Extremely low genetic variability within and among locations of the greenfish holothurian *Stichopus chloronotus* Brandt, 1835 in Okinawa, Japan

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The greenfish sea cucumber *Stichopus chloronotus* is an economically and ecologically important sea cucumber species throughout its range. This species is widely distributed, inhabiting coral reefs of the Indo-Pacific Ocean. Our study evaluated population genetic structure and levels of genetic diversity in southern Japan. A total of 180 individuals were collected from eight locations from Okinawa and Okinoerabu Islands and sequenced using mitochondrial 16S ribosomal DNA (16S) and nuclear histone H3 (H3) gene. Only three 16S haplotypes were detected with 518 bp with haplotype diversity ranging from 0 to 0.56 and nucleotide diversity from 0 to 0.1%. H3 showed no variation among the studied locations. It is plausible that such results could be due to a shift to asexual reproduction. Additionally, the presence of the species on the east coast of Okinawa could only be detected in one location and all individuals consisted of a single haplotype. Genetic differences between the east and west coasts of Okinawa have been noticed in other coral reef organisms, and attributed to either ecological or biogeographical historical differences between the coasts due to differing levels of isolation during Pleistocene ice ages. Results from the present study should inform management and conservation policies of *S. chloronotus* in southern Japan.
Introduction

Sea cucumbers (holothurians) belong to class Holothuroidea of phylum Echinodermata and play an important and essential part in maintaining and structuring the marine ecosystem. They are deposit feeders of carbonate sand and rubble through their digestive tracts and dissolve CaCO$_3$ as part of their digestive process (Schneider et al. 2011). As ‘ecosystem engineers’, healthy populations of sea cucumbers are important for maintaining the continuing health of many marine ecosystems.

However, sea cucumbers of the order Aspidochirotida (families Holothuroidea and Stichopodidae) are suffering from overexploitation and overfishing in several countries (Anderson et al., 2011; Friedman et al., 2011; Purcell, 2010; Toral-Granda et al., 2008) due to high demand from Chinese and Southeast Asian markets, where they are consumed as food and utilized for traditional medicine (Bordbar et al., 2011). Unfortunately, statistics on fisheries/fishing activities for sea cucumbers are unavailable for many countries as most fisheries organizations still record invertebrate catch as ‘others’. However, some organizations such as the Secretariat of the Pacific Community (SPC) and the Food and Agriculture Organization of the United Nations (FAO) have made efforts to make these records available for invertebrates including sea cucumbers.

The species *Stichopus chloronotus* Brandt, 1835 (greenfish sea cucumber), belonging to the family Stichopodidae, is a member of such highly commercial marine sea cucumber fisheries. This species is a widely distributed across most of the tropical Indo-Pacific region, from the Western Indian Ocean to the Central Pacific, and from southern Japan to northern Australia (Purcell et al., 2012). This species inhabits shallow waters from the intertidal zone to depths of 15 m and can be found on coral reef flats and slopes in dense numbers. Conand et al. (2002) reported on the sex ratios of *S. chloronotus* in some areas of the Indian and Pacific Oceans and
found a very high number of males compared to females. *S. chloronotus* is known to reproduce by two methods; sexual reproduction through indirect development (pelagic auricularia larvae) and asexually by transverse fission (Uthicke, 1997, 1999, 2001). In some marine invertebrates the pelagic larval stage, produced by sexual reproduction, has a large and long-term effect on gene flow via pelagic larval dispersal (Ellingson and Krug, 2016; Pechenik, 1999). The duration of the pelagic larval stage (PLD) of *S. chloronotus* is not well known, although it may be approximately 20 days as seen in other Stichopodidae (Soliman et al., 2013). In the northwestern Pacific, larval dispersal of *S. chloronotus* is influenced by the Kuroshio Current, which is a major ocean current bringing tropical water north from tropical regions along of the Ryukyu Archipelago and Japan's southeast Pacific coast (Barkley, 1970).

In Okinawa, Japan, similar to many other regions of the world, sea cucumbers have been under increasing commercial pressure due to high demand. A recent study showed that some populations of *Holothuria edulis* Lesson, 1830 around Okinawa Island had lower than expected genetic diversity, indicating potential overharvesting and/or ecosystem degradation (Soliman et al., 2016). Prefectural regulations have been put in place to try and limit further damage to sea cucumber populations, but basic genetic and abundance data are lacking for all other sea cucumber species for the entire Ryukyu Archipelago, and for Okinawa Island in particular. In order to properly conserve sea cucumbers around areas of large-scale human populations and coastal development like Okinawa Island, genetic population data are urgently needed.

Population genetic structure studies of *S. chloronotus* on the Great Barrier Reef have previously been carried out through polymorphic allozyme electrophoresis (Uthicke et al., 1999) and amplified fragment length polymorphism (AFLP) markers (Uthicke and Conand, 2005). In addition, there has been one study on the characterization of microsatellite markers for *S.
chloronotus utilizing specimens from the Ryukyu Islands, Japan, although there was no
discussion of biogeographical findings (Taquet et al., 2011). The allozyme and AFLP studies
suggested that population structure and genetic diversity of *S. chloronotus* is affected by asexual
reproduction, with lower than expected genetic diversity due to the presence of clones.

In the present study we assess the genetic diversity and population structure among
eight *S. chloronotus* locations in southern Japan (Okinawa and Okinoerabu Islands) using
sequences of mitochondrial DNA (16S ribosomal DNA) and nuclear DNA (histone H3), in order
to provide genetic connectivity. We examine whether *S. chloronotus* on the east and west coasts
of Okinawa are genetically differentiated as has been observed in a number of recent studies on
other sea cucumber (Soliman et al. 2016) and other invertebrate species (White et al., 2016). This
information will be useful for future conservation and fisheries management. Based on recent
increasing commercial pressure in Okinawa, we also expect that locations near to large human
populations may show reduced *S. chloronotus* genetic diversity, as well as a potential shift to
asexual reproduction due to reduced water quality, as was previously observed in *H. edulis*
(Soliman et al. 2016).

**Materials and methods**

A total of 180 specimens of the greenfish sea cucumber *S. chloronotus* were collected
from February to April 2015 by snorkeling and SCUBA diving at seven locations around
Okinawa Island (Itoman, Oyama, Zanpa, Ryugu, Motobu, Yona, Teniya) and one location at
Okinoerabu Island, Kagoshima, Japan (Table 1, Fig. 1). During our surveys, *S. chloronotus* was
not found at any locations on the eastern coast of Okinawa Island aside from at Teniya (Soliman
et al., unpublished data). Samples of *S. chloronotus* were obtained non-lethally through body
wall-tissue biopsy (1 cm²) as in our previous study (Soliman et al. 2016) and preserved in
absolute ethanol in the field. Live animals were released back to the same location from where they were collected.

Total genomic DNA was extracted from a small piece (0.25 g) of body wall tissue using a DNeasy Blood & Tissue extraction kit (Qiagen, Tokyo, Japan) following the manufacturer’s protocol. Polymerase chain reactions (PCR) were used to amplify fragments of two DNA markers; mitochondrial 16S and nuclear H3 gene, using the following primers: 16SA-R 5’- CGC CTG TTT ATC AAA AAC AT -3’; 16SB-R 5’- GCC GGT CTG AAC TCA GAT CAC GT -3’ (Palumbi et al., 1991), and H3a: 5’- ATG GCT CGT ACC AAG CAG ACV GC -3’; H3b: 5’- ATA TCC TTR GGC ATR ATR GTG AC -3’ (Colgan et al., 1998), respectively. Amplification reactions were carried out in a 15 µl volume containing 5 to 20 ng of template genomic DNA, 0.5 µM of each primer, and 10 µl of HotStarTaqTM Master Mix (Qiagen, Tokyo, Japan), in deionized water. PCR reactions were performed on an Eppendorf thermocycler (Eppendorf AG, Hamburg, Germany) with an initial denaturation step at 95°C for 15 min, 35 cycles of denaturation at 94°C for 1 min, an annealing at 46°C (16S) and 48°C (H3) for 1 min, extension at 72°C for 1 min, and a final extension step at 72°C for 10 min. The amplification products were tested using electrophoresis (1.5% agarose gel) and purified with Exonuclease I and Alkaline Phosphatase Shrimp (Takara) and incubated at 37°C for 20 min, followed by deactivation at 83°C for 30 min. Purified PCR products were sequenced using an ABI Prism automated sequencer at Fasmac Co., Kanagawa, Japan (http://www.fasmac.co.jp/index.html), in both forward and reverse directions.

Consensus sequences of 16S and H3 were aligned and edited using Geneious v.8.1.3 (http://www.geneious.com, (Kearse et al., 2012), and MEGA v.6.0 (Tamura et al., 2013). The H3 sequences showed no variation and no additional analyses were conducted. For 16S, number of haplotypes ($h$), haplotype diversity ($h_d$) and nucleotide diversity ($\pi$) were estimated with DNASP
v.5.10.01 (Librado and Rozas, 2009). The best-fit model of DNA sequence evolution was estimated using MEGA v.6.0 (Tamura et al., 2013). The AIC (Akaike Information Criterion) indicated that Jukes-Cantor (JC) was the best-fit model. We computed pairwise $\Phi_{ST}$ and average of pairwise differences (Nei’s distance, within and between locations) for all pairs of locations using Arlequin 3.5.1.2 (Excoffier and Lischer, 2010; Weir and Cockerham, 1984). The simultaneous tests correction was adjusted using the modified false discovery rate (FDR) method (Narum 2006). To investigate genetic partitioning among regions, among locations within regions, and between all locations we applied a hierarchical analysis of molecular variance (AMOVA), using Arlequin, and tested the following hypothesis; A – Three groups: Okinoerabu Island as group 1, Teniya (east) as group 2 and all other western locations (Itoman, Motobu, Oyama, Ryugu, Yona, Zanpa) as group 3; and B – Two groups: Teniya from east as group 1 and western locations of Okinawa Island (Itoman, Motobu, Oyama, Ryugu, Yona, Zanpa) plus Okinoerabu Island as group 2. Principal Coordinates Analysis (PCoA) based on 16S haplotype frequencies among locations were calculated and visualized using PAST 3.11 (Hammer et al., 2001).

Results

We obtained sequences of a 518 bp fragment of 16S from 175 specimens (GenBank Accession numbers KX522950-KX523124) from eight total locations around Okinawa and Okinoerabu Islands (Table 1, Figure 1). The alignment of 16S sequences resulted in three haplotypes: Hap_1 was common among seven locations (Itoman, Oyama, Zanpa, Ryugu, Motobu, Yona, Okinoerabu), Hap_2 was shared among five locations (Itoman, Oyama, Ryugu, Motobu, Teniya), and Hap_3 was exclusive to Oyama (Figure 1 and 2). Four locations showed no haplotype diversity for 16S; all 69 sea cucumbers sampled in Zanpa, Yona (both on
Okinawa’s west coast), and Okinoerabu were Hap_1 while all 15 specimens sampled at Teniya (Okinawa’s east coast) consisted of Hap_2 (Table 1 and Fig. 1). Overall, the haplotype diversity was low ($h_d = 0.36$), and was highest in the Oyama ($h_d = 0.558$) and Ryugu ($h_d = 0.5143$) locations. In addition, the nucleotide diversity of 16S was low for all locations combined ($\pi = 0.0008$; Table 1).

We also obtained a 307 bp fragment of the nuclear H3 gene from 180 individuals of *S. chloronotus* collected in the same locations around Okinawa and Okinoerabu Islands. Obtained H3 sequences showed no variation. Therefore, all subsequent population genetics analyses were based on 16S data only.

Significant pairwise $\Phi_{ST}$ values were found in 17 out of 28 comparisons after multiple tests correction ($P < 0.01$) (Table 2). Most of the significant comparisons were between locations not sharing haplotypes, Teniya (east coast of Okinawa Island) vs Yona and Zanpa (west coast of Okinawa Island) or with Okinoerabu Island. Although the null hypothesis of panmixia in the overall AMOVA test (global $\Phi_{ST} = 0.559$, $P < 0.001$) was rejected, results of the hierarchical AMOVAs did not support significant differences between east and west coasts or between north (Okinoerabu Island), east of Okinawa Island (Teniya) and west of Okinawa Island (Itoman, Motobu, Oyama, Ryugu, Yona, Zanpa) (Table 3).

Principle Coordinates Analyses (PCoA) using overall pairwise $\Phi_{ST}$ for 16S revealed three clusters; 1) Yona, Zanpa, Motobu, Itoman and Okinoerabu, 2) Ryugu and Oyama, and 3) Teniya (Figure 4).

**Discussion**

The present study revealed low levels of genetic diversity of the greenfish sea cucumber within and between the southern Ryukyu Islands of Japan, with only three haplotypes in 16S and one in
These low levels of polymorphism in *S. chloronatus* impose some caution in the overall interpretation of results. Various studies of *S. chloronotus* across regions using different genetic markers have also revealed low genetic diversity. Uthicke and Conand (2005) observed low genetic diversity in the southwestern Pacific (Great Barrier Reef, Torres Strait) compared to the western Indian Ocean using amplified fragment length polymorphism (AFLP) analyses, and suggested that AFLP markers are a suitable method to examine clonality in sea cucumbers. In addition, allozyme studies have showed a low number of polymorphic loci in *S. chloronotus* in some locations such as the Great Barrier Reef, Torres Strait, and La Reunion (Uthicke et al., 1999; Uthicke et al., 2001). The low genetic diversity results may stem from reported high rates of asexual reproduction in *S. chloronotus* in several other Indo-Pacific regions, as well as a biased male sex ratio (Conand et al., 2002). In addition, strong variation of rates of asexual reproduction over small geographic scales in *S. chloronotus* has been reported in genetic surveys and reproductive biology studies (Conand et al., 2002; Uthicke et al., 1999). This pattern seems to be common in other sea cucumbers: low genetic diversity has been observed in *Holothuria edulis* around Okinawa Island using mitochondrial (16S and COI) and nuclear H3 genes (Soliman et al., 2016), and in *Holothuria atra* in Okinawa and the Ogasawara Islands based on COI sequences (Skillings et al., 2011). We posit that our observations of low genetic diversity could be attributed to shifts to asexual reproduction in *S. chloronotus*.

Low habitat stability combined with high seawater concentrations of nutrients (eutrophication) and small individual sizes (caused by over-exploitation) have been identified as the most important factors that can lead to asexual reproduction in sea cucumbers (Conand et al., 2002). Over the past 40 years, coastal development and landfill have reduced and degraded shallow water habitats and water quality along the shoreline of Okinawa Island, and this has been
implicated in reductions in biodiversity of near-shore marine communities (Reimer et al., 2015). These anthropogenic impacts may also at least partially explain the reduction in genetic diversity among these locations. Overharvesting can also lead to reductions of genetic diversity (Smith et al. 1991; Pinsky et al. 2014) and from our field observations at different sites of Okinawa Island it appears that some sea cucumber species such as *H. edulis* and *S. chloronotus* are overexploited (Soliman et al., 2016). Similar anecdotal evidence on a reduction in sea cucumber numbers led Okinawa Prefecture to implement a harvesting licensing system in 2013 with further strengthening in 2016.

The haplotype frequency differences between the west coast of Okinawa and the single location on the east coast may be due to the existence of habitat differences, oceanographic barriers to dispersal promoting a reduction of larval exchange, and/or historical factors. We extensively searched for *S. chloronatus* individuals on the east coast, however, Teniya was the only location on this coast where the species could be found, further indicating the possibility of the above mentioned factors. Furthermore, our results indicated the presence of gene exchange among all Okinawa Island west coast locations and Okinoerabu Island. This pattern is consistent with the route of the Kuroshio Current flowing northward to the west of Okinawa Island (Figure 1), and suggests this current may play a key role in connecting locations of *S. chloronatus*. On the other hand, areas isolated from the Kuroshio Current, such as the east coast of Okinawa Island, may harbor genetically distinct populations. Extensive geographic fine grid surveys on the east coast of Okinawa Island and from regions more isolated from the Kuroshio Current (e.g. the Daito Islands 310 km to the east of Okinawa Island; the disputed Senkaku Islands 410 km west of Okinawa Island) in southern Japan are needed to confirm this assertion. Similar patterns of genetic differentiation between east and west Okinawa Island locations have been observed in recent studies on the sea cucumber *Holothuria edulis* (Soliman et al., 2016) and in the amphipod...
Leucothoe vulgaris (White et al., 2016). White et al. (2016) suggested this reduction caused by Pleistocene sea level changes (Ni et al., 2014) and/or geographic discontinuity (Castelin et al., 2012). In contrast, in one species of scleractinian coral (Acropora digitifera), Okinawa Island locations from both east and west coasts showed no separation and were treated as one panmictic population (Shinzato et al., 2015). Differences in the level of genetic structure could be explained by the shorter pelagic larval duration in the sea cucumber [approx. 20 days in Stichopodidae; Soliman et al. (2013)] than in A. digitifera [approx. 45 days; Nishikawa and Sakai (2005)]. However, multiple studies point to the complexity of making predictions on genetic connectivity based on the length of pelagic larval development (e.g. Selkoe et al., 2014; Shanks, 2009). Nevertheless, from the few population genetics studies published in sea cucumbers to date (Eastwood et al. 2016; Skillings et al., 2014; Soliman et al., 2016; Valente et al., 2015; Vergara-Chen et al., 2010) there is a generalized pattern of highly geographically structured genetic variation and reduced dispersal ability, which may compromise the persistence of sea cucumber populations and species when facing intense fishing pressure, habitat degradation, or other environmental threats.

In conclusion, the present genetic survey of S. chloronotus indicates low levels of genetic diversity and reduced genetic exchange among sites in Okinawa and Okinoerabu islands, and particularly between the east and west coasts of Okinawa Island. These results will be useful for fisheries management decisions regarding this species in southern Japan. Both Soliman et al. (2016) and the present study also suggest that more research on sea cucumbers is needed from the Ryukyu Islands in order to properly conserve and protect the genetic diversity of these economic and ecologically important animals.
Acknowledgments

S. Hisashi is deeply thanked for his help on Okinoerabu Island – he will be deeply missed. We also thank R. Diaz for help with sample collection and members of the MISE laboratory for their logistic and scientific support.

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Table legends

**Table 1** Molecular diversity of *Stichopus chloronotus* of mitochondrial 16S ribosomal DNA sequences from eight locations in southern Japan (*n*: Sample size; *h*: number of haplotypes; *h*d: haplotype diversity; *π*: nucleotide diversity).

**Table 2** Pairwise estimates of *Φ*ST among eight locations of *Stichopus chloronotus* in southern Japan inferred from a 518 bp DNA fragment of the mitochondrial 16S ribosomal gene from 175 specimens.

**Table 3** Hierarchical analyses of molecular variance (AMOVA) based a 518 bp fragment of the mitochondrial 16S ribosomal gene from 175 specimens of *S. chloronotus* testing different hypotheses of geographical groupings among eight locations in southern Japan. Hypothesis A:
Group 1 = Okinoerabu Island, Group 2 = Teniya, Group 3 = Itoman, Motobu, Oyama, Ryugu, Yona, Zanpa. Hypothesis B: Group 1 = Okinoerabu, Itoman, Motobu, Oyama, Ryugu, Yona, Zanpa, Group 2 = Teniya.

Figure legends

Figure 1. Map with sampling locations and frequency of 16S ribosomal DNA haplotypes of *Stichopus chloronotus* from Okinawa and Okinoerabu Islands, Japan. The arrows depict the prevailing path and direction of the Kuroshio Current.

Figure 2. Median-joining haplotype network inferred from 16S ribosomal DNA sequences of *Stichopus chloronotus*. Each circle represents a different haplotype and the size of a circle is proportional to the frequency of each haplotype. Colours denote the proportion of haplotypes collected at each location as indicated by the embedded key. Haplotypes separated by black crossbars differ by a single nucleotide, whereas hatch marks indicate unsampled haplotypes.

Figure 3. Average number of pairwise differences within and among locations of *Stichopus chloronotus* for 16S ribosomal DNA.

Figure 4. Principal Coordinates Analysis (PCoA) based on 16S ribosomal DNA haplotype frequencies of *Stichopus chloronotus* locations from Okinawa and Okinoerabu Islands.
Table 1 (on next page)

Molecular diversity of Stichopus chloronotus of mitochondrial 16S ribosomal DNA (16S) sequences from eight locations (= populations) in southern Japan (n: Sample size; h: number of haplotypes; hd: hap
| Locations               | Lat. and Long.                     | n  | h  | $h_d$ | $\pi$     | Fu's Fs | Fu's Fs p-value |
|-------------------------|-----------------------------------|----|----|-------|----------|---------|-----------------|
| Okinoerabu Island       | 27°25'05.2"N 128°39'00.6"E       | 24 | 1  | 0.0000| 0.0000   | 0.0000  | N.A.            |
| Yona, Okinawa Island    | 26°45'56.6"N 128°11'42.4"E       | 24 | 1  | 0.0000| 0.0000   | 0.0000  | N.A.            |
| Motobu, Okinawa Island  | 26°40'45.8"N 127°52'55.4"E       | 24 | 2  | 0.0833| 0.0002   | -1.0279| 0.079           |
| Ryugu, Okinawa Island   | 26°31'48.6"N 127°55'34.8"E       | 21 | 2  | 0.5143| 0.0011   | 1.4737  | 0.757           |
| Zanpa, Okinawa Island   | 26°25'58.8"N 127°42'57.2"E       | 21 | 1  | 0.0000| 0.0000   | 0.0000  | N.A.            |
| Oyama, Okinawa Island   | 26°17'01.2"N 127°44'20.5"E       | 22 | 3  | 0.5584| 0.0013   | 0.1995  | 0.504           |
| Itoman, Okinawa Island  | 26°07'15.1"N 127°39'34.7"E       | 24 | 2  | 0.1594| 0.0003   | -0.2489| 0.212           |
| Teniya, Okinawa Island  | 26°33'52.6"N 128°08'16.0"E       | 15 | 1  | 0.0000| 0.0000   | 0.0000  | N.A.            |
**Table 2** (on next page)

Pairwise estimates of ΦST among eight populations of *Stichopus chloronotus* in southern Japan inferred from a 518 bp DNA fragment of the mitochondrial 16S ribosomal gene from 175 specimens.
| Location | Okinoerabu | Yona | Motobu | Ryugu | Zanpa | Oyama | Itoman | Teniya |
|----------|------------|------|--------|-------|-------|-------|--------|--------|
| Okinoerabu | 0.000     |      |        |       |       |       |        |        |
| Yona     | 0.000     | 0.000 |        |       |       |       |        |        |
| Motobu   | 0.020     | 0.000 | 0.000  |       |       |       |        |        |
| Ryugu    | 0.569     | 0.569 | 0.486  | 0.000 |       |       |        |        |
| Zanpa    | 0.000     | 0.000 | 0.006  | 0.550 | 0.000 |       |        |        |
| Oyama    | 0.441     | 0.441 | 0.351  | 0.021 | 0.422 | 0.000 |        |        |
| Itoman   | 0.044     | 0.044 | 0.028  | 0.408 | 0.036 | 0.271 | 0.000  |        |
| Teniya   | 1.000     | 1.000 | 0.947  | 0.355 | 1.000 | 0.473 | 0.894  | 0.000  |

Significant $\Phi_{ST}$ at $P < 0.01$ following the False Discovery Rate (FDR) correction (Narum, 2006) indicated in bold.
Table 3  (on next page)

Hierarchical analyses of molecular variance (AMOVA) based a 518 bp fragment of the mitochondrial 16S ribosomal gene from 175 specimens of S. chloronotus testing different hypotheses of geographical gr
| Hypothesis                        | Variation | d.f. | SS  | Variation (%) | Fixation indices | P value | d.f. | SS  | Variation (%) | Fixation indices | P value |
|----------------------------------|-----------|------|-----|---------------|------------------|---------|------|-----|---------------|------------------|---------|
| Among groups                     |           | 2    | 17.22 | 47.08         | $\Phi_{CT} = 0.4781$ | 0.1691  | 1    | 16.09 | 70.85         | $\Phi_{CT} = 0.7085$ | 0.1202 |
| Among population within groups   |           | 5    | 11.56 | 20.28         | $\Phi_{SC} = 0.3885$ | <0.0001 | 6    | 12.7  | 11.56         | $\Phi_{MC} = 0.3966$ | <0.0001 |
| Within populations               |           | 167  | 22.06 | 31.91         | $\Phi_{ST} = 0.6782$ | <0.0001 | 167  | 22.06 | 17.59         | $\Phi_{ST} = 0.8241$ | <0.0001 |

d.f.: degree of freedom; SS: sum of squares
Figure 1

Map with sampling locations and frequency of 16S ribosomal DNA (16S) haplotypes of Stichopus chloronotus from Okinawa and Okinoerabu Islands, Japan. The arrows depict the prevailing path and direction.
Figure 2

Median-joining haplotype network inferred from 16S ribosomal DNA sequences of Stichopus chloronotus. Each circle represents a different haplotype and the size of a circle is proportional to the frequency.
Figure 3

Average number of pairwise differences within and among populations of Stichopus chloronotus for 16S ribosomal DNA.
Figure 4

Principal Coordinates Analysis (PCoA) based on 16S ribosomal DNA haplotype frequencies of Stichopus chloronotus populations from Okinawa and Okinoerabu Islands.