Genetic differentiation and connectivity of morphological types of the broadcast-spawning coral *Galaxea fascicularis* in the Nansei Islands, Japan

Yuichi Nakajima¹*, Yuna Zayasu², Chuya Shinzato², Noriyuki Satoh² & Satoshi Mitarai¹

¹Marine Biophysics Unit, Okinawa Institute of Science and Technology Graduate University, 1919-1 Tancha, Onna, Okinawa 904-0495, Japan
²Marine Genomics Unit, Okinawa Institute of Science and Technology Graduate University, 1919-1 Tancha, Onna, Okinawa 904-0495, Japan

**Abstract**

Population connectivity resulting from larval dispersal is essential for the maintenance or recovery of populations in marine ecosystems, including coral reefs. Studies of species diversity and genetic connectivity within species are essential for the conservation of corals and coral reef ecosystems. We analyzed mitochondrial DNA sequence types and microsatellite genotypes of the broadcast-spawning coral, *Galaxea fascicularis*, from four regions in the subtropical Nansei Islands in the northwestern Pacific Ocean. Two types (soft and hard types) of nematocyst morphology are known in *G. fascicularis* and are significantly correlated with the length of a mitochondrial DNA noncoding sequence (soft type: mt-L; hard type: mt-S type). Using microsatellites, significant genetic differentiation was detected between the mitochondrial DNA sequence types in all regions. We also found a third genetic cluster (mt-L+), and this unexpected type may be a cryptic species of *Galaxea*. High clonal diversity was detected in both mt-L and mt-S types. Significant genetic differentiation, which was found among regions within a given type (*F*$_{ST}$ = 0.009–0.024, all *P* ≤ 0.005 in mt-L; 0.009–0.032, all *P* ≤ 0.01 in mt-S), may result from the shorter larval development than in other broadcast-spawning corals, such as the genus *Acropora*. Nevertheless, intraspecific genetic diversity and connectivity have been maintained, and with both sexual and asexual reproduction, this species appears to have a potential for the recovery of populations after disturbance.

**Introduction**

Scleractinian reef-building corals constitute the framework of reef ecosystems and are especially endangered due to a multitude of rapid, oceanic environmental changes caused by anthropogenic disturbances, including extreme thermal stress from increasing sea surface temperatures (e.g., Hoegh-Guldberg 1999; Cantin et al. 2010), ocean acidification from increasing atmospheric CO$_{2}$ (e.g., Hoegh-Guldberg et al. 2007), and various local problems such as sediment pollution or nutrient influx from runoff (e.g., McCulloch et al. 2003). Population connectivity resulting from larval dispersal is essential for the maintenance or recovery of populations in marine ecosystems. Fertilized larvae of the broadcast-spawning corals disperse after spawning, and dispersal distances are dependent upon larval characteristics, such as the duration of the larval period, or time until settling (e.g., Nishikawa et al. 2003; Ayre and Hughes 2004). Therefore, broadcast-spawning corals are likely to maintain high genetic connectivity among populations. While various studies have reported high genetic connectivity in broadcast-spawning taxa such as the genus *Acropora* (e.g., Nishikawa et al. 2003; Ayre and Hughes 2004; Nishikawa and Sakai 2005a; Under-
wood et al. 2009; Nakajima et al. 2010, 2012a; Davies et al. 2015), some broadcast-spawning species have lower genetic connectivity than expected, due to the short duration of larval dispersal (e.g., *Platgyra daedalea*: Miller and Ayre 2008). Population genetic analysis in another genus will provide further insights into the genetic connectivity of the broadcast-spawning corals in this region. We conducted population genetics of the broadcast-spawning coral, *Galaxea fascicularis* (Linnaeus, 1767). This is the first report of a population genetic study of the genus *Galaxea* focusing on a large geographic range in an island reef system. *Galaxea fascicularis* is mainly distributed in reef areas in the Indo-western Pacific region (Veron 2000). It is a gonochoric, broadcast-spawning species; female colonies produce egg bundles, and male colonies form bundles consisting of sperm and infertile pseudo-eggs (e.g., Harrison 1988; Hayakawa et al. 2007; Keshavmurthy et al. 2012). This species is classified into soft and hard types based on nematocyst morphology (Hidaka 1992), and this morphological characteristic is correlated with the length of the noncoding region between the mitochondrial genes *cyt b* and *nad 2* (mt-Long: soft type; mt-Short: hard type) (Watanabe et al. 2005; Abe et al. 2008a). Significant genetic differentiation between these mt-Long (mt-L) and mt-Short (mt-S) types was shown at Zampa, on Okinawa Island using polymorphic nuclear microsatellite markers (Nakajima et al. 2015). However, significant genetic differentiation between mt-L and mt-S has not been demonstrated in other regions. Studies of boundaries between coral species are needed to estimate species diversity, which is essential for coral conservation. This species is one of the dominant, easily identified species at some reef sites in the Nansei Islands located in southwestern Japan. These islands are subtropical in nature, and a strong current, the Kuroshio Current, flows from southwest to northeast along the islands (Fig. 1). This current is considered a major factor in the expansion and maintenance of coral reefs and reef-dwelling organisms in the Nansei Islands (Nishihira and Veron 1995; Nishihira 2004).

In broadcast-spawning corals in this region, the genetic connectivity of *Acropora digitifera* (Nishikawa and Sakai 2005a; Nakajima et al. 2010), *Acropora tenuis* (Nishikawa et al. 2003), and *Coelastrea aspera* (formerly *Goniastrea aspera*) (Nishikawa and Sakai 2003, 2005b) has been analyzed using allozyme or microsatellite markers. The genetic differentiation index among regions for *A. digitifera* and *A. tenuis* (Nishikawa and Sakai 2005a) is not correlated with maximum pelagic larval duration until settlement (*A. digitifera*: 54 days, $F_{ST} = 0.021$, $P < 0.01$; *A. tenuis*: 69 days, $F_{ST} = 0.048$, $P < 0.01$). Also, local genetic differentiation ($F_{SC}$ among populations = 0.039, $P < 0.001$) was detected among sites in *C. aspera*, which employs planula brooding as well as broadcast spawning (Nishikawa and Sakai 2005b). *Galaxea fascicularis* appears to maintain the genetic diversity and connectivity within types by the virtue of the broadcast spawning. On the other hand, asexual reproduction probably contributes to the population maintenance because multiple colonies are often distributed locally (Nishihira and Veron 1995). Further research is needed to understand the dispersal patterns and connectivity in this species.
thermore, Nakajima et al. (2015) confirmed extensive clonality, although clonal diversity also has not been determined at other sites. We assessed genetic differentiation between mt-L and mt-S types in the Nansei Islands using cross-type microsatellite markers developed by Nakajima et al. (2015). Furthermore, we analyzed clonal diversity, genetic diversity, and differentiation within types at four regions in the Nansei Islands. In addition, we searched for novel genetic clusters (other types) in this species because two types had been identified based only on tentacle characters and a mitochondrial DNA noncoding region.

Materials and Methods

Study areas and collection of coral specimens

Sampling sites were located around the Nansei Islands (17 sites in four regions; Fig. 1, Table 1). The maximum distance between sites is greater than 750 km, between Ayamaru at Amami Island (A-Aya) and Amitor in the Yaeyama Islands (Y-Ami). We randomly collected a single branch from each colony of G. fascicularis. Specimens were preserved in ethanol and brought to the laboratory. The dataset from Zampa is from Nakajima et al. (2015).

Table 1. Geographic location information and population genetic parameters for each site. N: the number of analyzed specimens. G: the number of multilocus genotypes (MLGs). MLL: the number of multilocus lineages (MLLs), considering somatic mutation and scoring error and including identical genotypes resulting from sexual reproduction by chance, estimated by PSEX values for each site. R = (MLL−1)/N−1. Because sampling strategies were not necessarily identical at all locations, clonal diversity cannot be strictly compared between sites; however, it is apparent that clonality varies between populations, based upon environmental factors, for example, wave action, which can cause increased fragmentation.

| Region      | Location | Code       | Latitude (N) | Longitude (E) | Ntotal | N | G | MLL | R   | N | G | MLL | R   |
|-------------|----------|------------|--------------|---------------|--------|---|---|-----|-----|---|---|-----|-----|
| Amami       | Ayamaru  | A-Aya      | 28°28’34”    | 129°43’00”   | 15     | 10| 9 | 9   | 0.89| 5 | 5 | 5   | 1.00|
| Katetsu     | A-Kat    | 28°08’09”  | 129°20’39”   | 6            | 0   | – | – | –   | –   | – | – | –   | –   |
| Kuniano     | A-Kun    | 28°22’29”  | 129°24’14”   | 33            | 14 | 14| 14 | 1.00| 19 | 19| 19 | 1.00|
| Okinawa     | Zampa    | O-Zam      | 26°26’20”    | 127°42’40”   | 97    | 53| 10| 10 | 0.17| 44 | 10| 7  | 0.14|
| Odo         | O-Odo    | 26°05’17”  | 127°42’27”   | 13            | 6  | 5 | 5  | 0.80| 7  | 6 | 5  | 0.67|
| Kume        | O-Kum    | 26°19’15”  | 126°51’24”   | 39            | 20 | 20| 18 | 0.89| 9  | 9 | 8  | 0.88|
| Miyako      | Ikema    | M-Ike      | 24°56’02”    | 125°13’50”   | 14    | 9 | 9 | 9  | 1.00| 5  | 5 | 5  | 1.00|
| Yoshino     | M-Yos    | 24°44’52”  | 125°26’41”   | 27            | 24 | 2 | 1  | 0.00| 3  | 2 | 2  | 0.50|
| Ueno        | M-Uen    | 24°43’05”  | 125°20’30”   | 53            | 6  | 6 | 5  | 0.80| 46 | 8 | 8  | 0.16|
| Yaeyama     | Hirakubo | Y-Hir      | 24°35’35”    | 124°18’29”   | 12    | 5 | 2 | 2  | 0.25| 7  | 7 | 7  | 1.00|
| Ohama       | Y-Oha    | 24°20’24”  | 124°11’55”   | 38            | 36 | 3 | 3  | 0.06| 2  | 1 | 1  | 0.00|
| Kannon      | Y-Kan    | 24°21’55”  | 124°06’40”   | 17            | 10 | 9 | 9  | 0.89| 7  | 7 | 6  | 0.83|
| Taketomi    | Y-Tak    | 24°20’40”  | 124°05’21”   | 30            | 14 | 10| 9  | 0.62| 16 | 14| 14 | 0.87|
| Kuroshima   | Y-Kur    | 24°18’05”  | 124°06’56”   | 13            | 4  | 4 | 4  | 1.00| 9  | 9 | 8  | 0.88|
| Nakano      | Y-Nak    | 24°25’52”  | 123°47’26”   | 37            | 35 | 10| 7  | 0.18| 2  | 2 | 2  | 1.00|
| Amitori     | Y-Ami    | 24°19’47”  | 123°41’46”   | 39            | 27 | 27| 16 | 0.42| 2  | 2 | 2  | 1.00|
| Haemida     | Y-Hae    | 24°16’06”  | 123°49’47”   | 27            | 12 | 7 | 3  | 0.18| 15 | 15| 15 | 1.00|
| Total       |          |            |              |               | 510  | 295| 147| 124| 204| 127| 120| 11  | 11  | 11|

Genetic Differentiation and Connectivity of Coral

Scoring genotypes of microsatellites and identification of mitochondrial DNA sequence type

Genomic DNA was extracted using a DNeasy Blood & Tissue kit (Qiagen, Hilden, Germany). To analyze polymorphism and amplification of designed primer sets and to identify the mitochondrial DNA sequence type (mitochondrial type), we used the tailed primer method to perform PCR. We developed 11 microsatellite markers from G. fascicularis mt-L type, which are available for both mt-L and mt-S types (Nakajima et al. 2015). Three of 11 loci frequently indicated complex genotypic patterns showing a 1-bp shift among some alleles, complicating the process of determining genotypes. As a result, eight microsatellite markers were used for scoring genotypes in this study (see Table S1). The reaction mixture (5 μL) contained template DNA (<100 ng), AmpliTaq Gold 360 Master Mix (Thermo Fisher Scientific, Waltham, MA, USA), and three primers for each locus: a non-tailed forward primer (0.2 μM), a reverse primer with a U19 sequence tail (0.2 μM), and a U19 (5’-GGTTTTCAGTCACGCACGCGA-3’) primer (0.5 μM) fluorescently labeled with FAM, VIC, or NED, based on the method of Schuelke (2000). Furthermore, 188-1 (5’-GAATAGGTATACTAGCAGTGTC-3’, see Watanabe et al. 2005), 188-R3-U19 (5’-GGTTTTCAGTCACGCACGCACTATTATCTCTTCAA-3’) were used.
GG-3′), and U19 primers fluorescently labeled with VIC were used to identify the mitochondrial type (mt-L: 460 bp or mt-S: 170 bp) of the noncoding region between cyt b and nad 2. Amplifications of all microsatellite loci and the noncoding region between cyt b and nad 2 were performed with the following PCR conditions: 95°C for 9 min; followed by 35 cycles at 95°C for 30 s, 54°C for 30 s, and 72°C for 1 min; and a final extension at 72°C for 5 min. Amplified PCR products with the internal size standards GeneScan 600 LIZ (Thermo Fisher Scientific) were analyzed using an automated capillary-based DNA sequencer (ABI 3130xl Genetic Analyzer, Thermo Fisher Scientific) and GeneMapper ver. 3.7 (Thermo Fisher Scientific).

To compare the sequence of the mitochondrial non-coding region between cyt b and nad 2 to the one reported previously, PCR products amplified using 188-1 and 188-R3 (5′-CATCATTACCTTCTTCAAG-3′) for ten colonies (composed of mt-L, mt-S, and mt-L+ collected in O-Kum, see Results about mt-L+) were purified using ExoSAP-IT (Affymetrix, Santa Clara, CA, USA) and were sequenced bidirectionally using a BigDye Terminator Kit ver. 3.1 (Thermo Fisher Scientific). Procedures for sequencing were purified by ethanol precipitation, and sequences were analyzed using an ABI 3730d Genetic Analyzer (Thermo Fisher Scientific). Colonies with both mitochondrial types by fragment analysis (see Results) were also sequenced to confirm the sequences of the mitochondrial noncoding region.

Detection of clonal replicates

Multilocus lineages (MLLs) were employed to classify clonal replicates (Arnaud-Haond et al. 2007a). Genotyped colonies from the same site, displaying differences at no more than one locus, were considered clones derived from fragmentation (asexual reproduction), to avoid the misidentification of multilocus genotypes by incorrect genotyping caused by somatic mutation or scoring error. When slightly different genotypes were determined to belong to the same MLL, the most common genotype was used to represent the MLL, following the method of Arnaud-Haond et al. (2007b). However, if the most common genotype could not be determined (e.g., if two genotypes occurred in equal numbers), the genotype was set to zero so that this locus in the MLL was excluded from further analyses. MLLs were counted to estimate clonality using GenClone ver. 2.0 (Arnaud-Haond and Belkhir 2007). We removed one or two loci for the calculation of exact PSEX values if a somatic mutation or a scoring error appeared in the target MLL. We retained the replicated MLG in an MLL if two or more MLGs were the same, but occurred by chance as a result of sexual reproduction. However, if a population contained only one or several genotypes, even with a high sampling effort, and no further MLGs were detected, the statistical power associated with PSEX could be low (Arnaud-Haond et al. 2007b). In such a case, we did not consider the PSEX value to accurately estimate the replicated MLG and assumed that clonal reproduction accounted for such similarities.

Clonal diversity was estimated with the following index proposed by Dorken and Eckert (2001): \[ R = \frac{\text{MLL} - 1}{N - 1} \] (MLL: the number of multilocus lineages, N: the number of colonies analyzed). We removed clonal replicates from the dataset, and MLLs were retained for the following population genetic analyses.

Statistical analyses for population genetic indexes

The number of alleles, values of observed and expected heterozygosity (H_S and H_E, respectively), and a deviation index (F_IS) from Hardy–Weinberg equilibrium (HWE) for each type were evaluated with GenAlEx for each region. INEST ver. 2.0 (Chybicki and Burczyk 2009) was used to estimate the null allele frequency and the extent of inbreeding in each region. Using a Bayesian approach (individual inbreeding model), INEST was run using nbf (null alleles, inbreeding coefficients, and genotyping failures) and nb (null alleles and genotyping failures) models to detect the existence of inbreeding effects in our dataset with 50,000 burn-in cycles and 500,000 cycles overall. Postprocessing was conducted to calculate the mean null allele frequency of eight microsatellite loci, the mean value of the inbreeding coefficient, and the limit of the highest density posterior interval. The deviance information criterion (DIC) was used to determine the best model. For the estimation of genetic diversity, we calculated allelic richness (A_E) at each region using FSTAT ver. 1.2 (Goudet 1995). Furthermore, the H_E value was also used as the index of genetic diversity. Genetic differentiation was estimated by a hierarchical analysis of molecular variance (AMOVA; Excoffier et al. 1992) using GenAlEx. The genetic differentiation index between regions, pairwise F_ST, was also calculated using GenAlEx. The significance of each F_ST value was tested with 999 permutations. To estimate the effect of null alleles to the F_ST values, we calculated F_ST with both including null alleles (INA) and excluding null alleles (ENA) effects using FreeNA (Chapuis and Estoup 2007; Chapuis et al. 2008). The number
of replicates was 1000 for the computation of the bootstrap 95% confidence intervals.

The population structure for all MLLs based on Bayesian clustering was inferred using STRUCTURE ver. 2.3.4 (Pritchard et al. 2000). We ran STRUCTURE on the data for 10 independent chains for each K value (K = 1 to 11). A burn-in of 100,000 iterations followed by 1,000,000 Markov chain Monte Carlo (MCMC) replications was used for population clustering without prior information under the admixture model and assuming correlated allele frequencies (Falush et al. 2003). In the admixture model, individuals were assumed to have drawn exclusively from the K genetic clusters and were allowed to have a mixed ancestry (Pritchard et al. 2000; Falush et al. 2003). After calculating the mean log probability, Ln P(D), the number of clusters (K) that best fit the data, was determined using the method of Evanno et al. (2005), as implemented in STRUCTURE HARVESTER (Earl and von Holdt 2012). ΔK is an ad hoc quantity for predicting the number of possible clusters (Evanno et al. 2005). These data were also merged and outputted using CLUMPACK (Kopelman et al. 2015). To infer the population genetic structure within type, another Bayesian clustering algorithm was implemented in InStruct (Gao et al. 2007), which is an extension of the approach of STRUCTURE for simultaneous inference of inbreeding or selfing rates and population-of-origin classification. This procedure was used because inbreeding was suggested as a possible reason for the high inbreeding coefficients in our dataset (see Results). This algorithm analyzed and eliminated HWE assumptions within possible genetic clusters and calculated the expected genotype frequencies based on inbreeding or selfing rates (Gao et al. 2007). We ran InStruct for 10 independent chains for each K value (K = 1 to 5) and type. Each chain was iterated 1,000,000 times including burn-in with 100,000 iterations. Determination of the number of K clusters that best fit the data was conducted using the method of Evanno et al. (2005). These data were also merged and outputted using CLUMPP ver. 1.1.2 (Jakobsson and Rosenberg 2007) and distruct ver. 1.1 (Rosenberg 2004), respectively.

Results

Determination of mitochondrial type

Of 531 colonies of G. fascicularis collected from 17 sites at four regions in the Nansei Islands, we successfully determined the multilocus microsatellite genotypes and mitochondrial type in all but 21 colonies. When both peaks were indicated from the mitochondrial noncoding region between cyt b and nad 2, the main peak was used to decide the mitochondrial type if the other peak was extremely small (less than one-tenth the height of the main peak). Fourteen of these 21 are indeterminate colonies, and distinct and clear fragment peaks for both mitochondrial types were detected, although only one peak is usually derived from the mitochondrial noncoding region between cyt b and nad 2. Both peaks were derived from both types of mitochondrial noncoding region by sequencing. We excluded these 21 colonies and used genotypes from the remaining 510 colonies for all other analyses. Detailed site data are provided in Table 1. Of 510 colonies, 295 were identified as mt-L, while 204 were mt-S. Both mt-L and mt-S were found at 16 sites, excluding A-Kat (Table 1). In ten colonies collected at O-Kum and one at M-Uen, we found an unexpected mitochondrial type that showed 3 bp (TGG) more than the usual mt-L fragment length from the noncoding region. We named the unexpected type mt-L+. Haplotype networks from sequencing the mitochondrial noncoding region are shown with the sequence output in the study by Watanabe et al. (2005) (Fig. 2). As a result, this mt-L+ type is similar to the mt-L type E lineage collected at st.6 in Yaeyama in the previous study.

Multilocus lineages and inbreeding coefficients

Multilocus genotypes were determined from 295 mt-L, 11 mt-L+, and 204 mt-S colonies. Eight microsatellite loci utilized for scoring genotypes displayed 124, 11, and 120 MLLs, respectively (Table 1). The same MLG occurred by chance as a result of sexual reproduction in an mt-S MLL at M-Uen, based upon the PSEX value. We added this doubled MLL as a different lineage for subsequent analyses. In these MLLs, there is a large variation in clonality at each site. Values ranged from 0 to 1 for both mt-L and mt-S types, although MLLs were few at some sites because of high clonal diversity within types. Three MLLs (corresponding to three colonies analyzed) in mt-L and two MLLs (corresponding to two colonies analyzed) in mt-S showed the opposite genetic cluster in STRUCTURE (see black triangles in Fig. 3A). These five MLLs were excluded for InStruct and further statistical analyses to avoid misleading genotypic data.

Our dataset displayed high FIS values (FIS = 0.122 to 0.256 for each region), and this result was supported by a relatively high frequency of null alleles (Null freq. = 0.061 to 0.140 among regions) and inbreeding influences (Avg Fi = 0.021 to 0.133 among regions) from the results of INEST (Table 2). Of nine populations including mt-L+, the nb model is the best in two populations and nb is the best in seven populations; however, the difference in DIC values between models is <1.0 in seven populations, except for mt-L in Miyako and mt-L+.
Genetic differentiation among mitochondrial types

Significant genetic differentiation among types was detected based on AMOVA ($F_{RT} = 0.054$, $P < 0.001$, Table 3). The genetic differentiation between mt-L and mt-S was significant in all cases (pairwise $F_{ST} = 0.034$ to 0.068, all $P_s = 0.001$, Table 4). Genetic differentiation between mt-L+ and other types was also significant, and these values are higher than values between mt-L and mt-S (between mt-L and mt-L+: pairwise $F_{ST} = 0.158$ to 0.179, all $P_s = 0.001$; between mt-S and mt-L+: pairwise $F_{ST} = 0.158$ to 0.183, all $P_s = 0.001$). These tendencies are identical with the results of pairwise $F_{ST}$ excluding the effect of null alleles, and no large influence of null alleles for the calculation of these genetic differentiation indexes (Table S2). The results from STRUCTURE, based on Bayesian statistical model-based clustering, also indicated that these three types are genetically isolated (Fig. 3A). Two peaks are observed in the graph of $\Delta K$ value, which is an index to estimate the greatest possible number of assumed clusters ($\Delta K = 187.96$ at $K = 3$ and $\Delta K = 48.49$ at $K = 5$, Fig. S1). This application of the method of Evanno et al. (2005) indicated that $K = 3$ offers the best explanation for the genetic data.

Genetic diversity and differentiation within mitochondrial types

Allelic richness and the expected heterozygosity of mt-L and mt-S at northern regions were comparable to those of southern regions (Table 2). Allelic richness values ranged from 10.57 to 11.94 in mt-L and from 10.14 to 11.14 in mt-S. In mt-L+ collected at O-Kum and M-Uen, the
value was 7.95. For the expected heterozygosity, values ranged from 0.886 to 0.917 in mt-L and from 0.843 to 0.889 in mt-S. The value of mt-L+ collected at O-Kum was 0.621. Compared with the results of genetic differentiation among types, genetic differentiation within type (mt-L or mt-S, but not mt-L+) showed significant variation among regions. Values ranged from 0.009 to 0.024 in mt-L and from 0.009 to 0.032 in mt-S (Table 4). The extent of genetic differentiation was similar in both mt-L and mt-S. These tendencies are also identical with the results of pairwise $F_{ST}$ excluding the effect of null alleles (Table S2). STRUCTURE results (Fig. 3B) suggest slight genetic structuring among regions of each type. A slight genetic discontinuity occurs between Miyako and Yaeyama in mt-L. On the other hand, a slight genetic discontinuity is also observed between Miyako and Yaeyama in mt-S, resembling that cluster composition is found in Amami and Yaeyama. The results of InStruct, considering inbreeding coefficients, also showed the same tendency. A peak was observed in the graph of the $\Delta K$ value at $K = 2$ in both mt-L and mt-S ($\Delta K = 311.28$ in mt-L and 729.49 in mt-S, Fig. S2).

**Discussion**

For the broadcast-spawning coral, *G. fascicularis*, in the Nansei Islands of Japan, this study analyzed mitochondrial types and compared these types with genotypes of...
Table 3. The three mitochondrial DNA sequence types are genetically well differentiated, showing low variation within types. Analysis of molecular variance (AMOVA), showing degrees of freedom (df), sum of squares (SS), variance components (Var.), percentage of variances (%), and F-statistics between types, among regions within types and within regions.

| Source          | df | SS     | Var. (%) | F-statistics (P value) |
|-----------------|----|--------|----------|------------------------|
| Among types     | 2  | 69.331 | 0.209    | $F_{RT} = 0.054$ (0.001) |
| Among regions   | 6  | 40.454 | 0.057    | $F_{SR} = 0.016$ (0.001) |
| Within type     |    |        | (1.5%)   |                        |
| Within regions  | 491| 1758.089| 3.581    | $F_{ST} = 0.069$ (0.001) |
| Within type     |    |        | (93.1%)  |                        |
| Total           | 499| 1867.874| 3.847    | (100.0%)               |

Microsatellites to estimate the species diversity of *Galaxea*. Mitochondrial type is related to microsatellite genotype in all regions. Values of clonal diversity are variable among sites for each type. Genetic differentiation is significant, but low among regions, and genetic diversity has been maintained for each type.

Genetic differentiation among mitochondrial types

Although mt-L colonies were more numerous than mt-S colonies, the ratio of types varies widely among the sites that we sampled. This pattern agrees with the previous studies (Watanabe et al. 2005; Abe et al. 2008a). All sites, excluding A-Kat, possessed both types, and the ratio is unrelated to latitude in the Nansei Islands. Colonization may be more influenced by evolutionary processes than by latitudinal factors such as temperature. Evolutionary divergence of tropical coral species is likely to occur in the center of the tropical region, known as the Coral Triangle. However, the genus *Galaxea* has an extensive range in the Indo-western Pacific, and it is not clear what sort of divergence has occurred in the Coral Triangle itself. In Okinawa, mt-L and mt-S types can be distinguished not only by morphology, but also by the timing of spawning (Watanabe et al. 2005); mt-L spawns a few weeks earlier than mt-S in some cases. With this slight but significant genetic differentiation between mt-L and mt-S, segregated spawning may explain genetic differentiation between types. However, the spawning season of mt-L+ is not known. Further observation will be needed to understand the reproductive strategies contributing to genetic isolation among types in this species.

The mitochondrial type, mt-L+, found at O-Kum and M-Uen, was unexpected, indicating greater genetic differentiation; however, the sequence of the mitochondrial noncoding region is interestingly similar to the E lineage of mt-L reported by Watanabe et al. (2005) (Fig. 2). Although *G. fascicularis* is easily distinguished from other *Galaxea* species by morphology, we cannot exclude the possibility that this unexpected type is a known *Galaxea* species that has previously been reported in the Nansei Islands, such as *Galaxea astreata* or *Galaxea pauciradiata* (formerly *Galaxea paucisepala*) (see Nishihira and Veron 1995; Veron 2000). However, we could not differentiate morphological types in the field (see Fig. 2). Morphological characterization of *Galaxea* appears to be more difficult than we anticipated, as in some other corals. For example, the genus *Pocillopora* also shows interspecific ambiguity with high levels of morphological variability in colony shape and branch size, but these well-known variants do not appear to correlate with genetic variation (Pinzón and Lajeunesse 2011). Nevertheless, the morphs are distinguished by microsatellite markers and mitochondrial DNA regions (ORF and control region) (Pinzón and Lajeunesse 2011; Pinzón et al. 2013; Schmidt-Roach et al. 2013, 2014). Thus, genetic identification has contributed

Table 4. Pairwise $F_{ST}$ values for all mitochondrial DNA sequence types as an index of genetic differentiation between pairs of regions. The lower left table shows pairwise $F_{ST}$ values, and the upper right shows $P$ values. Larger $F_{ST}$ values denote greater genetic differentiation. Because the number of mt-L+ specimens was small (MLL = 11), all specimens are combined into a single population.

| Type | Region   | Amami | Okinawa | Miyako | Yaeyama |
|------|----------|-------|---------|--------|---------|
| mt-L | Amami    | 0.009 | 0.001   | 0.001  | 0.001   |
|      | Okinawa  | 0.024 | 0.013   | 0.002  | 0.002   |
|      | Miyako   | 0.016 | 0.009   | 0.019  | 0.001   |
|      | Yaeyama  | 0.037 | 0.042   | 0.053  | 0.043   |
| mt-S | Amami    | 0.037 | 0.034   | 0.046  | 0.039   |
|      | Okinawa  | 0.037 | 0.042   | 0.053  | 0.043   |
|      | Miyako   | 0.050 | 0.052   | 0.054  | 0.053   |
|      | Yaeyama  | 0.058 | 0.057   | 0.068  | 0.055   |
| mt-L+|         | 0.179 | 0.165   | 0.171  | 0.158   |

© 2016 The Authors. Ecology and Evolution published by John Wiley & Sons Ltd.
to species classification and estimates of species diversity of corals. We tried to compare differences between G. fascicularis and other Galaxea species using gene sequences from GenBank. We found six gene sequences commonly shared by G. fascicularis and G. astreata. However, large clear differences between types cannot be confirmed using the haplotype network by TCS ver. 1.21 (Clement et al. 2000). Some registered sequences may actually be derived from other genera because some of the haplotypes from the same nominal species are separated (Fig. S3). Understanding the phylogenetic relationship of Galaxea, more phylogenetic analyses (e.g., genome-wide single nuclear polymorphism (SNP) markers, or amino acid sequences derived from all coding genes on mitochondrial genome) are needed for the exact estimation of species diversity. With nematocyst types, fine-scale investigations of the corallite and subcorallite features observed with scanning electron microscopy (SEM) and thin sections may reveal defining characters each type.

### Interbreeding and ambiguous mitochondrial type

Intermediate genetic lineages were also detected with STRUCTURE. Interbreeding is less likely to occur between the two types than within types, suggesting that spawning seasons do not overlap in some cases (Watanabe et al. 2005). Abe et al. (2008b) reported that the seasons are not completely separated because they managed to conduct fertilization experiments between types. Therefore, a reproductive isolation may not be complete between mt-L and mt-S, which may have resulted from a recent evolutionary speciation process. Although crosstype hybrids occurred as a result of fertilization experiments between types (Abe et al. 2008b), the incubation time (4–5 h) with higher sperm concentration appears to be longer than those that occur in nature (see Iguchi et al. 2009; Nozawa et al. 2015). Therefore, as Watanabe et al. (2005) suggested, actual hybridization between types may be rare, even if spawning seasons partly overlap in some colonies. Different spawning times appear to maintain the genetic divergence between types. Two sympatric Acropora species, A. digitifera and a cryptic species Acropora sp. 1 aff. digitifera (Hayashibara and Shimoike 2002), also show significant genetic differentiation (Nakajima et al. 2012b). The octocoral species, Helioipsa coerulea, is also separated into two genetic clusters that are related to branch morphotype (Yasuda et al. 2014). Compared with G. fascicularis, these Acropora and Helioipsa species are influenced by the difference in temporal reproductive isolation (Acropora: Hayashibara and Shimoike 2002; Ohki et al. 2015; Helioipsa: Villanueva 2015).

We confirmed the existence of some colonies with both mt-L and mt-S peaks in the fragment analysis, although we excluded MLLs with undefined types from our analyses (see Results). Both mitochondrial types were confirmed in these colonies from the sequencing results. Although we suspected possible contamination in these specimens, repeated experiments did not reveal experimental errors. Furthermore, among microsatellite genotypes, there is no evidence of chimeric or triploid colonies, which are occasionally detected in genetic experiments (e.g., Baums et al. 2005; Puill-Stephan et al. 2012). We believe that mitochondrial types in a colony may be derived from heteroplasmy by introgression of both parental mitochondria via hybridization between types in previous generations. Further organelle genome studies of corals may eventually resolve this question.

### Population maintenance of Galaxea within mitochondrial type

Species that comprise the frameworks of ecosystems, such as corals and seagrasses, often reproduce clonally (Baums et al. 2006). Small-scale disturbances, such as waves during local storms, may increase the rates of clonal reproduction in coral populations as a result of fragmentation (Baums et al. 2006; Aranceta-Garza et al. 2012). In G. fascicularis, both sexual and asexual reproductions contribute to population persistence and recovery from anthropogenic disturbances, showing a high degree of clonal diversity. Although we employed nonstandardized sampling among sites and did not standardize the sampling strategy due to the magnitude of the study area, spatial genetic and clonal structure within individual reefs is also informative (see Gorospe and Karl 2013). Further collection and analyses of specimens standardized among sites will provide the detailed clonal structure among populations, and spatial correlation based on locality data should elucidate kinship among colonies on a regional scale.

It is generally accepted that broadcast-spawning corals have longer larval durations and higher connectivity among regions than brooding species. We expected that within types, G. fascicularis would also show high connectivity among regions and reduced genetic differentiation. Populations in high-latitude areas and islands isolated from the central habitat of the species show decreased genetic diversity, and some other coral species reflect this tendency (e.g., Ayre and Hughes 2004; Ridgway et al. 2008; Davies et al. 2015). However, high genetic diversity was shown in high-latitude peripheral populations of Acropora species (Nakajima et al. 2010; Noreen et al. 2013). Some Acropora species, classified as long-term winner species, which increase in their relative contribution to the total habitat cover in survey site at Okinawa, are
also pioneers following local extinctions due to disturbances such as mass bleaching events (van Woesik et al. 2011).

Two possible reasons can explain the significant genetic differentiation among regions within type. First, the larval duration may be shorter than for broadcast-spawning corals, such as Acropora. Although larval duration is not certain for G. fascicularis, the first larval stage lasts only ~18 h before larvae start swimming (Okubo et al. 2013). Embryonic development of G. fascicularis is significantly faster than in Acropora muricata (Keshavmurthy et al. 2012). Second, we can explain high clonal diversity at some sites. Populations of G. fascicularis are largely maintained by fragmentation, which promotes the deviation from HWE by preventing random mating. In Bayesian clustering, slight genetic discontinuity occurs between Miyako and Yaeyama in mt-L. On the other hand, it also exists between Miyako and Yaeyama in mt-S, although similar cluster composition is found in Amami and Yaeyama. The difference between these two types may have resulted from physiological characteristics and historical transitions. Differences between types should be resolved in future studies to clarify the population dynamics and evolutionary history of each type.

In the Nansei Islands, this species maintains high genetic diversity and connectivity, as do other broadcast-spawning coral species, such as A. digitifera (Nakajima et al. 2010). In the Nansei Islands, longer larval duration of the broadcast-spawning corals probably contributes to high genetic diversity and connectivity, which may be enhanced by the Kuroshio Current and its branches. Galaxea fascicularis is also distributed in nonreef areas in the temperate zone of Japan (Nishihira and Veron 1995), but population sizes are small because that environment is harsh for tropical species. If the sea surface temperature gradually increases due to climate change, northern populations of this species may increase and its range may extend northward. As a result, other reef-associated fauna also may extend their ranges as well.

Conclusions

This population genetic study using microsatellite markers revealed different ratios of mt-L and mt-S types and clonal diversity among regions in G. fascicularis. Clonal diversity is related to geographic factors rather than to type or latitude. Further study may provide a picture of clonal structure within single reefs. We also defined genetic differentiation among types using multiple analyses. An unexpected mitochondrial type was found, and interestingly, this third type is genetically quite different from both the mt-L and mt-S types. The sequence of the mitochondrial noncoding region in this type is similar to that of a colony reported by Watanabe et al. (2005). This unexpected type is possibly a novel Galaxea species, although we cannot exclude the possibility that this type is a known Galaxea species with an ambiguous morphotype. Genetic differentiation among regions within types is lower, but still significant. With oceanographic factors as the Kuroshio Current and branch currents, both larval dispersal via sexual reproduction and fragmentation via asexual reproduction as biological characters appear to contribute to the maintenance of Galaxea populations in the Nansei Islands.

Acknowledgments

This study was supported by the Japan Society for the Promotion of Science (JSPS) KAKENHI (25840152 to Yuichi Nakajima, 25660172 and 26290065 to Chuya Shinzato) and a research grant for the natural conservation of Nansei Islands to Yuichi Nakajima by Pro Natura Foundation (http://www.pronaturajapan.com/). We express our gratitude for financial support from the Okinawa Institute of Science and Technology Graduate University. The collection of corals was approved by Kagoshima (permission number not assigned) and Okinawa (No. 25-53 and 26-62) prefectures. We thank Kazuhioko Sakai and Go Suzuki for field assistance and Steven D. Aird for editing the manuscript.

Conflict of Interest

None declared.

References

Abe, M., T. Watanabe, Y. Suzuki, and M. Hidaka. 2008a. Genetic and morphological differentiation in the hermatypic coral Galaxea fascicularis in Okinawa, Japan. Plankton and Benthos Res. 3:174–179.
Abe, M., T. Watanabe, H. Hayakawa, and M. Hidaka. 2008b. Breeding experiments of the hermatypic coral Galaxea fascicularis: partial reproductive isolation between colonies of different nematocyst types, and enhancement of fertilization success by the presence of parental colonies. Fish. Sci. 74:1342–1344.
Aranceta-Garza, F., E. F. Balart, H. Reyes-Bonilla, and P. Cruz-Hernández. 2012. Effect of tropical storms on sexual and asexual reproduction in coral Pocillopora verrucosa subpopulations in the Gulf of California. Coral Reefs 31:1157–1167.
Arnaud-Haond, S., and K. Belkhir. 2007. GENCLONE: a computer program to analyze genotypic data, test for clonality and describe spatial clonal organization. Mol. Ecol. Notes 7:15–17.
Arnaud-Haond, S., M. Migliaccio, E. Diaz-Almela, S. Teixeira, M. S. van de Vliet, F. Alberto, et al. 2007a. Vicariance
patterns in the Mediterranean Sea: east-west cleavage and low dispersal in the endemic seagrass *Posidonia oceanica*. J. Biogeogr. 34:963–976.

Arnaud-Haond, S., C. M. Duarte, F. Alberto, and E. A. Serrão. 2007b. Standardizing methods to address clonality in population studies. Mol. Ecol. 16:5115–5139.

Ayre, D. J., and T. P. Hughes. 2004. Climate change, genotypic diversity and gene flow in reef-building corals. Ecol. Lett. 7:273–278.

Baums, I. B., C. R. Hughes, and M. E. Hellberg. 2005. Mendelian microsatellite loci for the Caribbean coral *Acropora palmata*. Mar. Ecol. Prog. Ser. 288:115–127.

Baums, I. B., M. W. Miller, and M. E. Hellberg. 2006. Geographic variation in clonal structure in a reef-building Caribbean coral, *Acropora palmata*. Ecol. Monogr. 76:503–519.

Cantin, N. E., A. L. Cohen, K. B. Karnauskas, A. M. Tarrant, and D. C. McCorkle. 2010. Ocean warming slows coral growth in the central Red Sea. Science 329:322–325.

Chapuis, M.-P., and A. Estoup. 2007. Microsatellite null alleles and estimation of population differentiation. Mol. Biol. Evol. 24:621–631.

Chapuis, M.-P., M. Lecoq, Y. Michalakis, A. Loiseau, G. A. Sword, S. Piry, et al. 2008. Do outbreaks affect genetic population structure? A worldwide survey in *Locusta migratoria*, a pest plagued by microsatellite null alleles. Mol. Ecol. 17:3640–3653.

Chybicki, I. J., and J. Burczyk. 2009. Simultaneous estimation of null alleles and inbreeding coefficients. J. Hered. 100:106–113.

Clement, M., D. Posada, and K. Crandall. 2000. TCS: a computer program to estimate genealogies. Mol. Ecol. 9:1657–1660.

Davies, S. W., E. A. Treml, C. D. Kenkel, and M. V. Matz. 2015. Exploring the role of Micronesian islands in the maintenance of coral genetic diversity in the Pacific Ocean. Mol. Ecol. 24:70–82.

Dorken, M. E., and C. G. Eckert. 2001. Severely reduced sexual reproduction in northern populations of a clonal plant, *Decodon verticillatus* (Lythraceae). J. Ecol. 89:339–350.

Earl, D. A., and B. M. vonHoldt. 2012. STRUCTURE HARVESTER: a website and program for visualizing STRUCTURE output and implementing the Evanno method. Conserv. Genet. Resour. 4:359–361.

Evanno, G., S. Regnaut, and J. Goudet. 2005. Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study. Mol. Ecol. 14:2611–2620.

Excoffier, L., P. E. Smouse, and J. M. Quattro. 1992. Analysis of molecular variance inferred from metric distances among DNA haplotypes: application to human mitochondrial DNA restriction data. Genetics 131:479–491.

Falush, D., M. Stephens, and J. K. Pritchard. 2003. Inference of population structure using multilocus genotype data: linked loci and correlated allele frequencies. Genetics 164:1567–1587.

Gao, H., S. Williamson, and C. D. Bustamante. 2007. A Markov chain Monte Carlo approach for joint inference of population structure and inbreeding rates from multilocus genotype data. Genetics 176:1635–1651.

Gorospe, K. D., and S. A. Karl. 2013. Genetic relatedness does not retain spatial pattern across multiple spatial scales: dispersal and colonization in the coral, *Pocillopora damicornis*. Mol. Ecol. 22:3721–3736.

Goudet, J. 1995. FSTAT (version 1.2): a computer program to calculate F-statistics. J. Hered. 86:485–486.

Harrison, P. L. 1988. Pseudo-gynodioecy: an unusual breeding system in the scleractinian coral *Galaxea fascicularis*. Proc. 6th Int. Coral Reef Symp. 2:699–705.

Hayakawa, H., T. Andoh, and T. Watanabe. 2007. Identification of a novel yolk protein in the hermatypic coral *Galaxea fascicularis*. Zool. Sci. 24:249–255.

Hayashibara, T., and K. Shimoike. 2002. Cryptic species of *Acropora digitifera*. Coral Reefs 21:224–225.

Hidaka, M. 1992. Use of nematocyst morphology for taxonomy of some related species of scleractinian corals. *Galaxea* 11:21–28.

Hoegh-Guldberg, O. 1999. Coral bleaching, climate change, and the future of the world’s coral reefs. Mar. Freshwater Res. 50:839–866.

Hoegh-Guldberg, O., P. J. Mumby, A. J. Hooten, R. S. Steneck, P. Greenfield, E. Gomez, et al. 2007. Coral reefs under rapid climate change and ocean acidification. Science 318:1737–1742.

Iguchi, A., M. Morita, Y. Nakajima, A. Nishikawa, and D. J. Miller. 2009. In vitro fertilization efficiency in coral *Acropora digitifera*. *Zygote* 17:225–227.

Jakobsson, M., and N. A. Rosenberg. 2007. CLUMPP: a cluster matching and permutation program for dealing with label switching and multimodality in analysis of population structure. Bioinformatics 23:1801–1806.

Keshavmurthy, S., C.-M. Hsu, C.-Y. Kuo, V. Denis, J. K.-L. Leung, S. Fontana, et al. 2012. Larval development of fertilized “pseudo-gynodioecious” eggs suggests a sexual pattern of gynodoecy in *Galaxea fascicularis* (Scleractinia: Euphylliidae). Zool. Stud. 51:143–149.

Kopelman, N. M., J. Mayzel, M. Jakobsson, N. A. Rosenberg, and I. Mayrose. 2015. CLUMPAK: a program for identifying clustering modes and packaging population structure inferences across K. Mol. Ecol. Resour. 15:1179–1191.

McCulloch, M., S. Fallon, T. Wyndham, E. Hendy, J. Lough, and D. Barnes. 2003. Coral record of increased sediment flux to the inner Great Barrier Reef since European settlement. Nature 421:727–730.

Miller, K. J., and D. J. Ayre. 2008. Population structure is not a simple function of reproductive mode and larval type: insights from tropical corals. J. Anim. Ecol. 77:713–724.

Nakajima, Y., A. Nishikawa, A. Iguchi, and K. Sakai. 2010. Gene flow and genetic diversity of a broadcast-spawning
Genetic Differentiation and Connectivity of Coral

Y. Nakajima et al.

coral in northern peripheral populations. PLoS ONE 5: e11149.
Nakajima, Y., A. Nishikawa, A. Iguchi, and K. Sakai. 2012a. Regional genetic differentiation among northern high-latitude island populations of a broadcast-spawning coral. Coral Reefs 31:1125–1133.
Nakajima, Y., A. Nishikawa, A. Iguchi, and K. Sakai. 2012b. Population genetic approach delineates species boundary of reproductively isolated corymbose acroporid corals. Mol. Phylogen. Evol. 63:527–531.
Nakajima, Y., C. Shinzato, N. Satoh, and S. Mitarai. 2015. Novel polymorphic microsatellite markers reveal genetic differentiation between two sympatric types of Galaxea fascicularis. PLoS ONE 10:e0130176.
Nishihi, M. 2004. Hermatypic corals of Japan. Pp. 10–14 in Ministry of the Environment of Japan, Japanese Coral Reef Society, ed. Coral reefs of Japan. Japan Wildlife Research Center, Tokyo. (in Japanese).
Nishihi, M., and J. E. N. Veron. 1995. Hermatypic corals of Japan. Kaiyusha, Tokyo. (in Japanese).
Nishikawa, A., and K. Sakai. 2003. Genetic variation and gene flow of broadcast spawning and planula brooding coral, Goniastrea aspera (Scleractinia) in the Ryukyu Archipelago, southern Japan. Zool. Sci. 20:1031–1038.
Nishikawa, A., and K. Sakai. 2005a. Settlement-competency period of planulae and genetic differentiation of the scleractinian coral Acropora digitifera. Zool. Sci. 22:391–399.
Nishikawa, A., and K. Sakai. 2005b. Genetic connectivity of the scleractinian coral around the Okinawa Islands. Coral Reefs 24:318–324.
Nishikawa, A., M. Katoh, and K. Sakai. 2003. Larval settlement rates and gene flow of broadcast-spawning (Acropora tenuis) and planula-brooding (Stylophora pistillata) corals (Scleractinia). Mar. Ecol. Prog. Ser. 256:87–97.
Noreen, A. M. E., M. J. H. van Oppen, and P. L. Harrison. 2013. Genetic diversity and differentiation among high-latitude broadcast-spawning coral populations disjunct from the core range. Mar. Ecol. Prog. Ser. 491:101–109.
Nohzawa, Y., N. Isomura, and H. Fukami. 2015. Influence of sperm dilution and gamete contact time on the fertilization rate of scleractinian corals. Coral Reefs 34:1199–1206.
Ohki, S., R. K. Kowalski, S. Kitanobo, and M. Morita. 2015. Changes in spawning time led to the speciation of the broadcast spawning corals Acropora digitifera and the cryptic species Acropora sp. 1 with similar gamete recognition systems. Coral Reefs 34:1189–1198.
Okubo, N., T. Mezaki, Y. Nohzawa, Y. Nakano, Y.-T. Lien, H. Fukami, et al. 2013. Comparative embryology of eleven species of stony corals (Scleractinia). PLoS ONE 8:e84115.
Peakall, R., and P. E. Smouse. 2006. Genalex 6: genetic analysis in Excel. Population genetic software for teaching and research. Mol. Ecol. Notes 6:288–295.
Pinzón, J. H., and T. C. Lejeunesse. 2011. Species delimitation of common reef corals in the genus Pocillopora using nucleotide sequence phylogenies, population genetics and symbiosis ecology. Mol. Ecol. 20:311–325.
Pinzón, J. H., E. Sampayo, E. Cox, L. J. Chauka, C. A. Chen, C. R. Voolstra, et al. 2013. Blind to morphology: genetics identifies several widespread ecologically common species and few endemics among Indo-Pacific cauliflower corals (Pocillopora, Scleractinia). J. Biogeogr. 40:1595–1608.
Pritchard, J. K., M. Stephens, and P. Donnelly. 2000. Inference of population structure using multilocus genotype data. Genetics 155:945–959.
Puill-Stephan, E., M. J. H. van Oppen, K. Pichavant-Rafini, and B. L. Willis. 2012. High potential for formation and persistence of chimeras following aggregated larval settlement in the broadcast spawning coral, Acropora millepora. Proc. R. Soc. B 279:699–708.
Ridgway, T., C. Riginos, J. Davis, and O. Hoegh-Guldberg. 2008. Genetic connectivity patterns of Pocillopora verrucosa in southern African Marine Protected Areas. Mar. Ecol. Prog. Ser. 354:161–168.
Rosenberg, N. A. 2004. DISTRUCT: a program for the graphical display of population structure. Mol. Ecol. Notes 4:137–138.
Schmidt-Roach, S., K. J. Miller, G. Gerlach, A. M. E. Noreen, and N. Andreakis. 2013. Assessing hidden species diversity in the coral Pocillopora damicornis from Eastern Australia. Coral Reefs 32:161–172.
Schmidt-Roach, S., K. J. Miller, P. Lundgren, and N. Andreakis. 2014. With eyes wide open: a revision of species within and closely related to the Pocillopora damicornis species complex (Scleractinia; Pocilloporidae) using morphology and genetics. Zool. J. Linn. Soc. 170:1–33.
Schuelke, M. 2000. An economic method for the fluorescent labeling of PCR fragments. Nature Biotech. 18:233–234.
Underwood, J. N., L. D. Smith, M. J. H. van Oppen, and J. P. Gilmour. 2009. Ecologically relevant dispersal of corals on isolated reefs: implications for managing resilience. Ecol. Appl. 19:18–29.
Veron, J. E. N. 2000. Corals of the world. Australian Institute of Marine Science and CRR Qld Pty Ltd., Townsville.
Villanueva, R. D. 2015. Cryptic speciation in the stony octocoral Heliopora coerulea: temporal reproductive isolation between two growth forms. Mar. Biodivers. doi: 10.1007/ s12526-015-0376-y.
Watanabe, T., M. Nishida, K. Watanabe, D. S. Wevengkang, and M. Hidaka. 2005. Polymorphism in the nucleotide sequence of a mitochondrial intergenic region in the scleractinian coral Galaxea fascicularis. Mar. Biotech. 7:33–39.
van Woesik, R., K. Sakai, A. Ganase, and Y. Loya. 2011. Revisiting the winners and the losers a decade after coral bleaching. Mar. Ecol. Prog. Ser. 434:67–76.
Yasuda, N., C. Taquet, S. Nagai, M. Fortes, T.-Y. Fan, N. Phongsuwan, et al. 2014. Genetic structure and cryptic speciation in the threatened reef-building coral Heliopora coerulea along Kuroshio Current. Bull. Mar. Sci. 90:233–255.
Supporting Information

Additional Supporting Information may be found in the online version of this article:

**Figure S1.** The optimal number of genetic clusters is three, using STRUCTURE from STRUCTURE HARVESTER. These three clusters correspond very closely to the three mitochondrial DNA sequence types identified from noncoding mitochondrial DNA sequences (mt-L, mt-S, mt-L+). Mean Ln P(D) values ($K = 1$ to $11$) across 10 iterations per $K$, and $\Delta K$ values ($K = 2$ to $10$) using the method of Evanno et al. (2005).

**Figure S2.** The most likely number of genetic clusters is two for both mt-L (a) and mt-S (b) by estimation of the optimal number of genetic clusters of InStruct. Mean Ln P(D) values ($K = 1$ to $5$) across 10 iterations per $K$, and $\Delta K$ values ($K = 2$ to $4$) using the method of Evanno et al. (2005).

**Figure S3.** Haplotype networks based upon sequences of mitochondrial genes (12S rRNA, 16S rRNA, atp 6, cox 1, cyt b) and a nuclear gene (28S rRNA) from GenBank split nominal species, suggesting that some *Galaxea* species may either be misidentified or may contain cryptic species. These networks were constructed by TCS ver. 1.21 (Clement et al. 2000). Sequences of these six loci are registered in both *Galaxea fascicularis* and *Galaxea astreata*.

**Table S1.** Characteristics of eight polymorphic microsatellite loci in this study: locus name, repeat motif, primer sequence, size range of amplification products including U19 sequence, and GenBank accession number. U19, 5'-GGTTTTCAGTCACGACG-3'.

**Table S2.** Pairwise $F_{ST}$ values calculated using FreeNA (Chapuis and Estoup 2007; Chapuis et al. 2008) to estimate the effect of null alleles. The values above the diagonal are pairwise $F_{ST}$ by ENA (i.e., excluding null alleles) model and lower diagonal are by INA (i.e., including null alleles) model.