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The community profiles of symbiotic bacteria at the different life stages of coral *Dipsastraea favus*

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Corals live in a symbiotic relationship with various bacteria that are fundamental to host fitness, health, and survival. Though the diversity of symbiotic bacteria has been revealed in the early life stages of some corals, the dynamic bacterial community profiles of one coral are still poorly characterized, particularly the stage-specific bacteria. In this study, the bacterial communities in the patent, eggs, and 4-day-old planula larvae of a hermaphrodite coral *Dipsastraea favus* were investigated by high-throughput sequencing of 16S ribosomal RNA gene. As a result, dynamic profiles of bacterial diversity in the planula larvae was a bit higher than that in the *D. favus* parent, and distinct stage-specific symbiotic bacteria were detected, e.g., Oceanospirillaceae, *Kordia*, and Legionellaceae in *D. favus* larvae, and Kiloniellales and *Prosthecochloris* in adult *D. favus*. The dynamic change of bacterial community in coral adults and larvae may expand our understanding of the complex relationship between coral host and its symbiotic microbiota.

KEYWORDS

Dipsastraea favus, life stage, bacterial community, high-throughput sequencing, horizontal transmission

Introduction

Coral holobionts harbor a complex consortium of microorganisms, including zooxanthellae, bacteria, archaea, fungi, viruses, and protists (Knowlton and Rohwer, 2003). Coral microbial flora has been attracting attention for their potentials to protect corals from environmental stresses and their relevance with coral hosts, such as defense against pathogens, promotion of coral health, development, nutrition-based symbioses, and environmental adaption (Krediet et al., 2013; Soffer et al., 2015; Lema et al., 2016; Yang et al., 2016; Zhou et al., 2017; Bernasconi et al., 2019; Babbin et al., 2021; Gavish et al., 2021; Kitamura et al., 2021). For example, the confirmation of anaerobic nitrogen metabolism in coral microorganisms sheds new light on coral and reef productivity (Babbin et al., 2021). Pollock et al. (2018) found evidence of coral-

microbe phylosymbiosis, in which coral microbiome composition and richness reflect coral phylogeny.

The majority of coral microbiology studies focus on coral adults (Rohwer et al., 2002; Ceh et al., 2011). To date, the coral-bacteria symbiosis particularly the relationship between the coral host and its intracellular bacteria is just beginning to be acknowledged (Kitamura et al., 2021; Maire et al., 2021). Bernasconi et al. (2019) indicated that the uptake and structure of bacterial communities were developmental, supporting the idea that microbial communities are likely to play specific roles within the distinct life history stages of the coral host. The establishment of coral microbial community in the early developmental stages of corals is fundamental to coral fitness, health, and survival (Apprill et al., 2009; Lema et al., 2014; Lema et al., 2016). However, though the diversity of symbiotic bacteria has been revealed in early life stages of some corals (Sharp et al., 2010; Sharp et al., 2012; Lema et al., 2014; Zhou et al., 2017; Epstein et al., 2019), we still rarely know about the dynamic profiles of bacteria at the different development stages of one coral because it is difficult to get enough male and female gametes and larvae at the early development stage of corals. Thus, investigation on the dynamic community profiles of symbiotic bacteria at the different life stages of one coral is with great importance for our understanding of the possible roles of bacterial symbionts, particularly stage-specific bacteria, in the host's development and coral-bacteria symbioses (Sharp et al., 2010; Lema et al., 2014).

Dipsastraea favus is a hermaphrodite, broadcast-spawning species coral belonging to the family Merulinidae (Shlesinger and Woesik, 2021), which is able to release gamete bundles with both sperms and eggs during its spawning period. In this study, with the seawater as control, the bacterial diversity in the coral *D. favus* parent, eggs and larvae was investigated, respectively, using high-throughput sequencing of 16S ribosomal RNA gene, with the aim to reveal the coral's dynamic profiles of symbiotic bacteria and consequently find stage-specific bacteria for the future understanding of *D. favus*-microbe symbioses and utilization of *D. favus* symbiotic microbes.

Materials and methods

Coral spawning and larval cultivation

Eight colonies of hermaphrodite *D. favus* (ca. $30 \times 40 \times 5$ cm³), 10 m away from each other, were collected at the Egong bay (114°29'E,

22°29'N), at a depth of 5 m, Shenzhen, China, approximately 1 month before spawning, and cultivated in pools with circulating natural seawater, and the temperature was kept at 26°C. The spawning occurred on 9 June 2017. The gamete bundles from each of the eight coral parents were captured and transferred to separate experimental tanks. After floating to the surface, the gamete bundles ruptured and released eggs and sperm. The eggs were collected and fixed immediately after being washed by sterile seawater. The seawater in tanks was not changed after spawning to avoid loss of sperm and kept at 26°C. Approximately 4 days after fertilization, most larvae tended to swim close to the substrate and searched for settlement locations, indicating that they were ready to settle and metamorphose soon; thus, we collected the 4-day-old planula larvae before they became settled juveniles. Because of the number limitation of eggs, no juveniles were collected in this study. The coral eggs, 4-day-old larvae, and parent fragments (Figure 1) were washed by sterile seawater several times to remove the surface microbes and fixed in RNALater. In order to provide enough biomass for successful PCR amplification, at least 50 eggs and 20 individuals of 4-day-old larvae were collected as one replicate for each coral parent. Approximately 2,000 ml of seawater in tanks was filtered by a 0.22µm filter as one replicate. All samples with three duplicates were frozen in -80°C for further processing.

DNA extraction and high-throughput sequencing

For coral adult DNA extraction, fragments of approximately 1 cm² including mucus, tissue, and skeleton were ground in liquid nitrogen by mortar and pestle. Proteinase (20 ml) was added to egg using Qiagen DNeasy Plant Mini Kit (cat. no. 69104), following the manufacturer's instructions. The extracted DNA was preserved at -20° C. DNA samples, including eggs, larvae, adult clones, and tank seawater (3 replications for each), were PCR-amplified using the following primers: forward primer (CCTACGGRRBGCASCAGKVRVGAAT) and reverse primer (GGACTACNVGGGTWTCTAATCC), which were designed by GENEWIZ (Suzhou, China), targeting the hypervariable regions V3–V4 of 16S ribosomal RNA gene for bacteria. The PCR solution contained 2.5 µl of TranStart Buffer, 2 µl of dNTPs, 0.5 µl of TransStart Taq DNA polymerase, 20 ng of total DNA, 1 µl of each primer, and ddH₂O in a final volume of 25 µl. The PCR amplification



FIGURE 1 The eggs (left), larvae (middle), and adult (right) of *D. favus*.

reaction was performed with the following procedure: initial denaturing at 94°C for 3 min; 35 cycles at 94°C for 5 s, 57°C for 1.5 min, and 72°C for 10 s; final extension at 72°C for 5 min. The library quality of PCR products was validated using Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA). The library concentration was quantified by Qubit2.0 Fluorometer (Invitrogen, Carlsbad, CA). All the amplicon products were sequenced on an Illumina Miseq platform according to the manufacturer's instructions, using 2X300 bp mode at GENEWIZ (Suzhou, China).

Sequence analysis

Analysis of the 16S rDNA sequences was conducted using the bioinformatics package QIIME 1.6 (Caporaso et al., 2010). A total of 74,9631 raw reads were generated from the nine samples (three replications for coral parent, eggs, and larvae, respectively). The sequences containing the adapter were discarded by Trimmomatic (Bolger et al., 2014). The paired-end sequence was merged using Pandaseq (Masella et al., 2012) with the parameter (minimum sequence length of 430 bp and maximum length of 470 bp). The chimera sequences detected by Usearch 61 were abandoned. The clean reads were clustered using UCLUST. Operational taxonomic units (OTUs) were selected at 97% identity by pick_open_reference_otus.py with the greengenes database (DeSantis et al., 2006). The OTUs annotated as chloroplast, mitochondria, and unassigned were discarded. After quality control, an average of 27,839 sequences were selected from each sample, ranging from 11,299 to 46,463. For a fair comparative analysis, samples were subsampled to an equal depth of 11,229 and were conducted by core_diversity_analyses.py. Summarized taxonomic abundance at the phylum, class, order, family, and genus levels was analyzed by QIIME.

Statistical analysis

Alpha diversity statistics including OTU, Chao1, and Shannon index were calculated in QIIME by alpha_diversity.py. The differences among groups were analyzed by one-way ANOVA with Tukey's HSD test. OTU count data were square-root-transformed. Bray-Curtis distance matrix was built to examine the patterns of community structure and was visualized using non-metric multidimensional scaling (NMDS) analysis. Pearson correlation vectors were overlaid to demonstrate which taxon has strong correlation with group difference. Multidimensional statistical analysis was performed in PRIMER v7 with the PERMANOVA+ add-on (PRIMER-e, Quest Research Limited, Auckland, NZ).

Results and discussion

A total of 250,555 high-quality 16S rRNA gene sequences (clean reads in Table 1) were obtained from all of the nine larva and adult samples of *D. favus* and seawater sample (three replications each). The rarefaction curves showed that the diversity reached a plateau (Figure S1). It was a pity that no 16S rRNA gene sequences were retrieved from the egg samples by egg DNA sequencing. The number of 16S rRNA gene reads, number of OTUs, and alpha diversity (Shannon and Chao) estimate were calculated with OTUs at 3% dissimilarity (Table 1). The alpha diversity of the bacterial community of D. favus larvae is a bit higher than that of adult D. favus based on the Shannon index, and the bacterial diversity shows statistical difference among the coral parent, larvae, and seawater samples (Table 1). The main shared bacterial OTUs among the three samples belong to the class of Rhodobacteraceae and Alteromonadaceae (Table S1). The shared OTUs in the coral parent and its larvae mainly belong to Bacteriodaceae and Flavobacteriales, despite the fact that these bacteria were not detected in the seawater (Table S2). In the case of seawater control, its bacterial community is composed of Alphaproteobacteria, Gammaproteobacteria, and Bacteroidetes (Figure 2).

The dominant 16S rRNA sequences from 4-day-old coral larvae are Gammaproteobacteria and Bacteroidetes (Figure 2). In contrast, Alphaproteobacteria dominates the 16S rRNA gene sequences recovered from adult F. favus, representing ~71% of the total sequences (Figure 2). The most abundant OTUs in the larvae are unidentified Oceanospirillaceae (OTU2) and Kordia sp. (OTU5), while the unidentified Kiloniellales (OTU1) and Prosthecochloris sp. (OTU10) are abundant in the adults (Figure 3). Based on non-metric multi-dimensional scaling analysis and permutational multivariate analysis of variance (PERMANOVA), we found obvious variations in the bacterial community composition among the D. favus adults, larvae, and seawater (PERMANOVA: p = 0.03) (Figure 4), e.g., Oceanospirillaceae, Kordia, and Legionellaceae in D. favus larvae, Kiloniellales and Prosthecochloris in adult D. favus, while Oceanospirillaceae, Rhodobacteraceae, and Sediminicole in seawater. Clearly, Figure 4 shows that the D. favus adults and larvae host different predominant bacteria. Similarly, significant stagespecific changes were detected in bacterial profiles during the

TABLE 1 Alpha diversity of bacteria in coral and the environmental seawater samples.

Sample	Raw reads	Clean reads	Observed OTUs	Chao1	Shannon
Coral egg	/	/	\	\	١
Coral larva	99,108 ± 16,883.19	29,069 ± 11,217.87	557.76 ± 32.69(a)	717.72 ± 31.632(a)	5.79 ± 0.14(b)
Coral adult	100,181 ± 5,852.11	12,723 ± 1,281.38	613.8 ± 26.94(a)	752.14 ± 20.46(a)	5.42 ± 0.07(a)
Seawater	50,588 ± 35,24.03	41,725 ± 5,064.39	917.2 ± 12.55(b)	1,729.39 ± 28.83(b)	6.14 ± 0.03(c)

Data are the average of three replicates. Letters indicate the significant differences based on the analysis of one-way ANOVA and Tukey's HSD test: same letter indicates that there is no significant difference.



development of scleractinian coral *Acropora gemmifera* (Zhou et al., 2017). Lema et al. (2014) suggested dynamic overall bacterial communities associated with the coral *Acropora millepora* at early life stages (larva, 2-week-old juveniles, and 3-, 6-, 12-month-old



Heat map reflecting the relative abundance and distributions of the top 20 OTUs. The scales 0.01 and 0.1 show the abundance at 5% and 10%, respectively.

juveniles). Damjanovic et al. (2020) also suggested that *Pocillopora acuta* recruits harbored dynamic and diverse bacterial assemblages.

We detected abundant Oceanospirillaceae in the larvae of D. favus rather than in adult D. favus (Figures 3, 4). Oceanospirillaceae has been identified from some stony corals (Morrow et al., 2012; Rodriguez-Lanetty et al., 2013). Oceanospirillaceae was only detected in 1-week old juveniles of A. millepora and disappeared in the later development juveniles (Lema et al., 2014). The same pattern occurred in the Caribbean coral Porites astreoides, which harbors high abundant Oceanospirillaceae in 4-day-old larvae while less Oceanospirillaceae is presented in 11-day-old larvae (Sharp et al., 2012). Previous studies demonstrated that the Oceanospirillales, belonging to Oceanospirillaceae, are able to metabolize DMSP in the coral A. millepora, which indicates that Oceanospirillaceae may play an important role in the coral-microbiome interactions because DMSP is a carbon and sulfur source for bacteria and produced by symbiotic dinoflagellates (Bourne et al., 2013). Moreover, the genus Kordia, which was abundant in the D. favus larvae (Figures 3, 4), was previously detected in the sea urchin and the sea anemone (Becker et al., 2009; Herrera et al., 2017). The genus Kordia is one of many genera affiliated with the family Flavobacteriaceae of the phylum Bacteroidetes, well known for its degradation of high-molecularweight organic matters. The genome of Kordia antarctica IMCC3317T encodes many carbohydrate-active enzymes, some of which are clustered into approximately seven polysaccharide utilization loci, thereby demonstrating the potential for polysaccharide utilization (Lim et al., 2020).

In contrast to *D. favus* larvae, we detected abundant bacteria Kiloniellales (Figures 3, 4) in the adult *D. favus*, which were previously detected in sponges and coral larvae (Sharp et al., 2012; Cleary et al., 2013). In the coral *Montastraea annularis*, Kiloniellales is significantly more abundant in healthy corals than the diseased corals and able to utilize nitrates through denitrification, indicating its positive function on coral's health (Soffer et al., 2015). The future research on the metabolic function of Kiloniellales associated with corals may explain its high abundance in *D. favus*. Another abundant bacterium in adult *D. favus* is *Prosthecochloris* (Figures 3, 4). *Prosthecochloris* sp. is a green sulfur bacterium, frequently detected in corals *Platygyra carnosa* and *Porites lutea* (Li et al., 2014; Cai et al., 2017).



Prosthecochloris sp. may provide organic and nitrogenous nutrients to coral host (Yang et al., 2016).

In this study, the most abundant bacteria shared by the larvae and adults are the unidentified Rhodobacteraceae (Table S1), which has been detected at the early development stages of corals (Sharp et al., 2012; Zhou et al., 2017). Despite the fact that Rhodobacteraceae is generally associated with diseased corals (Roder et al., 2014), the OTUs belonging to Rhodobacteraceae are also observed in healthy corals, such as the Oculina patagonica and Isopora palifera (Chen et al., 2011; Rubio-Portillo et al., 2016). The genus Roseobacter belonging to the family Rhodobacteraceae is reported to have an important function, that is, supplying fixed organic nitrogen to Symbiodiniaceae or defending against pathogenic bacteria (Nissimov et al., 2009). The Alteromans (OTU836759) and HTCC2207 (OTU805719) belonging to the Alteromonadaceae were also detected in both D. favus larvae and adults in this study (Table S1). A similar bacterial community was detected in the larvae of D. favus like other corals, indicating that these bacteria may be important at the early stages of corals.

Based on this study, some core bacteria were shared among the coral larvae, adults, and seawater despite their small proportion, suggesting that the coral D. favus is able to obtain bacteria from environmental seawater. The fact that abundant bacterial population in the coral larvae, adults, and the seawater is different (Figure 4) indicates that the selection of bacterial partners by the coral host is not random. Similarly, some studies also suggest the possible horizontal transmission of symbiotic bacteria from the environmental seawater to corals during their development because of the dynamic microbial community change. For instance, Lema et al. (2014); Lema et al. (2016) demonstrated horizontal (environmental) uptake of coral bacteria, e.g., nitrogen-fixing bacteria. Apprill et al. (2009) revealed that the composition of bacteria differed between life stages, i.e., prespawned oocyte bundles, spawned eggs, and week-old planulae, and concluded that the onset of association between microorganisms and the coral P. meandrina appeared to occur through horizontal uptake by planulae older than 79 h. Epstein et al. (2019) found that only a small number of bacterial strains were shared between offspring and their respective parents, revealing bacterial establishment as largely environmentally driven at the early life stages and suggested that recruits harbored more variable microbiomes compared to their parents, indicating that winnowing occurs as corals mature (Epstein et al., 2019).

On the contrary, Sharp et al. (2010) suggested a bacterial vertical transmission strategy based on the finding of bacteria in the ectoderm layer in newly released planula larvae of P. astreoides, but it was not known whether there were bacteria in the gametes of P. astreoides. Leite et al. (2017) found bacteria from the gamete bundles to the planula larvae and parental colony of M. hispida based on sequencing, and suggested that spawning coral *M. hispida* could vertically transfer microbes to the offspring based on the similar bacterial community in different stages of M. hispida, but the presence of bacterial cells in association with the (female and male) gametes of M. hispida through microscopy was not verified. Zhou et al. (2017) found that the microbial communities in the eggs of A. gemmifera were highly similar to those in parental colonies, indicating a transmission from coral parents to eggs. However, the possibility that some bacteria are eventually transmitted inside the gametes or larvae from the environments cannot be excluded. Most likely, the coral offspring can be partly colonized postspawning through the uptake of microbial associates released by the parental colonies into the surrounding seawater. In the case of coral D. favus in this study, no bacteria were detected by the 16S rRNA gene amplification from the egg samples, but we could not exclude the presence of bacteria in D. favus eggs because the failure in detecting the bacterial sequences in eggs may result from the very low abundance of bacteria or the primers and PCR procedure used. Meanwhile, in this study, some of the bacteria shared between the coral parents and larvae, and stage-specific bacteria were not detected in the seawater, suggesting the possible vertically transfer. Whether there is a vertical transmission of bacteria by coral D. favus needs future in-depth study, e.g., the confirmation of bacteria in the eggs or gametes of D. favus using electron microscopy and fluorescence in situ hybridization (FISH). Meanwhile, the characteristics of the symbiotic community profiles of D. favus in different regions need to be wholly investigated using abundant

samples from all life stages, e.g., gametes, larvae, and the early juvenile.

According to the results from this study, the larvae of D. favus harbor a bit higher number of diverse bacteria in comparison to adult D. favus. This may be explained by the different selective ability of corals at different life stages. It has been proposed that morphology change at different life stages may provide different complex niches, which may lead to a significant shift in the microbial community structure (Zhou et al., 2017). An initial higher diversity of bacterial community at the early life stage of corals, followed by low species diversity associated with the adults, has been observed (Franzenburg et al., 2013; Lema et al., 2014). The same situation occurs in the coral Acropora gemmifera, which has higher bacterial diversity at the larvae and 6-day-old juvenile stages than in adults (Zhou et al., 2017). The higher diversity of bacteria at the early life stage (e.g., larvae period) of corals provides a higher possibility for nutrient cycling due to the presence of special bacteria, such as Alteromonas and Roseobacter, which have been suggested to help coral larvae acquire additional nutrients in the oligotrophic environment and consequently increase the survival rate and fitness of corals at the early life stage (Lema et al., 2016). Moreover, the higher diversity of bacteria in the larvae may increase the possibility for more production of antibiotics, which can protect coral larvae from pathogens (Sharp et al., 2012; Lema et al., 2014).

In summary, we investigated the dynamic profiles of bacteria in the coral *D. favus* adult and its eggs and larvae by the high-throughput sequencing method, with the seawater as control. The *D. favus* parent and its larvae host variant bacterial community with larvae have a higher diversity. The shared bacteria among *D. favus* adult and its larvae, and seawater suggest that *D. favus* is able to acquire specific bacteria from the environment. The observed stage-specific bacteria, e.g., Oceanospirillaceae, *Kordia*, and Legionellaceae in *D. favus* larvae, and Kiloniellales and *Prosthecochloris* in adult *D. favus*, could be an important clue to explain the biological or ecological functions of bacteria in *D. favus* holobionts.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession

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Author contributions

JX, ZX, and XY finished the experiment. JX, GC, and YX analyzed the data. BL and BX participated in the experiment design. JX and ZL designed the study plan and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2023.1055848/ full#supplementary-material

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