

# [Phylogeographic analysis suggests a recent population bottleneck in the rare red ...](https://assignbuster.com/phylogeographic-analysis-suggests-a-recent-population-bottleneck-in-the-rare-red-sea-tridacna-squamosina/)

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## Introduction

The recognition of giant clams (subfamily Tridacninae) as a contributor to net primary productivity and biomass of Red Sea coral reefs is driving a growing research interest in their abundance, ecological roles and population genetic structure ( [Roa-Quiaoit, 2005](#B30) ; [Richter et al., 2008](#B29) ; [Pappas et al., 2017](#B26) ; [Rossbach et al., 2019a](#B33) , [b](#B34) ; [Fauvelot et al., 2020](#B8) ). They are a common food source within the coral reef community, with 75 known predators, including fishes such as wrasses, triggerfishes, and emperor fishes ( [Neo et al., 2015](#B23) ). Among these predatory fishes, 11 species naturally occur in the Red Sea ( [Golani and Bogorodsky, 2010](#B12) ), suggesting that giant clams are an important food source for coral reef food webs in this region. Three species of giant clams, specifically *Tridacna maxima* , *T. squamosa* , and *T. squamosina* are found in the Red Sea. The first two species are listed in the International Union for Conservation of Nature (IUCN) Red List of Threatened Species while the latter is a relatively recent resurrected species that has not been assessed for IUCN Red Listing, which requires information on their ecology, habitat, and density ( [Neo, 2020](#B22) ).

To date, at least 31 reported sightings of *T. squamosina* have been documented from the Red Sea since their first description during the Pola Red Sea Expeditions 1895/96 and 1897/98 ( [Huber and Eschner, 2011](#B14) ). The observations from the literature include 16 individuals from [Roa-Quiaoit (2005)](#B30) and [Richter et al. (2008)](#B29) , seven individuals from [Huber and Eschner (2011)](#B14) , two individuals from [Fauvelot et al. (2020)](#B8) and six individuals from [Rossbach et al. (2020)](#B32) . The past and present known distribution of *T. squamosina* are Egypt, Jordan, Yemen, Saudi Arabia, and possible sightings in Israel based on photographic anecdotes ( [Richter et al., 2008](#B29) ; [Huber and Eschner, 2011](#B14) ; [Neo et al., 2017](#B24) ; [Fauvelot et al., 2020](#B8) ; [Rossbach et al., 2020](#B32) ). Although a recently published analysis using genome skimming has provided highly credible phylogenetic relationship between *T. squamosina* and the other Tridacninae ( [Tan et al., 2021](#B38) ), information about their population genetic structure across the Red Sea is still lacking.

Cryptic diversity within giant clams was evident through the recent rediscovery of *T. noae* ( [Su et al., 2014](#B36) ; [Borsa et al., 2015](#B5) ) and *T. elongatissima* ( [Fauvelot et al., 2020](#B8) ) with the aid of molecular tools. *Tridacna squamosina* is another example of a cryptic species, having previously been identified to be a local variant of *T. squamosa* in the Red Sea. This species has been recently resurrected by [Richter et al. (2008)](#B29) using mitochondrial gene marker (16S) with detailed descriptions on its morphological characteristics. Mitochondrial cytochrome c oxidase subunit I (mtCOI) gene sequencing is a rapid and powerful tool for accurate identification of giant clam species in the Indo-Pacific ( [Nuryanto et al., 2007](#B25) ). However, to our knowledge, there was no reference mtCOI sequences of *T. squamosina* available prior to the study conducted by [Fauvelot et al. (2020)](#B8) . Although the authors have successfully amplified the mtCOI sequences of two *T. squamosina* individuals collected from the Red Sea, only short fragments (i. e., 234 base pairs) were obtained. Given the scarcity of reference sequences for this species, our study aims to amplify the mitochondrial sequences of the rare *T. squamosina* using the *Tridacna* specific primers, to improve barcode reference libraries and investigate their population structure. This paper highlights the limited genetic diversity of *T. squamosina* in the Red Sea, which indicates a need for local conservation programs to maintain and even enhance their low populations.

## Materials and Methods

### Study Method

Six specimens of *Tridacna squamosina* from a reef in Farasan Banks, as part of the large-scale giant clam survey that covered 58 unique reefs (∼1, 300 km) along the eastern Red Sea. For each *T. squamosina* , we recorded the water depth and shell length. *In situ* images (camera model: Canon G7x) with their mantles exposed were taken to catalog species ( [Figure 1](#F1) ). Biopsies of the mantle tissues of each clam were cut using surgical scissor and forceps, the samples were then stored in seawater at −80°C. We combined our results with those of [Richter et al. (2008)](#B29) and [Fauvelot et al. (2020)](#B8) , to document the known distribution of *T. squamosina* in the Red Sea ( [Figure 2](#F2) ).

FIGURE 1

Specimens of the(a) smallest and(b) largest *T. squasomina* specimens that were found in Farasan Banks. Scale bar = 5 cm.

FIGURE 2

Fifty-eight reefs studied during a large-scale giant clam survey along the eastern Red Sea (black and red dots). The red dot indicates the sampling site where the specimens of *Tridacna squamosina* were found and sampled for this study while yellow dots indicate the other three locations where *T. squamosina* were reported in the previous studies.

### DNA Extraction, Amplification and Sequencing

DNA was extracted using the QIAGEN DNeasy ® Blood and Tissue Kit (QIAGEN, Hilden, Germany). Two mitochondrial gene markers (mtCOI and 16S) were amplified and sequenced. First, a fragment of the mtCOI gene for the giant clam was amplified via polymerase chain reaction (PCR) with SQUA-R1 (5′–ATG TAT AAA CAA AAC AGG ATC–3′) and SQUA – F3 (5′–CAT CGT TTA GAG TAA TAA TTC G–3′) ( [DeBoer et al., 2008](#B6) ). PCR was performed using the QIAGEN ® Multiplex PCR Kit (QIAGEN, Valencia, CA, United States) in a total volume of 25 μL containing 2. 5 μL of genomic DNA, 12. 5 μL of QIAGEN Multiplex PCR master mix, forward and reverse primers at a concentration of 10 mM (1. 25 μL each) and RNase-Free water to adjust the volume. The protocol for isolating the mtCOI gene locus was modified from [DeBoer et al. (2008)](#B6) as follows: initial denaturation at 95°C for 15 min, 35 cycles of 94°C for 1 min to denature the DNA, 45°C for 1. 5 min to anneal the DNA, 72°C for 1 min for elongation of the DNA, and a final elongation step at 72°C for 5 min. The second fragment used for barcoding was part of the large 16S ribosomal RNA gene which was amplified using the 16sar-L (5′–CGC CTG TTT ATC AAA AAC AT–3′) and 16sbr-H (5′–CCG GTC TGA ACT CAG ATC ACG T–3′) primers ( [Richter et al., 2008](#B29) ); PCR was conducted using the same conditions as the mtCOI gene except the annealing temperature and time were set to 43°C and 30 s respectively. The PCR products were checked using UV light after running in a 1% agarose gel (1 × TAE) pre-stained with SYBR Safe dye (Invitrogen Corp., Carlsbad, CA, United States) under 100 V for an hour. Amplicons were bead-cleaned using Agencourt AMPure XP (Beckman Coulter) and both forward and reverse strands were sequenced using Sanger sequencing at the KAUST Bioscience Core Lab.

### Sequence Alignment and Phylogenetic Inference

Sequences were aligned and trimmed to a common length of 500 base pairs (bp) for mtCOI and 410 bp for 16S, using Geneious R9 (Biomatters Ltd., Auckland, New Zealand). To infer the phylogenetic relationship of Red Sea *T. squamosina* , six 16S sequences and 18 reference sequences (MN068557, MN068558, and AM909726–AM909741; [Supplementary Table 1](#TS1) ) were included in the analysis. The dataset was exported to DnaSP to estimate the genetic diversity. A 16S haplotype network was constructed in PopArt [1](#footnote1) using TCS network approach. Using the same 16S dataset, we estimated the value of π and θ from the Tajima’s test of neutrality to gain insight into the species’ population history (see [Templeton, 1993](#B39) ) using MEGA 7. 0. 14. We used Arlequin v. 3. 5 ( [Excoffier and Lischer, 2010](#B7) ) to test for genetic structure with Analysis of MOlecular Variance (AMOVA). As there is a genetic break around 19°N reported among various reef species due to the heterogeneous environmental conditions in the region ( [Froukh and Kochzius, 2007](#B9) ; [Nanninga et al., 2014](#B21) ; [Giles et al., 2015](#B10) ), we calculated the AMOVA by grouping the samples into two geographic clusters based on the proximity of their environmental variables. The northern population consists of populations from Jordan and Egypt (SST: ∼26°C, salinity: ∼40) and the southern population consists of Farasan Banks population (SST: ∼22°C, salinity: ∼37). As we had only one individual from the Farasan Island, this locality was omitted from population analyses. We used the same software to calculate Wright Fisher’s *F ST* to infer the relative level of genetic difference between the northern and southern populations. As the low number of mtCOI sequences (i. e., less than 20 samples) obtained in current study can cause incomplete estimates of genetic diversity within species ( [Phillips et al., 2019](#B28) ), the mtCOI sequences from current study were combined with the short reference sequences (MN068808 and MN068809; [Supplementary Table 1](#TS1) ) to visualize the haplotype network and no further analysis is done.

## Results

All individuals were found shallower than 3 m water depth, with shell lengths ranged from 15 to 30 cm (mean = 22. 5 ± 4. 6 cm). Twelve sequences from six individuals were successfully amplified using the *Tridacna* specific primers (six for mtCOI and 16S, respectively), and they were deposited in GenBank with accession numbers MW411954–MW411959 (mtCOI), MW407155–MW407160 (16S). The genetic diversity of *T. squamosina* based on the 16S sequences from the pooled dataset was 0. 163 ± 0. 099. From the 16S haplotype network, three unique haplotypes, representing the 24 individuals of Red Sea *T. squamosina* were detected. Five out of six samples collected in the current study shared the same haplotype with other samples found in the northern Red Sea and Farasan Islands, in the southern Red Sea (H1) ( [Figure 3A](#F3) ). The Tajima estimator (π) and coalescent estimator of genetic diversity (θ) values were 0. 001 and 0. 003, respectively. AMOVA analysis showed that 99. 81% of variations measured in the overall population are found within individuals of the same populations ( [Table 1](#T1) ). The fixation index ( *F ST* ) among populations was 0. 00189, which was not significant. One individual (G6) from current study shared the same point mutation with the Egypt specimen (MN068808) at the 51st position based on the mtCOI alignment (i. e., 234 bp) in Geneious R9 ( [Supplementary Figure 1](#FS1) ), congruent with the haplotype network as depicted in [Figure 3B](#F3) .

FIGURE 3

Haplotype networks based on(A) 24 16S sequences and(B) eight mtCOI sequences of *T. squamosina* in the Red Sea. Each circle corresponds to one haplotype (H for 16S; Hap for mtCOI) and the size of each circle is proportional to its frequency. Connecting lines between haplotypes represent one mutational step with additional mutational steps represented by the hatches.

TABLE 1

Analysis of molecular variance of three populations in the Red Sea. Group refers to the pooling of Jordan and Egypt together (northern population) and Farasan Banks alone (southern population).

## Discussion

This study confirms the occurrence of *Tridacna squamosina* in the South-eastern Saudi Arabian Red Sea, based on DNA sequences. [Roa-Quiaoit (2005)](#B30) reported that *T. squamosina* is geographically restricted at the marginal tip of the Gulf of Aqaba, where the cooler water temperatures are assumed to favor this species. However, our results combined with the finding of a single individual from Farasan Island by [Fauvelot et al. (2020)](#B8) , confirm that this species also occurs in the warmer, productive waters of the southern Red Sea. Apart from the environmental factors, their current distribution pattern could be affected by the settlement behavior of giant clam larvae and the geographical events. The disjointed distribution of *T. squamosina* under two extreme environmental conditions in the Red Sea is perplexing, given that the environmental conditions in the central Red Sea should also be suitable for this species. The most parsimonious explanation for this distribution pattern is a local extinction in the central Red Sea as suggested by [Pappas et al. (2017)](#B26) from their field survey data. The ecology and habitat of *T. squamosina* we describe here is similar to that of the bear paw clam ( *Hippopus hippopus* ), which has suffered substantial range contraction and local extinction in Pacific Islands as a result of human harvesting ( [Seeto et al., 2012](#B35) ) as well as glacio-eustatic sea level fluctuations ( [Paulay, 1996](#B27) ). Unlike most other Tridacninae species, *T. squamosina* is not permanently attached to a hard substrate, thus making it more susceptible to manual harvesting by hand [see [Neo et al. (2017)](#B24) and references therein]. This species may have been the main target for human collection in the Red Sea coast during the last interglacial period (> 1, 250, 000 years ago) ( [Richter et al., 2008](#B29) ), but the ongoing exploitation of *Tridacna* stocks in the northern, central, and southern Red Sea was also evident in the past decades ( [Bodoy, 1984](#B4) ; [Gladstone, 2000](#B11) ; [Ashworth et al., 2004](#B3) ). The decline in wild *Tridacna* stocks in the Red Sea could also be linked with coastal development around the major cities. In early 2000s, reduced coral cover (i. e., 20%, at 5 m) was observed in central Red Sea (e. g. Jeddah) due to the urban tourism and city development, in comparison to less disturbed reefs in the North and South ( [Kotb et al., 2004](#B15) ). High sedimentation can hinder the settlement and recruitment of giant clam larvae ( [Guest et al., 2008](#B13) ), thus jeopardizing their recruitment success in the central Red Sea. Hence, we hypothesize that the disjoint distribution of *T. squamosina* is a result of extirpation of populations between the Gulf of Aqaba in the North and the Farasan regions in the South by the combined effects of historical overexploitation and coastal development.

The overall size of *T. squamosina* found in this study (15–30 cm; six individuals) was generally smaller than reported by [Richter et al. (2008)](#B29) (27–34 cm; five individuals), but meaningful conclusions from this remains limited. The sexual maturation of giant clams can be inferred from their gonad status ( [Richter et al., 2008](#B29) ) and hence, biopsies of the gonadal tissue can help to estimate their age. However, we did not perform gonad biopsies due to logistic limitations and further investigations are needed to verify clam age. [Mohammed and Yassien (2008)](#B20) reported that high coral cover plays a major role in the bivalve assemblages in the Red Sea, it is however, the opposite for *T. squamosina* . All individuals were found free-living on the sandy-rubble flats with extremely low coral cover (i. e., ∼5%, see [Anton et al., 2020](#B2) ), supporting the hypothesis that this species might have overcome the competitive advantages of its sympatric congeners by inhabiting shallow areas with low coral cover. With respect to our genetic results, we obtained *T. squamosina* mtCOI sequences that were twice the length (i. e., 500 bp) of those from [Fauvelot et al. (2020)](#B8) . The reasons for this large difference are unclear. Our results confirm that the use of SQUA-primers for DNA barcoding is successful for amplification, identification and discrimination of the rare species from its congeners in the Red Sea. Low mitochondrial genetic diversity inferred from the pooled 16S dataset suggests that this population is exposed to an increased risk of extinction ( [Markert et al., 2010](#B19) ), but the data should be treated with caution as low genetic diversity could also result from a small population size. Low levels of genetic diversity observed in marine organisms is often associated with overexploitation and habitat degradation in the past ( [Rodrigues et al., 2008](#B31) ), suggesting that *T. squamosina* has experienced a recent population bottleneck. The number of haplotypes ( *n* = 3) and variable sites were extremely low ( *n* = 5/410, 1%). This level of variation is comparable to the rare bear paw clams ( *Hippopus hippopus* ) in the Philippines using the same marker ( *n* = 1/498, 0. 2%; [Lizano and Santos, 2014](#B17) ). As the surveyed population is too small to make a comparison, it is challenging to distinguish between a species having naturally low levels of genetic variation and a significant loss in variation as the result of a population bottleneck. To overcome this limitation, we looked into the species’ population history. The coalescent estimator of genetic diversity (θ = 0. 003) ( [Watterson, 1975](#B41) ) was larger than Tajima estimator (π = 0. 001) ( [Tajima, 1983](#B37) ), suggesting historically higher diversity for *T. squamosina* .

The genetic structure of *T. squamosina* based on 16S mtDNA marker revealed in this study is important for ecological conservation and restoration. The fact that major proportion of the variability (99. 81%) is present within populations inferred from AMOVA analysis implies that outbreeding depression is not a concern, and sourcing donor clams for future restocking programs does not have to be restricted to certain sites or environmental conditions. The low, non-significant *F ST* showing weak mitochondrial DNA (mtDNA) differentiation between northern and southern populations of *T. squamosina* in the Red Sea could be attributed to high gene flow. Such a weak population structure noticed in the present study is in accordance with [Lim et al. (2020)](#B16) , where a panmictic population was identified for its sister species, *T. maxima* in the same region. However, the limited sample size of *T. squamosina* specimens collected in this study did not allow us to fully address the true population structure. Thus, the genetic differentiation could be overestimated in our scenario. While mtDNA is one of the popular choices of genetic markers in conservation genetics, it has a few drawbacks in interpreting population connectivity as discussed in [Zhang and Hewitt (2003)](#B43) and [Wan et al. (2004)](#B40) . Firstly, the maternal inheritance of mtDNA restricts it to exploring events at the maternal angle, which may underrepresent the species’ true dispersal potential. Secondly, it has limited use in investigating recent loss of genetic variation and any individual-level events such as identity, individual dispersal, and mating systems. Future studies should include the use of microsatellite markers to study the demographic connectivity of Red Sea giant clams at these spatial scales, which was tested in other endangered marine bivalves based on the isolation by distance (IBD) hypothesis ( [Wesselmann et al., 2018](#B42) ).

In conclusion, our study suggests that *T. squamosina* is an endangered species in the Red Sea. Given their already low population abundance and genetic variability, this species deserves an enhanced monitoring effort in Saudi Arabia. Overharvesting of Red Sea giant clams was probably one of the main drivers for their historical population decline ( [Gladstone, 2000](#B11) ; [Ashworth et al., 2004](#B3) ). Nowadays, fishing pressure seems to have a rather negligible impact on the present giant clams stocks in the region ( [Rossbach et al., 2020](#B32) ). The particularly sharp decline in *T. squamosina* densities, however, may be caused by its weak attachment to the substratum by its byssus threads, making it more vulnerable to collection by hand than the other Tridacninae species in the same region. Nonetheless, we call for actions to protect and conserve the remaining population of *T. squamosina* along with biobanking and fertility preservation programs to safeguard remnant populations in the Red Sea, paralleling successful efforts in cryopreserving the gametes and embryos of other marine bivalve molluscs elsewhere in the world ( [Marina et al., 2020](#B18) ). Lastly, integrated coastal zone management plans need to be implemented in the coastal cities to mitigate the impacts of coastal developments and wastewater discharges on nearby reefs. Hence, we urge the relevant authorities to work closely with the coastal communities toward conservation of giant clams, such has been proposed on listing *Tridacna* spp. as a key taxa of high conservation priority in Saudi Arabia ( [AbuZinada et al., 2004](#B1) ).

## Data Availability Statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/ [Supplementary Material](#S9) .

## Author Contributions

SR collected the samples. KL conducted the barcoding with assistance from NG and ES. KL took the lead in writing the manuscript. All authors discussed the results and commented on the manuscript.

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## Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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## Supplementary Material

The Supplementary Material for this article can be found online at: https://www. frontiersin. org/articles/10. 3389/fmars. 2021. 628142/full#supplementary-material

Supplementary Figure 1 | Alignment of eight *T. squamosina* mtCOI sequences showing the two specimens sharing the same point mutation site on the 51st position.

Supplementary Table 1 | Summary information of Red Sea *Tridacna squamosina* sequences downloaded from GenBank used in the phylogenetic analysis and comparison purpose.

## Footnotes

1. [^](#footnote1a) [http://popart. otagoac. nz](http://popart.otagoac.nz/)

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