus from the Florida Keys, Belize and Honduras were collected by aspiration with needle-less sterile syringes that were uncapped immediately before the sample collection. A total of 60 samples were collected, including 30 BBD consortia, 20 surface mucus layer samples overlying healthy tissue on corals with BBD and 10 mucus samples from asymptomatic corals (Supplementary Table S1). Some of the corals in Belize were tagged so that the same coral was sampled at up to three time points, although not all tagged corals were resampled at every time point because of difficulties in locating tags on some of the subsequent samplings. Surface mucus from healthy tissue on corals with BBD was collected at least 10–15 cm away from an active lesion. Samples were spun in a microfuge at 4000 r.p.m. for 5 min, seawater was decanted and the pelleted mucus was preserved with ~5–10 volumes of RNAlater (Qiagen, Germantown, MD, USA) and stored at –20 °C until extraction of nucleic acids. Seawater was collected by boat off Carrie Bow Cay, Belize from an area adjacent to but not overlying the

reef and transported to shore. Seawater was filtered with a 0.2- μm filter in duplicate, with a total of 3 L of seawater for each filter. The filters were placed in a 15-ml conical tube, immersed in RNAlater and stored $-20\text{ }^{\circ}\text{C}$ until extraction of nucleic acids.

Coral-associated vibrios were isolated in April 2014 from healthy mucus and from the black band layer of *Montastraea cavernosa* from Florida on marine agar or TCBS (thiosulfate-citrate-bile salts-sucrose) agar. Individual colonies were sequentially streaked to isolation a minimum of three times and identified through sequencing of the V6-V8 region of the 16S rRNA gene with primers F968 and R1401 (Nübel *et al.*, 1996). A total of 29 coral-associated vibrios were screened for the production of CAI-1 autoinducer in AB medium (Supplementary Table S4). Of them, two bioluminescent isolates *Vibrio* spp. BBD33 (from black band) and BBD69 (from healthy mucus) were selected for subsequent assays with lyngbic acid.

QS assays

Cultures of *V. harveyi* and *Escherichia coli* reporters (Supplementary Table S2) were grown overnight with antibiotics in AB medium (for *V. harveyi*; Bassler *et al.*, 1994) or in LB (for *E. coli*) with kanamycin ($50\text{ }\mu\text{g ml}^{-1}$), chloramphenicol ($10\text{ }\mu\text{g ml}^{-1}$) or ampicillin ($200\text{ }\mu\text{g ml}^{-1}$) as appropriate to the genotype. To determine whether coral vibrios make CAI-1 signal, overnight AB cultures were spun, filtered through a 0.2 micron filter and mixed (at $\sim 30\%$) with a diluted culture of JMH626 ($\Delta luxN luxQ::Tn5 cqsA::Cm$), which only responds to CAI-1 (Supplementary Table S2 and Henke and Bassler, 2004). For the bioassays with lyngbic acid, a panel of the *V. harveyi* QS reporters and *Vibrio* spp. BBD33 and BBD69 were grown overnight, diluted at least 5000-fold, essentially as in Henke and Bassler (2004) and then added to the test wells containing fivefold serial dilutions of lyngbic acid. Dilutions of lyngbic acid were made in ethanol, and before adding reporter suspensions, the solvent was completely evaporated in a laminar flow hood. Control wells contained residue from the dried carrier solvent. Suspensions of the reporters were mixed with the dried lyngbic acid residue in each well by vigorous pipetting. Luminescence of the reporters was quantified over time with a Victor-2 multimode microtiter plate reader (Perkin-Elmer, Waltham, MA, USA).

'Indirect' assays were conducted to determine whether the addition of the cognate CAI-1 signal overrides the effect of lyngbic acid. Serial dilutions of lyngbic acid were prepared as above. Thirty microliters of filter-sterilized culture filtrates of *V. harveyi* KM413 ($\Delta luxS \Delta luxM$), which produces only the CAI-1 signal (Henke and Bassler, 2004) were added to all wells with lyngbic acid and then mixed with $90\text{ }\mu\text{l}$ of the diluted suspension of *V. harveyi* JMH626 (which only responds to CAI-1) (Supplementary Table S2). Luminescence was

measured every 30 min using the multimode microtiter plate reader.

16S Illumina tag sequencing

Genomic DNA for 16S Illumina tag sequencing was extracted with a PowerSoil DNA Isolation Kit (MoBio, Carlsbad, CA, USA). The V6 region of 16S rRNA genes was amplified in triplicate for each sample with previously reported Illumina-compatible primers designed for bacteria (Eren *et al.*, 2013). Samples were amplified in $25\text{ }\mu\text{l}$ reactions containing 0.5 U Phusion High-Fidelity Polymerase (New England Biolabs, Ipswich, MA, USA), $1\times$ Phusion HF Reaction Buffer, $0.75\text{ }\mu\text{l}$ dimethyl sulfoxide and 0.2 mM each dNTP. Triplicate PCR amplifications were pooled for each sample, cleaned with a MinElute Kit (Qiagen, Germantown, MD, USA), visualized on an ethidium bromide stained 1% agarose gel and quantified by NanoDrop (ThermoScientific, NanoDrop Products, Wilmington, DE, USA). Two hundred nanograms of each cleaned amplicon library was submitted to the Genomics Core Facility at Pennsylvania State University where the pooled libraries were size selected with a 2% agarose PippinPrep cassette to produce a narrow range of fragment sizes from 200 to 240 bp for sequencing (confirmed by a bioanalyzer) and cleaned again to remove agarose. Sequencing was performed on an Illumina MiSeq with a 150-bp paired-end protocol, using single indexing.

Sequencing reads were parsed by Illumina index at the sequencing center. Reads were then further parsed by the inline barcode, paired reads merged and primers and adaptors removed using a combination of tools in cutadapt (Martin 2011), Galaxy (Giardine *et al.*, 2005; Blankenberg *et al.*, 2010; Goecks *et al.*, 2010) and eautils (Aronesty, 2011). Parsed raw sequencing reads are publicly available through NCBI's Sequence Read Archive under the BioProject ID PRJNA269585. Sample names were added to the definition lines of sequencing reads using sed and concatenated into one fasta file, to make them compatible for analysis in QIIME v.1.8 (Caporaso *et al.*, 2010). Clustering of OTUs at 97% similarity was performed with the subsampled open-reference OTU picking method (Rideout *et al.*, 2014), with no removal of singletons. The Greengenes reference data set version 13.8 (DeSantis *et al.*, 2006) was used as the reference for OTU picking and for taxonomy assignment with uclust (Edgar, 2010). OTUs that were not assigned taxonomy were manually queried with BLASTn (Altschul *et al.*, 1990) against the non-redundant nucleotide collection of the NCBI (Mizrachi, 2002). OTUs identified as mitochondrial DNA or as chloroplasts were removed from further analyses. Community structure was analyzed in R with phyloseq (McMurdie and Holmes, 2013) and plotted with ggplot2 (Wickham, 2009). Analysis of similarities (ANOSIM) was performed in R using VEGAN v.2.0-8 (Dixon, 2003).

Network analysis of interactions between genera was conducted on a subset of 52 of the 60 coral microbiomes, focusing on the Florida and Belize locations that had both healthy and diseased samples from *Orbicella* and *Montastraea* hosts. Significant positive and negative correlations between genera with a minimum occurrence of 10 (out of 52 microbiomes) were determined by support of four separate measures: Pearson's correlation, Spearman's correlation, Bray–Curtis dissimilarity or Kullback–Leibler dissimilarity, using the CoNet app (Faust *et al.*, 2012) in Cytoscape v.3.0.2 (Shannon *et al.*, 2003). Networks from the four measures were merged by intersection, keeping only interactions with support from all methods. Rows were shuffled and the measures were performed with 1000 iterations, Benjamini–Hochberg multiple test correction was applied and highly interconnected regions (clusters) were identified with MCODE (Bader and Hogue, 2003).

Isolation of lyngbic acid

Lyngbic acid was isolated and completely characterized from two separate collections of BBD, one from the Florida Keys and one from Belize. The cyanobacterial mat from the black band consortium on *M. cavernosa* corals on the Looe Key Reef, Florida was collected on 23 April 2014. The sample was lyophilized to give a dry weight of 3.672 g and then extracted two times with 50 ml of MeOH. The extract was concentrated by rotary evaporation at 45 °C under reduced pressure to give 1.03 g of the MeOH-soluble extract. This extract was partitioned between ethyl acetate (EtOAc) and H₂O, and the EtOAc-soluble fraction was rotary evaporated. The EtOAc-soluble fraction (0.105 g) was chromatographed on a C₁₈ column using a MeOH:H₂O step gradient system to give six subfractions. A portion (0.003 g) of subfraction 4 (0.012 g), eluted with MeOH:H₂O (4:1) was further separated by reversed-phase HPLC (semi-prep, 5 µm, RP-C₁₈) using MeOH:H₂O (19:1) to give 1.2 mg of lyngbic acid (yield 0.13% dry wt, 4.8% of EtOAc-soluble fraction). Similarly, cyanobacterial mats from the BBD zones of coral species of *Orbicella* and *Pseudodiplora* in Belize were collected on July 13–17, 2013. The combined freeze-dried material (9.43 g) was extracted with MeOH saturated with helium gas. This MeOH extract (1.82 g) was partitioned between EtOAc and H₂O to give the EtOAc-soluble fraction (0.237 g) and H₂O-soluble fraction. The EtOAc extract was chromatographed on a C₁₈ column (16 g) using MeOH–H₂O step gradient system to give six subfractions. A portion (0.004 g) of the subfraction 4 (0.023 g), eluted with MeOH–H₂O (4:1), was further separated by reversed-phase HPLC using the same conditions to give 0.9 mg of lyngbic acid (yield 0.10% dry wt, 4.1% of EtOAc-soluble fraction). The structure of lyngbic acid was determined using the proton nuclear magnetic resonance spectroscopy,

high-resolution mass spectrometry and optical rotation data and confirmed by comparison with published data and data from our previously isolated samples of lyngbic acid (Cardellina *et al.*, 1978; Kwan *et al.*, 2010; Soares *et al.*, 2015). The presence of lyngbic acid in other collections of BBD (Belize: 1 March 2013; FL Keys: Looe Key 17 May 2013, 12 September 2013, 20 November 2013; and Wonderland Reef 10 June 2013) was determined by identifying the presence of the key methoxy and unsaturation proton signals in the nuclear magnetic resonance spectroscopy spectra of the extracts and fractions.

Results

Community structure of healthy and BBD coral microbiomes

The bacterial community structure and diversity of both healthy corals and those with BBD was determined for 60 coral surface microbiomes from Carrie Bow Cay, Belize, Looe Key in the Florida Keys, and four locations in Honduras, as well as for two background seawater samples from Carrie Bow Cay, Belize. A total of 14 571 218 bacterial small subunit rRNA gene sequences were analyzed after quality filtering, with 7497 to 750 674 sequences per sample (Supplementary Table S1). Distinct bacterial communities were detected in the surface microbiomes of healthy corals (epibiomes), in seawater and in the BBD consortia (Figure 1). Epibiomes were remarkably conserved and consisted primarily of Gammaproteobacteria (in particular, the genera *Halomonas* and *Moritella*) and *Renibacterium* (Figure 2). Epibiomes exhibited the lowest OTU richness and α -diversity compared with bacterial communities in seawater and BBD (Figure 3). Overall, the polymicrobial BBD consortia displayed more variable bacterial community structure than epibiomes (e.g., Figure 1 and Supplementary Figure S1), but this variability did not correspond with sample location (ANOSIM R statistic = 0.0867, significance from permutations = 0.044) (Figure 1a) or with coral species (ANOSIM R statistic = 0.1456, significance from permutations = 0.014) (Figure 1b), where an ANOSIM R statistic near zero indicates random variation between groups. However, a hypothesis that the composition of BBD varied seasonally was supported. BBD communities collected in Belize in cooler times of the year when the disease is thought to be less active were distinct from the BBD microbial communities sampled in warm times of the year when the disease was active (Figure 1). Furthermore, winter BBD samples tended to cluster with the seawater samples (Figure 1). Seawater samples exhibited the highest OTU richness and α -diversity (Figure 3).

Overall, bacterial communities in healthy mucus had high proportions of *Halomonas*, whereas black band consortia had high proportions of *Roseofilum*

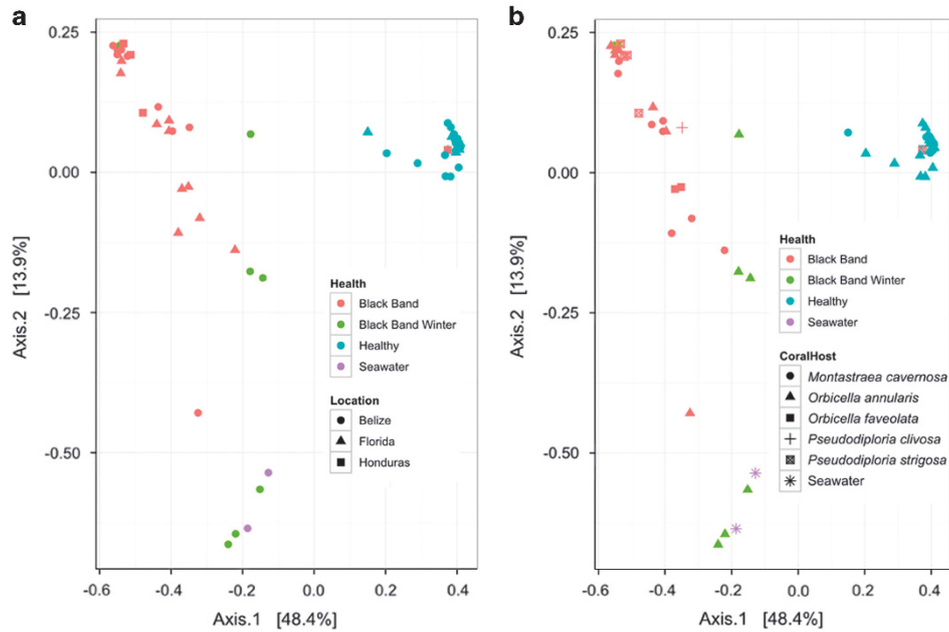


Figure 1 Comparison of bacterial community structure in healthy and diseased coral microbiomes. Bacterial community similarity based on Morisita–Horn β -diversity of Illumina MiSeq 16S rRNA gene libraries in the surface microbiomes of corals with and without BBD, in comparison with overlying seawater, with health status indicated by color and collection location (a) or coral host (b) indicated by symbol shape. Black Band communities sampled in February were consistently different from summer Black Band communities. Healthy surface microbiomes clustered together regardless of collection location or coral species.

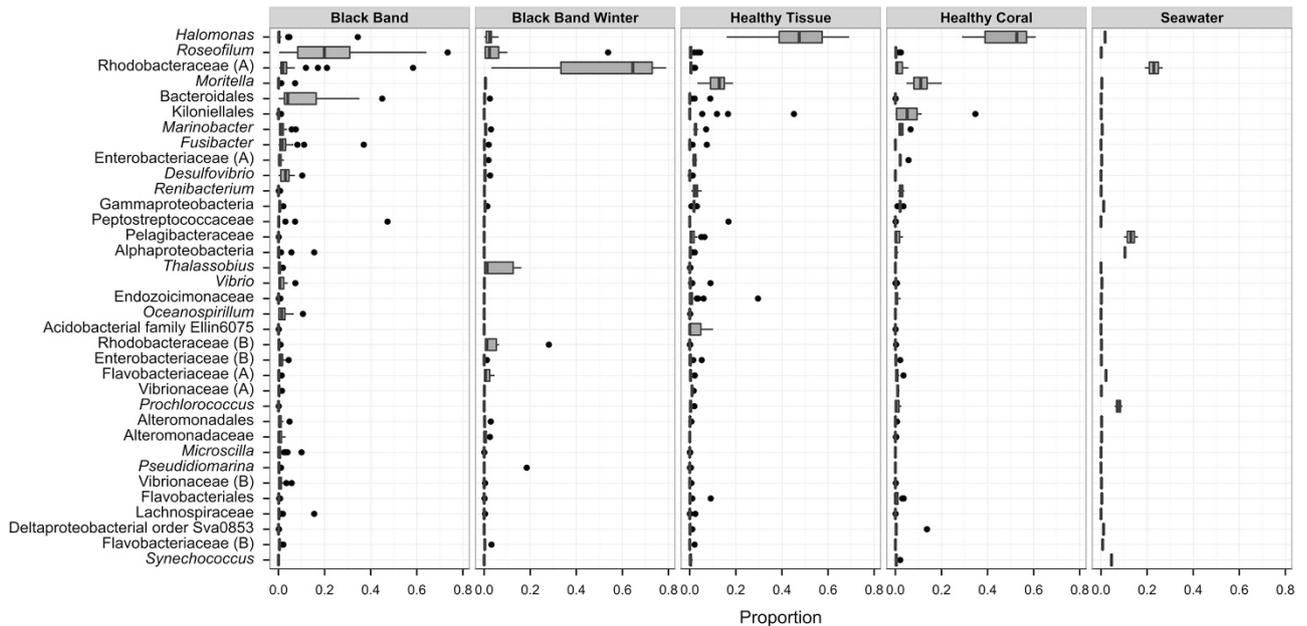


Figure 2 Distribution of abundant bacterial genera in healthy and diseased coral microbiomes. Box and whisker plot of the relative proportion of sequencing reads assigned to 35 most abundant bacterial taxa of surface microbiomes from corals with BBD ($n=23$), with BBD in winter ($n=7$), in healthy tissue on corals with BBD ($n=20$), in healthy coral heads ($n=10$), and in comparison with seawater above the reef ($n=2$). Points outside of the box and whiskers represent data outliers. Taxa were classified to genus when possible; taxa labeled with higher taxonomic classifications did not match named lineages within the database. Families labeled ‘(A)’ or ‘(B)’ represent separate lineages, which were not classified to more specific levels and match taxa labels in Figure 4 and Supplementary Figure S1.

(Figure 2 and Supplementary Figure S1). The filamentous, nonheterocystous, phycoerythrin-rich cyanobacterium *Roseofilum reptotaenium* was detected in every BBD sample, although great

variation in its abundance was observed (Figure 2 and Supplementary Figure S2). Although the dominant cyanobacterial OTUs in the Illumina libraries were classified as ‘*Leptolyngbya*’ by

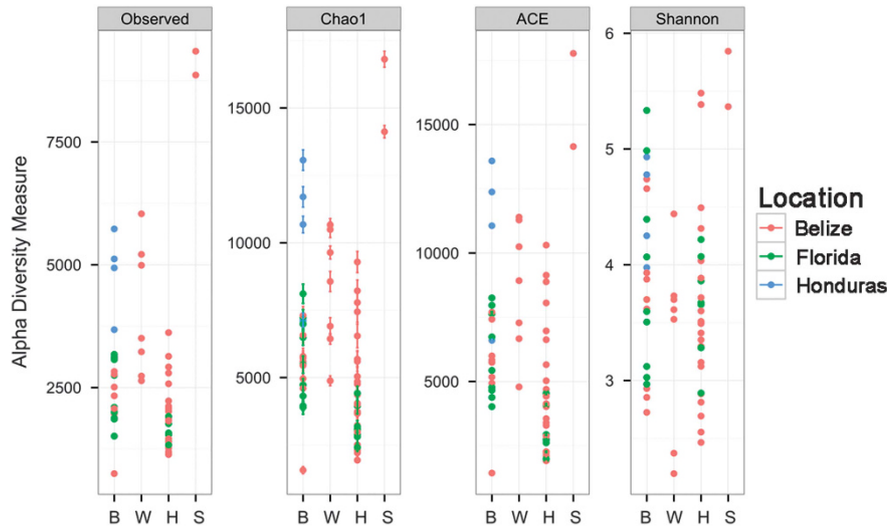


Figure 3 α -Diversity in healthy and diseased coral microbiomes. Comparison of observed taxonomic richness (number of OTUs clustered at 97% similarity), and the Chao1 (with s.e.), ACE and Shannon diversity indices in ‘B’: summer BBD consortia; ‘W’: winter BBD consortia; ‘H’: healthy surface microbiomes; and ‘S’: seawater. The background seawater samples were most diverse, and diseased microbiomes tended to be more diverse compared with healthy microbiomes.

Greengenes v.13.8, comparison of full-length cyanobacterial 16S rRNA genes recovered from Belize and Florida Keys metagenomic libraries prepared from the same DNA extractions and from a *Roseofilum* enrichment culture (KP689103) (Meyer *et al.*, in preparation) were >99% similar to sequences from the uncultured cyanobacterium (AY038527) identified in the first molecular characterization of Caribbean BBD mats (Frias-Lopez *et al.*, 2002), the cultured BBD cyanobacterium *Roseofilum reptotaenium* strain 101-1 (EU743965) from Caribbean *Pseudodiploria strigosa* (Casamatta *et al.*, 2012), and *Pseudoscillatoria coralii* (FJ210722) from *Favia* corals in the Red Sea (Rasoulouniriana *et al.*, 2009). Collectively, these results support the hypothesis that *R. reptotaenium* is the main cyanobacterium present in BBD mats.

In addition to *Roseofilum*, the disease consortia were dominated by uncultured genera of Rhodobacteraceae and Bacteroidales, *Fusibacter*, and *Desulfovibrio* (Figure 2). The dominant OTUs classified as uncultured Rhodobacteraceae closely match (≤ 2 nucleotide differences along the 60-bp length of the V6 amplicon sequence) uncultured Alphaproteobacteria recovered from *Siderastrea* corals with BBD (DQ446102) (Sekar *et al.*, 2006), from healthy *Orbicella annularis* corals (FJ952692) (Rypien *et al.*, 2010), and from corals with Porites White Patch Syndrome (KF179798) (Sere *et al.*, 2013). The dominant OTUs classified as uncultured Bacteroidales were not closely related to any Bacteroidetes detected in coral microbiomes.

Healthy coral microbiota are stable over time, depth and location

To test the hypothesis that the epibiome of healthy corals is stable over spatial and temporal gradients,

we sampled healthy surface microbial communities from *M. cavernosa*, *O. annularis*, *O. faveolata* and *P. strigosa* in Belize and Florida Keys over 2 years in February, April, May, June, July and August. Epibiomes generally clustered together (Supplementary Figure S3). Healthy microbiomes from *O. annularis* collected over a range of depths were indistinguishable from those collected in shallow waters (Supplementary Figure S3C). Importantly, all samples (including those collected from depths ranging from ~12 to 21 m, where BBD is not known to occur) contained *R. reptotaenium*, the cyanobacterium associated with BBD. Up to 0.3% of sequencing reads from healthy corals were identified as *Roseofilum*. These observations clearly establish that coral epibiomes are stable across temporal and spatial gradients. Furthermore, these observations support the hypothesis that *R. reptotaenium* is a normal, albeit minor, member of the coral microbiota, even at depths where BBD does not occur.

Microbiome shifts and BBD progression

In addition to the overall characterization of healthy and black band microbiomes, we examined spatial patterns in the distribution and progression of BBD. First, we fortuitously located adjacent colonies of *O. annularis* and *P. strigosa*, which both displayed signs of BBD (Supplementary Figure S4). A comparison of their microbiomes revealed that the bacterial communities from healthy tissues were conserved and clustered together, whereas BBD communities on adjacent corals were distinct (Supplementary Figure S4). Second, examination of the surface microbiome at 5 cm increments away from the BBD front in three separate corals of *M. cavernosa* showed a rapid transition from the BBD consortium, dominated by *R. reptotaenium*, to the conserved epibiome

consisting primarily of Gammaproteobacteria, in particular, the genera *Halomonas*, *Moritella* and *Marinobacter* (Supplementary Figure S5). It is important to note that in the asymptomatic transition zone (5 cm away from the lesion front), *R. reptotarium* was significantly represented (up to 4.5% of the microbiome), and the cyanobacterium was present as a minor ($\leq 0.1\%$) component of the stable microbiomes 10 and 15 cm away from the lesion front.

Interactions between healthy and BBD coral microbiota

As the black band layer migrates across the surface of infected corals, *Roseofilum* and its associated heterotrophs continually invade the adjacent healthy surface mucus layer. Our deep-sequencing results show that *Roseofilum* and other members of the polymicrobial disease consortia are present at low levels in healthy tissues and uninfected corals and that 63% of genera were shared between the two sample types. Taken together, this suggests that interactions between healthy microbiota and BBD consortia members may ultimately determine the overall health of the coral host. To test this hypothesis, we examined co-occurrence patterns of bacterial genera to identify microbe–microbe interactions that potentially influence the progression of BBD. A total of 97 significant interactions (multiple corrected P -value < 0.05) were detected between 23 bacterial genera in a matched subset of *Montastraea* and *Orbicella* corals from Belize and Florida, containing 23 black band consortia and 29 healthy samples (Figure 4a). All 23 genera with non-random co-occurrence patterns were detected in both healthy and diseased samples. Overall, dominant genera within either healthy or diseased microbiomes had significant positive interactions (co-occurrences) with other microbiome members of the same health state and negative interactions (mutual exclusions) with dominant microbiome members of the opposite health state. *Halomonas* had a total of 17 significant interactions, including negative interactions with all of the most abundant members of the BBD consortium: *Roseofilum*, uncultured Rhodobacteraceae, uncultured Bacteroidales, *Fusibacter* and *Desulfovibrio*. *Halomonas* also had significant positive interactions with six genera, four of which were other Gammaproteobacteria, as well as the single-celled cyanobacterium *Synechococcus* and the actinobacterium *Renibacterium*. In contrast, *Roseofilum* had six significant negative interactions, including *Halomonas* and *Moritella*, the two most abundant genera detected in the healthy microbiomes and only one positive interaction, with *Desulfovibrio*, a sulfate-reducing Deltaproteobacterium. The fewer significant interactions between *Roseofilum* and other genera may reflect the more variable community composition in black band consortia. Further analysis of the interaction networks revealed two large clusters of highly

interconnected regions of interactions. Cluster 1 involved 22 of the 23 genera with significant interactions in Figure 4a (all except *Marinobacter*) and revealed that many of the dominant Gammaproteobacteria, including *Halomonas*, *Moritella*, *Vibrio*, uncultured Enterobacteriaceae and *Oceanospirillum*, are hubs of connections, interacting both positively and negatively with many other taxa (Figure 4b and Supplementary Figure S6). Cluster 2 consisted of bacteria that are over-represented in BBD samples (*Roseofilum*, *Thalassobius*, *Vibrio* spp., uncultured Rhodobacteraceae and *Fusibacter*) (Figure 4c). Notably, while *Roseofilum* had only seven significant interactions, it was part of both clusters and several of its significant interactions were with taxa that are highly interconnected hubs (such as *Vibrio* spp.).

Lyngbic acid is abundant within BBD

To better understand interactions within the BBD consortium, secondary metabolites present within the extracts of the BBD mats were purified. Lyngbic acid was identified as one of the major secondary metabolites in BBD extracts and represented 0.13% (dry wt) from the Florida sample and 0.10% (dry wt) from the Belize sample, and over 4% of all hydrophobic substances (which includes many primary metabolites such as pigments and lipids) extractable into ethyl acetate. It was identified based on the HRMS data (HRESI/APCIMS m/z 257.2127 $[M+H]^+$ (calculated for $C_{15}H_{29}O_3$, 257.2117)) and was further confirmed by comparing the 1H nuclear magnetic resonance spectroscopy data with an authentic sample of lyngbic acid previously isolated from other benthic filamentous marine cyanobacteria. Lyngbic acid extracted from BBD gave a specific rotation value of $[\alpha]^{25D} -12.0$ (c 0.06, $CHCl_3$; Supplementary Figure S7), similar to the specific rotation value of lyngbic acid reported earlier from other cyanobacteria $[\alpha]^{26D} -9$ (c 7.3, $CHCl_3$) (Cardellina *et al.*, 1978); $[\alpha]^{20D} -12.6$ (c 0.8, CH_3OH) (Kwan *et al.*, 2010). When BBD mats were collected in quantities that were suitable for chemical extractions and subsequent analyses, the presence of lyngbic acid in BBD samples was ascertained by nuclear magnetic resonance spectroscopy and/or HRMS. It was detected in all seven tested Caribbean BBD samples, including those collected from Carrie Bow Cay in Belize and from Wonderland and Looe Keys in Florida (Supplementary Table S2). Lyngbic acid did not have a strong effect on growth of culturable coral-associated bacteria, such as *Halomonas* spp. and *Vibrio* spp. (data not shown). Therefore, it is unlikely to function as a general biocide.

Lyngbic acid inhibits CqsS-mediated QS in Vibrio spp.

With the subsequent experiments we tested the hypothesis that lyngbic acid interferes with QS in *Vibrio* spp. The rationale for this hypothesis was

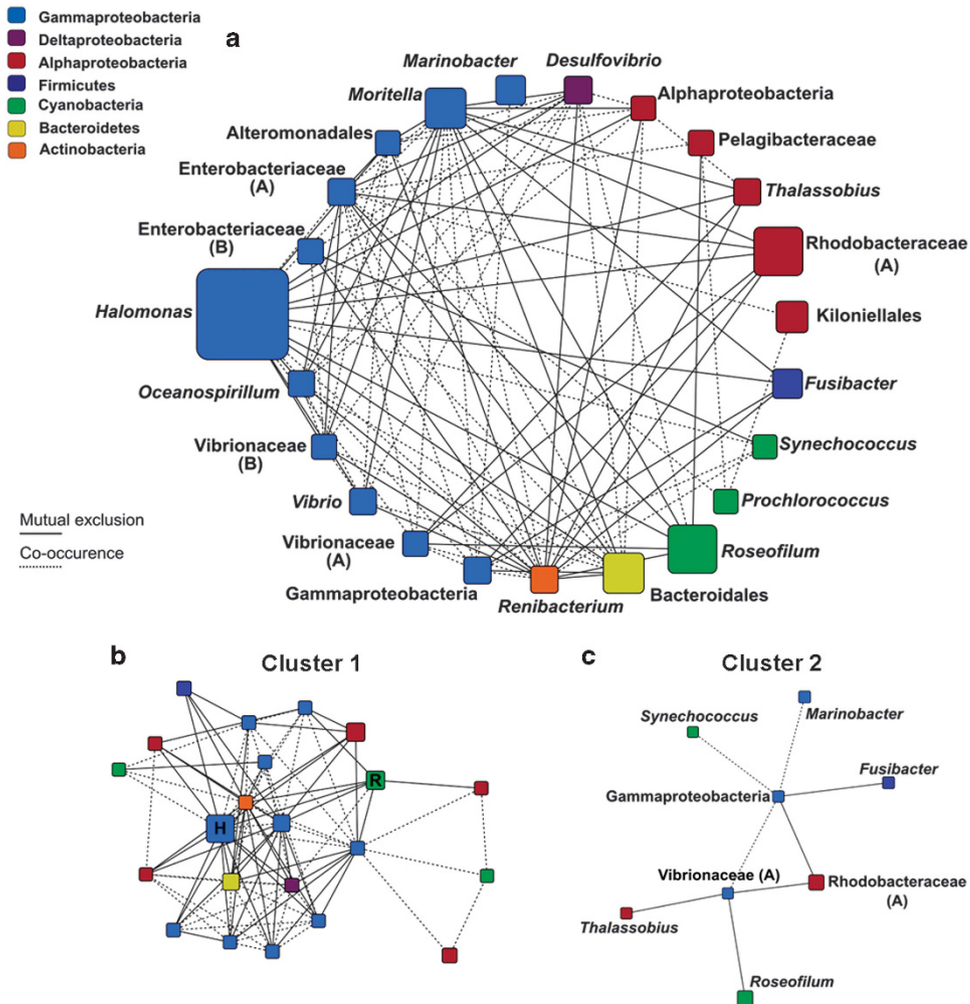


Figure 4 Network analysis of interactions between abundant bacterial genera. Significant interactions between bacterial genera in 52 *Orbicella* and *Montastraea* corals from Belize and Florida with and without BBD. Solid lines indicate negative interactions (mutual exclusions) and dotted lines indicate positive interactions (co-occurrences). The size of the nodes reflects the relative abundance of the genus in the entire data set, and the nodes are sorted and colored by phylum. (a) All 97 significant interactions, involving 23 genera, sorted by class and densely interconnected regions, Cluster 1 (b) and Cluster 2 (c).

based on the observation that *R. reptotaenium* and *Vibrio* spp. co-occurred in the BBD consortia and were predicted to have a direct negative interaction (Figure 4 and Supplementary Figure S6). *Vibrio* spp. form highly connected hubs in the cluster representing stable microbiome (Supplementary Figure S6). Cross-species manipulation of QS is a well-characterized mechanism of microbe–microbe interactions with community-wide consequences (Dobretsov *et al.*, 2009; Teplitski *et al.*, 2011). Therefore, lyngbic acid extracted from BBD was subjected to a series of bioassays to elucidate its effect on QS in *Vibrio* spp. A suite of *V. harveyi* mutants, each lacking either synthases or receptors of its three QS regulatory cascades, were exposed to serial dilutions of lyngbic acid. Despite deletions in QS genes, these mutants are still luminescent, although they produce light at levels lower than those of the wild type. If lyngbic acid acted as a signal agonist, then luminescence of the strains

lacking either of the QS signal synthase genes (*luxM*, *cqsA* and *luxS*, which are involved in synthesis of AHL, hydroxyketone CAI-1 and furanosyl borate diester AI-2 QS signals, respectively) would have increased. Instead, strong inhibition of luminescence in these mutants was observed (Figure 5 and Supplementary Table S3). Lyngbic acid did not inhibit luminescence of a constitutively luminescent *E. coli* DH5 α pTIM2442 (Alagely *et al.*, 2011b), indicating that it does not function as a generic toxin or a nonspecific inhibitor of luminescence at the tested concentrations (Supplementary Figure S8). To determine exactly where lyngbic acid affects QS cascades in *V. harveyi*, its effects on luminescence of mutants in *luxN*, *cqsS* and *luxPQ* (encoding receptors of AHLs, CAI-1 and AI-2 signals, respectively) were tested. Absence of a strong response to lyngbic acid in a particular receptor mutant indicates that the corresponding missing protein is critical to the signal transduction. The mutant lacking the *cqsS*

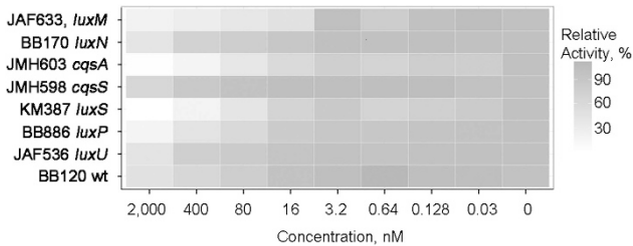


Figure 5 Effect of lyngbic acid on QS pathways. Luminescence assays were performed in seven mutant strains and one wild-type strain of *Vibrio* with concentrations up to 2 μ M lyngbic acid to determine its effect on several different pathways in the *Vibrio* model. *luxM*, *cqsA* and *luxS* encode synthases of the AHL, furanosyl borate diester AI-2 and hydroxyketone CAI-1 signals; *luxN*, *cqsS* and *luxPQ* encode receptors of these signals. LuxU is a phosphorelay protein through which inputs of all three QS systems are integrated into downstream regulatory cascades. Assays were performed in duplicates in three independent experiments. Heatmaps were constructed using data from a representative experiment after 8–10 h incubation with lyngbic acid.

receptor gene was the least responsive to lyngbic acid (Figure 5 and Supplementary Table S3), whereas strains in which *cqsS* was intact remained responsive to the cyanobacterial compound. A deletion of *luxU*, which functions as an ‘integration point’ for the input of all three QS signaling cascades, also strongly reduced sensitivity to lyngbic acid, consistent with the model in which effects of lyngbic acid require a QS receptor positioned below LuxU in the QS cascade (such as CqsS). Collectively, these observations suggest that the effect of lyngbic acid on QS in *V. harveyi* is likely mediated by the CAI-1 QS signal receptor CqsS, and lyngbic acid may function as a natural antagonist of the CqsS/CAI-1 QS system.

To more robustly test the hypothesis that lyngbic acid interferes with CqsS-mediated QS in *V. harveyi*, its ability to inhibit perception of the CAI-1 signal was determined using ‘indirect assays’. *V. harveyi* JMH626, which only contains a functional CqsS receptor (and lacks genes encoding receptors of the AHL or AI-2 signals, *luxN* and *luxQ*, as well as the CAI-1 synthase gene *cqsA*) (Henke and Bassler, 2004), was used as a reporter. *V. harveyi* JMH626 is normally dark, unless it is exposed to CAI-1 (Henke and Bassler, 2004). CAI-1 was provided in the cell-free culture filtrates of *V. harveyi* KM413, which lacks AHL and AI-2 synthases *luxM* and *luxS*, respectively, but has a functional Cqs QS system (Henke and Bassler, 2004). Dilutions of lyngbic acid were dissolved in the 1:3 mix of the CAI-1-containing culture filtrate of KM413, and the suspension of the CAI-1-specific reporter JMH626. This ratio of the culture filtrate to the reporter was shown to elicit maximum responses (Henke and Bassler, 2004). Increasing concentrations of lyngbic acid inhibited stimulatory effects of CAI-1 (Figure 6), consistent with the postulated function of lyngbic acid as a naturally occurring inhibitor of the Cqs-mediated QS.

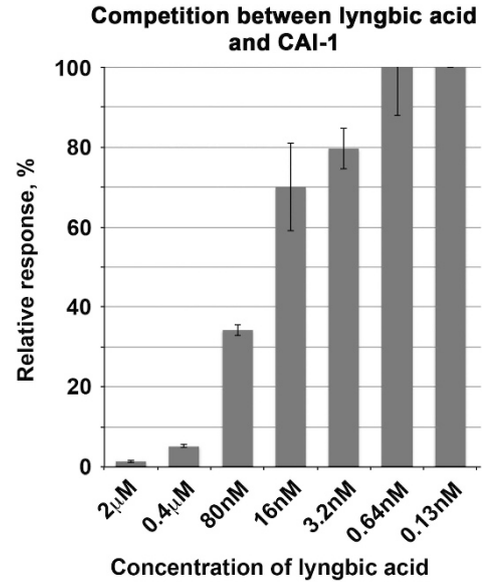


Figure 6 Competition between lyngbic acid and CAI-1. Luminescence of *V. harveyi* JMH626 (*luxN luxQ cqsA*, which does not respond to the AHL or AI-2 signals, does not make its own CAI-1 signal and only luminesces in response to the exogenous CAI-1) was measured with microtiter plate reader Victor-2 (Perkin-Elmer). Serial dilutions of lyngbic acid (indicated on x axis) were mixed with 30 μ l of filter-sterilized culture filtrates of *V. harveyi* KM413 ($\Delta luxS \Delta luxM$, which produces only the CAI-1 signal) before adding 90 μ l of the diluted suspension of JMH626. Inhibition of luminescence is indicated as the percentage of full activation of the JMH626 reporter with culture filtrates from KM413.

CAI-mediated signaling is common in coral vibrios

In addition to establishing that lyngbic acid can antagonize QS in a well-characterized reporter system, it was important to determine whether such interactions are possible *in situ*. Therefore, we tested the hypothesis that lyngbic acid interferes with QS in native coral-associated *Vibrio* spp. in a two-step experiment on vibrios isolated from both BBD mats and from healthy surface mucus layers. First, culture filtrates of coral *Vibrio* isolates were surveyed for their ability to induce luminescence in *V. harveyi* JMH626. Extracts of 27 out of 29 coral vibrios stimulated luminescence of this CqsS reporter by ~10–10 000-fold (Supplementary Table S4), indicating that CAI-mediated signaling is highly prevalent in coral vibrios. The observed differences in the levels of luminescence induced by culture filtrates are consistent with similar surveys in other *Vibrio* spp. (Ng *et al.*, 2011), and may be a consequence of different amounts or structures of CAI-1-like signals produced by the isolates.

Lyngbic acid inhibits luminescence in coral vibrios

To determine whether lyngbic acid impacts native *Vibrio* spp., two bioluminescent strains of coral vibrios that were shown to possess the CAI-1-mediated signaling were exposed to dilution series of lyngbic acid. Similarly to the observation with the

V. harveyi reporters, lyngbic acid inhibited bioluminescence of coral vibrios (Supplementary Table S3).

Discussion

Coral diseases, such as BBD, have been devastating reef ecosystems worldwide for decades (Antonius, 1973; Bourne *et al.*, 2009; Rosenberg, 2002). Even though disease signs and ecosystem-wide consequences have been cataloged for over a dozen coral diseases, much remains to be learned about the initiation of coral diseases, as well as interactions between native microbiota and disease consortia. More generally, to what extent host-associated microbiota are involved in diseases also remains an open question, even in the best-characterized model systems.

This study focused on the characterization of BBD microbiomes and coral epibiomes in distinct ecosystems throughout the Caribbean. Healthy epibiomes were characterized by a relatively low microbial diversity and were dominated by *Halomonas* spp., with *Moritella* spp., *Renibacterium* spp., *Kiloniella* spp. also prominent in healthy epibiomes. Dominance of *Halomonas* spp. in commensal coral microbiomes is consistent with observations in some other corals (McKew *et al.*, 2012); however, the closely related genus *Endozoicomonas* tends to be dominant in others (Morrow *et al.*, 2012; Bayer *et al.*, 2013; Meyer *et al.*, 2014).

BBD microbiomes were more diverse than healthy coral epibiomes; similarly, an increase in bacterial diversity was characteristic of other coral diseases (Sunagawa *et al.*, 2009; Closek *et al.*, 2014; Roder *et al.*, 2014a). Even though α -diversity of BBD microbiomes was generally higher compared with that in coral commensal epibiomes, the same key members (*R. reptotaeneum*, Bacteroidales, Rhodobacteriaceae, *Fusibacter* spp. and *Desulfovibrio* spp.) were the most prominent BBD members in the samples from the Florida Keys, Belize and three locations in Honduras. This suggests that the assembly of the BBD consortia likely involves a combination of environmental factors and interactions within the microbiomes. Interestingly, there were seasonal differences in the BBD microbiome composition. BBD communities in winter had a significantly higher representation of Rhodobacteraceae, and a relative decrease in abundance of *R. reptotaenium*. This relationship could be owing to the fact that these microorganisms are capable of both photosynthesis and nitrogen fixation, and that these biochemical processes could be required for the formation and progression of BBD. Seasonal representation of these organisms in BBD communities may reflect differences in their adaptation to water temperature, light intensity or other environmental conditions.

Mechanisms and signals associated with coral microbiome transitions to BBD remain unknown,

although the detection of bacterial cell-to-cell QS signals (*N*-acyl homoserine lactones, AHLs and AI-2) within the BBD consortia (Zimmer *et al.*, 2014) indicates that QS likely has a role in BBD communities. The ability to manipulate QS in associated microbiota is a common trait of marine organisms (Dobretsov *et al.*, 2009). Therefore, it is, perhaps, not surprising that one of the most abundant cyanobacterial metabolites within BBD is a compound capable of manipulating QS in *Vibrio* spp.

By examining species co-occurrence patterns, we uncovered a negative correlation between *Roseofilum* and a number of bacteria that are present mostly in epibiomes. Negative co-occurrence patterns were also observed for *Roseofilum* and *Vibrio* species. Vibrios have previously been postulated to have a role in BBD (Barneah *et al.*, 2007; Arotsker *et al.*, 2009), although their contribution to the initiation and progression of the disease has yet to be characterized. Although vibrios were detected at moderately low levels in both healthy coral mucus and in the black band, vibrios are readily cultivable from both sample types and have QS activity through both the AHL-based autoinducer 1 system and the furanosyl borate diester-based autoinducer 2 system (Tait *et al.*, 2010; Alagely *et al.*, 2011a; Golberg *et al.*, 2011, 2013; Zimmer *et al.*, 2014). In addition to AHLs and AI-2, vibrios also produce hydroxyketone CAI-1 signals (Henke and Bassler, 2004; Ng *et al.*, 2011; Wei *et al.*, 2012), and with this study we demonstrate that coral vibrios also produce CAI-1-like activities. Furthermore, this CqsS/CAI-1 QS system appears vulnerable to manipulation by lyngbic acid, a cyanobacterial secondary metabolite.

Unlike previously characterized naturally occurring antagonists, which target AHL- or AI-2-based QS systems, activity of lyngbic acid required the presence of a functional CqsS, a receptor for the hydroxyketone CAI-1 signal. Synthetic agonists and antagonists of the CqsS-mediated signaling have been previously identified (Bolitho *et al.*, 2011); however, lyngbic acid is the first characterized naturally occurring inhibitor of CqsS-mediated signaling. Even though lyngbic acid did not fully inhibit activity of the QS reporters (as was achieved by the some synthetic analogs of CAI-1), its activity at 16 nM is more potent than those reported for many of the synthetic antagonists (Bolitho *et al.*, 2011). We note that deletion of the gene coding for the AHL receptor LuxN also reduced sensitivity of the strain to lyngbic acid at 2 μ M, although not to the same extent as the deletion of the *cqsS* gene (Figure 5). It is possible that in addition to strongly inhibiting the CqsS-mediated cascades, lyngbic acid may also modestly affect the LuxN/AHL QS system, perhaps similarly to the ability of CAI-1 to antagonize AHL-based QS systems in other bacteria (Ganin *et al.*, 2012). The consequences of manipulation of *Vibrio* QS with the cyanobacterial compound lyngbic acid are not yet known. In *Vibrio* spp., QS signals act as timers, with the CqsS/CAI-1 system active at low population

densities (Henke and Bassler, 2004; Anetzberger *et al.*, 2012). In concert with other QS signals, the CqsS/CAI-1 system regulates toxin and metalloprotease production, as well as type 3 secretion system (Henke and Bassler, 2004; Higgins *et al.*, 2007). Therefore, functions of the cyanobacterial metabolite lyngbic acid may include manipulation of these behaviors to control interactions within BBD and its interactions with the host coral. This fundamental role for lyngbic acid as a QS inhibitor against vibrios may also explain its presence in so many different benthic filamentous marine cyanobacteria (Cardellina *et al.*, 1978; Kwan *et al.*, 2010; Soares *et al.*, 2015).

Deep sequencing conducted in this study clearly establishes that *Roseofilum reptotaenium*, the cyanobacterial pathogen linked to BBD of corals, is a minor but ubiquitous member of healthy Caribbean coral microbiota. This observation has important implications for understanding how BBD initiates and spreads. Knowing that both *Roseofilum* and potentially pathogenic heterotrophs are in low abundance in healthy corals suggests that the disease consortium is held in check by currently unknown constraints exerted by either the coral polyp or the symbiotic algae, or the commensal Gammaproteobacteria (including *Halomonas* spp., *Moritella* spp.) and actinomycetes (*Renibacterium* spp.) characteristic of the healthy state. Release from these constraints is associated with the loss of the commensals (*Halomonas* spp., *Moritella* spp. and *Renibacterium* spp.) and an increase in the diversity and species richness of the microbiome. Even though *R. reptotaenium* is known to be the major member of the BBD consortium, BBD is the only example of a disease associated primarily with a cyanobacterium (Richardson *et al.*, 2014). Given that cyanobacteria are not generally known for encoding genes involved in animal virulence, it is reasonable to hypothesize that the main function of *R. reptotaenium* in BBD is to act as an ‘engineer’ of the disease consortium: while not virulent on its own, through photosynthesis, nitrogen fixation and the biosynthesis of secondary metabolites, it creates an environment where opportunistic pathogens attack the coral host and degrade its tissues. To more robustly test this hypothesis, it will be important in future studies to characterize functions and chemicals that are involved in structuring of BBD.

Conflict of Interest

The authors declare no conflict of interest.

Acknowledgements

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