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## Genomic analysis reveals multiple mismatches between biological and management units in yellowfin tuna (*Thunnus albacares*)

Rachel B. Mullins<sup>1,†</sup>, Niall J. McKeown<sup>2,\*†</sup>, Warwick H. H. Sauer<sup>1</sup>, and Paul W. Shaw<sup>1,2</sup>

<sup>1</sup>Department of Ichthyology & Fisheries Science, Rhodes University, Grahamstown, South Africa

<sup>2</sup>Institute of Biological, Environmental and Rural Sciences, Aberystwyth University, Aberystwyth SY23 3DA, UK

\*Corresponding author: tel: 0044 1970622323; e-mail: [njm2@aber.ac.uk](mailto:njm2@aber.ac.uk).

†Equal contribution.

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The South African (SAF) yellowfin tuna (*Thunnus albacares*) fishery represents a potential example of misalignment between management units and biological processes. The SAF fishery spans an operational stock with a boundary at 20°E, either side of which fish are considered part of Atlantic or Indian Ocean regional stocks. However, the actual recruitment of fish from Atlantic and Indian Ocean spawning populations into SAF waters is unknown. To address this knowledge gap, genomic analysis (11 101 SNPs) was performed on samples from Atlantic and Indian Ocean spawning sites, including SAF sites spanning the current stock boundary. Outlier loci conferred high discriminatory power to assignment tests and revealed that all SAF fish were assigned to the Indian Ocean population and that no Atlantic Ocean fish appeared in the SAF samples. Additionally, several Indian Ocean migrants were detected at the Atlantic spawning site demonstrating asymmetric dispersal and the occurrence of a mixed-stock fishery in Atlantic waters. This study highlights both the spatial inaccuracy of current stock designations and a misunderstanding of interactions between the underlying biological units, which must be addressed in light of local and global declines of the species. Specifically, the entire SAF fishery must be managed as part of the Indian Ocean stock.

**Keywords:** adaptation, assignment, conservation, dispersal, genetic differentiation, stock, sustainability

### Introduction

The worldwide depletion of fish communities (Myers and Worm, 2005) with evidence of fishery induced economic (Botsford *et al.*, 1997) and biological extinctions (Jackson *et al.*, 2001) highlights the importance of identifying biologically distinct units within marine fishes for both sustainable management and conservation of marine biodiversity (Ruzzante, 2006). The ability to monitor the dynamics of such components within systems involving seasonal migration and potential spatial overlap is beneficial, as more easily exploited and/or less productive units may be susceptible to overharvesting, contributing to loss of diversity and adaptive potential (Iles and Sinclair, 1982) and producing negative

effects on recruitment potential and population/fishery viability (Ryman *et al.*, 1995). Recent advances in population genomic methods offer considerable potential as tools to meet the many challenges associated with sustainable fishery management (Allendorf *et al.*, 2010). However, this potential is yet to be fully harnessed for a variety of reasons. The integration of genomics data and fishery management is particularly hampered in developing countries where threats to fishery sustainability may be most concentrated (Waples *et al.*, 2008; Willette *et al.*, 2014; Bernatchez *et al.*, 2017).

Yellowfin tuna, *Thunnus albacares* is globally distributed in tropical waters and supports fishery stocks that extend across

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boundaries of national exclusive economic zones (EEZs) and into the high seas. The species accounts for the second largest worldwide catch of tuna and tuna-like species (after skipjack tuna *Katsuwonus pelamis*) in terms of catch weight and volume (Miyake *et al.*, 2010; Juan-Jordá *et al.*, 2013; FAO, 2017), with an average annual catch of ~1.25 million metric tons over the past decade (Pecoraro *et al.*, 2017). At present, four regional tuna management units are described, Atlantic, Indian, Eastern Pacific, and West Central Pacific, and each is managed as a single stock by the respective tuna Regional Fisheries Management Organisations (tRFMOs) (Pecoraro *et al.*, 2017). Recent genomic studies have confirmed genetic differences and restricted interbreeding among regional groups (Pecoraro *et al.*, 2016; Barth *et al.*, 2017). As previous studies have been macrogeographic in scale, there is still considerable uncertainty over the fine-scale boundaries and interactions among genetic groups and their congruence with current operational stock boundaries. Mismatch between biological and management units is recognized as a major threat to global fishery sustainability (Reiss *et al.*, 2009), and clarifying any such inconsistencies for yellowfin tuna is an urgent consideration as the management units are currently described as fully exploited and may even be overexploited (Majkowski, 2007), with the species being described as “near threatened” by the IUCN (Collette *et al.*, 2011).

The South African (SAF) yellowfin tuna fishery provides a striking case where management units may be incongruent with biological population processes of dispersal, interbreeding and adaptation. The present management boundary between the Atlantic and Indian Ocean operational stocks lies off South Africa at 20°E (FAO, 2017), and the two stocks are assessed and managed by the International Commission for the Conservation of Atlantic Tunas (ICCAT) and the Indian Ocean Tuna Commission (IOTC), respectively (Figure 1). Yellowfin tuna catches west of the 20°E are reported to ICCAT for inclusion in the Atlantic Ocean stock assessment, whereas catches east of this line are reported to the IOTC. However, this boundary is not based on any recognized biogeographic boundary or species-specific biological information but is based on a geographic

feature, the southernmost tip of Africa at Cape Agulhas at the confluence of the Atlantic and Indian Oceans. Yellowfin tuna do not spawn off the southern coast of South Africa, as the local environment does not provide conditions for optimal survival of early life stages (Pecoraro *et al.*, 2016), and thus adults occurring there represent allochthonous feeding migrants. Barth *et al.* (2017) suggested that the Benguela current system (BCS) along southwest African coast may be a barrier to dispersal of individuals from Atlantic spawning areas north of the BCS into SAF waters. It could therefore be hypothesized that the SAF fishery is being sustained solely by individuals from Indian Ocean spawning population(s). If so, the accuracy of current stock monitoring methods is fundamentally compromised. These inaccuracies need to be urgently resolved, because the Standing Committee on Research and Statistics of ICCAT has reported the Atlantic stock as over-fished (ICCAT, 2017), and the Scientific Committee of the IOTC has declared the Indian Ocean stock as overfished and has proposed an interim rebuilding plan (IOTC, 2016).

The present research aimed to build on and extend previous studies of yellowfin tuna by using RAD-Seq genotyping to assess the membership of yellowfin tuna adults sampled in SAF waters, from sites spanning the operational stock boundary, to Atlantic and Indian Ocean genetic stocks. A specific hypothesis being tested was that, in line with current management unit delineation, Western Cape and Southern/Eastern Cape yellowfin are derived from distinct Atlantic and Indian Ocean spawning populations, respectively. This study reports robust genetic patterns that both corroborate previous studies and reveal new aspects of yellowfin biocomplexity that collectively highlight both the spatial inaccuracy of current stock designations and a misunderstanding of dispersal patterns and population boundaries. These discrepancies prevent the accurate assessment of population productivity and dynamics, and so undermine the effectiveness of management actions that may compromise resilience and sustainability of the resource.

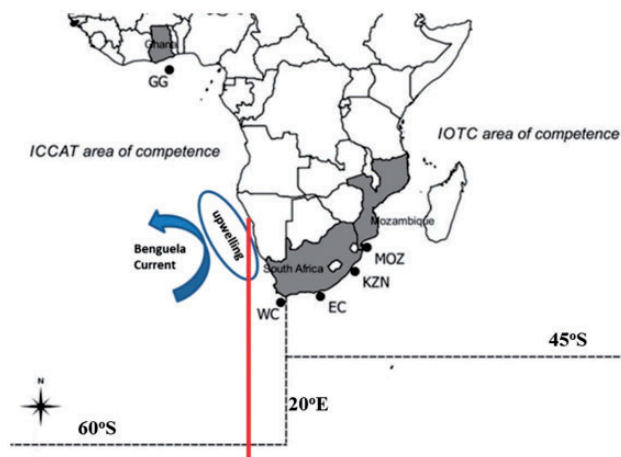
## Material and methods

### Sample collection, DNA isolation, and species verification

Our sampling strategy (Table 1 and Figure 1) was devised to include fish off the Western (Atlantic Ocean waters) and Eastern (Indian Ocean waters) Cape Provinces, as well as fish from putative spawning areas of regional stocks in the Atlantic Ocean [Gulf of Guinea (CG), ICCAT, 2016 and Indian Ocean (KwaZulu-Natal), IOTC, 2014]. Fin clips from individual yellowfin tuna were fixed in 95% ethanol. DNA was extracted from fin clips using a phenol/chloroform/isoamyl alcohol (PCIA) method following Winnepeninckx *et al.* (1993). A segment of the mitochondrial Control Region was amplified with the polymerase chain reaction (PCR) and sequenced in both directions using the species-specific primers (5'-TCCTACCCCTAACTCCCAAAG-3'; and reverse primer: 5'-AAACTGTGGGGATTCTCAC-3'). Sequences were used to confirm species identity using the BLAST function in GenBank. MtDNA summary statistics and estimates of inter-sample differentiation were calculated following McKeown *et al.* (2015).

### SNP genotyping

SNP genotyping by sequencing was performed using tunable Genotyping By Sequencing (tGBS), a modified version of RAD-



**Figure 1.** Geographical location of sample sites (codes correspond to Table 1) with approximate location of the Benguela Current System (BCS), and 20°E operational stock boundary (dashed line). Solid line denotes proposed location of new management boundary at ~13.35°E based on genetic results from this study.

**Table 1.** Sample information and summary indices of variation for mtDNA ( $N$  = number of individuals sequenced; NHap = number of distinct haplotypes resolved;  $h$  = haplotype diversity;  $\pi$  = nucleotide diversity) and nuclear SNP data including numbers of individuals genotyped ( $N$ ) and polymorphic SNPs (NpolySNP), observed and expected heterozygosities ( $H_O$  and  $H_E$ , respectively) and their standard deviations, and  $F_{IS}$  ( $F_{IS}$  values all non-significant).

| Sample (code)                     | Sample coordinates and date   | mtDNA |       |      |       | Nuclear SNPs |           |               |               |          |
|-----------------------------------|-------------------------------|-------|-------|------|-------|--------------|-----------|---------------|---------------|----------|
|                                   |                               | N     | N Hap | $h$  | $\pi$ | N            | Npoly SNP | $H_O$         | $H_E$         | $F_{IS}$ |
| Gulf of Guinea (GG)               | 2.5N/0.30W<br>August 2015     | 20    | 20    | 1    | 0.025 | 20           | 10 710    | 0.228 (0.160) | 0.256 (0.148) | 0.13     |
| Western Cape, South Africa (WC)   | 35.61S/18.70E<br>October 2014 | 16    | 16    | 1    | 0.026 | 13           | 9 831     | 0.241 (0.163) | 0.276 (0.145) | 0.11     |
| Eastern Cape, South Africa (EC)   | 34.72S/25.18E<br>June 2015    | 17    | 17    | 1    | 0.024 | 16           | 10 413    | 0.235 (0.164) | 0.264 (0.148) | 0.11     |
| KwaZulu-Natal, South Africa (KZN) | 30.83S/30.48E<br>July 2015    | 20    | 20    | 1    | 0.022 | 20           | 10 766    | 0.224 (0.225) | 0.254 (0.149) | 0.11     |
| Mozambique (MOZ)                  | 36.53S/34.50E<br>August 2015  | 20    | 19    | 0.99 | 0.026 | 20           | 10 755    | 0.227 (0.160) | 0.255 (0.148) | 0.11     |

Seq that incorporates digestion with two enzymes for genome reduction and results in an increased number of reads per site (Ott *et al.*, 2017). The tGBS libraries were sequenced on a Life Technologies' Ion Proton System. Sequenced reads were analysed using a custom Perl script (available at <https://github.com/orgs/schnablelab>; accessed 20 July 2018), which assigned each read to a sample and removed barcode sequences. Seqclean (sourceforge.net/projects/seqclean) was then used to remove adaptor sequences and chimaeric reads harbouring internal restriction enzyme sites. Retained reads were subjected to quality trimming in two phases using the software Lucy2 (Li and Chou, 2004) in which bases with PHRED scores <15 (of 40) were removed. In the first phase, sequences were scanned at each end; whereas in the second phase, sequences were scanned using overlapping 10 bp windows. Quality trimmed sequence reads were aligned to the reference genome of Pacific bluefin tuna, *Thunnus orientalis* (Nakamura *et al.*, 2013), using GSNAP (Wu and Nacu, 2010). Sequence alignments were then scanned for polymorphisms. A SNP was called homozygous if at least 15 reads supported the genotype at the site and at least 90% of all reads covering that site shared the same nucleotide. A SNP was considered heterozygous if each of the two nucleotide variants were reported at least 10 times, and each allele was represented in >30% of the total reads.

### SNP summary statistics and outlier analysis

Allele frequencies and observed ( $H_O$ ) and expected ( $H_E$ ) heterozygosities were estimated using ARLEQUIN 3.4.2.2 (Excoffier and Lischer, 2010). ARLEQUIN was also used to test for departures from expectations of Hardy–Weinberg Equilibrium (HWE). There are numerous challenges to detect SNP “outlier” loci that are putatively under selection (Narum and Hess, 2011). Even though the concept of balancing selection is well established, there remain methodological limitations for its identification in hitchhiking mapping (Hansen *et al.*, 2010). Therefore, we restricted our analysis to the detection of signals of directional selection using two conceptually different approaches. First, we used the FDIST2 outlier detection method (Beaumont and Nichols, 1996) implemented in LOSITAN (Antao *et al.*, 2008). We used the “force mean  $F_{ST}$ ” option to obtain the genome-wide mean (neutral)  $F_{ST}$  value, to reduce the rate of false-positive detection of outliers. We used a null model distribution of the mean

$F_{ST}$  as a function of  $H_E$  under an island model of population subdivision using 50 000 simulations. Positive outlier SNPs were identified as those falling above the 99% confidence intervals of the null distribution. Second, the Bayesian approach implemented in BAYESCAN 2.1 (Foll and Gaggiotti, 2008) was performed using default Markov Chain Monte Carlo (MCMC) parameters. Following the suggestion of Foll and Gaggiotti, (2008) to minimize false positives, we used the “decisive” criterion under Jeffreys’ scale of evidence (Jeffreys, 1961) to identify outliers. Loci were considered to be under directional selection if identified as outliers using both methods, as the detection of outliers through multiple methods increases the confidence that these loci are non-neutral. The results between global and pairwise sample tests were compared to confirm consistency of outlier designations. Functional significance of loci was investigated using BLAST in GenBank following Milano *et al.* (2014).

### Geographical structuring of genetic variation

Genetic differentiation among samples was quantified by global and pairwise  $F_{ST}$  (Weir and Cockerham, 1984) with statistical significances evaluated in ARELQUIN with 10 000 permutations. Genetic structuring was also investigated using the Bayesian clustering method in STRUCTURE 2.3.4 (Pritchard *et al.*, 2000), which identifies the most probable number of genetically distinct groups ( $K$ ) represented by the data and estimates assignment probabilities ( $Q$ ) for each individual (specifically their genomic components) to these groups. The analysis can be run without prior information; however, incorporating prior information using the LOCPRIOR model allows the clustering algorithm to assume that the probability of assignment varies among samples thereby increasing power while not biasing results (Hubisz *et al.*, 2009). The analysis was performed with and without the LOCPRIOR model, in both cases assuming admixture. Simulations were run 10 times for each proposed value of  $K$  (1–5) to assess convergence. Each run had a burn-in of 100 000 MCMC samples followed by 1 000 000 MCMC repetitions. Models were assessed using  $\Delta K$  (Evanno *et al.*, 2005). To complement the STRUCTURE analysis, classical assignment tests were performed in GENECLASS 2.0 (Piry *et al.*, 2004). The “detect migrants” function was used to identify first generation migrants born in a population other than the one in which they were sampled (Paetkau *et al.*, 2004; Piry *et al.*, 2004). The test

statistic  $L_h$  was used following Paetkau *et al.* (2004) and employing Nei's  $D_A$  (Nei *et al.*, 1983), which does not require conformance to HWE. The simulation method in GENECLASS (10 000 simulations) was also used to exclude individuals from a candidate population if that individual's probability of assignment to that population fell below a 0.05 threshold. Genetic structuring was also assessed using the discriminant analysis of principal components (DAPC) implemented in ADEGENET (Jombart *et al.*, 2010). Whereas STRUCTURE assigns cluster membership by minimizing Hardy–Weinberg and linkage disequilibria within clusters, DAPC employs fewer assumptions and simply maximizes differences between groups while minimizing differences within groups with the optimal number of groups identified using the Bayesian Information Criterion (BIC).

## Results

### MTDNA variation

Alignment of 633 bp sequences across 93 individuals revealed 137 variable sites that defined 91 haplotypes, of which 89 were found in a single individual. BLAST confirmed that all sequences were yellowfin tuna. Haplotype diversity was high ( $h=0.999$ ) across samples (Table 1). Global and pairwise tests of differentiation among samples yielded non-significant values (global  $\Phi_{ST}=0.001$ ;  $p=0.3$ ).

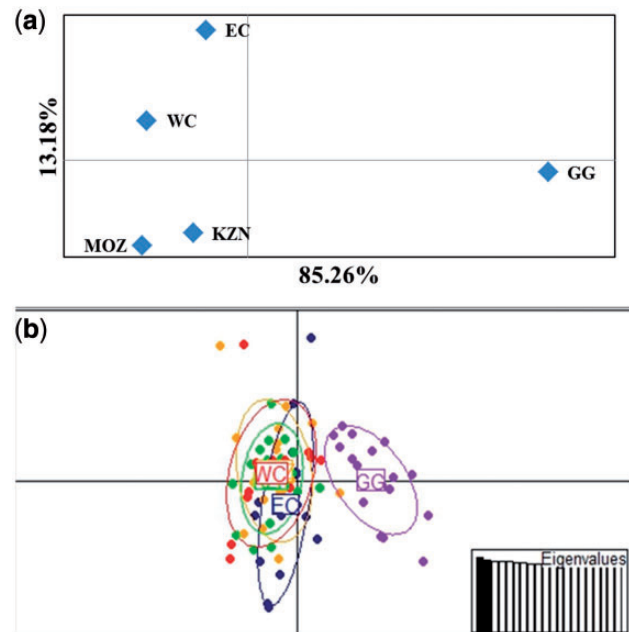
### Nuclear variation

A total of 179 521 855 raw sequenced reads across 96 yellowfin tuna were obtained with between 56 871 and 10 442 486 raw reads (average = 1 870 018) per individual. Seven individuals from the WC sample with >60% of identified SNPs missing were removed from the dataset, resulting in a dataset of ~30 000 SNP loci for 89 individuals. The number of genotypes was further reduced by removing SNP loci that were missing in >10% of individuals; this resulted in a final SNP genotype dataset of 11 101 SNP loci for 89 fish with a mean of 10 495 ( $SD=398$ ) SNPs per individual. Levels of genetic variability were similar across samples, and all samples conformed to HWE (Table 1). Global genetic structuring was weak ( $F_{ST}=0.002$ ) but statistically significant ( $p<0.0001$ ). Pairwise tests revealed this significant structuring largely resulted from the divergent GG sample, which showed significant  $F_{ST}$  values with all other samples (Table 2). All other pairwise  $F_{ST}$  values were not significant, and excluding GG global structuring was not significant ( $F_{ST}=-0.0002$ ).

The LOSITAN outlier analyses consistently reported a greater number of positive outlier loci than the BAYESCAN analysis in the various global and pairwise tests. For example, in the initial global outlier test LOSITAN identified 756 positive outliers compared with 11 by BAYESCAN. In all cases, outliers identified by BAYESCAN were also among the outliers identified by LOSITAN. Therefore, we restricted our discussion to the smaller number of outliers identified using BAYESCAN, as the consensus

outliers were ultimately defined by the BAYESCAN results. The results of outlier tests revealed a clear pattern that aligned with the differentiation of GG from the remaining samples. Firstly, no outliers were identified in global tests excluding the GG sample compared with the detection of 11 in global test including GG. Second, in tests between pairs of samples outliers were only identified in comparisons involving GG. All outliers identified in pairwise tests were among the 11 identified initially in the global test and certain outlier were common across different pairwise comparisons. BLAST analysis did not provide insight into potential functional associations of any of the outlier loci.

When outlier loci were excluded, the pattern of significant differentiation of the GG sample from all other (non-differentiated) samples was still evident in  $F_{ST}$  values (Table 2; Figure 2a) and PCA clustering (Figure 2b). Analysis of the 11 outlier loci reported considerably higher levels of differentiation of the GG sample while retaining the pattern of homogeneity among the remaining samples (Table 2; Figure 3a and b). Bayesian clustering analysis of outlier genotypes unanimously supported a model of  $K=2$ , with identical results for the analyses with and without LOCPRIOR. Under  $K=2$ , all non-GG fish strongly assigned to group B with high individual Q values, whereas the majority of GG fish had higher or intermediate (overlapping confidence



**Figure 2.** (a) Principal co-ordinate analysis of  $F_{ST}$  based on neutral loci. (b) Principal component analysis (PCA) clustering analyses of individuals based on neutral loci. For b ellipses summarize variation per site using ADEGENET's default "cellipse" settings.

**Table 2.** Pairwise  $F_{ST}$  values between samples with statistically significant values in bold.

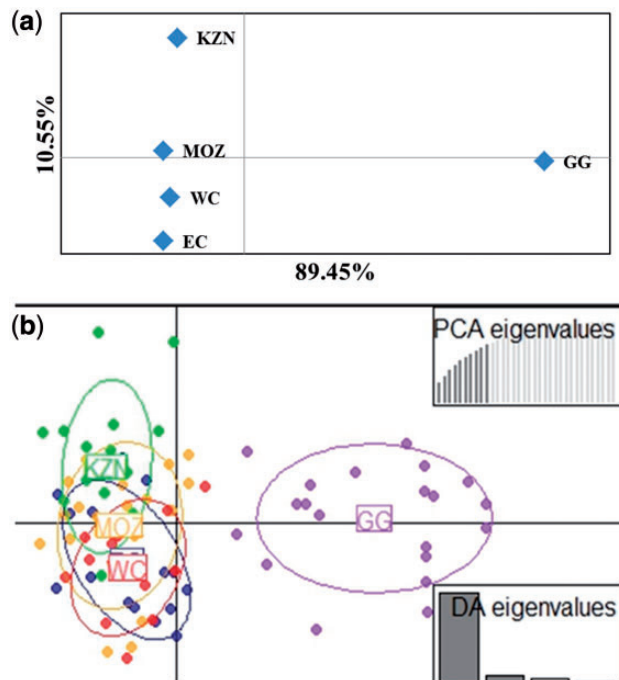
|                | GG                 | WC              | EC              | KZN            | MOZ          |
|----------------|--------------------|-----------------|-----------------|----------------|--------------|
| Gulf of Guinea | –                  | <b>0.330</b>    | <b>0.351</b>    | <b>0.341</b>   | <b>0.344</b> |
| Western Cape   | <b>0.005/0.005</b> | –               | –0.013          | 0.028          | –0.011       |
| Eastern Cape   | <b>0.004/0.003</b> | –0.0016/–0.0017 | –               | 0.056          | –0.005       |
| KwaZulu-Natal  | <b>0.005/0.003</b> | –0.0008/–0.0001 | –0.0004/–0.0006 | –              | 0.005        |
| Mozambique     | <b>0.005/0.004</b> | 0.0003/0.0002   | 0.0009/0.0007   | 0.0001/–0.0001 | –            |

Below diagonal: Estimates across all SNP loci/neutral SNP loci. Above diagonal: estimates based on outlier loci only.

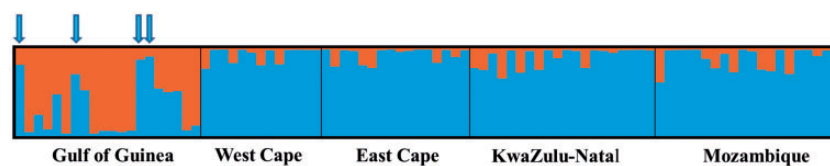
intervals for  $Q$  values) assignment probabilities of belonging to group A (Figure 4). Four individuals in the GG sample had larger  $Q$  membership values for group B. On the basis of the lack of differentiation among the SAF/Indian Ocean samples, the samples were pooled for the classical assignment tests between GG and the remaining samples. Assignment tests identified the same four GG individuals that had higher group B  $Q$  values as being first generation migrants from the pooled SAF/Indian Ocean group. No other migrants were detected with all individuals correctly assigned to the CG or Pooled group. Exclusion analysis using the strict 0.05 threshold further highlighted the genetic differentiation between both groups; first, of 16 non-migrants in the GG sample, 13 could be excluded from the Atlantic group, second, 42 of 69 non-migrants in the Atlantic/Indian ocean group could be excluded from GG.

## Discussion

Mismatches between fishery management units and biological population processes of dispersal, interbreeding and adaptation are recognized as fundamental threats to fishery sustainability



**Figure 3.** (a) Principal co-ordinate analysis of  $F_{ST}$  based on outlier (non-neutral) loci. (b) Discriminant analysis of principal components (DAPC) describing variation among groups of individuals from different samples estimated from outlier loci. For b ellipses summarize variation per site using ADEGENET's default "cellipse" settings.



**Figure 4.** Bar graph illustrating average patterns of Bayesian clustering of individual under the optimum model of  $K = 2$ . Arrows above bar graph denote  $Q$  values from those individuals identified as migrants in the GENECLASS tests.

and to long-term resilience to climate change (Reiss *et al.*, 2009). In light of global declines in yellowfin tuna numbers and continued intensive harvesting, a primary objective of our study was to assess the biological validity of the current Atlantic Ocean and Indian Ocean operational stocks through genomic analysis of fish spanning the current stock boundary off the south coast of South Africa. The salient findings were (i) a lack of population structuring among the SAF samples, and (ii) genetic differentiation of the GG population from the SAF population. These patterns were evident in  $F_{ST}$  estimates from the entire SNP and neutral SNP data sets. A major hindrance to the integration of genetics into fishery management has been the inability of traditional  $F_{ST}$ -based measures to infer short-term demographics because these measures reflect connectivity over multiple generations (Whitlock and McCauley, 1999; Hellberg, 2009; Berry *et al.*, 2012). Additionally, divergence is expected to be slow for large populations experiencing little genetic drift. Increasingly, individual-based kinship and assignment tests are being used to directly assess dispersal (Christie *et al.*, 2010). Combining assignment tests with outlier loci can provide high discriminatory power even with low levels of background differentiation among populations (Helyar *et al.*, 2011; Bekkevold *et al.*, 2015). This approach revealed two genetic clusters in which all SAF specimens were assigned to one cluster with the other comprising only GG fish. Collectively the neutral and non-neutral genetic data concurrently resolved two genetic groups.

On the basis of our sampling scheme, it can be assumed that the two genetic groups (GG vs. the rest) correspond respectively to the distinct Atlantic and Indian ocean genetic stocks previously suggested by allozyme variation (Ward *et al.*, 1997) and confirmed by genomic studies (Pecoraro *et al.*, 2016; Barth *et al.*, 2017). The results therefore demonstrate that Western Cape yellowfin tuna, currently assigned to the Atlantic operational stock, are derived from an Indian Ocean genetic stock. Tagging data corroborate the movement of yellowfin tuna individuals from the western equatorial Indian Ocean to the southern Benguela region off western South Africa (Eveson *et al.*, 2015; Murua *et al.*, 2015). The concordant results among the present and previous genetic studies and the temporal sampling range represented point to a temporally stable pattern in which the Atlantic population is not contributing to recruitment in SAF waters. Nevertheless, it cannot be ruled out that there may be seasonal variation in stock mixing or overlap.

The identification of isolating mechanisms is a crucial facet for inferring demographic independence with a central topic in fisheries genetics being the relative roles of environmental forcing and behaviour (Heath *et al.*, 2008; McKeown *et al.*, 2017). The spatial genetic structure reported here and in Barth *et al.* (2017) is compatible with a role for the cool upwelled waters of the BCS as a potential barrier to dispersal of Atlantic yellowfin tuna into SAF waters. Cold water barriers have been reported for several marine

fishes (Gwilliam *et al.*, 2018). Big eye tuna, *T. obesus*, is an interesting comparison because genetically distinct Atlantic and Indian Ocean fish mix in SAF waters. Yellowfin and bigeye tuna have similar geographical distributions but occupy different ecological niches and vertical habitats (Schaefer *et al.*, 2009); bigeye tuna are more tolerant of cooler water temperatures than are yellowfin tuna (Brill and Lutcavage, 2001). Hence, the cold water of the BCS may have less influence on the movement of bigeye tuna from Atlantic Ocean waters north of the BCS into SAF waters, than on yellowfin tuna.

Barth *et al.* (2017) reported evidence of asymmetric Indo-Pacific-to-Atlantic gene flow across the BCS in yellowfin tuna, and Reid *et al.* (2016) described a similar pattern in the cosmopolitan bluefish (*Pomatomus saltatrix*). In the present study, the Bayesian clustering analysis revealed patterns compatible with unidirectional gene flow from South Africa to GG. However, such gene flow is typically regarded as being historical and no studies have yet revealed recurrent dispersal of adults across the BCS in species for which the BCS is a barrier to contemporary gene flow. The multilocus identification of four Indian Ocean yellowfin tuna among GG fish is striking as it shows that Indian Ocean spawned yellowfin tuna are crossing the BCS and are mixing with members of the Atlantic genetic stock. These dispersals may be linked to Agulhas warm-core rings (Schouten *et al.*, 2000) that can facilitate the transfer of larvae/adults across the BCS and/or to active swimming. In the latter case, unidirectional dispersals may be facilitated by distinct genetic/plastic thermal tolerances of Indian Ocean spawned fish. This suggests that a cryptic mixed-stock fishery exists north of the BCS. Temporal studies are needed to assess the consistency with which Indian Ocean migrants cross the BCS.

The maintenance of genetic integrity of the Atlantic Ocean stock in the face of potentially high rates of dispersal of Indian Ocean genotypes implicates additional mechanisms that may be isolating both stocks. The capacity for active dispersal of adults highlights the potential importance of natal homing (Gaggiotti *et al.*, 2009; McKeown *et al.*, 2017). Indian Ocean fish may migrate into Atlantic spawning areas but return to the Indian Ocean to spawn. If this is occurring samples collected during spawning times should contain fewer migrants than at other times. In this study the GG sample was collected outside of the spawning period and so comparative analysis of a spawning sample from this area could be highly informative regarding the role of homing. The two genetic stocks may also be isolated by adaptation (Nosil *et al.*, 2009) with migrant offspring being selected against (Hendry, 2004). The increased level of genetic divergence between groups at outlier loci might be explained by directional selection, with the resolved groups corresponding to differentially adapted units (Milano *et al.*, 2014). Alternatively, outlier loci may reflect endogenous forces stemming from pre- and/or post-zygotic incompatibilities between populations with different genetic backgrounds to produce intrinsic incompatibilities, rather than reflecting adaptive environmental selection through genotype-environment associations (Bierne *et al.*, 2011). However, adaptation and intrinsic incompatibilities are not mutually exclusive. Along with natal homing, the identification of slowly changing isolating factors is important, as these factors may lead to resistance to hybridization, colonization of new habitats, and by extension, recovery of stocks (e.g. Svedäng *et al.*, 2007).

## Applications and management implications

The use of management units that include only a portion of a larger population can present problems with understanding population stock dynamics and environmental linkages (Frisk *et al.*, 2008). Conversely, management units containing multiple biological populations can lead to inaccurate descriptions of population-specific abundances and productivity, and may conceal the declines of vulnerable populations (Kell *et al.*, 2009; Ying *et al.*, 2011). Our study reveals that the management of the yellowfin tuna fishery around southern Africa presents both types of problems. At a local level, it is recommended that the South Africa Department of Agriculture, Forestry and Fisheries report all future yellowfin tuna catches in SAF waters to the IOTC for inclusion in Indian Ocean stock assessments. Our results indicate that the SAF catch is derived entirely from Indian Ocean spawned fish. It may be more accurate to move the boundary between the Atlantic and Indian Ocean management stocks to the western extent of South Africa's EEZ at 13.35°E (Figure 1). At a regional level, it is essential to understand the proportions of individuals derived from the Atlantic and Indian Ocean genetic stocks over space and time in the mixed-stock fishery north of the BCS. The proportion of "consensus" outliers identified here (0.9%) was generally smaller than the proportions of outlier loci reported by other studies (5.2%, Bradbury *et al.*, 2013; 4.5%, Milano *et al.*, 2014; 3.65%, Hess *et al.*, 2013; 0.99%, Guo *et al.*, 2015). These outlier loci represent a resource that can be used to identify migrants and to characterize mixed-stock dynamics. At present the indiscriminate harvesting of both stocks in the region means that the Atlantic spawning stock biomass is overestimated.

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## Data accessibility

Data will be deposited on Dryad.

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