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ORIGINAL ARTICLE



Population genetics of the hound needlefish *Tylosurus crocodilus* (Belonidae) indicate high connectivity in Tanzanian coastal waters

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ABSTRACT

The hound needlefish *Tylosurus crocodilus* (Belonidae) is a highly demanded fish in the local markets of Tanzania, but the growing coastal population threatens its sustainability. As belonids are highly migratory fishes utilising various parts of the seascape, increased fishing pressure may disrupt connectivity patterns on different spatiotemporal scales and disaggregate populations. Using the COI gene, this study assessed the genetic population structure, connectivity patterns, and historical demography of *T. crocodilus* collected in seven sites spread along Tanzanian coastal waters. Results showed fourteen haplotypes with low overall nucleotide and haplotype diversity. Pairwise F_{ST} comparisons revealed no significant differences among the sampled sites, except for the northernmost site (Tanga) and an island in the south (Songosongo). Analysis of molecular variance (AMOVA) revealed a non-significant genetic structure among populations ($F_{ST} = 0.01782$), suggesting the fishery across Tanzanian waters exploits the same population. Moreover, there was no correlative relationship between genetic and pairwise geographic distances, rejecting the isolation by distance hypothesis. However, neutrality tests and mismatch distribution analysis revealed that recent demographic expansion might exist. Empirical evidence of panmixia suggests high genetic connectivity. In combination with low genetic diversity, management should be directed to actions that prevent genetic diversity loss and the effect of genetic drift on populations.

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Introduction

The hound needlefish (*Tylosurus crocodilus*) is a marine epipelagic fish widely distributed in tropical oceans (Froese and Pauly 2022). It is a habitat generalist (Gaither et al. 2016) occupying a range of coastal marine ecosystems (Riede 2004) and is the most prominent member of the family Belonidae. It is estimated to reach a maximum length of 1.5 m and a maximum weight of 6.4 kg (IGFA 2001). The hound needlefish is mostly caught from seagrass beds, coral reefs and offshore waters (Silas 2022). This species is in high demand in the tropics and subtropics, including the Western Indian Ocean (WIO) markets, because of their tender flesh and few bones (Cervigón et al. 1992). It is caught in high numbers in artisanal fisheries and forms an important food source for coastal

communities (Froese and Pauly 2022), but as such, it is also subject to high fishing pressure. The fish is generally caught by casting and trolling from a boat and sometimes using seine nets. In Tanzania, however, it is mostly targeted using handlines and ringnets, with the latter often catching juveniles and spawning aggregations (Bett et al. 2021), which may impede recruitment (Alonzo and Mangel 2004; Vasilakopoulos et al. 2011) and potentially lead to a loss of genetic diversity (Pinsky and Palumbi 2014). The hound needlefish has long been managed as a single (randomly mating) population in Tanzanian waters. However, because there are no data on patterns of genetic connectivity, it is not known whether the current fisheries management approach is directed towards conserving the genetic population structure of the locally adapted

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population of this species as well as other unexplored key fishery species in the area.

The genetic population structures of marine fish species are largely determined by ocean currents and behavioural processes (White et al. 2010). The hound needlefish as many coral reef-associated species exhibits ontogenetic migration (Igulu et al. 2014). Like most belonids, it spawns in coastal waters and lays eggs with tendrils and sticky surfaces attached to floating objects (Breder and Rosen 1966; Bhatt and Bhargava 1973; Polte and Asmus 2006). Their eggs and pelagic larvae are dispersed by ocean currents to various locations, facilitating recruitment in the receiving populations (Dorenbosch et al. 2006; Riginos et al. 2014). With a planktonic nature and an egg incubation period of 8–10 days (Cole 2010), they have the potential to disperse across wide geographical distances (Riginos et al. 2014; Gaither et al. 2016), depending on local hydrodynamics and barriers. The fish are pelagic (Claro 1994), occupying the lagoons and seaward reefs either as individuals or in small groups.

Understanding the genetic connectivity of marine populations is crucial to be able to determine the recovery potential of marine species, especially when subjected to overfishing or other stressors. Failure to incorporate such information in management can lead to loss of genetic diversity (and genetic drift), potentially leading to a collapsing fishery (Reiss et al. 2009; Cadrin et al. 2014). This was experienced in Atlantic cod, where the fishery was subjected to overfishing and failed to recover because management approaches were not aligned with the genetic population structure of the fishery (Lage et al. 2004; Zemeckis et al. 2014).

In coastal East Africa, high fishing pressure and non-selective fishing gears (e.g. ringnet) characterise artisanal fisheries, which mostly catch juvenile pelagic fishes. Such fishing practice may interfere with gene dispersal and recruitment patterns of the caught species. This highlights the need to get information on genetic diversity and patterns of connectivity of marine fishes, which is important for successful conservation and management of coastal fisheries. Most population studies on East African marine fishes have reported extensive genetic connectivity (Okada and Sasaki 2005; Dorenbosch et al. 2006; Borsa et al. 2016; Huyghe and Kochzius 2017; Johnson et al. 2021). In contrast, limited gene flow has been reported using microsatellites and mitochondria markers in fiddler crabs (Nehemia and Kochzius 2017) and mangrove snails (Nehemia et al. 2017) mainly due to mangrove habitat degradation.

This study aimed to assess the genetic population structure, connectivity patterns and historical demography of the hound needlefish *T. crocodilus* in Tanzanian coastal waters. Given the strong currents in the area, we hypothesised genetic isolation by distance and not a well-mixed population.

Materials and methods

Field sampling was conducted in different locations spread along the Tanzanian coast (Figure 1). The region contains diverse and extensive seascapes containing mangrove forests, seagrass meadows, and

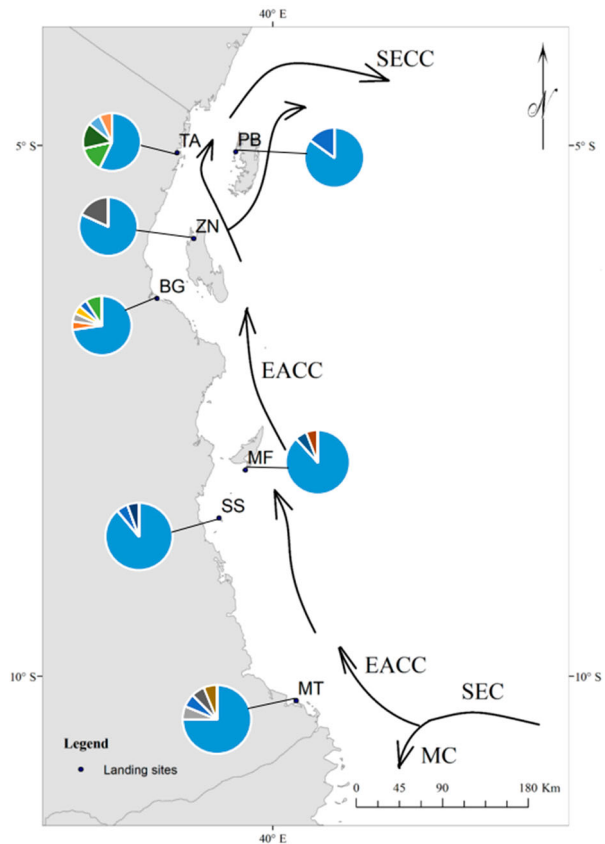


Figure 1. Sampling locations in Tanzania and pie charts showing the haplotype frequencies at each sampling site. The sampled sites include Zanzibar (ZN), Bagamoyo (BG), Tanga (TA), Mafia (MF), Songosongo (SS), Pemba (PB), and Mtwara (MT). The colours in the pie charts indicate the unique and/or shared haplotypes among the different sampling sites. The haplotype in blue (haplotype number one) was dominant and abundant in all sampling sites. Apart from the haplotype one dominating the sites, other haplotypes were at least shared by two or more sites, except the remainder two haplotypes at Songosongo (SS) that were uniquely not found at the highest haplotype diversity site in Tanga (TA). The arrows show major ocean currents in the region, which include the East African Coast Current (EACC), the South Equatorial Current (SEC), the South Equatorial Counter Current (SECC), and the Mozambique Current (MC).

coral reefs supporting marine biodiversity. The East African Coastal Current (EACC) is the major ocean current in the region, emerging from splitting the South Equatorial Current (SEC) near the border of Tanzania and Mozambique (Figure 1). The EACC transverses into the Somali Current along the coast of Kenya (Obura et al. 2019). The climate in the area is influenced by monsoon winds, with the southeast monsoon dominating from May to September and the northeast monsoon prevailing from October to April (McClanahan 1988). Therefore, the current speed varies depending on the monsoon and is higher during the SE monsoon period than in the NE monsoon period (Semba et al. 2019), which may have a seasonal effect on larval dispersal.

Sampling

Tissue samples of 118 hound needlefish (*Tylosurus crocodilus*) were collected between 2016 and 2021 from seven landing sites along the Tanzanian coast, including Bagamoyo (BG), Mafia (MF), Mtwara (MT), Pemba (PB), Songo-songo (SS), Tanga (TA), and Zanzibar (ZN) (Figure 1). On each site, specimens were collected from multiple fishers on different occasions to enhance the randomness and independence of sampling sites. Fishers in the studied sites utilise ringnet to catch pelagic species, which includes the hound needlefish. Fish specimens were identified based on local field guides (Bianchi 1985; Richmond 2011). The locations and overview of the COI sequences submitted to GenBank are presented in Table 1. A piece of the pectoral fin of each fish was excised and preserved in absolute ethanol at room temperature until further analysis.

DNA extraction, amplification of fragments of the COI, and gene sequencing

The total genomic DNA was extracted from the sampled tissues using the QIAGEN Blood & Tissue Kit

according to the protocol provided by the manufacturer. The quality and concentration of DNA extracts were checked using a NanoDrop Lite Spectrophotometer. Fragments of the COI gene were amplified using the primers FishF1-5'TCAACCAACCACAAA GACATTGGCAC3' and FishR2-5'ACTTCAGGGTGACCGA AGAATCAGAA3'. Each PCR reaction was conducted in a total volume of 50 µL containing 0.2 mM dNTP, 3 mM MgCl₂, 0.3 U of the *Thermus aquaticus* DNA polymerase, 1x standard Taq reaction buffer, 5 mg bovine serum albumin, 0.3 µM of each primer, and 2.5 µL of the DNA template. The temperature profile used in the PCR reaction included 94°C for 3 min, 35 cycles [94°C for 30 s, 54°C for 45 s, 72°C for 60 s], and 72°C for 15 min. Samples were sequenced using the Sanger sequencing technology on an ABI 3730 DNA Analyzer (Applied Biosystems).

Data analysis

Sequence editing was performed in the software CHROMASPRO (v. 2.1, Technelysium Ltd, Leicester, UK). Inspection and corrections were conducted to ensure the correct fitting of nucleotides to the right chromatography peaks. DNA barcoding for species identification was accomplished using the two Identification Systems (IDS) for COI; GenBank (<https://www.ncbi.nlm.nih.gov/genbank>, last consulted on 15/11/2021) and the data portal of Barcode of Life Data Systems (BOLD Systems, <http://v3.boldsystems.org/>, accessed on 16/11/2021). All samples were identified as the hound needlefish *T. crocodilus* at 99% similarity. Sequences were trimmed to the least common length, and 552 bp length was conserved for further analyses. Because the reference sequences were slightly shorter (518 bp), the sample sequences were reduced to 518 bp during the phylogeny construction. The quality control check was conducted where the COI sample sequences were translated into amino acids. This was done to rectify sequencing errors and ensure that the sample sequences represented the functional mitochondrial DNA and not pseudogenes. The sequences were uploaded to GenBank and given the accession numbers ON644875-ON644992. The phylogenetic reconstruction (see Appendix 1) was conducted in the MEGA X (Kumar et al. 2018) by applying the Maximum Likelihood method using the Kimura 2-parameter model (Kimura 1980). FaBox DNA Collapser (Villesen 2007) (<https://usersbirc.au.dk/palle/php/fabox/dnacollapser.php>, accessed on 16/11/2021) was used to reduce the sequences into haplotypes. Fourteen haplotypes were obtained in this process and used to generate an input file for population

Table 1. Summary of the amplified COI sequences submitted to GenBank, site locations and the sampled *Tylosurus crocodilus* specimen's n = sample size.

Site	Site code	Latitude	Longitude	n	GenBank accession numbers
Bagamoyo	BG	−6.436648	38.906450	22	ON644876 – 897
Mafia	MF	−7.900883	39.675290	17	ON644898 – 914
Mtwara	MT	−10.28533	40.187918	16	ON644915 – 930
Pemba	PB	−5.039935	39.785469	21	ON644931 – 950
Songosongo	SS	−8.518225	39.499065	17	ON644951 – 967 & ON644875
Tanga	TA	−5.081787	39.120887	14	ON644968 – 981
Zanzibar	ZN	−6.169694	39.199789	11	ON644982 – 992

structure analyses. Haplotype diversities were computed based on the methodology used by Nei (1987) in the Arlequin software version 3.5 (Excoffier and Lischer 2010). Additional reference sequences of *T. crocodilus* were retrieved from GenBank and added to the dataset during the phylogeny construction.

The genetic connectivity patterns were assessed using analysis of molecular variance, AMOVA (Excoffier et al. 1992), in the Arlequin software version 3.5 (Excoffier and Lischer 2010). The overall extent of genetic differentiation (F_{st}) and the pairwise F_{st} values were computed to determine the dataset structure and compare variations among different paired sites. The resulting pairwise F_{st} p-values were adjusted using the Bonferroni correction (Holm 1979). A Minimum Spanning Tree of the haplotypes was constructed using PopART version 1.7 (Bandelt et al. 1999) to explore relationships among the sampled populations. Furthermore, a correlation between genetic (pairwise F_{st} – values) and geographic distances was assessed using the Mantel test (Mantel 1967) in the Isolation by Distance (IBD) software version 1.52 (Bohonak 2002). The assessment examined if there was an association between genetic and geographic distances along the gradient coastline of Tanzania. In this case, the geographic distances among the sampled locations were estimated in Google Earth as the shortest marine pathways connecting different sampling sites. The final graphical

output was constructed using the R programming software (R Core Team 2019).

The historical demography was analysed using the programme Arlequin version 3.5. The analysis tested the marker's null hypothesis of neutral evolution by applying the Tajima's D (Tajima 1989) and Fu's FS (Fu 1997) tests. The demographic metrics were computed to detect if there was a recent population expansion or a bottleneck. Furthermore, a mismatch distribution analysis was performed to affirm the patterns in the previous tests and identify any signs of selective pressure and demographic growth (Excoffier and Schneider 1999). Finally, the sum of square deviation and Harpending's raggedness index were calculated to evaluate the sudden population expansion model (Rogers and Harpending 1992; Harpending 1994).

Results

Genetic connectivity patterns

AMOVA indicated a substantially higher genetic variation within sampling sites (98.22%) than between sites (1.78%; Table II). Furthermore, the analysis revealed low and non-significant F_{ST} values, suggesting that the null hypothesis of panmixia cannot be rejected ($F_{ST} = 0.018$, $p > 0.05$). Similarly, all pairwise F_{ST} values were non-significant, except for one pairwise test between Tanga (TA) and Songo-songo (SS) (Table III). The non-significant pairwise tests and the AMOVA results indicate high connectivity among populations in the different sampled sites. Furthermore, the isolation by distance analysis showed no correlation between genetic and geographic distances ($Z = 147.86$, $r = -0.05$, $p = 0.58$; Figure 2). The frequency relationships among the 14 haplotypes obtained in this study were demonstrated using the haplotype network (Figure 3), which showed a star-like shape with one dominant haplotype shared by 95 samples and several singletons containing up to three mutation steps.

Table II. Results of analysis of molecular variance (AMOVA) among sampled sites of the *Tylosurus crocodilus* from the Tanzanian coast.

Source of variations	Degree of freedom	Sum of squares	Variance component	Percentage of variance
Among sites	6	1.450	0.00336	1.78
Within sites	111	20.575	0.18536	98.22
Total	117	22.025		

Fixation Index F_{ST} : 0.01782.

Significance tests (1023 permutations).

Va and F_{ST} : P (rand. value > obs. value) = 0.14.

P (rand. value = obs. value) < 0.001.

P -value = 0.14 ± 0.013 .

Table III. Pairwise genetic (F_{ST} values, below diagonal) and geographic distances (km, above diagonal) between sampled sites of *Tylosurus crocodilus* along the coast of Tanzania. In bold are the significant values after sequential Bonferroni correction when $k = 21$ (significance level: $p < 0.05$). Where $k = 21$ represents the number of pairwise tests. For site abbreviations, see Table I.

	TA	PB	ZN	BG	MF	SS	MT
TA							
PB	0.103						
ZN	0.055	0.027					
BG	0.001	0.006	0.001				
MF	0.104	0.007	0.008	0.012			
SS	0.113	0.026	0.013	0.010	0.028		
MT	0.022	0.014	0.040	−0.032	−0.001	−0.007	

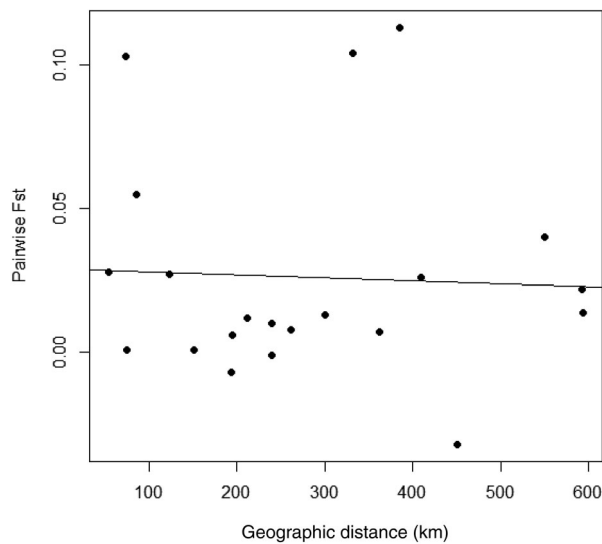


Figure 2. Graph showing the relationship between pairwise F_{ST} and geographic distance ($r = -0.0469$) from 21 pairwise comparisons of the seven *Tylosurus crocodilus* populations sampled along the coast of Tanzania.

Genetic diversity and historical demography

The sampled fish showed 14 haplotypes associated with low overall haplotype and nucleotide diversities ($h = 0.38 \pm 0.06$ and $\pi = 0.09 \pm 0.09$). The haplotype diversity ranged from 0.22 ± 0.12 at Songosongo (SS) to 0.67 ± 0.13 at Tanga (TA) (Table IV). The samples from Bagamoyo had the highest number of haplotypes ($N_h = 6$) followed by Tanga ($N_h = 5$), while the

specimens from Zanzibar and Pemba ($N_h = 2$) had the lowest number of haplotypes (Table IV). Nucleotide diversity (π) among samples was lowest at Songosongo and Mafia (both $\pi = 0.04 \pm 0.06$) and highest at Tanga ($\pi = 0.19 \pm 0.15$; Table IV).

The overall Tajima D and Fu's F_s values were negative and significant, suggesting purifying selection or recent population expansion (Tajima D = -2.08 , $p < 0.05$; Fu's $F_s = -15.26$, $p < 0.01$). The mismatch distribution analysis of the observed and expected pairwise differences produced a unimodal distribution typical for expanding populations. The raggedness index and sum of square deviations were low and insignificant, indicating a good fit of the observed and expected pairwise differences to the population expansion model (SSD = 0.0001, HRI = 0.16, $p > 0.05$; Table IV).

Discussion

Genetic connectivity patterns

The findings of this study indicate that the hound needlefish *Tylosurus crocodilus* (belonging to Belonidae) in Tanzania belongs to a single panmictic population, as evidenced by the constructed haplotype network and the results of the AMOVA. The haplotype network analysis displayed a distinct central dominant haplotype present at all sampled locations, indicating that the sampled fish from the different locations exhibit

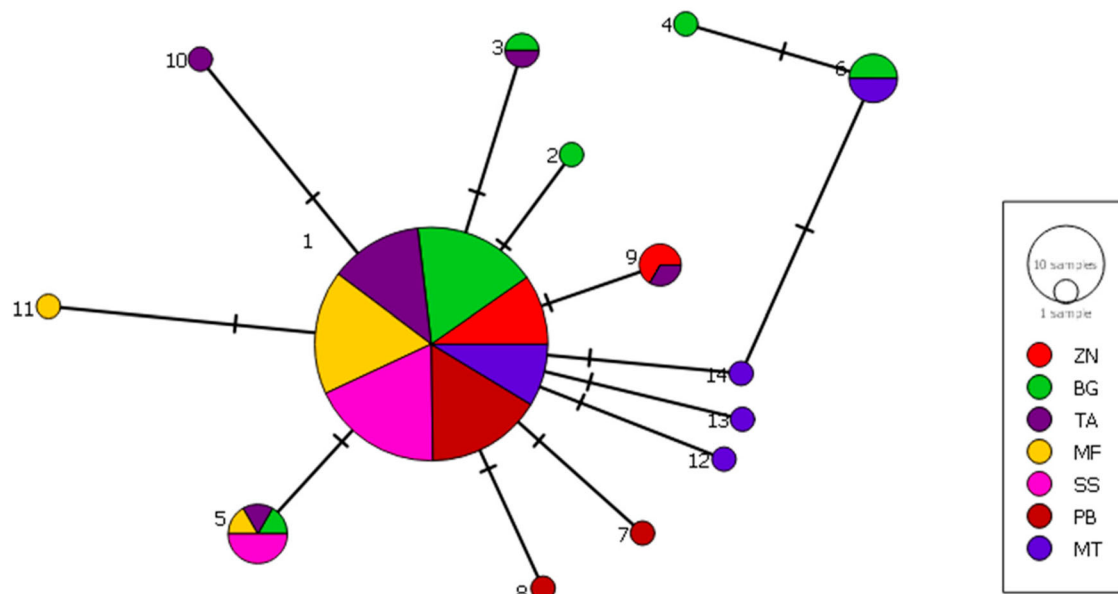


Figure 3. A haplotype network of sequences of the *Tylosurus crocodilus* samples from the coast of Tanzania. The circle size represents the number of sample sequences in that haplotype. The biggest inner circle equals 95 samples, while the smallest outer circles represent one sample each. The sampled sites include Zanzibar (ZN), Bagamoyo (BG), Tanga (TA), Mafia (MF), Songosongo (SS), Pemba (PB), and Mtwara (MT).

Table IV. Results of the genetic diversity, neutrality tests, and mismatch distributions of the *Tylosurus crocodilus* samples collected along the coast of Tanzania. n = sample size; N_{hp} = number of haplotypes; h = haplotype diversity (\pm SD); π = nucleotide diversity (\pm SD), Tajima D, Fu's Fs, SSD = sum of square deviation, HRI = Harpending's raggedness index. Significance levels: *0.05 > p > 0.01; **p \leq 0.01.

Population	Code	n	N _{hp}	Genetic diversity		Neutrality test		Mismatch distribution	
				h	π (%)	Tajima's D	Fu's Fs	SSD	HRI
Overall		118	14	0.38 \pm 0.06	0.09 \pm 0.09	−2.08	−15.26	0.0001	0.16
Zanzibar	ZN	11	2	0.33 \pm 0.15	0.06 \pm 0.07	−0.10	0.36	0.003	0.23
Bagamoyo	BG	22	6	0.48 \pm 0.13	0.16 \pm 0.13	−1.49*	−2.58*	0.306*	0.11
Mtwara	MT	16	5	0.45 \pm 0.15	0.09 \pm 0.09	−1.83*	−3.31**	0.007	0.15
Songosongo	SS	18	3	0.22 \pm 0.12	0.04 \pm 0.06	−1.51	−1.74**	0.002	0.37
Pemba	PB	20	2	0.27 \pm 0.11	0.05 \pm 0.06	−0.09	0.38	0.263	0.29
Mafia	MF	17	3	0.23 \pm 0.13	0.04 \pm 0.06	−1.50	−1.68*	0.003	0.35
Tanga	TA	14	5	0.67 \pm 0.13	0.19 \pm 0.15	−0.59	−1.55	0.001	0.05

extensive genetic connectivity in Tanzania. Further evidence of high genetic connectivity was revealed by AMOVA, which produced non-significant indices of genetic differentiation between the sampling sites, and no isolation by distance could be detected. These results align with other genetic studies conducted on mud crabs (Rumisha et al. 2018), prawns (Mwakosya et al. 2018) and octopus (Van Nieuwenhove et al. 2019) on the East African coast. In addition, the lack of genetic structure has also been shown for belonids due to the transport of larvae by strong and warm currents in the Indo-Pacific (Yu et al. 2016). However, a shallow but significant genetic population structure has been reported for the skunk clownfish *Amphiprion akallopisos* (Huyghe and Kochzius 2017) and the East African giant mud crab *Scylla serrata* (Rumisha et al. 2017), which are both known to have limited dispersal patterns.

Ocean currents are the important drivers for genetic structuring in the Western Indian Ocean (WIO) region (Jahnke et al. 2019; Obura et al. 2019). Further, evidence of genetic divergence in the WIO has been reported in stony corals (*Acropora tenuis*) (van der Ven et al. 2016). However, Tanzania's relatively narrow coastal area can explain this study's observed lack of genetic divergence. The strong EACC distributes larvae along the Tanzanian coastline connecting southern and northern populations in a south-to-north direction. During the planktonic stage of belonids, the relatively large eggs (4–4.6 mm in diameter) with threads like filaments and larvae are transported by ocean currents and dispersed along the coast, facilitating gene flow from the southern to the northern populations (Breder and Rosen 1966; Dorenbosch et al. 2006). Based on patterns of larval dispersal, a gradient in genetic population structure was expected along the coastline. However, the strong swimming ability of this species can also enable individual migration between populations at extensive

geographic distances, which could also explain the observed high genetic connectivity.

Although there was no clear pattern of isolation by distance and lack of distinguished population structures, we observed a significant difference in genetic population structure between the northernmost site (Tanga) and an island in the south (Songosongo). The reason for this separation could not be clarified but may result from currents and local eddies in the region (Baltazar-Soares et al. 2014).

Genetic diversity and historical demography

For marine animals, both haplotype (h) and nucleotide (π) diversities of less than 0.5 indicate low genetic diversity, although species-specific variation occurs (Grant and Bowen 1998). The hound needlefish *T. crocodilus* showed low haplotype diversity at all sites except at Tanga. On the other hand, the nucleotide diversity was low at all sites (Table IV). This may be because of the relatively high fecundity of *T. crocodilus* and the aggregate spawning and schooling behaviour. The values of haplotype diversity were within the range of the estimates from the WIO for the skunk clownfish (*A. akallopisos*) but rather small compared to the values reported for the same species in the Eastern Indian Ocean (EIO) region (Huyghe and Kochzius 2017). The records are comparable to the estimates documented for the giant mud crab *S. serrata* from East Africa (Rumisha et al. 2017) and the scalloped hammerhead shark *Sphyrna lewini* in the WIO (Hadi et al. 2020). The overall haplotype and nucleotide diversities (Table IV) were lower than the average diversity values reported for mangrove-associated crabs, e.g. the East African *Perisesarma guttatum*, the Indo-Pacific *Uca hesperiae*, and *Neosarmatium africanum* in the WIO (Silva et al. 2010; Fratini et al. 2016), and based on the mitochondrial DNA control region of the narrow-barred Spanish mackerel

Scomberomorus commerson sampled from the north of Tanzania (Johnson et al. 2021). The low overall values of haplotype and nucleotide diversity are typical for populations that are expanding after a historical bottleneck (Alves et al. 2001). This claim was well supported by the Tajima's D test, Fu's FS test, mismatch distribution analysis, and Rogers' test (Table IV) (Rogers and Harpending 1992; Rogers 1995). Furthermore, the estimates of genetic diversity were lower than the overall values reported in the EIO for the skunk clownfish *A. akallopisos*, particularly in the Palau Seribu archipelago and Karimunjawa National Park in Indonesia (Huyghe and Kochzius 2017). This is consistent with the same study's conclusion that the overall genetic diversities were far higher in the EIO than in the WIO, supporting the theory of fish distribution from the diversity-rich EIO to the WIO region (Huyghe and Kochzius 2017).

Concluding remarks

This study reported high genetic connectivity (based on overall low nucleotide and haplotype diversities) and recent demographic expansion of the hound needfish *T. crocodilus* in Tanzanian waters. The mitochondrial DNA (mtDNA), however, faces limitations such as assessing genetic variations at only a single locus, maternally inherited, and subject to selection pressure (Bazin et al. 2006). Therefore, to confirm the present findings, future work should apply multiple loci and high-resolution markers like microsatellites (Zink and Barrowclough 2008). Nevertheless, the strong connectivity found in this study using a single marker is coherent with other regional studies using high-resolution markers such as microsatellites (Jahnke et al. 2019). The empirical evidence of panmixia suggests that the fishery likely exploits a single population, which makes it less resilient to stressors such as overfishing and climate change. Future fisheries management should therefore consider genetic diversity (and genetic drift) as a priority in fish stock monitoring programmes to ensure healthy fisheries, sustainable fish communities and food security for coastal people.

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Author contributions

Conceived and designed the study: SM, MOS, PP, MS, MW and MG. Collected the samples: SM, MOS, MS, PP and MG. Performed the laboratory work: SM, MOS, CR and EV. Analysed the data: SM, CM, CR and EV. Developed the map: SM and MOS. Wrote the paper: All authors.

Disclosure statement

No potential conflict of interest was reported by the author(s).



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Data availability statement

The sequence data have been submitted to the GenBank database and given the accession numbers ON644875 – ON644992.

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Figure A1. Evolutionary relationships of sampled sequences of the hound needlefish (*Tylosurus crocodilus*) were inferred using the Maximum Likelihood method and Kimura 2-parameter model. All samples clustered phylogenetically with the reference *T. crocodilus* sequences (1000 bootstraps) at 99% similarity.