Phylogeography of Blue Corals (Genus Heliopora) Across the Indo-West Pacific

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Species delimitation of corals is one of the most challenging issues in coral reef ecology and conservation. Morphology can obscure evolutionary relationships, and molecular datasets are consistently revealing greater within-species diversity than currently understood. Most phylogenetic studies, however, have examined narrow geographic areas and phylogeographic expansion is required to obtain more robust interpretations of within- and among- species relationships. In the case of the blue coral Heliopora, there are currently two valid species (H. coerulea and H. hiberniana) as evidenced by integrated genetic and morphological analyses in northwestern Australia. There are also two distinct genetic lineages of H. coerulea in the Kuroshio Current region that are morphologically and reproductively different from each other. Sampling from all Heliopora spp. across the Indo-Pacific is essential to obtain a more complete picture of phylogeographic patterns. To examine phylogenetic relationships within the genus Heliopora, we applied Multiplexed inter simple sequence repeat (ISSR) Genotyping by sequencing (MIG-seq) on > 1287 colonies across the Indo-West Pacific. Maximum likelihood phylogenetic trees indicated the examined Heliopora samples comprise three genetically distinct groups: H. coerulea group, H. hiberniana group, and a new undescribed Heliopora sp. group with further subdivisions within each group. Geographic structuring is evident among the three species with H. hiberniana group found in the Indo-Malay Archipelago and biased toward the Indian Ocean whilst Heliopora sp. was only found in the Kuroshio Current region and Singapore, indicating that this taxon is distributed in the western Pacific and the Indo-Malay Archipelago.
**INTRODUCTION**

Coral reef ecosystems have the highest biodiversity of all marine ecosystems (Fisher et al., 2015) and are important in terms of their scientific, economic, and social value (Moberg and Folke, 1999; Spalding et al., 2001; Cesar et al., 2003). However, reef-building corals, the foundation organisms in coral reefs, have been rapidly deteriorating (Hughes et al., 2003; Altieri et al., 2017; Eyre et al., 2018) due to coral mass bleaching and other threats associated with the on-going climate change (e.g., global warming and ocean acidification) (Hoegh-Guldberg et al., 2007; Hoegh-Guldberg and Bruno, 2010; Hoegh-Guldberg, 2011; Hughes et al., 2018); local anthropogenic impacts (e.g., red-soil runoff, nutrient enrichment, and marine plastic pollution) (Halpern et al., 2008; Omori, 2011; Huang et al., 2020; de Oliveira Soares et al., 2020); outbreaks of corallivorous animals (Birkeland and Lucas, 1990; Yasuda, 2018; Montalbetti et al., 2019) and coral disease (Bruno et al., 2003, 2007; Burge et al., 2014). Approximately one-third of the world’s reef-building corals are facing an elevated risk of extinction (Carpenter et al., 2008; Wilkinson, 2008), but many more cryptic species may be undescribed (Bickford et al., 2007) and if the natural range of species is misunderstood, their threatened status may be under-estimated (Richards et al., 2016).

The classification of reef-building coral species has traditionally been based mainly on morphological characteristics (Veron, 2000; Fabricius and Alderslade, 2001). In some cases, however, morphological classification has been difficult due to a high level of phenotypic plasticity across geographic and depth gradients, and in some genera, there is a lack of discrete morphological features that can be used to underpin species identifications (e.g., Forsman et al., 2009; Martí-Puig et al., 2014; McFadden et al., 2017). More recently, genetic information based on several molecular markers has shown morphology can conceal cryptic evolutionary relationships (Richards et al., 2013) and it is increasingly common for genetically distinct cryptic species or lineages to be found within widespread coral species (e.g., Nakajima et al., 2012; Pinzon et al., 2013; Warner et al., 2015). Such discordance between morphology and genetics has led to confusion regarding the species boundaries in many groups (e.g., Acropora, van Oppen et al., 2001; Marquez et al., 2002, Pocillopora, Flot et al., 2008; Souter, 2010; Schmidt-Roach et al., 2013). Such ambiguity threatens to undermine effective biodiversity conservation efforts and prevents the true complexity of coral reef ecosystems from being properly understood.

Molecular data is not, however, always able to provide a definitive answer about species boundaries within reef-building corals. Molecular systematic studies have been partly hindered by the lack of appropriate genetic markers and the traditionally used mitochondrial markers have suffered from inadequate analytical resolution (Shearer et al., 2002; Ridgway and Gates, 2006; Bilewitch and Degnan, 2011). The ability to interpret nuclear datasets is also hampered by the multi-copy nature of genes (Vollmer and Palumbi, 2004) even those that were originally thought to be single copy (e.g., Pax-C, Rosser et al., 2017), introgressive hybridization events (Veron, 1995; Richards et al., 2013) and/or incomplete lineage sorting (e.g., van Oppen et al., 2001; Souter, 2010; Yasuda et al., 2015). Recent advances in High-Throughput Sequencing (HTS) have enabled the use of genome-wide polymorphisms such as restriction site associated DNA sequencing (RAD-seq, Miller et al., 2007; Baird et al., 2008; Rowe et al., 2011; and Dart-seq, Rosser et al., 2017) and Multiplexed inter simple sequence repeat (ISSR) genotyping by sequencing (MIG-seq, Suyama and Matsuki, 2015) to infer phylogenetic relationships. Such techniques often successfully delimit species of non-model organisms including corals with higher resolution than traditional genetic markers (RAD-seq, e.g., Pante et al., 2015; Herrera and Shank, 2016; Quattrini et al., 2019 and MIG-seq, e.g., Tamaki et al., 2017; Park et al., 2019; Hirai, 2019).

MIG-seq is an easy, cost-effective, novel method to obtain a moderate number of single nucleotide polymorphisms (SNPs) of non-model organisms with polymerase chain reaction (PCR) and HTS technology. The number of SNPs from MIG-seq analysis is generally less than those obtained using other techniques such as RAD-seq. However, MIG-seq has several advantages, since the method can be performed with small amounts and/or low-quality DNA and is relatively easy and cheap. Indeed, MIG-seq successfully revealed species boundaries of octocoral species that could be undetectable by traditional genetic markers (Richards et al., 2018; Takata et al., 2019). While some studies examined genetic relationships of closely related reef-building corals using HTS techniques in geographically restricted regions (e.g., Forsman et al., 2017; Johnston et al., 2017), only a few studies have analyzed speciation and/or genetic divergence patterns covering the Indo-Pacific scale (but see Warner et al., 2015; Nakajima et al., 2017; Arrigoni et al., 2020; Wepfer et al., 2020). For a more comprehensive picture of the phylogeographic structure of coral species in the Indo-Pacific, and to understand the species diversity of corals, it is necessary to obtain more extensive representation of species across their geographic range.

Increasingly recognized is the importance of integrating information about reproductive timing and physiology with genetic and morphological data to obtain more robust estimates of coral species (see Oliki et al., 2015; Rosser, 2015; Villanueva, 2016; Luzon et al., 2017; Richards et al., 2018). Timing of reproduction in broadcast spawning species is particularly...
important because gametes are viable for only a few hours, so individuals that spawn more than a few hours apart are unlikely to cross-fertilize (Levitan et al., 2011) and in this regard, reproductive timing heavily influences evolutionary patterns. Whilst integrated datasets that include reproductive data and cover a wide geographic area are emerging as the gold standard in phylogenetic studies, practical and logistical constraints often mean it is not possible to obtain complete sampling coverage in every region (Keyse et al., 2014; Wepfer et al., 2021). Nevertheless, examining the species boundaries, ecological preferences, and distribution of reef-building corals is fundamental for promoting effective conservation and management strategies.

The blue corals (genus *Heliopora*) are members of octocorals, though it has a well-developed aragonite skeleton like those of scleractinians (Quattrini et al., 2020). Owing to their hard skeleton, *Heliopora* spp. play an important role in coral reef accretion in various Indo-West Pacific reefs (e.g., Zann and Bolton, 1985; Planck et al., 1988; Abe et al., 2008; Takino et al., 2010), and therefore is an important reef-building coral taxon. *Heliopora coerulea* (Pallas, 1766) has mainly three growth forms, namely lobate, columnar, and encrusting (Dana, 1846), but with high morphological plasticity and continuous variations among growth forms (Eguchi, 1948; Colgan, 1984; H. Taninaka pers. obs.). *Heliopora coerulea* has long been regarded as a living fossil, being the sole extant member of the family Helioporidae (Colgan, 1984) until the recent description of *Heliopora hiberniana* Richards et al., 2018, a morphologically, genetically and reproductively distinct species from northwestern Australia in the Indian Ocean (Richards et al., 2018). To date, *H. coerulea* is known to be distributed throughout the Indo-Pacific realm (Wells, 1954; Bouillon and Houvenaghel-Crévecoeur, 1970; Zann and Bolton, 1985), whereas *H. hiberniana* is known from the north-west shelf of Western Australia, the Maldives, and the Wakatobi and Gili Islands in Indonesia (Richards et al., 2018, 2020).

*Heliopora* spp. are gonochoric, brooding corals (Babcock, 1990) with short larval dispersal durations (Hariri et al., 2002; Hariri and Kayanne, 2003). Due to such a low larval dispersal ability, genetic connectivity among geographically separate populations may be limited, a pattern that was evident within the scale of several hundred square kilometers (Taninaka et al., 2019). Therefore, it is possible that *Heliopora* includes further genetically distinct lineages in the Indo-Pacific. Previous genetic studies have indicated that additional cryptic species may indeed be present within *H. coerulea* along the Kuroshio Current region of the Philippines, Taiwan and Japan (Yasuda et al., 2014). Study from Yasuda et al., 2014 hypothesized that there are at least two genetically distinct lineages (HC-A and HC-B) of *H. coerulea* in the Kuroshio Current region that can also be distinguished by different gross morphologies (small branch for HC-A and flat shapes for HC-B, Yasuda et al., 2014, 2015; Eguchi et al., 2019). Subsequent observations (Saito et al., 2015; Villanueva, 2016) and histological examination (Taninaka et al., 2018) revealed that the reproductive timing of the two lineages is different by almost one month even under similar environmental conditions. However, the phylogeographical relationships among HC-A and B in the Kuroshio Current regions, and with *H. hiberniana* and *H. coerulea* in northwestern Australia are still unknown. The aim of this study was to clarify the phylogenetic relationships and distributions of the genus *Heliopora* across the Indo-West Pacific to gain insights on the speciation patterns of Indo-Pacific coral species. To achieve this goal, we collected *Heliopora* samples widely from the Indo-West Pacific region and applied genome-wide phylogenetic, phylogeographic, and population genetic analyses using the MIG-seq method.

**MATERIALS AND METHODS**

**Sample Collection and DNA Extraction**

A total of 1287 *Heliopora* samples from a total of 73 sites across three western Pacific regions (Japan, Taiwan and Guam) and Singapore, and three Indian Ocean regions (northwestern Australia, Thailand, and the Maldives) were collected using SCUBA from 2008 to 2019 (Supplementary Table 1). Additionally, we used previously collected samples from Japan and northwestern Australia (Supplementary Table 1). The samples include those from the northernmost distribution site (Yaku Island in Japan; Nakabayashi et al., 2017), and the known deepest depth (from 50 m depth from Green Island, Taiwan). All coral fragments were preserved in 99.5% EtOH and kept at −20°C until DNA extraction. During sampling, colonies were photographed, and GPS information was acquired wherever possible. All overseas sample collection was conducted through legal procedures and collection cooperation with the support of local research collaborating institutions; Academia Sinica in Taiwan, University of Guam in Guam, National University of Singapore in Singapore, Western Australian Museum and Curtin University in Western Australia, Phuket Marine Biological Center in Thailand, University of Milano-Bicocca and Marine Research and High Education Center in Maldives (see also FIELD STUDY PERMISSION). Genomic DNA was extracted from the preserved samples using the hot alkali solution method (Meeker et al., 2007) or QIAGEN DNeasy Blood & Tissue kit (QIAGEN, Hilden, Germany) on site. The first PCR procedure of MIG-seq analysis was carried out at each overseas laboratory except for Thailand samples from which ethanol preserved coral fragments were imported in 2011 (CITES number: AC.0510.2/407). We used them in this study under an additional permit from the Department of Marine and Coastal Resources in Thailand.

**MIG-seq Library Preparation and Sequencing**

All the DNA extraction and first PCR procedure except for domestic samples were carried out locally, and the first PCR products were transported to Miyazaki, Japan. The first PCR step was performed by using 8 pairs of multiplex ISSR primers of Suyama and Matsuki (2015). The fragments were amplified with the Multiplex PCR Assay Kit Ver.2 using a total volume of 7 μL reaction in a thermal cycler with the following modified profile: 94°C for 1 min followed by 29 cycles at 94°C for 30 s, 38°C for 1 min, 72°C for 1 min, and a final extension at 72°C for 10 min. The 50-fold diluted first PCR product was used as the template.
DNA for the second, tailed-PCR to add the individual index and the Illumina adapter sequence to each sample. PrimeSTAR GXL DNA polymerase (TaKaRa Bio Inc., Otsu, Shiga, Japan) was used for second PCR with a total volume of 12 µL reaction in a thermal cycler with a following profile: 20 cycles at 98°C for 10 s, 54°C for 15 s, 68°C for 1 min. The second PCR products were pooled as a single mixture library by using 1µL of each product. The mixed sample was electrophoresed on 0.1% agarose gel to verify amplification. The size ranged from 350 to 800 bp and DNA was manually extracted from the gel on a transilluminator. After extracting genomic DNA from the gel using FastGene Gel/PCR Extraction Kit (Nippon Genetics, Tokyo, Japan), library concentration was measured with a Qubit fluorometer (Thermo Fisher Scientific, Waltham, MA, United States). Finally, the DNA library was sequenced on a MiSeq Sequencer (sequencing control software v2.0.12, Illumina, San Diego, CA, United States) using a MiSeq Reagent Kit v3 (150 cycles; Illumina). Image analysis and base calling were performed using real-time analysis software v1.17.21 (Illumina). DarkCycle option was changed “Amplicon-dark 17-3” to “Amplicon-dark 17-17” on the “Chemistry” line (see also Suyama and Matsuki, 2015).

Sequence Processing and SNP Genotyping

The FASTX-toolkit version 0.0.14 (Gordon and Hannon, 2012), with a fastq-quality-filter setting of –Q 33 –q 30 –p 40, was used to eliminate low-quality reads and primer sequences from the MiSeq raw data. Adapter sequences for Illumina MiSeq were removed from both 5’ end and 3’ end using Cutadapt version 2.10 (Martin, 2011). Short reads less than 80 bp were excluded using an in-house python script. To further exclude contamination from Symbiodiniaceae-DNA, and to obtain larger numbers of loci (Shafer et al., 2017), the filtered raw reads were mapped onto the reference genome of Heliopora coerulea obtained from Symbiodiniaceae-free larvae (10.6084/m9.figshare.14356418) using Stacks version 2.2.0 (Catchen et al., 2011; Rochette et al., 2019) with the reference-aligned pipeline for single nucleotide polymorphism (SNP) discovery and genotyping. As a first step, the unpaired reads were removed using the repair.sh software tool within BBTools software package (Bushnell, 2017). Next, the retained reads were aligned to the reference genome using Burrows-Wheeler Aligner (BWA) version 0.0.17-r118 (bwa index and mem with default parameters) (Li and Durbin, 2009). Alignments were then compressed, sorted, and indexed with SAMTools version 1.10 (Li et al., 2009). Finally, SNP genotyping for each individual was carried out using the gstacks program in Stacks with default setting. SNP calling for each genetic analysis was carried out using the populations program in Stacks with the option -r to obtain different minimum proportions of genotyped individuals per locus (0.1 for phylogenetic analyses, 0.9 for assigning clones and population genetic analyses), and the option -ordered_export to ensure that only one of the overlapping SNPs is output from reference aligned data. All the output files from Stacks were converted to the appropriate formats for each genetic analysis using PGDSpider version 2.1.1.5 (Lischer and Excoffier, 2012).

Assign Clones

We used the software GenoDive version 3.04 (Meirmans, 2020) to assign possible clones within each population based on the infinite allele model (IAM) omitting all missing data. Less than 12 differences of multi-locus genotype within the same population were identified as possible clone mates (estimated by inspection of the pairwise distance histogram in GenoDive). All clone mates except for one individual with the least missing data were removed from this study. After removing possible clones, we selected up to three least missing data samples per sampling site. In case multiple different gross morphologies and/or genetically different lineages were found within the same site, up to three samples per morphology were selected. Finally, we rerun Stacks using this selected dataset excluding possible clones to obtain new SNPs for phylogenetic and population genetic analyses.

Phylogenetic Analysis

We estimated phylogenetic trees based on the Maximum likelihood (ML) method. The program IQ-TREE2 version 2.0.6 (Minh et al., 2020), using the TVM + F + R5 model selected based on the Bayesian Information Criterion (BIC), was used to reconstruct a phylogenetic hypothesis for Heliopora spp. The bootstrap analysis was performed with 1000 replicates using UFBoot, Ultrafast Bootstrap Approximation (Minh et al., 2013). The final tree was drawn using FigTree version 1.4.4 (Rambaut, 2012).

Population Genetic Analysis

We conducted Principal Coordinates Analysis (PCoA) to examine genetic relationships among Heliopora individuals using GenAlEx version 6.502 (Peakall and Smouse, 2012). In addition, Discriminant Analysis of Principal Components (DAPC) (adegenet package version 2.1.3 in R version 4.0.2) (Jombart, 2008; Jombart et al., 2010; Jombart and Ahmed, 2011) was performed to visualize the genetic structure. The lowest BIC was used to detect the optimal number of K clusters.

RESULTS

Number of Reads and SNPs From MIG-seq Analysis

In total 339,818,358 raw reads with an average of 264,039 reads per sample were obtained for 1,287 individuals. After filtering low-quality reads, 335,164,915 reads with an average of 260,423 reads per sample were obtained. After excluding index, adapter, and short reads less than 80 bp, 139,837,592 reads with an average of 108,654 reads were obtained. In total 67,119,478 reads were mapped onto the reference-genome with an average of 52,152 reads per sample. We used 795 SNPs across 1,287 individuals (r = 0.9) to find possible clones, and then excluded 576 individuals from the subsequently genetic analyses. We finally selected 245 individuals from 73 sampling sites from 7 regions (DRA Accession No. DRA012077) for phylogenetic (ML tree; 24,741 SNPs, r = 0.1) and population genetic analyses.
Taninaka et al. Phylogeography of Genus Heliopora (PCoA and DAPC; 1,092 SNPs, $r = 0.9$). For the latter, we selected 238 individuals from the final dataset after excluding seven individuals with a missing rate of more than 50% per 1,092 SNPs.

**Phylogenetic Relationships**

ML phylogenetic analysis revealed three major groups of *Heliopora* spp. in the Indo-West Pacific with 100% bootstrap values (Figure 1). The phylogenetic hypothesis presented here indicated that there is one new undescribed group (hereafter *Heliopora* sp. group) genetically different from the other two groups, each of which includes typical morphologies of the previously described species (slender-branching growth form *H. hiberniana* whose type locality is in the northwestern Australia; lobate growth form *H. coerulea* whose type locality is in the Indian Ocean). The “*Heliopora* sp.” group (shown in blue in Figure 1) comprises the HC-A lineage found in Kuroshio Current regions including Japan and Taiwan (Yasuda et al., 2010, 2014, 2015; Taninaka et al., 2019), and Singapore. The “*H. hiberniana*” group (shown in green in Figure 1) includes the holotype of *H. hiberniana* found in northwestern Australia (Richards et al., 2018) and the eastern Indian Ocean samples from Thailand and the Maldives. The “*H. coerulea*” group (shown in red in Figure 1) comprises mainly lobate growth morphs HC-B lineage found in Kuroshio Current regions including Japan and Taiwan (Yasuda et al., 2010, 2014, 2015; Taninaka et al., 2019), Guam, and *H. coerulea* found in northwestern Australia (Richards et al., 2018). In addition, there were genetically isolated subclades within each group that clustered together by different growth forms or geographic regions (Figure 1; sp.1–3, hib.1–2, and coe.1–3). The *Heliopora* sp. group was divided into three subclades: Singapore subclade (sp.1), HC-A with only columnar growth form subclade of Japan (sp.2), and HC-A with both encrusting and columnar growth forms subclade found in both Japan and Taiwan (sp.3). *Heliopora* sp.3 also included a sample collected from 50m depth in Green Island, Taiwan. A sample collected from Yaku Island in Japan, the northernmost distribution site, looks like an outgroup of sp.2 and sp.3 in the phylogeny, possibly because it had a hybrid genotype between *Heliopora* sp.2 and sp.3 based on STRUCTURE analysis (data shown in Supplementary Figure 1). The group “*H. hiberniana*” was divided into two subclades: northwestern Australia *H. hiberniana* subclade (hib.1) and a subclade consisting of Thailand and the Maldives samples (hib.2). The group “*H. coerulea*” was divided into three subclades; a
Guam subclade (coe.1), *H. coerulea* in northwestern Australia subclade (coe.2), and a subclade consisting of HC-B found in Japan and Taiwan samples (coe.3).

**Individual-Based Genetic Differentiation**

Consistent with the phylogenetic analysis result, population genetic analyses revealed three genetic isolated groups of *Heliopora* with eight subclades in the Indo-West Pacific (Figures 2A,B). The PCoA showed three clusters based on 11.73% (PC1) and 7.3% (PC2) of the total variability (Figure 2A). The DAPC also showed clear genetic separation of the three clusters and eight subclades (Figure 2B). K = 3 was selected supporting the ML phylogeny result, while K = 8 was determined as the optimal number of clusters by calculating the first 200 PCs for the maximum of 50 clusters based on BIC. The posterior DAPC assignments of both K = 3 and 8 were consistent with the prior clusters (all blue crosses were on red rectangles).

**Geographical Distribution**

The three groups of *Heliopora* indicated uneven geographical distribution (Figure 3A). The distribution of *Heliopora* sp. group was biased to western Pacific Ocean side including Kuroshio Current region (Japan and Taiwan) and the South China Sea (Singapore). On the contrary, the distribution of *H. hiberniana* group was mainly in the Indian Ocean side such as Kimberley region and Christmas Island in northwestern Australia, the Andaman Sea (Thailand), and the Maldives. The *H. coerulea* group was distributed both in the Indian Ocean and western Pacific ranging from the Guam and the Kuroshio Current region (Japan and Taiwan) to Kimberley and Christmas Island in northwestern Australia. The distribution of *H. coerulea* group...
FIGURE 3 | (A) Geographical distributions of the three Heliopora spp. groups. (B) Gross morphologies of the three Heliopora spp. groups. H. coerulea group: (a) lobate growth form in Guam (coe.1), (b) lobate growth form in northwestern Australia (coe.2), (c) lobate growth form in Japan and (d) Taiwan (coe.3), and (e) thick-branching growth in Guam (coe.1). H. hiberniana group: (f) slender-branching form in the type locality in northwestern Australia (hib.1), and (g) slender-branching and (h–j) lobate growth form in the Maldives (hib.2). Heliopora sp. group: (k) columnar growth form in Singapore (sp.1), (l) columnar growth form in Japan (sp.2), (m) columnar growth form in Japan (sp.3), (n) encrusting growth form in Japan (sp.3), and (o) encrusting growth form in the mesophotic zone (50 m) in Taiwan (sp.3).

was partly overlapping with the other two species groups in Japan, Taiwan, and northwestern Australia (Figure 3A, coe.2 and hib.1; coe.3 and sp.2-3).

Gross Morphology

The gross morphologies of the three groups of Heliopora were highly variable (Figure 3B). H. coerulea group included two typical morphs; lobate growth form (Figure 3B, a–d) and thick-branching growth form (Figure 3B, f–g) and lobate growth form (Figure 3B, h–j). Heliopora sp. group included two typical morphs; columnar growth form (Figure 3B, k–m) and encrusting growth form (Figure 3B, n–o). The typical growth forms of Heliopora sp. group were relatively distinct from other two species, while the morphologies of H. hiberniana group and H. coerulea group partly overlapped and were difficult to identify by gross morphology.

DISCUSSION

In the present study, we examined the phylogenetic relationships, gross morphological variations, and the geographical distributions of Heliopora spp. collected from the Indo-West Pacific region. Our phylogenetic reconstruction indicates that there are three genetically isolated species groups in the genus Heliopora and we provide evidence that there is substantial subcladal genetic structuring within each group.

We do not describe the newly found Heliopora sp. group or its subclades as new species of the genus Heliopora in this study. Cladistic analysis and formal species description are being undertaken separately.
**Phylogeography and Speciation Patterns of Heliopora**

Phylogenetic and population genetic analyses using genome-wide SNPs by MIG-seq showed that the Indo-West Pacific *Heliopora* was divided into three groups: *H. coerulea* group, distributed in the Indo-West Pacific and including HC-B in the Kuroshio Current region; *H. hiberniana* group distributed mainly in the Indian Ocean including the type locality of *H. hiberniana* in northwestern Australia; and a new undescribed *Heliopora* sp. group distributed mainly in the Western Pacific Ocean, including HC-A in the Kuroshio Current region (Figures 1, 2 and 3A; Supplementary Table 1).

The new undescribed *Heliopora* sp. group was distributed mainly in the Western Pacific (Japan and Taiwan) and Singapore in this study, while *H. hiberniana* group was distributed mainly in the Indian Ocean (northwestern Australia, the Andaman Sea in Thailand, and the Maldives in this study; distributed also in Indonesia, Richards et al., 2020). Such uneven geographical distributions imply the origin of *Heliopora* sp. group could be in the Pacific Ocean and that of *H. hiberniana* could be in the Indian Ocean, but this remains to be tested with a dated phylogeny. Cenozoic tectonic events (Hall and Holloway, 1998) and sea-level fluctuations associated with climate change (Pillans et al., 1998) in the area between the Indian and Pacific Oceans are thought to have promoted allopatric speciation in many coral reef organisms (Carpenter et al., 2011) including starfish (Benzie, 1999; Vogler et al., 2008) and reef fishes (Gaiter et al., 2011; Bowen et al., 2013). It is, therefore, possible that the genus *Heliopora* has also allopatrically speciated into the two groups, *Heliopora* sp. and *H. hiberniana*, due to the physical division between the Indian and Pacific Oceans caused by past tectonic and climatic changes. Morphological and molecular analyses on other Indo-Pacific reef-building coral species also found sibling species between the Indian and Pacific Oceans (Veron, 1995; van Oppen et al., 2001; Pinzon et al., 2013; Huang et al., 2014; Arrigoni et al., 2020; Wepfer et al., 2020).

The *H. coerulea* group is distributed in both the Indian and Western Pacific Oceans. The distribution of *H. coerulea* group partially overlaps with the *Heliopora* sp. group in the western Pacific and with the *H. hiberniana* group in northwestern Australia. Previous studies revealed sympatrically distributed *Heliopora* spp. have different reproductive timing: the reproductive timing of *Heliopora* sp. group (HC-A) and *H. coerulea* group (HC-B) in Japan (Saito et al., 2015; Taninaka et al., 2018) and the Philippines (Villanueva, 2016) differ by almost one month. *Heliopora hiberniana* and *H. coerulea* colonies in northwestern Australia have been found to have different reproductive timings even when found in sympathy (Richards et al., 2018). These patterns suggest that endogenetic regulation may contribute to differences in reproductive timing. A previous study revealed that there are fixed species-specific substitutions in the dopamine receptor 2-like gene (Isomura et al., 2013) and cryptochrome-1 (Oldach et al., 2017), both of which were identified as the genes that regulate reproductive timing in *Acropora* species (Iguchi et al., 2019). It is possible that allochronic speciation have occurred between these sympatrically distributed *Heliopora* spp. groups, or at least allochronism in reproductive timing plays a central role to keep their species boundaries after speciation. The difference in reproductive timing is an important mechanism as a prezygotic isolation for marine species that release gametes into the water column (Knowlton, 1993; Palumbi, 1994). Indeed, many of the sympatrically distributed broadcast spawning coral species have different reproductive timing in the same genus (e.g., *Orbicella*, Knowlton et al., 1997; Szmidt et al., 1997; *Acropora*, Fukami et al., 2003; Nakajima et al., 2012; Ohki et al., 2015). Generally, the reproductive timing of corals is also strongly associated with environmental factors (e.g., moon phases, moonlight, solar radiation, water temperature) in addition to genetic factors (e.g., circadian clock genes, photoreceptor proteins) that correlate with the environmental factors (e.g., Levy et al., 2007; Brady et al., 2009; Crowder et al., 2017; Oldach et al., 2017). Thus, future studies on reproductive timing of *Heliopora* spp. that include both environmental and genetic factors may provide further insights into speciation of sympatrically distributed *Heliopora* spp.

The *Heliopora* sp. group was found both in Yaku Island in Japan (N 30°16′21.45″, Nakabayashi et al., 2017), the northernmost site and at a depth of 50 m on Green Island in Taiwan, the deepest habitat records of the genus *Heliopora*. Therefore, contrary to previous knowledge on the ecological traits of *H. coerulea* that it prefers warm shallow water habitats (Zann and Bolton, 1985), *Heliopora* sp. group could be found in colder and deeper habitat; the northernmost known habitat of *H. coerulea* group (HC-B) is Miyako Island (N 24°51′52.10″, Yasuda et al., 2014) and that of *Heliopora* sp. group (HC-A) is Yaku Island (N 30°16′21.45″, Nakabayashi et al., 2017). In the field, contrasting ecological differences have been previously observed for *Heliopora* sp. group (HC-A) and *H. coerulea* group (HC-B). Colonies of *Heliopora* sp. group (HC-A) are relatively small and more abundant outside the well-developed fringing reef, where water temperature is lower, while colonies of *H. coerulea* group (HC-B) can be larger, forming micro-atolls, and dominating inner reef habitats where water temperature is higher (Yasuda et al., 2010; Taninaka et al., 2018), providing evidence of niche differentiation which further support the hypothesis of ecological speciation between these two groups.

**Gross Morphological Diversity of Heliopora**

There are some specific growth forms within each *Heliopora* group. Columnar and encrusting growth forms are relatively distinct from other morphologies and specific to *Heliopora* sp. group (Figure 3B, k–o). Slender-branching growth form with whitish skeleton (Figure 3B, f–g) is also specific to *H. hiberniana* group. However, lobate growth form (Figure 3B, a–d and h–j) can be found in both *H. coerulea* and *H. hiberniana* groups. Like
other reef-building coral species, young and/or juvenile stages are hard to distinguish among closely related species (Babcock, 1992; Babcock et al., 2003). Previous studies have also reported intermediate growth forms sometimes rendering it difficult to distinguish even between HC-A (Heliopora sp. group) and HC-B (H. coerulea group) possibly due to infrequent hybridization and phenotypic plasticity in the field (Yasuda et al., 2014; Taninaka et al., 2018). Therefore, not all colonies can be clearly classified into the Heliopora groups based on field observation alone.

Phenotypic plasticity and continuous morphological variation among closely related coral species are a common phenomenon and have hindered accurate morphological species identification (e.g., Forsman et al., 2009; Marti-Puig et al., 2014; McFadden et al., 2017). Light intensity and water movement are the most influential environmental factors responsible for coral phenotypic plasticity (Todd, 2008), while other physical environmental factors (e.g., dredging, sediment disturbances, turbidity, competition) are also correlated with morphological diversity, leading to continuous morphological variation of different species under similar environmental conditions (e.g., Darling et al., 2012; Erftemeijer et al., 2012; Swierts and Vermeij, 2016). However, it is still unclear what environmental factors influence gross morphological diversity in Heliopora spp.

Richards et al. (2018) previously reported the key morphological characteristics to distinguish H. coerulea from H. hiberniana, including gross morphology, the intensity of bluish skeletal color, and skeletal microstructure (e.g., highly elaborated coenenchymal echinulations and smaller and more numerous autopores), indicating skeletal color can be a key morphological characteristic. In the western Pacific, a few colonies with whitish skeletons have been found in the Heliopora sp. group in Japan (H. Taninaka and N. Yasuda pers. obs.). Contrary to the case in the northwestern Australian region, there was no genetic difference between the whitish and other bluish skeleton colonies, indicating skeletal color cannot be a universal key morphological character to distinguish Heliopora sp. from the other species.

Based on these findings, it is necessary to clarify the relationship between morphological diversity including fine skeletal structure and environmental and/or genetic factors among Heliopora spp. in more detail in the future.

Subclades Within Each Heliopora Group

Our phylogenetic tree showed substantial phylogenetic structure and multiple genetically isolated subclades exist within each Heliopora species group. These subclades partially correspond to typical growth forms and geographical distributions (Figures 1, 2 and 3A). In the Heliopora sp. group for example, the geographically distant Singapore subclade (sp.1) first diverged from Japan-Taiwan clades (sp.2 and 3). Then, the Japan-Taiwan clades were split into only columnar growth form subclade (sp.2) found only in Japan and encrusting and columnar growth forms subclade (sp.3) found in both Japan and Taiwan. These two subclades were genetically distinct despite being sympatric. In the field, some ecological differences can be observed between the two subclades: Heliopora sp.2 tends to be found in shallower inner reef areas with strong sunlight and relatively weak waves. Heliopora sp.3 tends to be found in relatively deep darker areas such as outer reef slope with fast water currents and turbidity (H. Taninaka pers. obs.). Further ecological and physiological differences between Heliopora sp.2 and Heliopora sp.3 would clarify the species status of the two subclades. The Heliopora sp.2 and Heliopora sp.3 subclades could not be distinguished in the previous studies using traditional genetic markers (e.g., mitochondrial DNA, ITS2, microsatellite and microsatellite flaking region sequences) (Yasuda et al., 2014, 2015; Taninaka et al., 2019), while this study corroborated the effectiveness of using genome-wide SNPs to elucidate detailed genetic boundaries between closely related lineages/species as in the case of other octocorals (Herrera and Shank, 2016; Quattrini et al., 2019; Takata et al., 2019).

In the H. hiberniana group, two subclades (hib.1-2) showed deep genetic divergence corresponding to geographical distance, namely the northwestern Australian subclade (hib.1) and the Maldives-Thailand subclade (hib.2). This genetic isolation could be the result of long-term restriction of gene flow between the two subclades. The current circulation system in the Indian Ocean mainly consists of currents that move from east to west that does not directly connect northern (Maldives and Thailand) and southern (northwestern Australia) sides of the mid-east Indian Ocean (Schott and McCreary, 2001; Schott et al., 2009). Due to such current patterns, previous studies on other marine organisms with pelagic larval phase also showed similar genetic isolation in the Indian Ocean (Benzie et al., 2002; Bay et al., 2004; Hui et al., 2016). On the other hand, geological research (Schettino and Turco, 2011; Keith et al., 2013; Obura, 2016) fossils, and phylogenetic data (Obura, 2012, 2016; Veron et al., 2015) propose a hypothesis that tectonic changes occurred during the Paleogene and the Neogene and may have promoted genetic isolation and speciation events in the Indian Ocean. Such historical geographic events might have promoted genetic divergence of Heliopora spp. in the Indian Ocean, but this requires further testing within a calibrated phylogenetic framework.

In the H. coerulea group, three subclades (coe.1–3) corresponding to geographically distinct regions were found, namely Guam subclade (coe.1), the northwestern Australia subclade (coe.2), and Japan-Taiwan subclade (coe.3). The most genetically isolated subclade that first split from the others is the Guam subclade (coe.1). It is notable that Japan-Taiwan subclade (coe.3) is genetically more closely related to the northwestern Australia subclade (coe.2) in the Indian Ocean than to the geographically closer Guam subclade (coe.1). It is likely that genetic connectivity has been restricted between Guam and Japan-Taiwan since the Guam subclade (coe.1) diverged from the other two subclades early in the history of H. coerulea. This is unusual because genetic connectivity between Mariana Islands region including Guam and the Kuroshio Current region is often strong in most of the marine invertebrate species with larval dispersal period (e.g., Palumbi et al., 1997; Williams and Benzie, 1998; Lessios et al., 2001; Pinzón et al., 2013; Arrigoni et al., 2020; Wepfer et al., 2020; but see exception...
in Wörheide et al., 2008). Although the reasons for the genetic break between Guam and Kuroshio Current region in Heliopora spp. are unclear, it could be caused by limited larval dispersal capability of Heliopora spp. (Taninaka et al., 2019) and/or local adaptation of each subclade. On the other hand, northwestern Australian populations of other marine species are genetically more similar to west Pacific populations than other Indian Ocean populations (e.g., Williams and Benzie, 1998; Lessios et al., 2001; Pinnzon et al., 2013; Arrigoni et al., 2020; Wepfer et al., 2020) but contrary examples also exist where Indian and Pacific Ocean coral populations are divergent (e.g., Richards et al., 2016). Williams and Benzie (1998) discussed that this might be due to Western Australia being recently colonized by recruits from the Western Pacific coastline after the end of the last ice age. Recent genome-wide phylogeographic analysis of reef-building corals have also found such large discrepancies between geographic and genetic distances (Arrigoni et al., 2020; Wepfer et al., 2020), which need to be examined in closer detail in the future.

**CONCLUSION**

The present study provides a comprehensive picture of phylogenetic relationships, distribution, and growth form patterns among closely related Heliopora species that gives new insights into speciation patterns of Heliopora spp. in the Indo-West Pacific. Our genome-wide genetic data of Heliopora spp. revealed three major groups with eight subclades. Heliopora sp. group was genetically distinct from the other two groups (H. coerulescens group and H. hiberniana group) containing the samples from type localities of the previously described species. Genetic diversification in the genus Heliopora could be attributed to allopatry, allochrony, and local and/or ecological adaptation. Further studies incorporating more genome-wide markers for phylogeographic analysis, geohistorical information, physiological and ecological data would more precisely delineate the boundaries of Heliopora species.

Our study also demonstrates the effectiveness of the MIG-seq method for clarifying the species boundaries of octocoral species that were indistinguishable using traditional genetic markers such as mtDNA and ITS2. In addition, we highlight the importance of wide geographic sampling to generate a more complete picture of the phylogeographic relationships and speciation patterns among closely related octocoral species and how they diversified across the Indo-Pacific. It is expected that the MIG-seq method can be applied to other coral species and non-model organisms to resolve longstanding questions in marine evolution.

**FIELD STUDY PERMISSION**

Domestic sampling except for Amami Oshima was conducted under each prefectural government permits; Yaku Island (Kagoshima Prefectural Government permit, No. 2–56), Okinawa including Okinawa Mainland, Kume Island, Ishigaki Island, Sekisei Lagoon, and Iriomote Island (Okinawa Prefectural Government permits, Nos. 18–34, 19–39, 19–60, 21–18, 22–15, 25–44, 26–10, 27–29, 27–81, 28–78, 30–42, and 31–18), and Amami Oshima was conducted under personal permit by Mr. Katsuki Oki. Foreign sampling was conducted under each local government permits; Taiwan (samples collected from Dabaisha and Gongguan, the Taitung County Government, No. 1074150258; a 50 m specimen collected in Chaoiku, Taitung County Government, No. 1040000285; and that is registered at the Zoological collection of the Biodiversity Research Museum in Academia Sinica AZIC0001313), Guam (SC-19-006 (cc), University of Guam Marine Laboratory), Singapore (National Parks Board Permit NP/RP16-156), Australia (samples in Western Australia were collected under permit number AU-COM2013-203 and Australian Government approval number 006-RRRW-130723-01), Thailand (the permission for collecting and sending the specimens is granted by the Department of Fisheries), Maldives (Permit No. (OTHR)30-D/INDIV/2017/123 released by the Maldivian Ministry of Fisheries and Agriculture). All the 1st PCR except for those from Thailand were conducted on site and only 1st PCR products were brought or sent to Japan. Ethanol-preserved coral fragment samples from Thailand were imported to Japan in 2011 (CITES number: AC.0510.2/407), and we used them in this study under an additional permit from the Department of Marine and Coastal Resources in Thailand.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: DDBJ (accession: DRA012077), FigShare (doi: 10.6084/m9.figshare.14356418 and doi: 10.6084/m9.figshare.14814231).

**AUTHOR CONTRIBUTIONS**

HT and NY conceived the study and drafted the manuscript. All authors except for TK and HY collected samples and conducted molecular genetic experiment. HT, TK, and HY conducted genetic analysis. All authors edited the draft and approved it for publication.

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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.2021.714662/full#supplementary-material

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