Quorum Sensing Interference and Structural Variation of Quorum Sensing Mimics in Australian Soft Coral

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Bacterial Quorum Sensing (QS), the indirect regulation of gene expression through production and detection of small diffusable molecules, has emerged as a point of interaction between eukaryotic host organisms and their associated microbial communities. The extracellular nature of QS molecules enables interference in QS systems, in many cases via mimicry. This study targeted QS induction and inhibition in soft coral holobionts, as many soft coral species commonly contain compounds with structural similarities to the well-studied bacterial QS molecules acyl homoserine lactones. Screening with two bacterial biosensors, Agrobacterium tumefaciens A136 and Chromobacterium violaceum CV026, demonstrated that QS interference differed between the two biosensor strains and extended across the soft coral families Alcyoniidae, Clavulariidae, Nephtheidae, and Xeniidae. Bioassay-guided fractionation revealed chemical activity patterns, particularly in the induction of QS. Cembranoid diterpenes from active fractions were purified and tested for QS interference activity. Interestingly, the type of QS activity (induction or inhibition) in A. tumefaciens A136 correlated with structural variability of the secondary oxygen ring; cembranoid diterpenes with a furan ring or five-membered lactone induced QS, while compounds with larger (six or seven membered) lactone rings inhibited QS. Addition of the dominant cembranoid diterpene in the soft coral Lobophytum compactum, isolobophytolide, to bacterial culture media increased the number and morphological diversity of bacteria recovered from the mucosal layer of this soft coral, demonstrating a selective effect on certain members of the soft coral bacterial community. The identity and QS activity of recovered isolates differed between the mucosal layers of L. compactum and Sinularia flexibilis. In conclusion, this study provides information on the complexity of the interaction between soft corals and their associated bacteria, as well as, a structural understanding of how QS mimic compounds are able to interfere with a bacterial communication system.

Keywords: quorum sensing, soft coral, bacterial isolates, cembranolide, quorum sensing mimic
INTRODUCTION

Quorum Sensing (QS) is one form of cell to cell signaling employed by bacteria to coordinate gene expression across entire populations through release and detection of extracellular signal molecules (Miller and Bassler, 2001). The aspects of multicellularity gained through QS, enable bacteria to perform many important ecological functions such as the ability to interact with their physical and biological environment (Miller and Bassler, 2001), form biofilms (Rice et al., 2005), and secrete virulence factors (Zhu and Mekalanos, 2003). The extracellular nature of QS signaling molecules facilitates their disruption and mimicry (Chhabra et al., 2005). Consequently, many bacteria possess the ability to detect and respond to QS signals of other species (Joint et al., 2007). Indeed, QS systems are more prevalent amongst bacteria associated with mixed bacterial biofilms and macro-organisms, suggesting that possession of QS systems confers an advantage in these habitats (Dudler and Eberl, 2006).

QS mimics are extrinsic signals that can interfere directly with QS gene expression (Bauer and Robinson, 2002). To be effective, QS mimic compounds must specifically interfere with the target QS system. Multiple QS systems have been discovered; in Gram-negative bacteria (Papenfort and Bassler, 2016) the most well studied system is the Auto Inducer One system (AI-1), which utilizes acyl homoserine lactones (AHLs) as signal molecules. Studies into the structure and functions of AHLs suggest that the γ-lactone ring is required for QS activity and that the length and functionality of the acyl side chain provides specificity (Parsk and Greenberg, 2000; Watson et al., 2002; Geske et al., 2008). For this reason, it has been hypothesized that AI-1 QS mimics would also contain a γ lactone ring or homologous functionality such as the furanones of the red alga Delisea pulchra. The furanones are one of the few QS inhibitors to have been structurally elucidated and the presence of an oxygenated ring was demonstrated to be essential to their activity (Manefield et al., 1999).

Disruption and mimicry of QS signals are increasingly recognized as mechanisms that are commonly employed by macro-organisms to regulate and manipulate their associated microbial communities (Bauer and Robinson, 2002; González and Keshavan, 2006). QS interference by host organisms can confer the ability to respond to the presence of certain pathogenic or mutualistic bacteria quickly and reliably (Kjelleberg et al., 1997; Mathesius et al., 2003), render a pathogenic species of bacteria non-pathogenic (Dong et al., 2007; Swem et al., 2009), and enable manipulation of the abundance and composition of its associated bacterial assemblies (Givskov et al., 1996). A host's microbiota can be a first line of defense against pathogen invasion (McFall-Ngai et al., 2013), therefore manipulation of QS could strengthen the resilience of the holobiont (González and Keshavan, 2006; Teplitski and Ritchie, 2009).

The ability of some bacterial species to detect non-native QS molecules has allowed the development of bacterial biosensor strains. QS bacterial biosensor strains are genetically modified bacterial isolates that require external addition of QS signal molecules. Their expression of QS-regulated genes is linked to reporter genes, which typically produce a pigment or bioluminescence (Steindler and Venturi, 2007). Two of the most commonly used AI-1 bacterial biosensor strains are based on the species Chromobacterium violaceum and Agrobacterium tumefaciens. In C. violaceum, QS regulates the production of the secondary metabolite violacein, which is purple in color (McClean et al., 1997). Bacterial biosensors based on this species utilize the LuxI/LuxR homolog genes CivI/CivR and are sensitive to AHLs with C4–C8 carbon chains as well as 3-oxo-C6 and -C8 carbon chains (McClean et al., 1997; Steindler and Venturi, 2007). A. tumefaciens QS biosensors have a genetically modified QS plasmid such that QS by these strains results in an enzymatic breakdown of X-gal and the formation of an indigo colored product (Zhu et al., 1998; Farrand et al., 2002; Zhu and Mekalanos, 2003). A. tumefaciens QS biosensors often utilize the TraI/TraR genes that provide sensitivity to AHLs with C6–C14 acyl side chains as well their equivalent 3-oxo-acyl side chains and C6–C10 hydroxyl acyl side chains (Zhu et al., 1998; Farrand et al., 2002; Zhu and Mekalanos, 2003; Steindler and Venturi, 2007). Despite the partly overlapping acyl chain lengths detected by these two sensors, previous studies have observed differences in their responses during screening of isolates (Chong et al., 2012).

Widespread QS inhibitory activity has been observed to occur in the marine benthos, particularly in sponges and soft corals (Taylor et al., 2004; Skindersoe et al., 2008; Hunt et al., 2012). Soft corals contain a number of secondary metabolites with the structural potential to mimic QS, including furanocembranes (cembranoid diterpenes with fused 5-membered ether rings) and cembranolides (cembranoids that possess a fused second ring in the form of a lactone). Cembranoid diterpenes are most commonly, but not exclusively, found within the family Alcyoniidae. Furthermore, cembranoid diterpenes are inherently variable in the presence, position and size of oxygenated ring systems (Wahlberg and Eklund, 1992). Variation of substituents, direction of cyclisation and the corresponding position of the isoprenoid double bonds of different diterpenes from different species of soft coral are also encountered (Wahlberg and Eklund, 1992) making cembranoid diterpenes a natural pool of compounds to investigate QS mimic structure-activity relationships.

Soft corals have a high incidence of QS interference; however, so far all screened soft corals have been from a single family, Alcyoniidae, and the presence of QS induction has not yet been investigated. While QS and QS interference have been implicated in the regulation of mixed bacterial communities including those found in the surface mucosal layer (SML) of hard corals (Tait et al., 2010; Golberg et al., 2011, 2013), it is unknown whether these communities may be regulated by QS mimics. Currently very little known about bacterial communities that associate with soft corals. In hard corals, bacteria in the SML, the first and largest point of interaction between a coral and the environment, are considered important to the health and resilience of the holobiont (Reshef et al., 2006; Rosenberg et al., 2007; Zilber-Rosenberg and Rosenberg, 2008; Shtit-Orland and Kushmaro, 2009).

This study aims to assess the potential of soft corals to interact with their associated microbes via QS, with a focus on their associated secondary metabolites such as cembranolides and furanocembranes. Crude extracts of 15 soft coral species, representing 4 families of soft corals, were tested for their
ability to induce or inhibit QS in 2 bacterial biosensors, *A. tumefaciens* A136 and *C. violaceum* CV026. Cembranolides and furanocembranes were isolated from soft corals and the effect of their structural variability on QS interference was assessed. To gain a better understanding of the role that QS can play in regulating eukaryote associated bacterial communities, bacterial strains isolated from the surface mucosal layer (SML) of two soft corals, *Sinularia flexibilis* and *Lobophytum compactum* were similarly evaluated for QS activity. The possible role that isolobophytolide, the major secondary metabolite of *L. compactum*, might play in bacterial selection within the soft corals mucus was also assessed by supplementing culture media with this secondary metabolite.

**MATERIALS AND METHODS**

**Soft Coral Collection**

Twenty four specimens of soft coral, representing 15 species (Supplementary Tables 1–3), were collected at a depth of 1–3 m near Orpheus Island (Great Barrier Reef, Australia; latitude 18°36′5.878″S; longitude 146°29′9.996″E). All specimens except *Cespitularia* sp. were photographed (Supplementary Figure 1) and sampled underwater, with samples placed directly into plastic bags filled with seawater. Samples were frozen (−80°C) within 1 h of collection and stored until freeze drying. Additional colonies of *L. compactum*, *S. flexibilis*, and *Pachyclavularia violacea* were collected for fraction analysis and isolation of pure compounds (Supplementary Tables 2,3). 1H NMR spectra of the crude extracts of all soft coral samples were compared between collections and with reference spectra from the Bowden laboratory to ensure consistency in the metabolites and species tested.

**Soft Coral Extract Preparation**

Dried soft coral tissue was weighed and homogenized before extraction. Three extracts of different polarity were generated for each soft coral sample. Solvents used for extraction were, in order, dichloromethane (DCM), methanol (MeOH), and water (H2O). Each extract was the result of three successive applications of solvent. Extracts were concentrated by rotary evaporation before being dried under a stream of nitrogen (N2) and stored at −20°C until analysis. The DCM and MeOH extracts were dissolved in ethanol and the aqueous extracts in H2O to a concentration of 20 mg/ml. All soft coral extracts were tested at two concentrations (4 µg/well and 40 µg/well) three times for the presence of QS induction and QS inhibition activity in *A. tumefaciens* A136 and *C. violaceum* CV026 in at least two independent experiments.

**Soft Coral Extract Fractionation**

To further investigate the patterns of QS activity observed in crude extracts, nine soft coral species from five genera (three families) were chosen for further fractionation (Supplementary Table 2). The chosen species displayed five patterns of crude extract activity (Supplementary Table 4). In addition to four species with strong crude extract activities, five species were selected to investigate observed within genera variation in activity patterns or because they represent common genera on the Great Barrier Reef (GBR). Ten fractions of decreasing polarity were generated from the dichloromethane extracts of the chosen soft corals. Extracts were fractionated using flash column chromatography on RP-C18 silica cartridges (Phenomenex Strata C18-E 55 µm 70 Å, 1,000 mg) eluted with a stepwise 20–100% MeOH: H2O gradient followed by a 1:1 DCM: MeOH wash. The resulting fractions were concentrated to dryness under a stream of N2 gas and re-dissolved in ethanol for QS screening as described above. 1H-NMR spectra (600 MHz) of active fractions were recorded with a Bruker 600 Avance spectrometer in deuterated chloroform (CDCl3).

**Isolation of Purified Metabolites From the Corals**

Ten cembranoid diterpenes with variable secondary ring structures were assessed in this study (Figure 1, Table 1). Isoneocembrene A (1) represents the base cembrane backbone, without an additional ring (Figure 1, Table 1). Compounds 2–4 are γ-lactones (5-membered rings). Two δ-lactones (6-membered rings) and one ε-lactone (7-membered ring) are also included in compounds 6–8 respectively (Figure 1, Table 1). The final two compounds are furanocembranes (Figure 1, Table 1). Pure samples of isoneocembrene A (1), lobolide (4) and sarcophine (5) were acquired from the Bowden Laboratory, Townsville Australia. Isolobophytolide (2), and isolobophytolide monoacetates (3) were isolated from freshly collected and extracted *L. compactum* (see below). Flexibilide (6), dihydroleflexibilide (7) and sinulariolide (8) were isolated from freshly collected and extracted *S. flexibilis* (see below). The furanocembranes, pachyclavulariadiol (9) and pachyclavulariadiol diacetate (10), were isolated from freshly collected and extracted *P. violacea*. Extracts of each species were generated as described previously (section Soft Coral Extract Preparation).

For the *L. compactum* extract, vacuum liquid chromatography of active crude DCM extract (2g) was performed over reverse phase C18 silica gel (Phenomenex Gemini 3 μm NX-C18 110 Å, LC Column 30 × 4.6 mm). Isolobophytolide (2) eluted at 15 min and the two isomers of isolobophytolide monoacetate (3) eluted at 17 min.

For the *S. flexibilis* extract, flexibilide (6) and dihydroleflexibilide (7) were isolated as reported by Kazlauskas et al. (1978). In brief, the DCM extract was subjected to normal phase flash column chromatography with combinations of hexane, DCM and EtOAc. Flexibilide was eluted by 6:1 DCM: EtOAc, 4:1 DCM: EtOAc yielded a mixture of 6 and 7 and 3:1 DCM: EtOAc afforded pure dihydroleflexibilide. Sinulariolide (8) was isolated separately by a method adapted from Tursch et al. (1975) by direct crystallization of the DCM extract after dissolution in diethyl ether. Purification of compounds was performed by HPLC as described above.
Due to instability of the furanocembranes from *P. violacea*, the pure compounds were generated semi-synthetically as per Bowden et al. (1979). In brief, the DCM extract was prepared at 4°C then partitioned between hexane and 10% aqueous MeOH. After removal of the solvent, the aqueous MeOH fraction was subjected to normal phase flash chromatography with a hexane:EtOAc gradient. All fractions containing (by TLC and $^1$H NMR) pachyclavulariadiol, diacetyl pachyclavulariadiol and the two monoacetyl pachyclavulariadiols were combined and hydrolyzed to pachyclavulariadiol ($^9$) by incubation for 24 h at room temperature with MeOH containing 1% (w/v) potassium hydroxide. Methanol was removed under vacuum and the residue was partitioned between diethyl ether and water. The ether fraction was evaporated and the residue was dissolved in hexane. Diacetyl pachyclavulariadiol (10) was acquired by acetylation of half of the obtained pachyclavulariadiol (9). Acetylation was affected by incubation for 24 h with 1:1 acetic anhydride in pyridine before evaporation of the solvent and retrieval via liquid a partition between hexane and water. The semi-synthesis of monoacetyl pachyclavulariadiols was not performed as both 9 and 10 exhibited similar activity, so it was considered unlikely that the activities of monoacetyl pachyclavulariadiols would be different.

Structure and purity of each extracted compound was confirmed by 1D and 2D NMR and comparison with literature values. $^1$H-NMR (600 MHz) and $^{13}$C-NMR (150 MHz) spectra were recorded with a Bruker 600 Avance spectrometer in CDCl$_3$, with tetramethylsilane (TMS) as internal standard. High
resolution mass spectra were collected using an unmodified Bruker BioAPEX 47e mass spectrometer equipped with an Analytica model 103426 (Branford, CT) electrospray ionization (ESI) source. Analytical thin layer chromatography (TLC) was performed on Merck Kieselgel 60. Spots were visualized by UV light or by spraying with a 1% vanillin in acetic acid solution. Pure compounds were re-solubilized in ethanol and serially diluted to generate five different concentration solutions (1 × 10⁻² mM to 1 × 10⁻⁵ mM) for each compound.

Collection, Culture and Extraction of Bacterial Isolates From S. flexibilis and L. compactum Surface Mucosal Layer

Surface mucosal layer samples (SML) were collected from three healthy replicate colonies of S. flexibilis and L. compactum from the same depth and location. SML samples were collected underwater from the mid-capitulum region of the coral colony using 50 ml needleless sterile syringes. Samples were maintained at ambient temperatures and processed within 3 h of collection. At the same time as SML samples were retrieved, tissue samples of each replicate were collected and their metabolite ³¹H NMR spectral profiles were compared with samples from S. flexibilis and L. compactum collected previously to ensure correct species identification.

SML samples were serially diluted (10⁻², 10⁻³, 10⁻⁴) using autoclaved artificial seawater (Instant Ocean; Spectrum Brands, Madison, WI, USA). One hundred microliters of each dilution were spread plated in triplicate on two types of media commonly used for studies of marine bacteria: 50% Marine Agar (50MA; BD) and Glycerol Artificial Seawater (GASW) agar (Smith and Hayasaka, 1982). Additionally, Thiosulfate Citrate Bile Salts (TCBS; BD) agar, which specifically selects for members of the family Vibrionaceae, was included. L. compactum SML dilutions were additionally plated onto 50MA and GASW agar supplemented with L. compactum's major secondary metabolite, isolobophytolide. All plates were incubated at 28°C and sampled after 48 h, 72 h, 1, and 2 weeks. Representatives of each colony morphotype from each plate were sub-cultured to purity for identification.

Where possible, two representatives of each morphotype were selected for QS activity screening. Where bacteria had initially been isolated using media embedded with isolobophytolide, growth was attempted on the equivalent medium without isolobophytolide. Strains that could not be cultured without isolobophytolide were not included in the screening. Screening was performed on acidified ethyl acetate (EtOAc) extracts of the cell free supernatant of soft coral isolates. These were acquired by transferring single colonies from 50MA plates to liquid culture (10 ml 50% Marine broth culture, 28°C at 170 rpm) and grown to late exponential phase. Cultures were centrifuged for 10 min at 4°C at 10,000 g to obtain the cell free supernatant (CFS). Each CFS was subjected to exhaustive extraction with acidified EtOAc (1% acetic acid) and concentrated to dryness under a stream of N₂ gas. Extracts were then dissolved and diluted to a concentration of 20 mg/l with ethanol.

Bacterial QS Biosensor Strains and Culture Medium

The biosensor strains A. tumefaciens A136 (Fuqua and Winans, 1996) and C. violaceum CV026 (McClean et al., 1997) were used for detection of QS induction and inhibition in soft coral extracts. A. tumefaciens A136 was grown on ABt media (Clarke and Maaloe, 1967) and C. violaceum CV026 was grown on Luria Bertani (LB) media (Bertani, 1951). In order to ensure that the QS plasmid was intact and functional, QS biosensor strains were grown in the presence of the appropriate antibiotic (Ravn et al., 2001). A. tumefaciens A136 was grown on media supplemented with 4.5 µg/ml of tetracycline and 50 µg/ml of spectinomycin, whereas, C. violaceum CV026 was grown on media supplemented with 20 µg/ml of kanamycin (Ravn et al., 2001).

QS Screening Assays

The presence of AHL type QS induction activity in soft coral extracts, fractions and pure compounds was detected by performing an agar diffusion assay as described in detail by Ravn et al. (2001). The QS biosensor strain, either A. tumefaciens A136 or C. violaceum CV026, was embedded within the agar and the sample being tested was added to a well cut or formed in the agar. For induction of QS, N-hexanoyl homoserine lactone was used as a positive control and extraction solvents were used as negative controls. Positive results were read as a blue coloration surrounding the wells of A. tumefaciens A136 and a purple coloration surrounding the wells of C. violaceum CV026 (see above and Supplementary Figure 2). The intensity of the response was measured as the diameter of the colored zone and normalized to the response of the positive control.

The agar diffusion assays described above were modified in order to detect QS inhibition. Briefly, as A. tumefaciens A136 and C. violaceum CV026 are not able to QS without the exogenous addition of AHLs, 8.5 μmol n-hexanoyl homoserine lactone was added into the agar embedded with the biosensor strain in order to test for QS inhibition. Two positive controls were chosen based on their previously reported ability to inhibit QS: n-dodecanoyl-DL-homoserine lactone (McClean et al., 1997) and vanillin (Choo et al., 2006). These controls proved effective for both biosensors. The extraction solvents were once again used as negative controls. Positive results in the inhibition assay were read as inhibition of blue or purple coloration of the plates containing A. tumefaciens A136 and C. violaceum CV026 (respectively) (Supplementary Figure 2). The intensity of the response was measured as the width of the inhibition zone surrounding the well and normalized to the positive control.

To generate dose response curves for pure compounds, agar containing the respective biosensor was poured into custom built molds with 28 preformed wells 4 mm in diameter. After solidification of the agar, 20 µl of sample was added to each well. Pure compounds were only tested with the A. tumefaciens A136 strain as it was the only strain to have both QS induction and inhibition activity.
**Bacterial DNA Extraction, PCR and Sequencing**

Genomic DNA of bacterial isolates was extracted using the Promega Wizard Genomic DNA Isolation kit (Promega, Madison WI USA) according to the manufacturer’s directions. PCR amplification of 16S rRNA gene fragments was performed using the primers 27F and 1492R (Lane, 1991). The PCR reactions contained the following reagents: 0.4 mM of each primer, 1x MyTaq buffer (Bioline, Australia), 1.25 U MyTaq (Bioline, Australia), 1 µL DNA extract (final volume of 50 µL). Cycling conditions were an initial denaturing step of 94°C for 5 min, followed by 30 cycles at 95°C for 1 min, 56°C for 45 s, 72°C for 60 s, and a final elongation step at 72°C for 10 min. PCR products were verified by agarose gel electrophoresis and purified for sequencing using the Qiaquick PCR purification Kit (Qiagen, Valencia, CA) according to company supplied directions. Sanger sequencing was carried out at Macrogen (Seoul, South Korea) using both 27f and 1492R as sequencing primers. 

**Phylogenetic Analysis of Bacterial Isolates**

Sequence fragments were assembled using Sequencher (Version 5, Gene Codes, Ann Arbour, USA). For each isolate, the 16S rRNA gene sequence was aligned with sequences in the nr and Ref_Seq database at the NCBI using the megablast tool (RRID:SCR_001598; Altschul et al., 1990) to identify closely related database sequences. Sequences of isolates and database matches were imported into MEGA6 (MEGA Software, RRID:SCR_000667) and aligned using ClustalW (Larkin et al., 2007). A Maximum Likelihood-based phylogenetic tree was constructed using the Maximum Parsimony algorithm for the starting tree, the Tamura-Nei model for nucleotide substitution, and 500 bootstrap replicates (Supplementary Figure 3). The 16S rRNA gene sequences for the 72 bacterial isolates were deposited into the NCBI Genbank database, under accession numbers KM360403-KM360473. Quantification and statistical analysis of CFUs and isolate morphotypes.

Colony forming unit (CFU) concentrations were estimated based on dilutions yielding between 30 and 300 colonies per plate. Differences in the number of CFUs between samples were determined based on three replicates for the corresponding dilution and media type. Statistical differences were determined using the non-parametric Kruskal-Wallis statistic, as the data violated both normality and homogeneity of variances required for ANOVA. Colony morphology profile analysis was conducted on the variables color, size and texture and were compared using a nonmetric Multidimensional Scaling (nMDS) analysis in the vegan package (version 2.5-1; Oksanen et al., 2018) in R. The nMDS was chosen based on its suitability for spatial representation of complex data sets containing multiple variables, large numbers of zeroes and non-normal distributions (Rabinowitz, 1975). The statistical analyses were performed using Graphpad PRISM (GraphPad Prism, RRID:SCR_015807).

**RESULTS**

**Quorum Sensing Activities of Soft Coral Crude Extracts**

Crude soft coral extracts demonstrated the ability to both induce and inhibit QS in the biosensors tested. While both induction and inhibition of QS were observed for the biosensor A. tumefaciens A136, none of the extracts, regardless of source coral species, polarity, or concentration, were able to induce QS in C. violaceum CV026 (Figure 2A). Inhibition of QS was more prevalent than induction across all polarity extracts (Figure 2A). For both biosensors, the number of active soft coral extracts was reduced at lower extract concentration, with none of the aqueous extracts retaining their activity at the lower dosage (Figure 2B).

Crude extracts of five species induced QS in A. tumefaciens A136; two species from the family Alcyoniidae (L. compactum, Lobophytum sarcophytoide), one from the family Nephtheidae (Nepthia chabroli), one from the family Clavulariidae (P. violacea), and one from the family Xeniidae (Cespitularia sp.) (Figure 3). The highest incidence and strength of induction activity was seen for DCM crude extracts, with the largest haloes of coloration produced by DCM extracts of L. compactum and P. violacea (Figure 3).

Low level QS inhibition of A. tumefaciens A136 was demonstrated for DCM crude extracts of most species, with the only exceptions being L. microlobulatum, Simularia polydactyla, P. violacea, Clavularia sp., and N. chabroli (Figure 4A). The same trend was seen for MeOH crude extracts, with the additional exception of Sarcophyton sp. 2 (Figure 4A). Inhibition of QS in C. violaceum CV026 was present in DCM crude extracts of
all species except P. violacea and Clavularia sp. (Figure 4B). Of the five species that induced QS in A. tumefaciens A136, three (L. compactum, L. sarcophytoides and Cespitularia sp.) also inhibited QS in both biosensors (Figure 4). This contrasted with the other two species capable of QS induction; N. chabroli extracts inhibited QS only in C. violaceum CV026 (Figure 4) while P. violacea demonstrated no QS inhibitory activity with either biosensor (Figure 4).

Quorum Sensing Activities of Soft Coral Fractions
For all fractionated species, at least two fractions induced QS in A. tumefaciens A136 (Figure 5), regardless of whether their corresponding crude extracts were active (Figure 4). All species produced fractions that induced QS and inhibited QS in at least one biosensor strain. The only exception was P. violacea, which did not inhibit QS in either strain (Figure 5) consistent with crude extract results (Figure 4). The two largest inductive haloes were observed for N. chabroli and L. compactum, which also retained their activity at the higher (1:10) dilution level (Figure 4). The two major bands of induction activity seen in the A. tumefaciens A136 bioassay occurred for fractions eluted at 60% MeOH and at 80–90% MeOH (Figure 5). Many of the latter fractions (80–90% MeOH) also showed activity in the corresponding QS inhibition bioassay (Figure 5). The distinct patterns of QS activity observed for A. tumefaciens A136 contrasted strongly with the broad C. violaceum CV026 inhibition activity (Figure 5). The presence of cembrene diterpenes correlated well with the QS active fractions (data not shown).

Quorum Sensing Activities of Pure Compounds
The response to pure compounds also depended on the biosensor strain utilized. None of the tested cembranoid diterpenes induced QS in C. violaceum CV026. In contrast, both QS induction and inhibition activity was observed against A. tumefaciens A136 over three to four orders of magnitude, with a loss of activity at higher concentrations (Figure 6). No QS interference was observed for isoneocembrene A (Compound 1; Figure 6). The strongest induction of QS in A. tumefaciens A136 was observed for pachyclavularadiol and diacetyl pachyclavularadiol (Compounds 9 and 10; Figure 6). Induction was also observed in isolobophytolide (Compounds 2 and 3), lobolide (Compound 4), and sarcophine (Compound 5). QS inhibition was strongest in the ε-lactone ring of sinulariolide (Figure 6). Peak QS interference for all compounds occurred at approximately 1 × 10⁻⁵ M (or 3 ppm; Figure 6).

Culturable Bacteria
A significantly higher number of colony forming units (CFUs) were estimated for mucus of S. flexibilis as compared to L. compactum (Kruskal-Wallis, H = 7.200, 2 d.f., P = 0.0036; Supplementary Figure 4). Interestingly, when growth media for L. compactum were amended with isolobophytolide, the estimated number of CFUs increased for this species but...
remained lower than the estimates for S. flexibilis (Supplementary Figure 4). The number and type of colony morphotypes also differed between L. compactum and S. flexibilis (Figure 7). S. flexibilis showed little variation in the morphotype profiles of GASW or 50MA media, forming a tight cluster on the nMDS biplot (Figure 7). In comparison, the morphotype profiles generated from L. compactum showed higher variation in the same culture media. This trend was also consistent when isolobophytolide was added as a selection agent (Figure 7).

**Sinularia flexibilis Bacterial Isolates**

In total, 20 bacterial isolates from S. flexibilis were identified through 16S rRNA gene sequencing followed by BLAST searches and construction of phylogenetic trees (Supplementary
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FIGURE 5 | Active soft coral fractions by QS biosensor assay. Fractions were generated using a C18 flash column with a stepwise MeOH:H₂O gradient. Fraction numbers reflect elution order with increasing MeOH percentage. Light blue bars represent the *A. tumefaciens* A136 induction assay, dark blue bars represent the *A. tumefaciens* A136 inhibition assay and the purple bars represent the *C. violaceum* CV026 inhibition assay.

Table 5, Figures 8, 9). The isolates were dominated by Gammaproteobacteria belonging to the family *Vibrionaceae* both for the non-*Vibrionaceae* targeted media (GASW and 50MA) as well as the *Vibrionaceae* targeted medium (TCBS). Other Gammaproteobacteria included two isolates whose closest relative was "*Spongiobacter nickelotolerans*" (hereafter referred to
as *Endozoicomonas*, see below); and three *Alteromonas*-related strains. Finally, one isolate was identified with 99% sequence identity to *Bacillus megaterium* and *Bacillus aryabhattai* (phylum *Firmicutes*).

The potential of soft coral isolates from *S. flexibilis* to participate in AHL-type QS communication systems was investigated using the same reporter bioassays as used for coral extracts. Quorum sensing activity under the used test conditions was demonstrated for 52.4% of the tested *S. flexibilis* isolates. Both tested *Alteromonas* strains exhibited QS induction activity. The *Alteromonas* SFB10_2 strain triggered QS induction in both sensors strains, whereas, the *Alteromonas* YSF strain only triggered QS induction in *C. violaceum* CV026 (*Figure 8*). Both *Endozoicomonas*-related strains triggered QS induction in *C. violaceum* CV026 only. In addition, both strains triggered QS inhibition in *A. tumefaciens* A136, while QS inhibition in *C. violaceum* CV026 was only triggered by strain SF102 (*Figure 8*). We note that out of the 62 bacterial strains screened in this study, the two *Endozoicomonas*-related strains from *S. flexibilis* were the only strains with both induction and inhibition QS activity. Of the tested *Vibrio* strains, 6 of 14 strains showed QS activity and this was evenly split between induction (3 strains) and inhibition (3 strains) activity.

### Lobophytum compactum Bacterial Isolates

The isolates cultured from *L. compactum* demonstrated a number of similarities to the bacteria isolated from *S. flexibilis* (Supplementary Table 5, *Figures 10, 11*). Firstly, the majority of *L. compactum* isolates were gammaproteobacteria of the genus *Vibrio* (30/51). Secondly, strains related to the genera *Endozoicomonas* and *Bacillus*, and the order *Alteromonadales*, were isolated also from this soft coral species. In this instance, however, the diversity of *Alteromonadales*-related strains was higher with strains related not only to the genus *Alteromonas* (seven strains) but also to the genera *Pseudoalteromonas* (six strains), *Paramoritella*, *Ferrimonas*, and *Shewanella*. In contrast to the *S. flexibilis* isolates, the *L. compactum* isolates also included strains belonging to the genera *Psychrobacter* (class *Gammaproteobacteria*), *Erythrobacter* (class *Alphaproteobacteria*), and *Micrococcus* (class *Actinobacteria*).

The potential of soft coral isolates from *L. compactum* to participate in AHL-type QS communication systems was also investigated (*Figure 10*). Of the tested isolates from *L. compactum*, 47.5% demonstrated QS activity at the growth conditions tested and activity was mixed between induction and inhibition. Three of the tested *Vibrio* strains (LC111, LC103, and LC105) showed inhibitory activity against both biosensors. Of the strains that were initially isolated with media containing isobutyloyltaide, 41% were unable to be cultured in the absence of this compound and consequently were not tested for QS activity.

### DISCUSSION

#### Soft Coral Extracts

This study has demonstrated that QS interference extends across at least four soft coral families (*Alcyoniidae*, *Clavulariidae*, *Neptheidae*, and *Xeniidae*). Further, it was shown that both induction and inhibition QS activity extends across both polar and non-polar fractions, indicating that QS interference capability is widespread in soft corals from the central Great Barrier Reef, Australia.

Widespread activity, across not only species that are known to contain different metabolite types but also across a range polarities (indicated by the activities of extracts obtained by use of solvents with different polarities), is probably indicative of active compounds of more than one structural type. This is further supported by the finding that most (except *P. violacea*) of the soft coral species that induced QS in *A. tumefaciens* A136, also inhibited both biosensors. The widespread prevalence of QS inhibition as well as the presence of QS induction in the soft corals screened here is consistent with QS activity found across a range of marine invertebrates (Taylor et al., 2004; Skindersoe et al., 2008; Hunt et al., 2012). The dual presence of induction and inhibition of QS is similar to that found previously in gorgonian coral extracts (Hunt et al., 2012) but contrasts with the sole QS inhibition activity that was identified in *D. pulchra* (Kjelleberg et al., 1997). QS induction was also established in extracts of marine sponges and sponge associated bacteria (Taylor et al., 2004). Results from this study highlight the need to examine both induction and inhibition of QS to generate a realistic understanding of the complexity of ecological interactions between a host organism and its associated bacteria.

#### Soft Coral Fractions

In contrast with the initial crude extract testing, all the fractionated soft corals displayed at least one active fraction in the *A. tumefaciens* A136 QS induction assay. This may reflect an inherent increase in concentration of the active components or a decrease in complexity of the samples being tested. Soft coral extracts and fractions may be highly complex mixtures of compounds with contrasting QS regulatory activities. The potential for activity masking within extracts, a phenomenon previously observed in the QS screening of marine sponge extracts (Taylor et al., 2004), is high. This is particularly true if only one concentration or level of complexity is tested.

The QS induction pattern of the soft coral fractions was generally limited to one or two active fractions (80 and 90% methanol elution) for each species suggests that the inductive capability may be due to the presence of structurally similar compounds. Cembranoid diterpenes are well documented in eleven of the soft coral species tested (MarinLit Database, 2013) and correlated well to QS induction therefore could be responsible for the observed QS activity in these species. The same diterpene scaffolds, however, are not commonly known in the genera *Nepthea* (Amir et al., 2012a,b) *Cespitularia* (Elshamy et al., 2016) or in gorgonian corals (Changyun et al., 2008), so cembranoid diterpenes cannot fully explain the observed QS interference in these species. Isolation of cembranoid diterpenes was therefore required to understand the relative importance of these secondary metabolites to act as QS mimics.

A similarly discrete pattern was not reflected in the QS inhibition profile for these fractions. The broad QS inhibition profile of these fractions, might be due to multiple compounds,
or, the compound(s) responsible may not be suited to the method of fractionation used and the same compounds could be spread across several fractions. The presence of multiple QS compounds within a single holobiont would potentially enable a larger number of interactions with different bacterial strains and/or trigger different QS responses. This complexity may also reflect the capability of some bacteria to possess multiple QS systems (Reshef et al., 2006), with each system regulating a different process or interaction.

**Pure Compounds**

The strength and type of QS interference by cembranoid diterpenes was observed to correlate with the size of the oxygenated ring. Those cembranoid diterpenes that contained either a five-membered furan or lactone ring were capable of inducing QS in A. tumefaciens A136, whereas the cembranoid diterpenes with larger lactone rings (six or seven members) were seen to inhibit QS in A. tumefaciens A136. The type of oxygenated functional group also appears to impact the strength of QS interference, with QS interference only observed for cembranoid diterpenes possessing secondary oxygen rings (Fuqua et al., 2001; Watson et al., 2002; Geske et al., 2008). The presence of other minor functional groups (epoxides, acetates or level of saturation) had minimal discernible impact on the strength of QS interference observed and no obvious effect on the type of activity observed with respect to A. tumefaciens A136. QS mimics have previously been isolated that possess a γ lactone, however, these mimics (such as the furanones of D. pulchra) are often associated with QS inhibition rather than the inductive activity demonstrated here (Givskov et al., 1996; Defoirdt et al., 2013). In the case of the furanones from D. pulchra, bromine substituents are also present and may be influencing the type of activity. The presence of these metabolites in soft corals is strongly correlated to their taxonomy and may represent different strategies of interaction between species.

A common feature of QS mimic compounds previously identified from eukaryotic extracts is multiple forms of biological activity (Davies, 2006; Yim et al., 2007; Defoirdt et al., 2013). Cembranoid diterpenes appear to be no different with a number, including those identified in the current study, having previously been reported to demonstrate antibiotic (Aceret et al., 1995), cytotoxic (Maida et al., 1993) and algacidal properties. The antimicrobial activity identified in flexibilide (7) by Aceret et al. (1995), however, was exhibited at concentrations at least one order of magnitude higher than those that produced QS...
interference in this study. The peak QS active concentration occurred $1 \times 10^{-5}$ mM (or 3 ppm), reflecting the concentrations of flexibilide and sarcophytoxide in the mucous and water column surrounding *S. flexibilis* and *S. crassocaule* detected by Coll et al. (1982). Rather than being incompatible, the contrasting activities could be evidence of a hormetic response. Hormetic relationships have been previously observed in the QS mimics from garlic (Persson et al., 2005) and some antibiotic compounds, whereby growth stimulation or cell signaling properties are exhibited at concentrations below their minimum growth inhibitory concentration (Davies, 2006; Yim et al., 2007). A hormetic response could be relevant in soft corals with loosely packed sclerites, where uptake or release of water from the tissue can lead to large changes in volume over a matter of hours (Freckelton, 2015). As a result, associated metabolites will show a correspondingly dramatic change in concentration in the tissues on a volumetric basis over the same time period. More research is required to understand the potential of hormetic relationships.
in QS mimics and how such concentration changes could be manipulated in the defense of the coral.

Strong evidence for the ecological role of cembranolides and furanocembranoid diterpenes as QS mimics is further exhibited in the strong differences in the ability of the two biosensor strains to respond to QS mimics in the soft corals. The QS inductive compounds present within the soft corals were more readily detected by the *A. tumefaciens* A136 strain. In contrast, QS inhibition was observed more frequently for *C. violaceum* CV026. This could suggest that *C. violaceum* CV026 can be inhibited by a broader range of compounds or that it is more sensitive to a broader range of compound concentrations. This

| FIGURE 10 | A phylogenetic tree based on 16S rRNA gene sequences retrieved from an analysis of bacterial isolates from the mucus of the soft coral *L. compactum*. Details of the Vibrionaceae are shown in Figure 11. The tree is based on maximum-likelihood analysis, using a 50% conservation filter. The scale bar indicates 10% estimated sequence divergence. Thermoplasma acidophilum was used as the outgroup for analysis. Isolated sequences and their accession numbers are indicated in bold type. The nearest matches from the NCBI databases are included. T indicates that the sequence originates in the species type strain. |
FIGURE 11 | Vibrionaceae sub-tree (part of tree presented in Figure 10) based on 16S rRNA gene sequences retrieved from an analysis of bacterial isolates from the mucus of the soft coral *L. compactum*. The tree is based on maximum-likelihood analysis, using a 50% conservation filter. The scale bar indicates 10% estimated sequence divergence. *Thermoplasma acidophilum* was used as the outgroup for analysis. Isolated sequences and their accession numbers are indicated in bold type. The nearest matches from the NCBI databases are included. T indicates that the sequence originates in the species type strain.
difference in sensitivity is despite an overlap of AHL acyl chain length detection by the two biosensor strains (Steindler and Venturi, 2007). A. tumefaciens A136 utilizes the TraR QS response regulator system and responds to a broad range of acyl chain lengths in AHL molecules (Steindler and Venturi, 2007). C. violaceum responds to a shorter range of acyl chain lengths in AHL molecules and utilizes the CviR QS response regulator (Steindler and Venturi, 2007). The differential responses of these two biosensors highlights the advantage of using multiple biosensors when screening for QS mimics where chain length sensitivities may have little applicability.

**Isolated Bacteria**

This study strongly suggests that isolobopytolide, the major secondary metabolite in L. compactum, is an important selection factor regulating the microbial community of this soft coral. Firstly, we demonstrated that isolobopytolide can interfere with the QS activity of sensor strains, and secondly we demonstrated that addition of isolobopytolide to culture media increased the number and morphological variation of colonies produced from L. compactum. Moreover, the latter result suggests that the inclusion of secondary metabolites in growth media can improve the success of culturing soft coral associated bacterial isolates.

Most isolates generated in this study had high sequence identity with bacterial sequences sourced from the marine environment, including marine invertebrate hosts (Supplementary Table 5). Many of the recovered genera have also previously been isolated from coral mucus samples including Alteromonas, Bacillus, Endozoicomonas, Erythrobacter, Micrococcus, Pseudoalteromonas, Shewanella, and Vibrio (Lampert et al., 2006; Nithyanand and Pandian, 2009; Pootakham et al., 2017). The isolates were dominated by gammaproteobacteria belonging to the family Vibrionaceae, a result that is consistent with previous observations in scleractinian corals (Kvennefors et al., 2010, 2012). For both coral species, several Vibrio strains were isolated whose sequences clustered together and separately from the most closely related database sequences and hence may represent novel species. Scleractinian corals have previously been recognized as harboring a number of novel bacterial taxa (Rohwer et al., 2002; Sunagawa et al., 2010). This situation still remains, with amplicon-based studies of coral microbiomes returning many unassigned OTUs (Blackall et al., 2015). The bacteria of alcyonacean corals are less well studied and it is reasonable to assume that a similar situation could exist.

Multiple strains capable of inducing and/or disrupting QS in the bacterial biosensors were isolated from L. compactum and S. flexibilis (57.5 and 57.8% of tested strains, respectively). The genera of all the bacteria isolated, regardless of activity detected in this study, have previously been reported to possess or interact with QS systems (Ansaldi et al., 2002; Long et al., 2003; Waters and Bassler, 2005; Case et al., 2008; Tait et al., 2009, 2010; Nithya et al., 2010; Albuquerque and Casadevall, 2012; Lade et al., 2014), providing support for the hypothesis that QS is one of the mechanisms regulating coral associated microbial communities.

There was no clear taxonomic pattern of QS activity within the Vibrionaceae, which is consistent with a previous study that assessed QS activity in 29 Vibrionaceae strains (Tait et al., 2010). It is well recognized that QS in Vibrio spp. is tightly regulated by environmental conditions including host-released cues and nutritional status (Waters and Bassler, 2005). Vibrio spp. are ubiquitous in the marine environment (Urakawa and Rivera, 2006), however many Vibrio strains have been implicated in disease either as primary or opportunistic pathogens (Urakawa and Rivera, 2006). Given that QS is involved in the regulation of a number of the genes involved in pathogenicity (de Kievit and Iglewski, 2000; LaSarre and Federle, 2013), the presence of a wide range of Vibrio spp. with QS capabilities in otherwise healthy corals warrants further investigation to elucidate which genes are under QS control in these species.

“Spongiobacter,” now recognized as belonging to the genus Endozoicomonas (Neave et al., 2016), was originally recovered from a marine sponge (Pike et al., 2013) but is also present in many gorgonian (Sunagawa et al., 2010; La Rivière et al., 2013) and scleractinian corals (Raina et al., 2009; Blackall et al., 2015; Bourne et al., 2016). “Spongiobacter” strains have been attributed a number of ecological roles; “Spongiobacter” strains from A. millepora demonstrated a dependence on DMSP and consequently a role in the biogeochemical sulfur cycle was postulated (Raina et al., 2009), whereas, “Spongiobacter” strains from the sponge Suberites carnosus demonstrated antibacterial activity (Flemer et al., 2012). Of greatest interest to this study is the QS activity detected in “Spongiobacter” strains from the sponges Mycale laxissima and Ircinia strobilina (Mohamed et al., 2008). The Endozoicomonas-related strains SF102 and SF204 from S. flexibilis that were tested in this study induced QS activity in C. violaceum CV026 and not in A. tumefaciens A136, whereas Mohamed and coworkers found the opposite response (positive in A. tumefaciens and negative in C. violaceum).

In this study, QS activities were assessed for the culturable fraction of bacteria associated with the mucus of two soft coral species. In future, functional gene analysis and gene expression analysis may allow a more complete assessment of the genes that are responsible for and regulated by QS in these bacteria. Moreover, new –omics techniques will allow investigations of quorum sensing genes and their expression also in bacteria that cannot easily be cultured with standard methods. A combination of culture-independent studies and manipulative experiments using isolates holds great promise for further elucidation of QS mechanisms in soft coral holobios.

**CONCLUSION**

This research establishes a framework for the importance of QS and the identity of potential QS mimics within the soft coral holobiont, highlighting the potential value of soft corals as a model system for both structural and ecological investigations of QS mimics. The results presented here
clearly show that cembranolides and furancembrenes are partially responsible for previously observed QS interference in soft coral extracts. Their QS interference translates to a potentially new structural backbone for QS mimic compounds. The size of the oxygenated ring had more bearing on the activity expressed than the presence or position of epoxides, double bonds or acetate groups, an observation which extends the structural understanding of QS mimics. QS interference extended however also to soft coral species not known to contain cembranolides and furancembrenes, suggesting that new structural backbones with QS activity remain to be elucidated. The presence of both QS metabolites and QS bacteria within soft corals supports the role of QS as a way of mediating soft coral associated microbial communities. If the active compounds in these extracts are indeed produced by the soft coral, this interaction with QS could be important to the health and resilience of the host organism and may reflect a more widespread strategy of sessile marine invertebrates.

ETHICS STATEMENT

Soft coral samples were legally collected between 2009 and 2015 under Great Barrier Reef Marine Park Authority (GBRMPA) permits (G09/30327.1, G12/35236.1).

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this manuscript will be made available by the authors, without undue reservation, to any qualified researcher.

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AUTHOR CONTRIBUTIONS

MF experimental design, sample collection, lab work, data analysis and write-up. LH experimental design, data analysis and write-up. BB experimental design, sample collection, data analysis and write-up.

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SUPPLEMENTARY MATERIAL

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