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# Article Diversity and Biogeography of Coral Mucus-Associated Bacterial Communities: The Case of Acropora formosa

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Abstract: The role of microorganisms in coral health, disease, and nutrition has been demonstrated in various studies. Environmental factors including pH, temperature, and dissolved oxygen also play crucial roles in maintaining sustainable coral ecosystems. However, how geographical and environmental factors influence bacterial diversity and community composition is unclear. Here, bacterial communities associated with Acropora formosa coral were sampled from four different locations-Phu Quoc Islands (Vietnam), Nha Trang (Vietnam), Ujung Gelam (Indonesia), and Bourake (New Caledonia)—and compared using tagged 16S rRNA sequencing. We identified 24 bacterial phyla, 47 classes, 114 orders, and 495 genera from 18 samples. Overall, Proteobacteria (1039 distant amplicon sequence variants [ASVs]) and Firmicutes (589 ASVs) were predominant, while Verrucomicrobiota (75 ASVs) and Planctomycetota (46 ASVs) were minor taxa. Alpha diversity analyses revealed that the bacterial community associated with Acropora formosa from Ujung Gelam had the highest indexes (Observed and Chao1), while the figures for Bourake were the lowest. Non-metric multidimensional scaling analysis (NMDS) showed significant differences in bacterial communities among locations (ADONIS,  $p = 1 \times 10^{-4}$ ). Temperature was strongly correlated with the distribution of bacterial communities in Bourake, whereas pH and dissolved oxygen were significantly correlated with the presence of coral-associated bacterial communities in Phu Quoc and Nha Trang. Across all samples, 28 potential biological markers and 95 core ASVs were found, revealing significant differences in coral-associated bacterial communities. Collectively, these findings provide a comprehensive understanding of bacterial communities living in coral reefs across different geographic sites, which could be useful springboards for further studies.

Keywords: corals; bacteria; microbial diversity; barcoding; core microbiome

## 1. Introduction

Coral ecosystems are one of the most biologically diverse systems on Earth due to their remarkable adaptation to short-term natural phenomena [1]. Corals can persist in habitats with relatively extreme conditions, such as shallow pools, reef flats, and mangroves [2]. However, coral reefs are endangered by various environmental stressors such as ocean pH, increasing temperature, and solar radiation [3] as well as biotic stressors such as pathogenic bacteria and viruses, predators, and the loss of endosymbiotic microalgae [4,5].

Some coral-associated microbiota continually play both mutualistic and commensal roles by exhibiting pivotal functions in coral metabolism, including nitrogen fixation, carbon fixation, carbon degradation, and sulfur metabolism [6]. Other types of bacteria serve as coral-antibiotic agents by inhibiting the growth of potentially harmful microorganisms [7]. Besides the presence of commensal bacteria, accumulating evidence indicate strong correlations between bacterial pathogens and coral diseases. For example, *Vibrio* 



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**Copyright:** © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). *shiloi* is a bacterial pathogen that causes the bleaching of several scleractinian corals [8,9], whereas *Aspergillus sydowii* and *Serratia marcescens* are causative agents for aspergillosis [10] and white pox disease [11], respectively.

The composition of coral-associated bacterial communities has been demonstrated to be dependent on several contrasting factors, including geographic location, season, environmental conditions, and coral health [12,13]. Our earlier studies showed significant shifts in coral mucus-associated microbial composition in healthy and bleached corals [14,15]. Recent studies have revealed that *Endozoicomonadaceae* are the dominant bacterial group associated with corals in Korea and Japan; *Mycoplasmataceae* dominate in Taiwan [1]; and *Campylobacteraceae*, *Francisellaceae*, and *Pasteurellaceae* are the most likely primary pathogens to colonize the Caribbean *Acropora* coral reefs [16].

There have been increasing studies on the relationship between environmental factors and coral bleaching that elucidate the critical roles of environmental stressors such as pH, temperature, solar irradiance, and free radicals of oxygen [3,17]. In this study, we assumed that there were relationships between environmental factors and the composition of the microbiome of coral located in different sites worldwide. Thus, an investigation of the composition of bacterial communities associated with *Acropora formosa* was conducted on four geographic locations, including Phu Quoc Island (Vietnam), Nha Trang (Vietnam), Ujung Gelam (Indonesia), and Bourake (New Caledonia Islands). The association between the coral core microbiomes and geographical and environmental factors is discussed in our study, which provides a comprehensive view of the species *Acropora formosa* and its related microorganisms to create a premise for further studies.

#### 2. Materials and Methods

#### 2.1. Sample Collection and DNA Extraction

Colony fragments of Acropora formosa (n = 2), a widespread ubiquist species present in the global ocean, were collected between 3 and 5 m depth by scuba diving in shallow coral reefs of Hon Xuong Island of Phu Quoc islands, Vietnam (9°55'20.6" N 104°01'16.4" E). Coral mucus was collected from two Acropora formosa colonies in two different locations of this island. Next, the samples were taken out of the water with 3 min of air exposure. The mucus secretion that was triggered by this desiccation stress consisted of long gellike threads dripping from the coral surface. The first 30 s of mucus production was discarded to prevent contamination and dilution by seawater. Then, the mucus samples were collected using sterile syringes and transferred from syringes to sterile cryotubes, where they were immediately fixed with 30% glycerol solution in a ratio of 1:1 then stored at −20 °C until further use [14]. DNA extractions were processed using the Easy-DNA<sup>TM</sup> gDNA Purification Kit (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA) following the manufacturer's instructions, and 500  $\mu$ L bacterial DNA was extracted from different coral mucus samples. To assess whether bacterial DNA used for analysis was contaminated during the DNA extraction and polymerase chain reaction (PCR) stages, we used distilled water as the negative control in the PCR, as it contained all reaction components except for the template DNA.

In Phu Quoc, water pH, temperature, and dissolved oxygen were measured from a depth just above the corals (3 to 5 m) using handheld meters such as a Hach refractometer (USA). For other locations, we only used data that were published in NCBI, with the accession number listed in Table 1.

Sampling Sites	Accessions Numbers	Species	Sample ID	рН	Temperature	Dissolved Oxygen
Phu Quoc	This study	Acropora formosa	PQ1	<b>7</b> .0( ) 0.01	27.8 ± 0.09	6.96 ± 0.11
(9°55′20.6″ N 104°01′16.4″ E			PQ2	$7.96 \pm 0.01$		
Nha Trang (Whale Island (12°39.1′ N, 109°23.9′ E)	SRP116049 [14]	Acropora formosa	NT1		$29.5\pm0.4$	$5.92\pm0.15$
			NT2			
			NT3	$8.19\pm0.05$		
			NT4			
			NT5			
Ujung Gelam (S 05°49′57.7″, E 110° 22′50.5″)	SRP071125 [18]	Acropora formosa	UG1	7.3 ± 0.1	30.1 ± 0.4	6.4 ± 0.6
			UG2			
			UG3			
			UG4			
Bourake-Lagoon (21°56.915 S; 165°59.577 E)	PRJNA510614 [19]	- Acropora formosa -	BR1	7.3	33	2.3
			BR2			
			BR3			
			BR4			
			BR5			
			BR6			
			BR7			

Table 1. Sampling sites and environmental factors associated with Acropora formosa coral.

#### 2.2. PCR Amplification and 16S rRNA Gene Sequencing

Amplification of the partial 16S rRNA gene with the purified template DNA was established by using the universal bacterial primer sets 343F (S-D-Bact-0343-a-S-15, 5'-ACGGRAGGCAGCAG-3') and 802 R (5'-TACCAGGGTATCTAATCCT-3') [14], 2X PCR Taq Master Mix (Thermo Fisher Scientific, Waltham, MA, USA). These primers were attached with specific 6-bp barcode sequences at the 5' end and then used for DNA amplification to produce 16S rRNA amplicons. A ~460-bp fragment belonging to the V3–V4 region of the 16S rRNA gene was amplified. PCR amplifications were performed in an Eppendorf 6331 Nexus Gradient MasterCycler Thermal Cycler (Hampton, NH, USA) as follows: 30 cycles at 94 °C for 5 min, 94 °C for 30 s, 65 °C for 30 s, 72 °C for 1 min, and a final extension at 72 °C for 2 min. The pyrosequencing sequencing method of the 16S rRNA gene was performed on the 454 Titanium platform. For further analysis, the obtained files (FASTA and QUAL) were combined into the FASTQ format using the Galaxy server (http://usegalaxy.org (accessed on 27 August 2021)) [20].

#### 2.3. Data Cleaning and Analyses

The dataset used in this study comprised a combination of four sub-datasets: 16S rRNA gene sequences of bacteria living in *Acropora formosa* collected in Hon Xuong, Phu Quoc, Vietnam; Van Phong Bay, Nha Trang, Vietnam (accession number: SRP116049); Ujung Gelam, Indonesia (accession number: SRP071125); and Bourake, New Caledonia (accession number: PRJNA510614) utilizing the format of FastQ files [14,18,19]. In this study, metagenomic data were collected from different regions of the 16S rRNA gene. Accordingly, raw sequences of bacterial communities from Phu Quoc and Nha Trang were amplified from the 16S rRNA gene V3–V4 region, while those from Ujung Gelam and Bourake belonged to the V5–V6 region. To reduce the influence of differences in data between DNA regions as well as locations, only taxa from phylum to genus were selected by using the tax\_glom function in R.

The protocol for processing raw sequence data reads included filtering and trimming low-quality sequences, denoising, inferring sequence variants, constructing an ASV table, and assigning taxonomy, as described by Callahan in 2016 [21]. Data analysis was performed using the DADA2 pipeline (version 1.8)—R Studio (version 3.6.1) with some modifications to optimize access of the dataset. Chimera filtering was performed by the "removeBimeraDenovo" function of the "dada2" package, while taxonomy was assigned using the Silva taxonomic training data formatted for DADA2 (SILVA ribosomal RNA gene database project version 132) [22]. Microbiome relative abundance was assessed by the "phyloseq" package, and bacteria comprising high percentages in the communities were visualized by the "ggplot2" package at phylum and genus levels.

Alpha diversity metrics were calculated to compare the microbial diversity of the samples, which included the observed ASVs, the Chao1 richness estimator, and the Shannon– Weaver index. To minimize the impact of the difference in sample size, we conducted a rarefaction analysis based on resampling using the rarefy function (the vegan package) in R. Comparisons of the alpha diversity indices among samples were tested using the analysis of variance (ANOVA) test, and Tukey's honestly significant difference test (Tukey's HSD) was performed on the ANOVA results to compare alpha diversity among locations.

Beta diversity was assessed by calculating the compositional similarities of all 18 samples from four locations on the basis of environmental parameters such as pH, temperature, and dissolved oxygen. Before performing the beta diversity analysis, the centered log ratio (CLR) transformation was used for the normalization of data distribution. A non-metric multidimensional scaling (NMDS) plot (based on the computed Bray-Curtis dissimilarity between samples) was visualized using the envfit function and Vegan package in R. To evaluate the statistical significance of differences among the four locations, ADONIS and betadisper (type = "centroid") functions were used to test the homogeneity of dispersion and similarity of composition among groups [23]. ANOVA was used to test for differences in beta dispersion among locations.

High-dimensional biomarkers were accessed using the linear discriminant analysis effect size, LEfSe [24], which emphasizes both biological relevance and statistical significance to identify bacterial taxa characterizing the differences between communities. Explaining the differences between biological classes and estimating the sizes of significant variations helped clarify relevant features.

The core ASVs in coral microbiomes were defined using different percentage cut-offs ranging from 30% to 100% [25]. In the current study, the presence of ASVs in at least 80% of samples was chosen as a conservative representation of the core microbiome. To detect core ASVs, we used the function "core\_members" in R. Appropriate taxa were chosen and displayed by using the functions add\_besthit and plot\_core in R, respectively. The core microbiomes of *Acropora formosa* from four locations were identified using the "microbiome" package and illustrated with an Upset diagram and a heatmap chart.

#### 3. Results

#### 3.1. Overview of the Sequencing Dataset and Data Preprocessing

The composition of bacterial communities living in *Acropora formosa coral* was investigated based on 16S rRNA gene sequences. A total of 1,390,711 raw reads were obtained from 18 coral mucus samples collected from four locations. Among these, 64,474 (2%) raw reads were collected from Phu Quoc. An ASV table was generated from non-chimeric sequences, including information about ASVs with corresponding abundances (the number of reads with number of ASVs). The total number of sequencing reads and filtered reads for all samples are presented in Table 2.

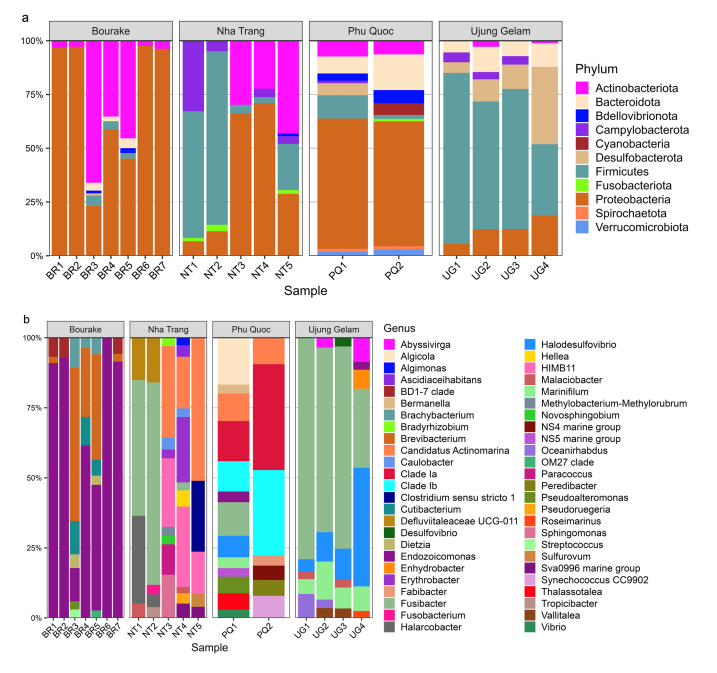
Sampling Sites	Species	Sample ID	No. of Raw Sequences	No. of Sequences after QC	Observed ASVs	Chao1	Shannon
Phu Quoc	Acropora formosa	PQ1	31,064	21,432	231	322.22	4.79
		PQ2	33,409	6008	174	210.67	4.38
Nha Trang	Acropora formosa	NT1	49,936	15,729	85	95.20	3.65
		NT2	52,510	28,100	77	96.46	3.33
		NT3	75,709	60,117	37	37.50	3.02
		NT4	65,007	52,221	54	54.50	3.47
		NT5	45,634	7970	123	143.32	3.67
Ujung Gelam	Acropora formosa	UG1	186,435	133,877	169	250.00	3.89
		UG2	195,464	128,950	236	426.83	4.56
		UG3	232,120	175,525	228	436.41	4.44
		UG4	145,104	75,380	142	266.25	3.81
Bourake	Acropora formosa	BR1	65,875	52,843	28	28.60	2.29
		BR2	55,164	43,544	37	41.00	2.62
		BR3	32,452	28,524	157	262.64	3.19
		BR4	10,782	9288	81	114.06	2.37
		BR5	4635	3764	65	70.35	2.55
		BR6	76,145	62,827	31	35.02	2.32
		BR7	33,266	28,381	24	29.00	2.05

Table 2. Alpha diversity estimates of bacterial communities associated with Acropora formosa.

#### 3.2. Taxonomic Assignment of Coral-Associated Bacterial Communities

In total, 24 bacterial phyla, 47 classes, 114 orders, and 495 genera were assigned to the coral mucus samples by DADA2 taxonomic classification. Overall, the surveyed bacterial communities residing in the corals were dominated by the phylum Proteobacteria (1309 distant ASVs), followed by Firmicutes (589 ASVs), Bacteroidota (564 ASVs), Desulfobacterota (151 ASVs), and Actinobacteria (151 ASVs). In addition, other phyla, including Bdellovibrionota (108 ASVs), Campylobacterota (84 ASVs), Verrucomicrobiota (75 ASVs), and Planctomycetota (46 ASVs), were shown in lower abundances (Figure 1a). At the genus level, *Endozoicomonas, HIMB11*, and *Vibrio* were majorly shown in Proteobacteria, whereas *Fusibacter, Halodesulfovibrio*, and *Marinifilum* were the predominating genera in the phyla Firmicutes, Desulfobacterota, and Bacteroidota, respectively (Figure 1b).

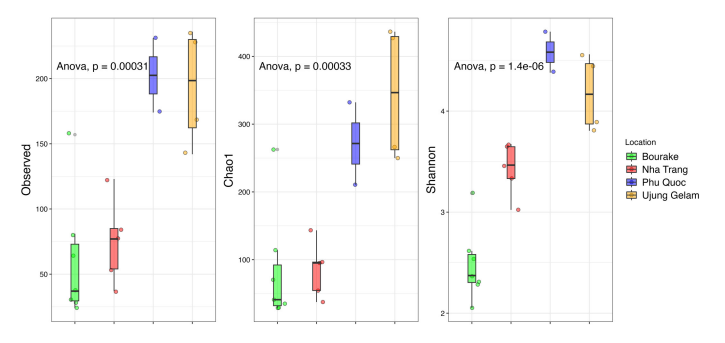
Comparative analysis based on locations indicated that the phylum Proteobacteria comprised the largest proportion of the bacterial communities from the most samples that were collected from Phu Quoc (56.6–59.6%) and Bourake (22.9–97.5%). Additionally, the relative abundance of the phylum Firmicutes displayed a significant increase in the coral bacteriome in Ujung Gelam (ranging from 32.8–79.1%). Similarly, the phylum Campylobacterota (Nha Trang, 32.7%) was found in greater abundance in bacterial communities at this location than at others. At a deeper taxonomic rank, significant variations of relative abundances in some bacterial genera were obtained depending on the coral sampling sites (Figure 1b). The numbers of assigned bacterial genera obtained from Ujung Gelam (276 genera) were higher than those of three other locations, thus demonstrating the higher diversity of bacterial communities at these locations. *Endozoicomonas* (96.5.%), *Fusibacter* (66.7%), *Candidatus\_Actinomarina* (39.9%), and *Clade\_ Ia* (23.33%) were the major bacterial genera across coral samples at Bourake, Ujung Gelam, Nha Trang, and Phu Quoc, respectively (Figure 1b).



**Figure 1.** Taxonomic classification of bacterial diversity at (**a**) phylum and (**b**) genus levels. The bar charts show the composition and abundance of bacterial communities at the phylum (relative abundance > 1%) and genus levels (relative abundance > 2%).

#### 3.3. Diversity and Biogeography of Bacterial Communities

Microbial alpha diversity indicated substantial variability among coral samples at different locations (Figure 2). After performing rarefaction analysis to remove rare ASVs (singletons and doubletons) or low-abundance ASVs, we retained 18 samples containing 98.83 ASVs on average, with a range of 24–236 per sample (Table 2). For each location, the total number of ASVs was Phu Quoc Island, 405; Nha Trang Island, 376; Ujung Gelam, 775; and Bourake, 303. The bacterial composition in Phu Quoc exhibited the largest number of bacterial species/ASVs (236), whereas Bourake exhibited the smallest number of species (24). Likewise, the Chao1 index indicated a significantly higher diversity of bacteria living in the *A. formosa* located at Ujung Gelam (436.41) than that of Bourake (28.60) and other



locations. The Shannon indices had the lowest and highest values in Bourake (2.05) and Phu Quoc (4.79) (Figure 2 and Table 2).

**Figure 2.** Alpha diversity of microbial communities associated with *Acropora formosa*. The line inside the box represents the median, while the whiskers display the lowest and highest values within the 1.5 interquartile range (IQR). Sample values are shown as dots.

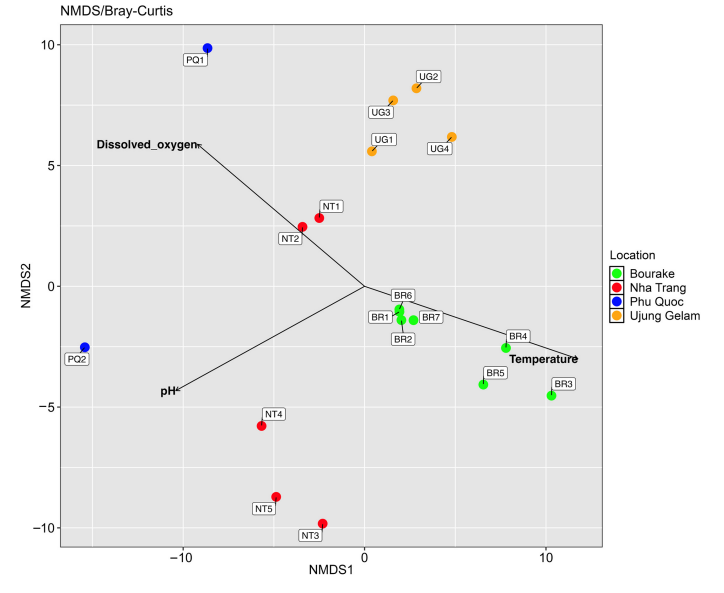
Alpha diversity was estimated by observed ASVs, Chao1, and the Shannon index. Calculations of alpha diversity showed that Ujung Gelam had higher species richness estimates (ANOVA, p < 0.01) than other locations, whereas that for Bourake was the lowest (Figure 2, Table S1). According to the Tukey's HSD test (Table S1), there were significant differences in the richness estimates (observed ASVs and Chao1) between pairwise comparations such as Phu Quoc–Bourake ( $p = 4.95 \times 10^{-3}$ ), Ujung Gelam–Bourake ( $p = 1.06 \times 10^{-3}$ ), Phu Quoc–Nha Trang ( $p = 1.50 \times 10^{-2}$ ), and Nha Trang–Ujung Gelam ( $p = 5.06 \times 10^{-3}$ ). Additionally, the Shannon diversity index did not change significantly between Phu Quoc and Ujung Gelam ( $p = 5.18 \times 10^{-1}$ , Table S1).

Rank-based methods such as NMDS, which is one of the best ways to visualize beta diversity, were used to illustrate the dissimilarities of the bacterial communities between sampled sites. Each symbol in the plot represents a bacterial community residing in the samples. When they were ordinated closer together, those communities were likely more similar, and vice versa. The non-metric multidimensional scaling analysis (NMDS) of microbial communities in four locations indicated that samples clustered based on their sites. In particular, Phu Quoc and Nha Trang samples were on the left side, while Bourake and Ujung Gelam samples were on the right side of the NMDS plot. Samples from Bourake or Ujung Gelam tended to be grouped closely together compared to other locations (Figure S1). In contrast, Phu Quoc or Nha Trang revealed a clear separation between samples and had more variance in bacterial community compositions (Figure S1).

Measures of the distance to the group centroid revealed that the locations Nha Trang (10.65) and Phu Quoc (10.12) had the highest heterogeneity (a greater distance from the centroid) compared to the others, while Ujung Gelam (6.48) had the lowest heterogeneity. This was supported by the NMDS plot (Figure S1), where the samples from Phu Quoc and Nha Trang showed more variance than Ujung Gelam and Bourake.

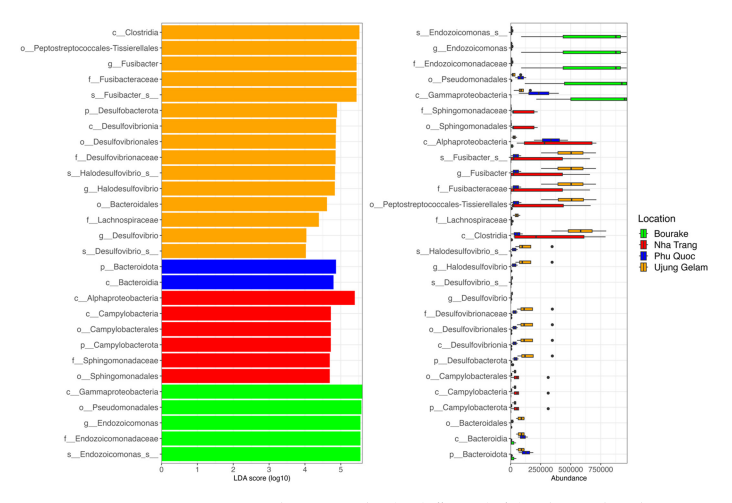
PERMANOVA indicated significant differences in bacterial composition among the sampling locations (ADONIS,  $R^2 = 0.55$ ,  $p = 1 \times 10^{-4}$ , Table S2). However, there were no significant differences in beta-dispersion by collection location (ANOVA, p = 0.09, Table S3).

Envfit analysis indicated that the temperature factor was significantly correlated (p < 0.05) with the ordination of microbial communities in Bourake, while pH and dissolved oxygen had significant influence on the distribution of bacterial communities in Phu Quoc and Nha Trang (Figure 3). However, in Ujung Gelam, the influence of environmental conditions such as pH, temperature, and dissolved oxygen on the bacterial composition was not observed.



**Figure 3.** Non-metric multidimensional scaling (NMDS) of microbial communities. Sample ordination based on bacterial taxonomic composition across samples is illustrated by an NMDS plot based on Bray–Curtis dissimilarity. Vectors represent the correlation between environmental variables and axes.

LEfSe was specifically designed to discover biomarkers in genomic data; thus, we applied the tool to the 16S rRNA gene to detect the differences in bacterial abundances across the four coral sampling sites. Bacterial taxa represented organisms comprising the majority in the communities, and colors indicate the sampling sites where a difference was observed. A total of 28 bacterial groups were differentially abundant (Linear discriminant analysis (LDA) score > 4.0) between the coral groups collected from the four different locations (Figure 4). Among these, 15 biomarkers were discovered in Ujung Gelam, 6 in



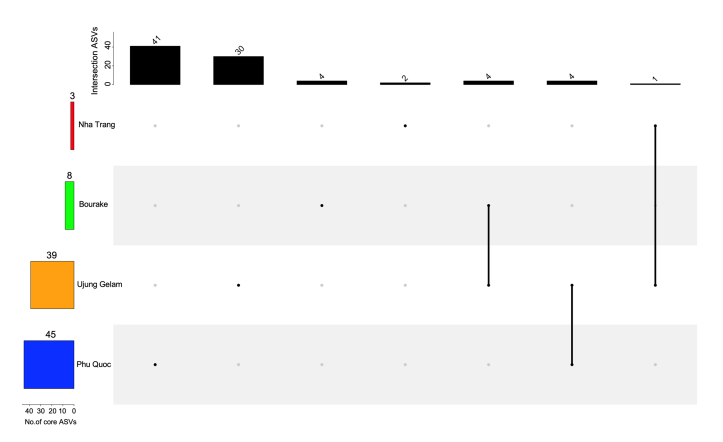
# Nha Trang, and 5 in Bourake, and the location with the lowest number of biomarkers was Phu Quoc (2).

**Figure 4.** Linear discriminant analysis (LDA) effect size (LEfSe) results on coral microbiomes. Taxonomic distribution of bacterial groups according to LDA score (on the left side) and abundance (on the right side).

As can be seen, there were considerable statistical differences among coral bacteriomes in the four studied sites. The coral samples from Ujung Gelam had the highest diversity of bacterial taxa (from species to phylum), of which most belonged to the phyla Desulfobacterota and Firmicutes, with LDA scores ranging from 4.52 to 5.52. Despite having less diversity than Ujung Gelam, Bourake had higher LDA scores, varying between 5.55 and 5.60 (Table S4).

### 3.4. The Coral Core Microbiome

Using the Upset plot (Figure 5), we identified a total of 95 ASVs, which are considered to be the core microbiome across all four sites. Phu Quoc contributed the highest number of coral core microbiome ASVs (45), while the figure for Ujung Gelam was lower, at 39 ASVs, and that for Nha Trang was lowest, at only 3 ASVs. Ujung Gelam had the largest number of shared ASVs, with a total of 9 ASVs (4 for Ujung Gelam and Phu Quoc, 4 for Ujung Gelam and Bourake, and 1 for Ujung Gelam and Nha Trang). Notably, none of the ASVs were shared across all sites.



**Figure 5.** Overview of the coral core microbiome composition in *Acropora formosa* coral. Upset diagram showing the number of shared and unique ASVs in four different locations.

A comparative analyis of core microbiome taxa revealed that each group harbors some specific taxa. In particular, genera *Clade\_Ia*, *Clade\_Ib*, and *Candidatus\_Actinoma* exist only in Phu Quoc (Figure S2), whereas *Endozoimonas* and *Brevibacterium* were considered as core genera in Bourake (Figure S3). Genus *Fusibacter* was found in both Nha Trang and Ujung Gelam (Figures S4 and S5, respectively).

#### 4. Discussion

One of the goals of this study was to investigate the bacterial diversity and composition in four different locations (Phu Quoc, Nha Trang, Bourake, and Ujung Gelam). Differences in the bacterial community composition across geographical locations have previously been described in Acropora species [6]. Although Proteobacteria were present in most of the locations, there was a notable shift in their abundance among locations, with the highest abundance recorded at Bourake and the lowest at Ujung Gelam, where Firmicutes was the most dominant phylum. At lower taxa, our results showed that the composition of the coral bacteriome strongly differed by location, where 28 potential biological markers were found by LEfSe. This suggests that microbial community dissimilarity arises with geographical distance, as was reported in recent studies [6,26–28]. In other aspects, however, environmental conditions can be considered main drivers of changes in coral bacterial communities. By comparing the beta diversity among four locations, we found that seawater dissolved oxygen, temperature, and pH also play important roles in the distribution of samples, except for samples from Ujung Gelam. This was in agreement with previous studies, as it showed the impact of environmental conditions on the bacterial community of the coral reef [29,30].

The influence of environmental variables upon bacterial diversity and community can also be examined through analyses of coral core microbiomes. *Fusibacter* and *Ery-throbacter* were the major bacterial genera in Nha Trang. According to Mhuantong et al., (2019), the rising abundance of *Fusibacter* was found in coral samples with white band

disease [18]. However, *Erythrobacter* is thought to play a crucial role in enhancing coral holobiont tolerance to heat stress [2]. Thus, the temperature was not a determining factor for the distribution of microbial composition in Nha Trang. Some genera, such as *Fusibacter*, *Halodesulfovibrio*, *Marinifilum*, and *vibrio*, were considered as core microbiome in Ujung Gelam (Figure S5). However, most of them were detected in diseased coral samples [18,31,32]. Thus, we thought that microbial composition and diversity in *Acropora formosa* were influenced not by a single determinant but resulted from a complex combination of different factors.

Another finding was the presence of *Endozoicomonas* and *Brevibacterium* in coral core microbiomein Bourake, which has the environmental conditions of high temperature and low dissolved oxygen. Generally, *Endozoicomonas* are recognized to play an important role within the coral holobiont due to their widespread predominance in many coral species [16,33–35] and apparent metabolic versatility [35,36]. Furthermore, the decrease in the abundance of *Endozoicomonas* in stressed, diseased, or bleached corals has been reported, which suggests that intensive abundance of *Endozoicomonas* is a potential indicator of habitat suitability [16,34,37–39]. *Brevibacterium* is thought to be capable of producing palytoxin, a chemical compound that offers nutrition for corals [40]. In Bourake, all samples were healthy. Thus, in addition to geographical and environmental factors, the health state of coral samples was an important factor, having a close correlation with bacterial community diversity.

### 5. Conclusions

In this study, we compared the composition and abundance of bacterial communities associated with the scleractinian coral *Acropora formosa* in four geographic sites. According to diversity analysis, there were 28 potential biological markers across all samples, which revealed that coral-associated bacterial communities exhibited significant differences among the studied sites. The results of the core microbiome analysis showed that 95 core ASVs were detected in different environmental conditions. Additionally, our findings indicated that the bacterial community composition in Nha Trang, Phu Quoc, and Ujung Gelam could be affected by environmental conditions such as pH, temperature, and dissolved oxygen.

Overall, our study provides new elements to confirm the strong sensitivity of the coral microbiome to environmental conditions and, therefore, its susceptibility to climate changes.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/jmse11010074/s1.

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