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Genetic population structure of the blue sea star (*Linckia laevigata*) and the boring giant clam (*Tridacna crocea*) across Malaysia

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Abstract Various studies on population genetics in the Indo-Pacific that include the Coral Triangle region have revealed phylogeographic patterns in marine species populations. However, little is known about the population structure and connectivity pattern among the coral reefs in four seas surrounding Malaysia: the Strait of Malacca (Andaman Sea), the South China Sea, the Sulu Sea, and the Sulawesi Sea. In view of this, we examined the genetic population structure of two invertebrate species: the blue sea star (*Linckia laevigata*) and the boring clam (*Tridacna crocea*) from seven localities across

Malaysia, including Pulau Layang-Layang in the Spratly Islands. Additional samples of *L. laevigata* were obtained from the Davao Gulf, the Philippines. The analyses were based on partial mitochondrial cytochrome oxidase I sequences of *L. laevigata* and *T. crocea*. Populations of *L. laevigata* and *T. crocea* showed departure from neutrality, indicating selection or population expansion. However, a mismatch analysis suggested population expansion. Lack of genetic population structure in *L. laevigata* ($\Phi_{ST} = -0.001$, $P = 0.399$) implies high connectivity among the sampled reef sites. There was subtle genetic structuring in populations of *T. crocea* ($\Phi_{ST} = 0.032$, $P = 0.009$), but AMOVA did not detect any structure in the a priori groupings. The genetic population structure and scale in connectivity are most likely attributed to the life history traits of the sea star and giant clam species,

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and also to the geological history and seasonal current patterns of the region.

Keywords Genetic diversity · Historical demography · Dispersal · Phylogeography · Conservation

Introduction

The Coral Triangle is paramount for its diverse marine habitats and species; yet this region is also faced with various threats, giving it high priority for conservation (Burke et al., 2011, 2012). Studies on population genetics and phylogeographic patterns in the Indo-Pacific including the Coral Triangle region (Crandall et al., 2008a; Carpenter et al., 2011; Bowen et al., 2014; Keyse et al., 2014) have been conducted to understand the processes driving and maintaining the extraordinary biodiversity in this region (Avice et al., 1987; Palumbi, 1997; Barber & Bellwood, 2005), and infer the connectivity among coral reefs. Information on population connectivity is essential in formulating effective management plans or designing MPA networks (Cowen et al., 2007; Cowen & Spangale, 2009). Different scales of restricted gene flow or barriers to dispersal have been discovered in the Indo-Pacific region (e.g. Barber et al., 2002; Ravago-Gotanco et al., 2007; Gaither et al., 2011; Crandall et al., 2019) and populations that are potentially isolated or dependent on other reef populations may be vulnerable, and thus would need protection tailored to the unique requirements of these reef areas.

Malaysia is composed of Peninsular Malaysia and East Malaysia separated by the South China Sea. East Malaysia makes up the north-northwest of Borneo and is bounded by the South China Sea in the West and the Sulu Sea and Sulawesi Sea in the East (Fig. 1). The Sulu-Sulawesi seas on the east coast of Sabah mark the boundary of the Coral Triangle for Malaysia, but its exact position there is disputed (Hoeksema, 2007; Veron et al., 2009; Lane & Hoeksema, 2016), and a review by Veron et al. (2015), including new data from the South China Sea (Huang et al., 2015) suggested a westward extension of this biogeographic boundary to include the South China Sea overlying the Sunda Shelf. Few reef areas have been sampled in Malaysia (Gaither & Rocha, 2013)

and very little is known about the connectivity patterns across this region of the South China Sea.

For population genetic studies, *Linckia laevigata* (Linnaeus, 1758) and *Tridacna crocea* Lamarck, 1819 are two of the top five model species with the highest number of locations sampled across the Indian and Pacific Oceans, and further studies were recommended to include mtDNA COI markers to address the gap in sampling localities (Keyse et al., 2014). *Linckia laevigata* is common in most coral reef environments from the Western Indian Ocean to south-eastern Polynesia (Clark & Rowe, 1971; Marsh, 1974; Yamaguchi, 1977). Populations from the Indian Ocean and the West Pacific are genetically dissimilar from each other, with the former predominantly being represented by an orange colour morph and the latter by a blue morph (Williams & Benzie, 1998). This sea star reproduces by broadcast spawning, releasing gametes into the water column for external fertilisation (Yamaguchi, 1973) and its larvae have a long pelagic duration of 22–28 days (Yamaguchi, 1977).

Tridacna crocea is the smallest giant clam in the genus *Tridacna* and can be found in shallow reef environments from the Eastern Indian Ocean to the West Pacific (Othman et al., 2010; Neo et al., 2017). Similar to *L. laevigata*, this *Tridacna* species reproduces by broadcast spawning, but has a pelagic larval duration of 7–14 days (Lucas, 1988; Shokita et al., 1991) and is able to settle on the reef in approximately 10 days (Copland & Lucas, 1988) by boring into the reef substrate.

Previous studies on the population structure of *L. laevigata* and *T. crocea* indicate varying degrees of gene flow among populations of these species over a wide geographical range (Crandall et al., 2008b; Otwoma & Kochzius, 2016; Hui et al., 2017; Keyse et al., 2018). There has been limited studies of these species in Malaysia. Therefore, in this study we aim to examine the genetic population structure of both species to infer the degree of connectivity among the coral reefs of Malaysia. This information is relevant for finding marine biogeographic boundaries across seas around Malaysia and in particular between sites in- and outside the Coral Triangle, with its westernmost boundary still under debate. Based on the possible westward extension of the Coral Triangle (Huang et al., 2015; Veron et al., 2015) and the course of major currents in the South China Sea (e.g. van der Ven et al., 2021, Fig. 1), we expect to find population

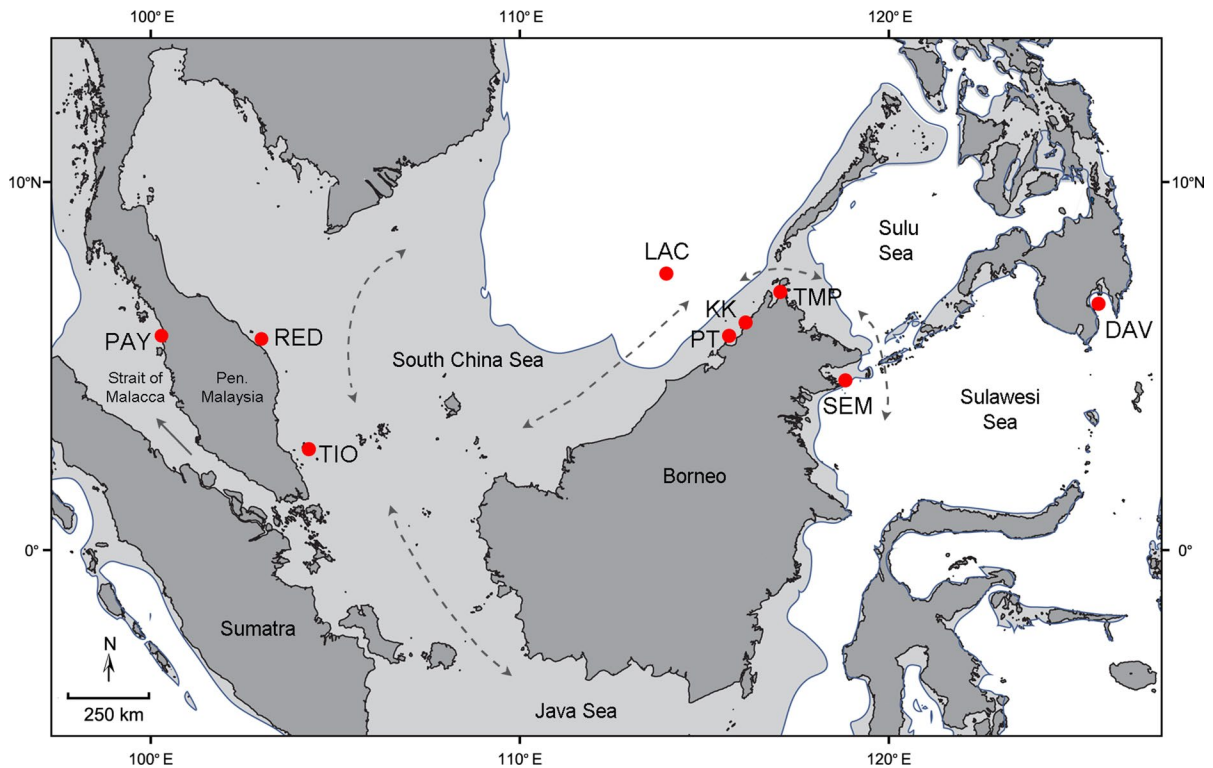


Fig. 1 Sampling sites across Malaysia: Payar (PAY), Redang (RED), Tioman (TIO), Pulau Tiga (PT), Kota Kinabalu (KK), Kudat (TMP), Semporna (SEM), and Pulau Layang-Layang, Spratly Islands (LAC). An additional locality is the Davao Gulf, the Philippines (DAV). Locality data is given in Table 1. Dark grey shading are present-day land formations; whereas

light grey shading are areas exposed during the Last Glacial Maximum (26,000–20,000 years ago) when the sea level stand was >130 m lower than at present (2000 2007, De Groeve et al., 2022). Dashed arrows indicate seasonal reversing current patterns

connectivity among the South China Sea, Sulu and Sulawesi Seas.

Materials and methods

Study area and sample collection

Linckia laevigata and *Tridacna crocea* were sampled in the years 2010–2013 from seven localities across Malaysia (Fig. 1, Table 1), including an atoll, Pulau Layang-Layang (LAC), in the southern Spratly Islands (Waheed et al., 2015a). Sampling was carried out within marine parks (Waheed & Hoeksema, 2013, 2014; Waheed et al., 2015b), except for 44 out of 59 sites in Semporna (SEM) and sites in Layang-Layang (LAC). Additional samples of *L. laevigata* were obtained from the Davao Gulf, the Philippines

(DAV). Samples of the bright blue morph of *L. laevigata* were collected in the form of a piece of arm cutting (~2–3 cm). For *T. crocea* some mantle tissue was collected. All samples were stored in 95% ethanol prior to DNA extraction.

DNA extraction, amplification and sequencing

DNA was isolated from the tissue using the DNeasy Blood and Tissue Kit (QIAGEN) following the standard protocol for animal tissue (version 07/2006). The mitochondrial COI gene was amplified using primers HCO2198 and LCO1490 (Folmer et al., 1994) for *L. laevigata*, and primers COI-Tricro-Frwd and COI-Tricro-Rev (Kochzius & Nuryanto, 2008) for *T. crocea*.

The PCR amplification was carried out in a 25 µl reaction volume, which contained 2.5 µl PCR

Table 1 Sampling localities, codes and coordinates

Site	Location	Code	Number of sampled sites	Latitude (N)	Longitude (E)
Pulau Payar, Peninsular Malaysia	Strait of Malacca	PAY	12	06° 03′–06° 05′	100° 02′–100° 03′
Pulau Redang, Peninsular Malaysia	South China Sea	RED	14	05° 43′–05° 49′	102° 59′–103° 04′
Pulau Tioman, Peninsular Malaysia	South China Sea	TIO	26	02° 42′–02° 56′	104° 03′–104° 13′
Pulau Layang-Layang, Spratly Islands	South China Sea	LAC	18	07° 22′–07° 23′	113° 47′–113° 52′
Pulau Tiga, East Malaysia	South China Sea	PT	6	05° 41′–05° 51′	115° 37′–115° 41′
Kota Kinabalu, East Malaysia	South China Sea	KK	28	05° 57′–06° 12′	115° 59′–116° 05′
Kudat, East Malaysia	Sulu Sea	TMP	35	06° 40′–07° 28′	116° 50′–117° 34′
Semporna, East Malaysia	Sulawesi Sea	SEM	59	04° 06′–04° 48′	118° 10′–118° 57′
Davao Gulf, The Philippines	Sulawesi Sea	DAV	5	06° 29′–07° 01′	125° 34′–125° 44′

CoralLoad Buffer (containing 15 mM MgCl₂) (QIAGEN), 1.0 µl of each primer (10 pmol), 0.5 µl dNTPs (2.5 mM), 0.5 µl Taq polymerase (15 units/µl) (QIAGEN), 18.5 µl of extra pure PCR water and 1.0 µl diluted DNA extract. The PCR cycles consisted of an initial denaturation step of 94 °C for 3 min, followed by 39 cycles of 94 °C for 1 min, annealing temperature of 5 °C for 1 min, an extension step of 72 °C for 1 min and a final elongation step of 72 °C for 5 min. The amplicons were analysed on a 1% agarose gel electrophoresis, stained with ethidium bromide and visualised on a Red™ Personal Imaging System. Successfully amplified samples were sent to MacroGen Europe for PCR cleaning and bidirectional sequencing on an ABI Automated Sequencer 3730xl.

Sequence alignment and phasing

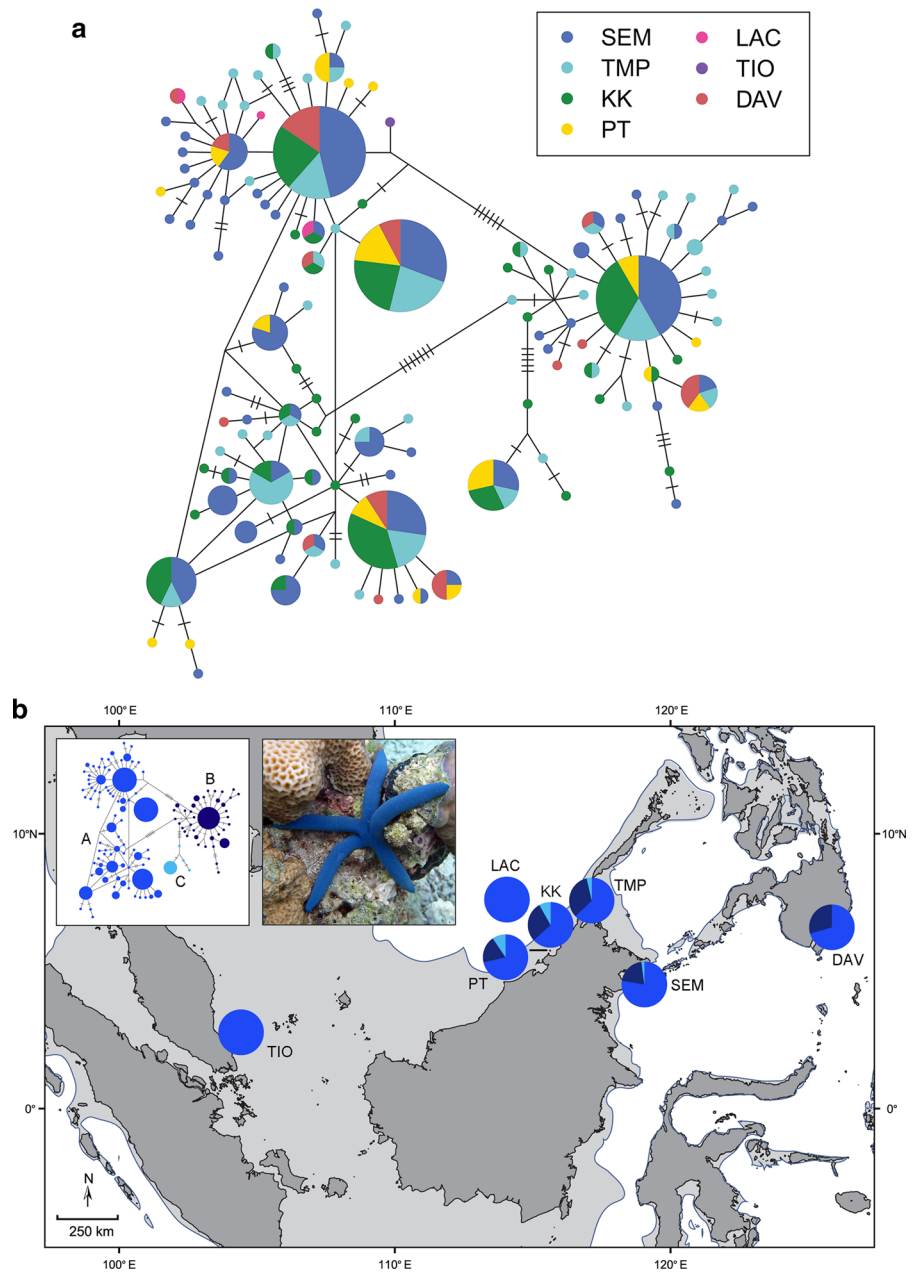
Sequences were assembled and edited with Sequencher 4.10.1 (Gene Codes Corporation), aligned on the GUIDANCE server using MAFFT algorithm (Penn et al., 2010a, b) and trimmed in BioEdit 7.2.5 (Hall, 1999). The sequences did not contain any gaps or ambiguous positions between base pairs. Protein translations were made in MEGA 6 (Tamura et al., 2013) to ensure that stop codons were not present.

Genetic diversity and historical demography

Genetic diversity indices such as haplotype diversity (h) and nucleotide diversity (π) (Nei & Li, 1979; Nei, 1987) were determined for all populations using Arlequin 3.5.1.3 (Excoffier & Lischer, 2010). Tajima's

D (Tajima, 1989) and Fu's F_s (Fu, 1997) were calculated to assess the neutrality of the marker. Significant negative values could indicate a selection sweep or recent population expansion, while positive values could reflect balancing selection or a population bottleneck (Tajima, 1989, 1993; Fu & Li, 1993). Fu's F_s is more sensitive in detecting population growth (Fu, 1997). To further investigate the demographic changes of each species, mismatch distributions were analysed to test the model of sudden population expansion (Slatkin & Hudson, 1991; Rogers & Harpending, 1992; Rogers, 1995). Populations that have undergone growth or expansion are expected to have unimodal and smooth distributions, whereas populations in demographic equilibrium are expected to have multimodal and ragged distributions. Sum of squared deviations (SSD) tests the fit between the observed and expected distribution (Schneider & Excoffier, 1999), whereas Harpending's raggedness index (HRI) evaluates the smoothness of the distributions (Harpending, 1994). Non-significant SSD and HRI values support a sudden population expansion (Harpending, 1994). In order to obtain accurate estimates of the demographic analyses, only locations with more than five samples were included in the analyses, which then excluded Layang-Layang and Tioman for *L. laevigata*. Additional analyses were run using only haplogroup A of *L. laevigata* (Fig. 2) to examine population demographic changes of this haplogroup. All demographic analyses were tested for statistical soundness with 10,000 permutations in Arlequin 3.5.1.3 (Excoffier & Lischer, 2010) with the final significance level determined by sequential Bonferroni corrections (Rice, 1989).

Fig. 2 **a** Haplotype network of *Linckia laevigata*. Each circle is a unique haplotype. The size of the circle is proportional to the haplotype frequency. The line between the circles represents one mutational step, and hatches represent additional mutational steps. **b** Distribution of haplogroups across *L. laevigata* populations



Population structure and gene flow

Population genetic structuring was examined by (1) creating haplotype networks to visualise the relationships among the unique haplotypes, (2) calculating Φ statistics and conducting analysis of molecular variance (AMOVA) to determine the degree of population differentiation among the locations, (3) performing hierarchical AMOVA to identify potential population

groupings based on geographical affinities, and (4) determining the association between genetic distance and geographical distance implemented with Mantel test to infer patterns of isolation by distance and gene flow among the populations.

A haplotype network for each species was constructed based on unique haplotypes using the statistical parsimony principle (Templeton et al., 1992) as implemented in TCS version 1.21 (Clement et al.,

2000). All haplotypes (of *L. laevigata* and *T. crocea*) were included in this analysis. Haplogroups were defined as clusters of haplotypes separated by the highest number of mutational steps (Kochzius et al., 2009). The haplogroups were depicted as pie charts and overlaid onto the map of the study area.

AMOVA was employed to calculate pairwise Φ_{ST} -values among all locations (excluding populations with less than five samples, i.e. Layang-Layang and Tioman for *L. laevigata*). Hierarchical groupings were made by considering the geography of the study site. Hierarchical AMOVA was conducted to calculate Φ_{ST} , Φ_{SC} and Φ_{CT} values to assess genetic variation of each grouping and determine the grouping with the most probable partitioning (largest Φ_{CT} difference). Statistical significance was assessed with 10,000 permutations in Arlequin and significance level was Bonferroni adjusted. The analyses were run using the Tamura and Nei model of nucleotide substitution, which is the best approximation of the TIM1 and TPM3uf models (as determined in jModelTest) available in Arlequin 3.5.1.3 (Excoffier & Lischer, 2010). For *L. laevigata*, a gamma distribution (0.108) was applied based on the most appropriate model of nucleotide substitution determined in jModelTest 2.1.6 (Darriba et al., 2012) based on Akaike Information Criterion (AIC). Additional pairwise Φ_{ST} and hierarchical AMOVA analyses were run using only haplogroup A of *L. laevigata* to account for effects of admixture of the Pacific and Indian Ocean haplogroups (see Crandall et al., 2008b).

For the Isolation by Distance (IBD) analysis, correlation between genetic (pairwise Φ_{ST}) and geographical distance (km) was evaluated using Mantel test (Manly, 1994) and reduced major axis (RMA) regression in Isolation by Distance Web Service (IBDWS) version 3.23 (Jensen et al., 2005) with 10,000 randomisations. Geographical distance was measured as the shortest path between two populations by sea, estimated on Google Earth version 7.1.5.1557.

Results

Sampling efforts were performed at 180 sites in seven localities in Malaysia, 18 sites in Layang-Layang and five sites in the Davao Gulf, the Philippines. Both species were absent from two localities: Pulau Payar in the Strait of Malacca, and Pulau Redang, Peninsular

Malaysia in the South China Sea. *Tridacna crocea* was also absent from Kota Kinabalu and Pulau Tiga at the time of the sampling.

Sequence data were obtained for 353 individuals from both species throughout nine sampling localities. Mitochondrial COI fragments of 601 bp from 233 *Linckia laevigata* and 439 bp from 98 *Tridacna crocea* resulted in 120 and 51 haplotypes, respectively. All unique haplotypes have been deposited in GenBank (*L. laevigata*: ON385154-ON385386; *T. crocea*: ON385440-ON385537).

Genetic diversity and historical demography

Genetic diversity within populations of *L. laevigata* and *T. crocea* was almost similar to each other, albeit slightly higher in the former (Table 2). High values of haplotype (h) and nucleotide (π) diversity were recorded (average values of *L. laevigata*: $h=0.98$, $\pi=1.8$; and *T. crocea*: $h=0.96$, $\pi=1.1$). Further analyses were not conducted for *L. laevigata* from Layang-Layang ($n=3$) and Tioman ($n=1$) due to the small sample size.

Tajima's D values were negative for all *L. laevigata* and *T. crocea* populations, and all values were not significant (Table 2). Fu's F_s values were negative for all *L. laevigata* and *T. crocea* populations, and only significant in Semporna, Kudat and Kota Kinabalu populations of *L. laevigata* and Semporna and Layang-Layang populations of *T. crocea* ($P<0.0001$). Haplogroup A of *L. laevigata* showed similar negative values for Tajima's D and Fu's F_s for all populations. Fu's F_s values were significant in all populations, while Tajima's D value was significant in the Semporna population only (Supplementary Table S1). The overall negative values for *L. laevigata* and *T. crocea* indicate an excess of rare haplotypes compared to what would be expected under neutrality, suggesting selection or population size expansion (Fu, 1997), especially for populations with supporting statistical significance in Fu's F_s test. The mismatch distributions showed multimodal patterns for *L. laevigata* with a significant SSD for the Semporna population (SSD=0.119, $P<0.002$), which indicates a constant population size or a declining population size. The non-significant values of HRI and SSD test for the Kudat, Kota Kinabalu, Pulau Tiga and Davao Gulf populations did not match the mismatch distributions under the population expansion model

Table 2 Summary statistics of *Linckia laevigata* and *Tridacna crocea* at each locality

Area	<i>Linckia laevigata</i>						<i>Tridacna crocea</i>									
	N	H	Genetic diversity		Neutrality tests		Mismatch distribution		N	H	Genetic diversity		Neutrality tests		Mismatch distribution	
			h	π (%)	F _s	D	SSD	HRI			h	π (%)	F _s	D		SSD
Semporna (SEM)	89	57	0.984	1.7	-24.75	-1.52	0.119	0.010	45	30	0.972	0.9	-24.59	-1.69	0.004	0.03
Kudat (TMP)	54	45	0.991	1.9	-24.72	-1.18	0.018	0.009	14	12	0.978	1.5	-4.35	-1.26	0.02	0.08
Kota Kinabalu (KK)	49	36	0.981	2.0	-18.90	-0.85	0.019	0.016	-	-	-	-	-	-	-	-
Pulau Tiga (PT)	20	17	0.984	2.0	-5.86	-0.45	0.016	0.017	-	-	-	-	-	-	-	-
Layang-Layang (LAC)	3	3	1.000	0.6	-	-	-	-	18	15	0.974	0.9	-9.83	-1.53	0.01	0.05
Pulau Tioman (TIO)	1	1	-	-	-	-	-	-	21	9	0.900	1.1	-0.21	-0.97	0.03	0.04
Davao Gulf (DAV)	17	14	0.978	1.7	-4.13	-0.19	0.024	0.024	-	-	-	-	-	-	-	-

Significant P values after Bonferroni correction are in bold

N number of samples, H number of haplotypes, h haplotype diversity, π nucleotide diversity, Fu's F_s, Tajima's D, sum of squared deviations (SSD) and Harpending's raggedness index (HRI)

(Supplementary Fig. S1). However, the mismatch distributions for *L. laevigata* haplogroup A showed unimodal patterns for all populations with a significant SSD and HRI for the Kota Kinabalu population (Supplementary Fig. S2). For *T. crocea*, raggedness indices from the mismatch distributions were all non-significant. Unimodal curves were observed for Sem-porna and Layang-Layang populations, supporting the model of recent population expansion for these two locations (Supplementary Fig. S3).

Population structure and gene flow

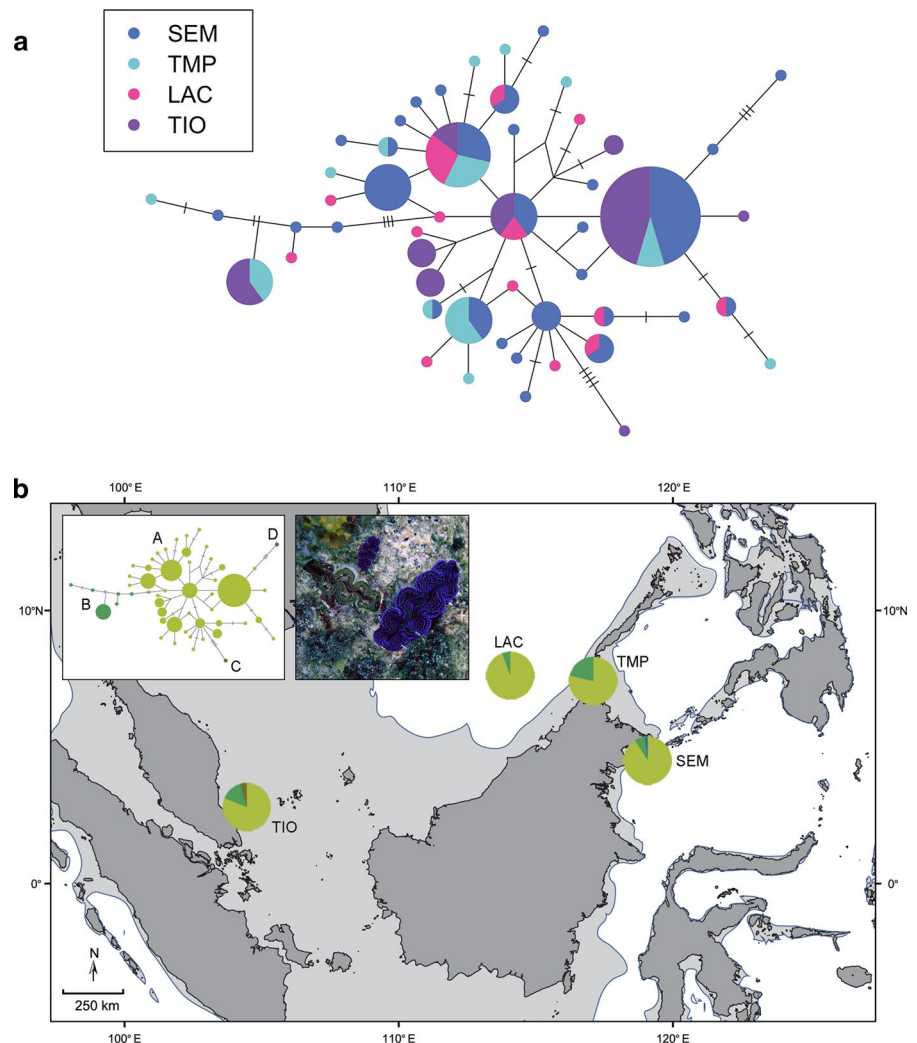
For *L. laevigata*, different haplogroups were separated by five or more mutations. Haplotypes from haplogroup A (163 samples) were shared by all

populations, and haplogroup B (60 samples) was only present in the populations from north Borneo and the Davao Gulf, whereas haplogroup C (10 samples) occurred exclusively in north Borneo, with a higher frequency in the west coast of Sabah (Fig. 2).

In *T. crocea*, the two main haplogroups were separated by three mutations. The distribution of the haplogroups on the map showed no clear pattern of subdivision. All sampled populations contained haplogroups A (86 samples) and B (10 samples), whereas the two singletons were from Tioman (C), and Sem-porna (D) (Fig. 3).

Overall genetic structure for *L. laevigata* was very low ($\Phi_{ST} = -0.001$, $P = 0.399$) and the pairwise Φ_{ST} -values between populations of *L. laevigata* were also very low, suggesting very little to no population

Fig. 3 **a** Haplotype network of *Tridacna crocea*. Each circle is a unique haplotype. The size of the circle is proportional to the haplotype frequency. The line between the circles represents one mutational step, and hatches represent additional mutational steps. **b** Distribution of haplogroups across *T. crocea* populations



differentiation. Similarly, low pairwise Φ_{ST} -values and no population differentiation were observed in populations of *L. laevigata* from haplogroup A. There was subtle genetic structuring in *T. crocea* ($\Phi_{ST}=0.032$, $P=0.009$) and significant genetic structure was detected between populations of Tioman and Semporna ($\Phi_{ST}=0.053$, $P=0.005$), and Tioman and Layang-Layang ($\Phi_{ST}=0.074$, $P=0.006$) (Supplementary Table S2).

Hierarchical AMOVA yielded very low support values in all a priori groupings and none were significant after Bonferroni correction. In both species, the percentage of variation within populations was much higher than the variation among populations within regions and among region groupings (Supplementary Table S3). AMOVA confirmed no significant structure in *L. laevigata* in the region. Furthermore, AMOVA did not detect structure in the assigned groupings for *T. crocea*, despite the significant pairwise Φ_{ST} values between populations of Tioman and Semporna, and populations of Tioman and Layang-Layang.

There was no correlation between genetic and geographical distances across all populations of *L. laevigata* ($r=-0.142$, $P=0.619$) and *T. crocea* ($r=0.067$, $P=0.165$) as indicated by Mantel test (Supplementary Fig. S4).

Discussion

In this study, sampling efforts were performed at seven localities across Malaysia, but both species were not encountered at two localities: Pulau Redang and Pulau Payar, at the east and west coast of Peninsular Malaysia, respectively. There have been anecdotal reports of *L. laevigata* from Redang, but no information is available for the Strait of Malacca, where Pulau Payar is located. Nevertheless, this species has been found in the Andaman Sea, situated northward from the Strait of Malacca (Putchakarn & Sonchaeng, 2004). Populations of *T. crocea* have been reported from the east coast of Peninsular Malaysia (Harborne et al., 2000; Wong & Arshad, 2011) in Pulau Tioman (Tan et al., 1998; Kee Alfian et al., 2005; Neo et al., 2015) and Pulau Redang (Lee et al., 2012), however specimens were only encountered and collected from Tioman. This species has also been reported from Lee Pae Island (=Koh Lipe) and Mu Ko Surin Marine

Park in Thailand (Chantrapornsyl et al., 1996), north of Payar. *Tridacna crocea* was also curiously absent from Kota Kinabalu and Pulau Tiga along the west coast of Sabah, although samples of this species were previously collected from Kota Kinabalu (Kochzius & Nuryanto, 2008).

Genetic variability of *L. laevigata* and *T. crocea* were comparable with other populations in the Indo-Malay Archipelago (*L. laevigata*: Crandall et al., 2008b; Kochzius et al., 2009; Alcazar & Kochzius, 2015, *T. crocea*: Kochzius & Nuryanto, 2008; DeBoer et al., 2008, 2014b; Neo & Todd, 2012; Hui et al., 2016; Kusnadi et al., 2022), which could suggest a similar demographic history among these population (e.g. Duda & Palumbi, 1999).

Population expansion in *L. laevigata* was less clear with conflicting outcomes between the neutrality tests and multimodal mismatch distributions in all populations, even though the neutrality tests indicated possible expansion of Semporna, Kudat and Kota Kinabalu. By contrast, the neutrality tests indicated expansion in populations of *T. crocea*, particularly for Semporna and Layang-Layang and this was further supported by the test of sudden population expansion as well as by the unimodal distribution for these two populations. Population expansion has been observed in populations of *T. crocea* (Neo & Todd, 2012; DeBoer et al., 2014a; Hui et al., 2016) and *L. laevigata* in the Indo-Pacific (Crandall et al., 2008b; Kochzius et al., 2009) including the Visayas in the Philippines (Alcazar & Kochzius, 2015).

An almost similar pattern of haplogroups was retrieved for both species with two main haplogroups (A and B) with shared haplotypes among all sampled populations. A smaller haplogroup (C) composed of three haplotypes was primarily found in north Borneo for *L. laevigata*. Similarly, a haplogroup exclusive to the eastern part of the Visayas, separate from the central Visayas, was detected in the Philippines (Alcazar & Kochzius, 2015). Nevertheless, more samples from other localities are needed to determine the presence of a distinct haplogroup in north Borneo.

Based on pairwise comparisons of populations and hierarchical ANOVA, no significant genetic population structure was detected for *L. laevigata* (in the whole dataset and the dataset containing only haplogroup A). Populations from north Borneo and the Davao Gulf, spanning across the South China Sea, Sulu Sea and Sulawesi Sea were not significantly

different from each other, indicating unrestricted gene flow. Connectivity between the seas is facilitated by north-easterly currents from the South China Sea entering the Sulu Sea through the Balabac Strait during the southwest monsoon between August and October, and the southward currents from the Sulu Sea into the Sulawesi Sea in October. Conversely, currents from the Sulu Sea enter the South China Sea during the northeast monsoon (Wyrтки, 1961; van der Ven et al., 2021; Fig. 1). The seasonally changing currents, possibly coupled with the long pelagic larval duration in *L. laevigata*, may enable larval dispersal across the reef communities of these seas. In addition, the series of reefs between these seas may facilitate larval dispersal (Kimura & Weiss, 1964; Juinio-Meñez et al., 2003).

Weak but significant genetic population structure was detected in *T. crocea*, especially between populations of Tioman and Semporna, and Tioman and Layang-Layang. However, there was no significant structure in the a priori groupings. The water circulation pattern along the east coast of Peninsular Malaysia reverses between the two monsoon seasons (Xu and Malanotte-Rizzoli, 2013). During the northeast monsoon a south-westerly current from the northern rim of the sea passes the coast of Vietnam and branches into a small current that turns north-westwards into the Gulf of Thailand, and a major current that flows southwards through the Karimata Strait into the Java Sea (Xu & Malanotte-Rizzoli, 2013), or turns north-easterly along the northwest coast of Borneo (Wyrтки, 1961; Morton & Blackmore, 2001). The current circulation pattern changes direction during the southwest monsoon, driving a northward current from the Karimata Strait through the central South China Sea towards the northern rim of the sea (Wyrтки, 1961; Morton & Blackmore, 2001; Xu & Malanotte-Rizzoli, 2013). This seasonally reversed circulation pattern creates a complex system of eddies and gyres in the South China Sea (Qu, 2000; Tangang et al., 2011; Xu & Malanotte-Rizzoli, 2013), which could aid in retaining or dispersing larvae across the reefs systems in this shallow sea, thereby connecting the reef populations between north Borneo and the east coast of Peninsular Malaysia. For example, Kochzius and Nuryanto (2008) suggested connectivity between *T. crocea* populations of Kota Kinabalu and Pulau Seribu in the Java Sea. Likewise, in the nearby Bohol Sea, Central Philippines, populations of *T. crocea*

exhibited genetic affinity as a result of prevailing current patterns in the region (Naguit, 2015).

Studies on a larger scale across the Indo-Pacific have revealed that populations of *L. laevigata* and *T. crocea* can be divided into three general regions, (1) Eastern Indian Ocean, (2) central Indo-Malay Archipelago, and (3) Western Pacific (Kochzius et al., 2009; Huelsken et al., 2013; Hui et al., 2016, 2017), with distinct divergence between the Indian Ocean and Pacific Ocean (Williams et al., 2002; Crandall et al., 2008b; Otwoma & Kochzius, 2016). The strongest genetic differentiation has been found to the east of Cenderawasih Bay along northeast New Guinea, thus segregating populations from the west: Cenderawasih Bay, Central Indonesia and Indian Ocean, and the east: Western and Central Pacific (Crandall et al., 2014). Our sampling localities within the South China Sea and Sulu and Sulawesi seas are located to the west of Cenderawasih Bay and are nested within the central Indo-Malay Archipelago, hence exhibited a homogenous population with connectivity among the coral reefs, consistent with the previous findings. Simulation models of connectivity structure distinguished the Spratly Islands (South China Sea), Palawan, the Philippines and the Sulu Archipelago as a region with high levels of connectivity, with the Spratly Islands as an upstream source of larvae for the Sulu Archipelago and the Coral Triangle (Kool et al., 2011) as well as other reefs areas within the South China Sea (Dorman et al., 2015). This underscores the importance of the Spratly Islands and the need to designate a conservation zone or have a concerted effort to establish a management regime for these groups of islands (McManus, 1994; Oakley & Pilcher, 1996; Ablan et al., 2002).

Approximately 26,000–20,000 years ago, when the sea level was >130 m lower than at present (De Groeve et al., 2022), Borneo and Palawan were separated by the Balabac Strait, but by a width of only 12 km, while Borneo and Sulawesi were separated by a narrow, deep oceanic trench (Voris, 2000). These narrow pathways may have allowed gene flow between these seas (but see McManus (1985) for review on the intermittent isolation between the South China Sea and Sulu Sea). According to Potts (1983, 1984, 1985) Plio-Pleistocene regressions occurred too frequently and lasted too shortly for speciation and perhaps this may also explain a lack of differentiation at population level for some species.

At a sea level 30 m lower than the present, land mass between Borneo and Sumatra (including Peninsular Malaysia) were very likely separated (Umbroge, 1947; Voris, 2000; Hoeksema, 2007). At present sea level, an approximately tenfold increase in shallow water habitat since the end of the Last Glacial Maximum around 20,000 years ago has allowed dispersal of marine species across the Sunda Shelf and between the Pacific and Indian Ocean (Bellwood & Wainwright, 2002; Crandall et al., 2012; Gaither & Rocha, 2013). This allows for marine species with moderate to long pelagic larval duration to establish in the Sunda Shelf, such as *T. crocea* and *L. laevigata*, albeit at a slower rate in the former than the latter (see Juinio-Meñez et al., 2003, and for other examples see Crandall et al., 2012), and to disperse under the influence of prevailing monsoonal current patterns.

Our results on the lack of genetic structuring in populations of *L. laevigata* and *T. crocea* in the South China Sea, Sulu and Sulawesi seas of Malaysia and the Davao Gulf in the Philippines suggest connectivity among the sampled reef populations, which is influenced by seasonal circulation patterns in the seas. Samples for this study in Malaysia were collected from marine park areas except for 44 out of 59 sites in Semporna, which are outside of the Tun Sakaran Marine Park boundary (Waheed & Hoeksema, 2013), and sites in Layang-Layang, the Spratly Islands. Semporna has the largest concentration of coral reefs in Malaysia and is known to have high species richness of hard coral, including a few endemic species (Waheed & Hoeksema, 2013). In spite of this, only small areas have been gazetted as the Tun Sakaran Marine Park, and the Sipadan Island Park. Reefs in Semporna should be assessed to determine appropriate conservation or management measures tailored to address the issues confronting these reefs, such as blast fishing (Kassem et al., 2012). Layang-Layang, together with the other islands and reefs in the Spratly Islands, is very likely an important source of larvae for other populations in the South China Sea, the Sulu Sea and the Coral Triangle, and should be part of a network of MPAs in the region (McManus, 1994; McManus & Meñez, 1997; Huang et al., 2016). As yet, no conservation measures have been afforded to these islands and reefs (McManus, 2017; Ablan-Lagman, 2019; Robles, 2020).

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Data availability Sampling locations and mtCOI sequence files are available in GENBANK.

Code availability Not applicable.

Declarations

Conflict of interest On behalf of all the authors, the corresponding author states that there is no conflict of interest.

Ethical approval All applicable international, national and/or institutional guidelines for sampling, care and experimental use of organisms for the study have been followed and all necessary approvals have been obtained. This study complied with all CITES regulations and the necessary permits.

Consent to participate Consent on behalf of all the authors.

Consent for publication Consent on behalf of all the authors.

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