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Comparative population genetics and evolutionary history of two commonly misidentified billfishes of management and conservation concern

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Abstract

Background: Misidentifications between exploited species may lead to inaccuracies in population assessments, with potentially irreversible conservation ramifications if overexploitation of either species is occurring. A notable showcase is provided by the realization that the roundscale spearfish (*Tetrapturus georgii*), a recently validated species, has been historically misidentified as the morphologically very similar and severely overfished white marlin (*Kajikia albida*) (IUCN listing: Vulnerable). In effect, no information exists on the population status and evolutionary history of the enigmatic roundscale spearfish, a large, highly vagile and broadly distributed pelagic species. We provide the first population genetic evaluation of the roundscale spearfish, utilizing nuclear microsatellite and mitochondrial DNA sequence markers. Furthermore, we re-evaluated existing white marlin mitochondrial genetic data and present our findings in a comparative context to the roundscale spearfish.

Results: Microsatellite and mitochondrial (control region) DNA markers provided mixed evidence for roundscale spearfish population differentiation between the western north and south Atlantic regions, depending on marker-statistical analysis combination used. Mitochondrial DNA analyses provided strong signals of historical population growth for both white marlin and roundscale spearfish, but higher genetic diversity and effective female population size (1.5-1.9X) for white marlin.

Conclusions: The equivocal indications of roundscale spearfish population structure, combined with a smaller effective female population size compared to the white marlin, already a species of concern, suggests that a species-specific and precautionary management strategy recognizing two management units is prudent for this newly validated billfish.

Keywords: Roundscale spearfish, White marlin, Genetic population structure, Genetic diversity, Effective population size, *Tetrapturus georgii*, *Kajikia albida*

Background

Identifying genetic conservation units of large-bodied, marine pelagic fishes remains challenging as a result of their often large population sizes, typically strong dispersal ability (via adult and/or larval phases) and few apparent physical barriers to gene flow. These parameters are generally associated with shallow levels of genetic differentiation across large geographic regions [1-3]. More

recently, however, low but statistically significant levels of genetic differentiation have been detected among populations of pelagic fishes, introducing exceptions to the traditional paradigm of little if any genetic structure across local and even broad spatial scales for such taxa [4,5]. Although the biological interpretation of such shallow genetic differentiation is sometimes unclear [2,6], defining genetic population boundaries remains essential for conservation of genetic legacies and adaptive potential. This issue is of particular interest in the case of apex predatory fishes given their likely important ecosystem role, and the fact that many are also exploited in highly

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valuable commercial and recreational fisheries. Istiophorid (Istiophoridae) billfish species fall in this category, and in several cases are known to have declined to levels low enough to cause international concern about their population status [7-10].

The recent validation of the roundscale spearfish (*Tetrapturus georgii*) and its routine misidentification as the morphologically very similar and overfished, sympatric white marlin, *Kajikia albida* (Figure 1), have raised new management and conservation challenges concerning the population status of both species [11-14]. While genetic analyses can readily differentiate these two species [11,15,16], only subtle morphological differences distinguish them, requiring either very close visual examination or the taking of morphometric measurements. These differences are: visual - shape of lateral torso scales; morphometric - ratio of the distance from the anus to the origin of the first anal fin to the maximum height of this fin, and branchiostegal to opercle length relationship [17]. Further exacerbating these challenges is that the roundscale spearfish and white marlin possess largely sympatric Atlantic-wide temperate and tropical distributions [13], and that misidentifications may also be occurring between these species and the longbill spearfish (*Tetrapturus pfluegeri*), a third morphologically similar species [7].

Two intertwined issues have complicated assessing the population status and planning of management strategies for these billfishes: (i) the widespread species misidentifications in the context of severe white marlin declines, and (ii) the current lack of almost any data for the roundscale spearfish. First, white marlin have undergone severe population declines over the past four decades, mostly as a result of offshore longline fisheries in the Atlantic [10]. This species is currently listed as Vulnerable on the IUCN Red List of Threatened Species [8]. In addition, it is now recognized that decades of unrealized species misidentifications have occurred between the white marlin, and the longbill and roundscale spearfish, and that management strategies for the nominal white marlin have unknowingly been based on catch information and stock assessments for a species-complex [7,18]. Potential

impacts of these species misidentifications may be severe [10]. Analysis of commercial catch data has suggested that the roundscale spearfish may comprise a significant proportion (27%) of the overall white marlin catch in the western North Atlantic and that both species may show contemporary evidence of over-exploitation and decline [18]. Further hindering management efforts, is the second relevant issue that almost nothing is known about the population dynamics of the roundscale spearfish, including its population genetic structure and demographic history.

Here, we provide the first population genetic assessment of the roundscale spearfish to inform management and conservation efforts for this enigmatic, recently recognized species. As part of this assessment, we utilize nuclear microsatellite and mitochondrial sequence markers to explore the population structure of this species in its western Atlantic range. Furthermore, we utilized existing white marlin mitochondrial DNA sequences to compare the genetic diversity and evolutionary history of the white marlin and roundscale spearfish, as the bulk of misidentifications are believed to occur between these two species [11,18]. As historical stock assessments of the white marlin were based on landings that were unknowingly comprised of a species-complex, this comparison allowed for a unique species-specific survey of the demographic history of two morphologically very similar, but evolutionarily distinct species.

Methods

Ethics statement

The roundscale spearfish tissue samples used in this study were obtained from fish harvested independently by commercial fisheries. This is not a CITES listed species and no permits or licenses were required to work with these samples. All laboratory work on these samples was performed in accordance with Nova Southeastern University guidelines.

Samples and collection sites

A total of 198 roundscale spearfish samples were obtained from animals incidentally caught in long-line fisheries



Figure 1 White marlin (top) and roundscale spearfish (bottom) showing strong morphological similarity. (Image credit: J. Foster/GHRI)

targeting other teleost species, including swordfish and tuna. The majority of individuals possessed a lower jaw fork length ranging from 126 to 186 cm (where data was available). To date, no information is available on the relationship between length and maturity for roundscale spearfish; however, assuming size at maturity of the roundscale spearfish and white marlin are comparable [19,20], our roundscale spearfish comprised mainly adult fish. Roundscale spearfish capture locations are shown in Figure 2, and comprise locations within the western Atlantic, both north and south of the equator [i.e., the western North Atlantic (WNA; n = 140) and western South Atlantic (WSA; n = 58)]. Samples were divided into *a priori* western North and South populations for population-level analyses for two reasons: (i) presence of the Amazon plume at equatorial latitudes, a known biogeographic barrier for numerous reef fishes [21,22], and (ii) previous genetic evidence of at minimum weak genetic differentiation between North and South collections of other billfishes as well as the presence of disjunct hemispheric spawning locations for some billfishes [23,24].

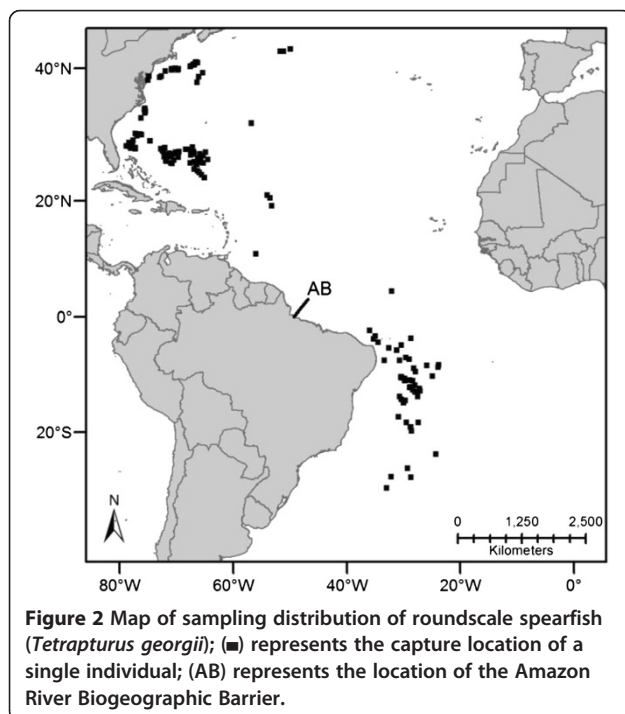
Tissue samples were stored in 95% ethanol until genomic DNA extraction. Prior to inclusion in this study, identities of all roundscale spearfish were verified using a multiplex species-specific primer test targeting the mitochondrial protein coding gene NADH dehydrogenase 4 (MS Shivji unpublished observation). Identities of a subset of samples (n = 43) were also confirmed by mitochondrial Cytochrome c oxidase I barcodes [16]. To investigate the genetic population characteristics of the roundscale spearfish, we

genotyped 198 individuals at 13 microsatellite loci, and sequenced ~580 base pairs (bp) of the mitochondrial control region (mtCR) locus of a subset of individuals (total n = 83: WNA n = 42; WSA n = 41). To assess the comparative genetic diversity and demographic history of the white marlin relative to roundscale spearfish, white marlin mtCR sequences (834 bp) from 99 individuals (91 haplotypes) were obtained from GenBank (Accession numbers DQ835191-DQ835281) comprising individuals sampled from the WSA, WNA, Caribbean Sea and the Eastern Atlantic [23]. As we only obtained partial mtCR locus sequences from roundscale spearfish, all white marlin sequences were cropped (to 601 605 bp) to ensure that the same region was analyzed for both species. Variation in length of mtCR sequences between species resulted from indels.

Mitochondrial DNA sequencing and microsatellite genotyping

Genomic DNA was extracted from ~25 mg of roundscale spearfish tissue using the DNeasy Kit (QIAGEN Inc., Valencia, CA) following manufacturer's instructions. To amplify and sequence the ~580 bp section of the mtCR, we used the primer pair Pro-5M13F (5'-CAC GAC GTT GTA AAA CGA CCT ACC YCY AAC TCC CAA AGC-3') and dLoopi (5'-CCA TCT TAA CAT CTT CAG TG-3') [15]. Total polymerase chain reaction (PCR) volumes were 50 µL and contained 1 µL of unquantified extracted genomic DNA. Final concentration of the remaining PCR reactants were 1 x PCR buffer (0.15 mM MgCl₂), 0.2 mM of each dNTP, 0.25 µM of each of the Forward and Reverse primers and 1.0 U of HotStar Taq DNA Polymerase (QIAGEN Inc.). PCR was performed in a Mastercycler Gradient (Eppendorf Inc., Westbury, NY) thermal cycler as follows: an initial denaturation at 95°C for 15 minutes (min), followed by 35 cycles of 94°C for a 1 min, 50°C for 1 min, 72°C for 1 min, and a 20 min final extension step at 72°C. A negative control (no genomic DNA) was included in each PCR set to check for reagent contamination. PCR products were purified using the QIAquick PCR Purification Kit (QIAGEN Inc.) and double-strand sequenced using standard protocols on an AB 3130 genetic analyzer (Applied Biosystems Inc., Foster City, CA). The mtCR sequences were aligned using MUSCLE as implemented in the program Geneious version 6.0.6 (Biomatters Inc., San Francisco, CA), and the alignment was subsequently refined and manually checked by hand.

The 13 microsatellite loci used for genotyping were those developed for roundscale spearfish by Bernard et al. [25] (*tge23*, *tge54*, *tge76*, *tge79*, *tge105*, *tge119*, *tge135*, *tge139*, *tge144*, and *tge151*) and blue marlin (*Makaira nigricans*) by Buonaccorsi and Graves [26] (Mn01, Mn10, and Mn60). The roundscale spearfish species-specific microsatellite loci were amplified as per Bernard et al.



[25]. The three blue marlin microsatellite loci, were cross-amplified in roundscale spearfish using a total PCR reaction volume of 25 μ L, containing 1 μ L (unquantified) genomic DNA. Final concentration of the remaining PCR reactants were 1 x PCR buffer (0.15 mM $MgCl_2$), 0.2 mM of each dNTP, 0.33 mM $MgCl_2$, 0.16 μ M of the Forward microsatellite primer which possessed a 5'-M13 tail [27], 0.4 μ M of the Reverse microsatellite primer, 0.4 μ M of the fluorescently labeled universal M13 primer (5'-TGTTAAACGACGGCCAGT-3') [27], and 0.5 U of HotStar *Taq* DNA Polymerase (QIAGEN Inc.). PCR was performed in a Mastercycler Gradient (Eppendorf Inc.) thermal cycler as follows: 95°C initial heating for 15 min, followed by 35 cycles of 94°C for 1 min, 1 min at the primer annealing temperature [$T_A = 60^\circ C$ (Mn01, Mn10, Mn60, *tge105*, *tge119*, *tge135*, *tge139*, and *tge151*), and 58°C (*tge23*, *tge54*, *tge76*, *tge79*, *tge144*)], 72°C for 1 min, and a final 20 min extension step at 72°C. Electrophoresis was performed on an AB 3130 (Applied Biosystems Inc.) genetic analyzer. All fragments were sized using LIZ 600 as the internal allele size standard and scored using the software GENEMAPPER 3.7 (Applied Biosystems Inc.).

Data analysis

Calculations of microsatellite allele frequencies, expected (H_E) and observed (H_O) heterozygosities, and tests for Hardy-Weinberg (HWE) and linkage equilibrium (LE) were performed using GENEPOP on the web (v.4.0.10) [28,29]. To estimate the significance of the above tests, we used an unbiased exact test, employing the Markov chain method (1000 dememorizations, 100 batches, 1000 iterations per batch) [30,31] as implemented in GENEPOP. Significance levels were adjusted using sequential Bonferroni correction [32] to accommodate multiple comparison testing. Microsatellite allelic richness (R_S) [33] for each collection site (*a priori* defined as samples from the WNA or WSA) was estimated using FSTAT 2.9.3.2 [34]. The frequency of null alleles was estimated using the program FreeNA [35].

Population genetic structure of roundscale spearfish: population-level analyses

To test for western Atlantic population subdivision with both mitochondrial and nuclear markers, we estimated divergence between WNA versus WSA samples. For mtCR sequence data, divergence was estimated using Φ_{ST} [Tamura and Nei (TN) model of evolution; 10 000 permutations] as implemented in Arlequin 3.1 [36], Jost's D statistic [37] as implemented in the program SPADE [38] (10 000 bootstrap iterations), and the nearest neighbor statistic (S_{nn}) [39] as implemented in DnaSP v5 [40] [significance of the S_{nn} test statistic was estimated using 10 000 permutations (sites with alignment gaps excluded)].

For microsatellite data, between population divergence was estimated using Jost's D (arithmetic mean of D_{est})

using DEMETics [41] within the statistical package R v2.15.1 [42] (significance estimated using 1000 bootstrap iterations), and F_{ST} as implemented in FSTAT. Gender information was available for a large sub-set of our roundscale spearfish samples (uncommon in billfish landings data), as such we tested for sex-biased dispersal (FSTAT: 1000 randomizations). Although sex-biased dispersal has not been documented previously in istiophorid billfish, Muths et al. [43] found support for this hypothesis in swordfish (*Xiphias gladius*), raising the potential for sex-biased dispersal in other migratory billfishes. All seven measures were utilized to test for sex-biased dispersal.

Population genetic structure of roundscale spearfish: individual-level analyses

The Bayesian multi-locus clustering program *Structure* v2.31 [44] was utilized to determine the most likely number of genetically discrete populations [$\ln Pr(X|K)$]. Two disparate *Structure* analyses (see below) were performed both consisting of ten replicates for the values $K = 1 - 5$ (MCMC chain length and burn-in consisted of 200 000 and 100 000 iterations, respectively), assuming correlated allele frequencies [45] and admixture. One analysis was performed without *a priori* sampling location information, while the second analysis implemented the model *locprior* [46], which incorporates *a priori* sampling location information (e.g., WNA versus WSA in this case).

Potential genetic spatial discontinuities were also assessed using the program Geneland 3.1.4 [47] as implemented in the statistical package R [42] to complement *Structure*'s individual-based analyses. All runs incorporated the Dirichlet distribution model of independent allele frequencies [47]. Geneland was run 10 times at $K = 1 - 5$ for 500 000 iterations (500 thinning; 50 000 burn-in) with zero uncertainty of geographical coordinates. Given the highly migratory nature of roundscale spearfish, additional Geneland analyses were performed assuming varying levels of coordinate uncertainty (10 km and 100 km). Geographic coordinates used represented the location of the start of the fishing long-line set on which each individual roundscale spearfish was captured.

To evaluate the hypothesis of whether genetic distance among roundscale spearfish individuals was correlated with geographical distance among their collection locations, Mantel tests were performed as implemented in GenAlEx [48]. The significance of correlations was assessed using 999 permutations.

Comparative mitochondrial DNA-based genetic diversity and demographic histories of the roundscale spearfish and white marlin

We used the software jModelTest 2.1.2 [49,50] to identify the most appropriate model of DNA evolution using the Akaike information criterion (AIC) for both mtCR

sequence datasets. Molecular diversity calculations for mtCR sequences (number of unique haplotypes, number of segregating sites, nucleotide composition, and haplotype (h) and nucleotide diversities (π) estimated with Neis corrected average genetic divergence [51]) were estimated in Arlequin.

To test for departures from a constant population size we estimated the summary statistics F_{IS} [52] in Arlequin (10 000 iterations) and R_2 [53] in DNAsp. Significance of the R_2 statistic was determined in DNAsp using 10 000 replicates. We note that 5 of the 91 white marlin mtCR sequences downloaded from NCBI (Accession numbers DQ835236, DQ835248, DQ835251, DQ835275, and DQ835277) contained ambiguous bases. Since DNAsp does not allow for ambiguous bases, these five sequences were not included in the R_2 analysis.

Demographic expansions were also assessed for both species by mismatch analyses [54] using the sudden demographic expansion model implemented in Arlequin. Model fit to our data was statistically tested using the sum of squared deviations (SSD) and the raggedness index (H_{ri}) [55] (1000 bootstrap replicates). Mismatch analyses were performed for each *a priori* roundscale spearfish population as well as for the overall (pooled) roundscale spearfish mitochondrial dataset. Population parameters τ , Θ_0 , and Θ_1 , where Θ_0 and Θ_1 are the expected pairwise differences before and after a change in population size, respectively [55], and τ is a relative measure of time since population expansion in generations, were also estimated. Using the above parameters, actual time since population expansion (t) was estimated as $t = \tau/2\mu$, where μ is the mutation rate per locus per generation. Reported estimates of teleost mitochondrial control region divergence rates have varied considerably across species and studies, but typically range between 3.6–9% per site per million years [56–58]. Given the absence of specific divergence rates for the istiophorid lineages, we adopted the provisional mutation rates of 1.8–4.5% per site per million years based on the typical divergence rates reported (note: mutation rate within a lineage = ? divergence time between lineages) [56,57], to determine time since population expansion from the mismatch distribution for the two billfishes. We estimated the long-term population parameters Θ ($2N_f\mu$, where N_f = the female historical effective population size, and μ = the mutation rate) and exponential growth rate (g) for roundscale spearfish and white marlin using the pooled sample set for each species, as well as separately for each of the WNA and WSA roundscale spearfish sample sets using the Bayesian method implemented in the program LAMARC 2.0 [59]. Assuming that the roundscale spearfish and white marlin share similar mtCR mutation rates, Θ will allow for a direct comparison of each species relative female effective population size and the relative genetic variability found

within species [60]. Analyses were performed using the GTR and F84 models of evolution (roundscale and white marlin samples, respectively) and three simultaneous chains implementing an adaptive heating scheme (1.0, 1.1, 1.3). A single chain comprising 10 to 20 million iterations was employed (10% burn-in). Convergence was assessed using the program Tracer 1.5 [61]; estimates were assumed to have converged once ESS scores exceeded 200 for all parameter estimates. Final parameter estimates and credibility intervals were the most probable estimates (MPE) determined after three replicates.

To assess the historical trajectories in female effective population size for roundscale spearfish and white marlin, we constructed coalescent-based Bayesian skyline plots (BSP) using the program BEAST v1.5.2 [62]. Plots were constructed for each *a priori* population of roundscale spearfish (i.e. WNA and WSA) and for pooled samples from each species. Priors included the implementation of the TVM + I + G and HKY + I + G models of substitution for roundscale spearfish and white marlin datasets, respectively (as defined by jModelTest), and the strict clock model. Priors for the site heterogeneity model [Gamma (G) and Invariant Sites (I)] were obtained from jModelTest. The piecewise-constant skyline model was selected and runs were fixed at 10 groups. The fixed substitution rate was set to correspond to the mutation rates utilized in the previous analyses (1.8–4.5% per site per million years). MCMC tests were run for 50 million generations and sampled every 5000th step (10% burn-in). Convergence was assessed using the program Tracer and estimates were assumed to have converged once ESS scores exceeded 200 for all parameter estimates.

Results

Population genetic structure of roundscale spearfish: population-level analyses

The genotypes of 198 roundscale spearfish were determined at 13 microsatellite loci. Sample sizes, basic genetic diversity statistics and the deviation from HWE for each locus and across all loci for each sampling location are listed in Table 1. Average heterozygosities and allelic richness for both roundscale spearfish populations across all loci ranged from 0.71–0.74 and 14.3–14.6, respectively. All loci met HWE expectations after sequential Bonferroni correction ($\alpha/26$; $\alpha = 0.05$); however, pairwise tests of LE within populations demonstrated significant disequilibrium between three locus pairs (WNA: Mn01 & *tge*105, *tge*23 & *tge*54; WSA: *tge*135 & *tge*139) after sequential Bonferroni correction ($P < 0.05$). We note, however, that where significant departures from LE were detected; they were not widespread, being restricted to only one of the two *a priori* defined populations (i.e., WNA or WSA). Furthermore, no evidence of linkage disequilibrium (LD) was found when all roundscale spearfish samples were

Table 1 Summary statistics of 13 microsatellite loci for roundscale spearfish (*Tetrapturus georgii*)

Sample	Locus													Average across loci
	MN01	MN10	MN60	tge23	tge54	tge76	tge79	tge105	tge119	tge135	tge139	tge144	tge151	
WNA														
<i>n</i>	137	137	127	140	135	139	139	139	134	135	136	134	128	
<i>a</i>	18	27	32	4	14	3	17	19	28	24	10	23	8	17.5
R_s	15.8	21.8	25.2	3.6	12.3	2.7	14.1	14.6	21.0	22.0	8.3	18.2	6.7	14.3
<i>as</i>	267-335	291-439	217-319	223-235	202-232	103-109	129-197	185-227	187-238	177-277	99- 139	110-212	191-205	
H_O	0.90	0.94	0.92	0.44	0.86	0.05	0.78	0.73	0.87	0.89	0.76	0.77	0.63	0.73
H_E	0.90	0.93	0.94	0.50	0.86	0.06	0.78	0.73	0.89	0.94	0.71	0.82	0.63	0.74
<i>HWE</i>	0.02	0.67	0.47	0.25	0.01	0.09	0.80	0.75	0.44	0.05	0.55	0.01	0.11	0.002
WSA														
<i>n</i>	58	55	57	58	58	57	58	58	55	56	57	57	58	
<i>a</i>	16	22	30	4	12	3	13	12	25	21	7	19	7	14.7
R_s	15.8	22	29.6	4.0	11.8	3.0	12.8	11.9	25	20.9	6.9	18.8	6.9	14.6
<i>as</i>	275-335	291-427	193-331	211-231	200-226	103-109	129-183	193-225	185-229	177-277	99- 127	110-198	189-207	
H_O	0.95	0.95	0.91	0.47	0.76	0.07	0.72	0.74	0.87	0.95	0.74	0.70	0.45	0.71
H_E	0.91	0.93	0.95	0.43	0.84	0.07	0.79	0.72	0.90	0.81	0.72	0.74	0.53	0.72
<i>HWE</i>	0.86	0.05	0.24	0.57	0.64	1.00	0.26	0.92	0.22	0.99	0.11	0.67	0.08	0.402

Abbreviations: WNA western North Atlantic, WSA western South Atlantic, *n* number of individuals, *a* number of alleles, R_s allelic richness, *as* size range of alleles, H_O observed heterozygosity, H_E expected heterozygosity, *HWE* probability of conformation to Hardy-Weinberg expectations.

pooled, suggesting that the surveyed microsatellite loci do sort independently. The frequency of null alleles estimated by FreeNA across all genotyped loci was < 5.0% and therefore considered negligible for analysis purposes [35,63].

Due to the significant LD between some loci, we utilized three post-hoc pairwise estimates of F_{ST} to assess population differentiation between WNA and WSA roundscale spearfish: fixation indices were computed for 1) each locus individually, 2) across all 13 loci, and 3) across just 10 loci (i.e., excluding loci Mn01, *tge54*, and *tge139* to eliminate any disequilibrium bias). Individual locus F_{ST} estimates of divergence between WNA and WSA roundscale spearfish populations ranged between -0.0046 to 0.0194, with five of 13 loci providing significant estimates of divergence at $P < 0.05$ (Mn1, Mn10, *tge119*, *tge144*, *tge151*). Across all 13 microsatellite loci, the overall F_{ST} was estimated at 0.0037 and was significant at $P = 0.05$. The 10 locus F_{ST} estimate was also 0.0037 and significant ($P = 0.05$). Estimates of the arithmetic mean of the D_{est} statistic were low but statistically significant: $D_{est} = 0.021$ ($P = 0.005$) and 0.019 ($P = 0.012$) for the 13 and 10 loci, respectively. No statistically significant, nuclear marker based evidence of sex-biased dispersal was detected in any of the parameters estimated across either suite of 10 or 13 loci.

Mitochondrial DNA analyses provided mixed evidence of differentiation between the WNA and WSA roundscale spearfish. The Φ_{ST} estimate of 0.0046 was non-significant ($P = 0.24$); in contrast, the S_{nn} statistic was 0.625 and significant ($P = 0.017$). Jost's D test statistic showed no divergence as D was estimated at -0.061 (95% confidence intervals 0.000, 0.146).

Population genetic structure of roundscale spearfish: individual-level analyses

Results from the 13- and 10-locus microsatellite data sets were congruent for all individual-based analyses. *Structure* identified a single, homogenous population of roundscale spearfish within western Atlantic waters. Mean Ln Pr ($X|K$) values across the ten runs peaked at $K = 1$ for both model-type analyses [without spatial model: Mean Ln Pr ($X|K$) = -10177.3; with *locprior* model: Mean Ln Pr ($X|K$) = -10177.3], and variances associated

with likelihood estimates increased at $K > 1$ (not shown). Geneland derived posterior distributions of the estimated number of populations (K) also produced a clear mode at $K = 1$ for all 10 runs. Log likelihoods ranged from -8805 (run 4) to -9002 (run 9) and no evidence of genetic subdivision among samples was detected (not shown). Coordinate uncertainty had no effect on the estimated number of populations (not shown). Individual-based Mantel tests revealed a lack of significant correlation between pairwise genetic and geographical distance among individuals for all comparisons ($R^2 = 0.00001$; $P = 0.487$).

Comparative mitochondrial DNA-based genetic diversity and demographic histories of the roundscale spearfish and white marlin

Sample sizes and population-level mitochondrial diversity indices for both billfish species are listed in Table 2. Sequencing 577 580 base pairs (bp) of the roundscale spearfish (total $n = 83$) mtCR resolved 69 haplotypes consisting of 17.83% cytosine, 32.70% thymine, 34.74% adenine, and 14.74% guanine (GenBank Accession no. KF441482-KF441550). The sequences revealed 154 polymorphic sites consisting of 134 transitions, 19 transversions, and 15 indels. Overall haplotype (h) and nucleotide (π) diversities were 0.993 ± 0.004 and 0.024 ± 0.012 , respectively, and were similar between the WNA and WSA samples. In comparison, white marlin mtCR sequences ($n = 99$) resolved 91 haplotypes consisting of 21.69% cytosine, 28.84% thymine, 30.48% adenine, and 18.99% guanine. A total of 225 polymorphic sites were identified, consisting of 208 transitions, 10 transversions, and 22 indels. Overall, mtCR genetic diversity estimates in white marlin were higher than roundscale spearfish (Table 2, see h , π , and Θ).

The TVM model of substitution plus invariable sites (I) and a gamma distribution (Γ) of rate heterogeneity across variable sites provided the best fit to the roundscale spearfish mtCR data set (jModelTest). The estimated parameters under this model were $\Gamma = 1.1990$, and $I = 0.51$. For white marlin, jModelTest identified the HKY + I + G model [64] ($\Gamma = 1.0830$, and $I = 0.3240$) as the most

Table 2 Mitochondrial control region sequence variability and population demographic parameters for roundscale spearfish and white marlin

Species and Sample	<i>n</i>	<i>nh</i>	<i>h</i>	π	MPE Θ (95% CI)	MPE <i>g</i> (95% CI)	F_s	R_2
RS WNA	42	36	0.992 0.007	0.024 0.012	0.391 (0.206, 0.895)	220.04 (129.93, 352.30)	-17.00	0.053*
RS WSA	41	37	0.994 0.007	0.024 0.012	0.219 (0.124, 0.447)	148.35 (68.13, 256.15)	-20.49	0.058*
RS overall	83	69	0.993 0.004	0.024 0.012	0.402 (0.267, 0.666)	193.47 (125.30, 284.18)	-24.21	0.044*
WM overall	99	91	0.998 0.002	0.037 0.018	0.772 (0.534, 1.226)	180.00 (130.55, 242.07)	-23.95	0.045*

Abbreviations: RS, roundscale spearfish; WM, white marlin; WNA, western North Atlantic; WSA, western South Atlantic; *n*, number of individuals; *nh*, number of haplotypes; *h*, haplotype diversity; π , nucleotide diversity; MPE Θ , most probable estimate of Kuhner's (2006) parameter theta; (95% CI), 95% credibility intervals; MPE *g*, most probable estimate of Kuhner's (2006) exponential growth rate; F_s , Fu's (1996) test statistic; R_2 , Ramos and Rozas (2002) test statistic. * indicates significance at $P < 0.01$; indicates significance at $P < 0.001$.

appropriate substitution model for the mtCR dataset. All demographic analysis results for roundscale spearfish and most (see below) for white marlin were consistent with a scenario of population expansions for both species. Estimates of F_{ST} were negative and significantly different from zero, whereas R_2 values were small, positive, and statistically significant (Table 2) for all three roundscale spearfish collections and for the pooled white marlin collection. Furthermore, for both species, mismatch analyses revealed large differences in Θ_0 and Θ_1 , also indicative of rapid population expansions (Suppl. online Additional file 1). Similarly, the mismatch distribution model fit statistics, Harpending's [55] raggedness index and SSD, failed to differ significantly from that expected under a model of sudden population expansion (Suppl. online Additional file 1). The mismatch distribution for the pooled roundscale spearfish samples appeared smooth and unimodal, consistent with a model of population expansion. In contrast, the mismatch distribution for white marlin was distinctly bimodal indicative of a largely stable population size (Suppl. online Additional file 2), but inconsistent with the demographic summary test statistics for this species. Estimates of time since expansion (τ) for both roundscale spearfish populations overlapped substantially, suggesting a similar timing of demographic events in the WNA and WSA regions (Suppl. online Additional file 1). The mean timing of this expansion was estimated as $\tau = 10.97$ for the pooled roundscale spearfish samples, and likely occurred approximately 211 000 - 530 000 years before present (ybp), assuming mutation rates of 4.5% and 1.8% per site per million years, respectively. Assuming the same mutation rates, the range of expansion times for the white marlin pooled samples was 235 000 - 585 000 ybp.

MPEs of Θ generated by LAMARC showed variation between the two species (and the two roundscale spearfish collections) (Table 2). The white marlin median estimate of Θ was approximately 1.9 times larger than that for roundscale spearfish, although substantial overlap among credibility intervals was found. Estimates of g were similar for both species and strongly positive, indicating substantial historical growth (Table 2).

Bayesian skyline plots provided a signal of mostly continuous historical population size growth for the roundscale spearfish and white marlin (Figure 3; Suppl. online Additional file 3). Credibility intervals (95%) around estimates of female N_E (female effective size \times generation time) showed substantial overlap between species, although the final median estimates of female N_E for white marlin were roughly 1.5-1.9 times higher than roundscale spearfish.

Discussion

Roundscale spearfish population structure

We provide the first examination of the genetic population structure of the roundscale spearfish, a large, pelagic

predator captured in international fisheries and whose existence has only recently been recognized. The mixed population structure inferences obtained from the different statistical approaches used here highlight some of the difficulties associated with identifying management units for pelagic teleosts with high vagility and contiguous distributions over large geographic scales. We address two issues relevant to deriving management and conservation inferences from our findings: (i) the discordance between population- and individual-level statistical analyses in the framework of the resolving power of these analyses, and (ii) the biological interpretation of the weak but significant genetic structure revealed by population-level statistical approaches.

Numerous studies have demonstrated that the use of highly polymorphic microsatellite markers in combination with population-level (pairwise) statistical tests have increased the ability to detect shallow genetic discontinuities between populations [65,66]. In contrast, individual, multilocus-based clustering or assignment methods may have lower power to resolve such weak genetic structure [66-68]. For example, rigorous testing of individual-based analyses suggests an inability to identify divergence below a threshold of $F_{ST} < 0.01 - 0.03$ [66-68]. While F_{ST} values below this magnitude often indicate low levels of genetic partitioning, biologically important differences between such mildly divergent populations may still be present, and should not be ignored as they may be relevant for the management and conservation of species of concern [69]. For roundscale spearfish, individual-based nuclear analyses (*Structure* and Geneland) failed to detect intra-specific genetic population structure between the northern and southern hemisphere sampling sites. In contrast, the significant F_{ST} ($P = 0.05$) obtained from population-level analyses supports the notion that at least shallow genetic differentiation exists between roundscale spearfish from the WNA and WSA. The individual-based analyses may not have resolved this shallow level of differentiation because it fell below their respective resolution thresholds.

Biological interpretation of the results of roundscale population-level analyses is complicated by the mixed outcomes obtained, which were dependent on the combination of marker and statistical test used. For example, even though population differentiation was not observed using individual-based analyses, the microsatellite pairwise statistical tests (F_{ST} and D_{est}) were notably congruent in suggesting very shallow but statistically significant divergence between WNA and WSA roundscale spearfish collections. While some controversy exists surrounding the relative utility of the estimators F_{ST} and D_{est} when paired with highly variable microsatellite genetic markers [37,41,70], the fact that both estimators provided congruent results, support the inference that shallow

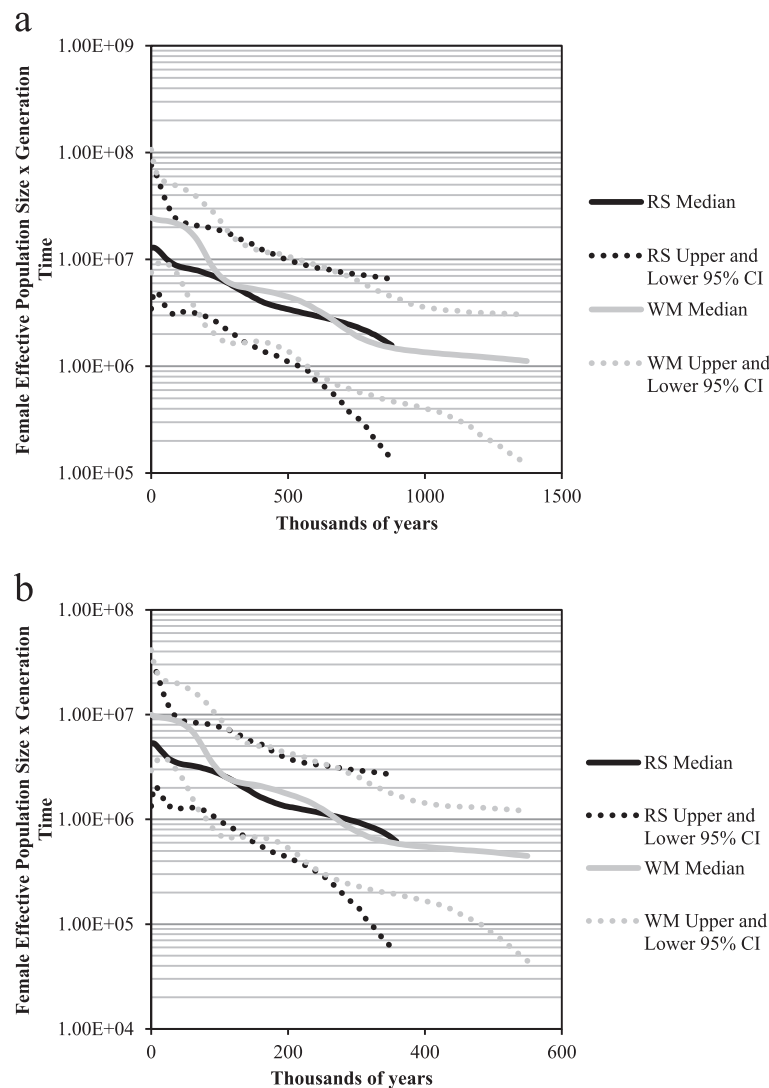


Figure 3 Bayesian skyline plot (BSPs) estimated using BEAST from pooled samples of roundscale spearfish (RS: *Tetrapturus georgii*) and white marlin (WM: *Kajikia albida*); BSPs derived using a mutation rate of (a) 1.8% per site per million years, and (b) 4.5% per site per million years.

genetic differentiation may exist between WNA and WSA populations.

Mitochondrial DNA analyses also revealed similar contradictions for inferences of roundscale spearfish population structure. Estimates of Φ_{ST} suggested an absence of differentiation, but the S_{nn} statistic identified significant differentiation among collections ($P = 0.017$). As roundscale spearfish haplotype diversity was quite high ($h = 0.992 - 0.994$), the S_{nn} test is likely a more powerful statistic than the traditional Φ_{ST} statistic to measure population-level differentiation [39].

Previous surveys of the genetic population structure of other istiophorid species (white marlin, blue marlin [*Makaira nigricans*] and sailfish, [*Istiophorus platypterus*]), have also found little, if any, support for population structure within the Atlantic [23,71-74]. However, notable

parallels may be found when comparing the population structure of roundscale spearfish to the sympatric and also Atlantic-limited white marlin. For example, analysis of white marlin genetic population structure utilizing microsatellite markers and mtCR sequences also provided mixed inferences, depending on marker class and analysis method used. Previous work [23], employed five microsatellite loci and found a small but statistically significant F_{ST} of 0.0041 ($P = 0.017$) between WNA and WSA collections, which is very similar to the level of differentiation we found for roundscale spearfish ($F_{ST} = 0.0037$). Furthermore, as was found with the roundscale spearfish, the white marlin mtCR data did not detect significant differentiation between the WNA and WSA samples based on the Φ_{ST} statistic [23]. However, in contrast to the roundscale spearfish results, the S_{nn} statistic did not differentiate

white marlin from the WNA and WSA, despite both species having similarly high haplotype diversities. These contrasting S_{nn} results may be due to the small white marlin sample sizes used for the mtCR analyses ($n = 20$ per collection), which likely reduced the power to detect shallow differentiation [23,75]. Based on the mixed results across divergent marker classes, along with the weak, albeit significant and nearly significant, spatial differentiation obtained with microsatellite markers (varied by analysis method), Graves and McDowell [23] recommended continued management of white marlin as a single, Atlantic-wide stock. However, given some indications of population heterogeneity, they also recommended that this issue be further investigated with more microsatellite markers, better planned sampling design and larger sample sizes.

Such shallow population differentiation in roundscale spearfish, despite sampling from geographically distant regions (northern vs. southern hemispheres), raises the question of whether these populations should be treated as separate management units (MUs, *sensu* [76]). While it is possible that such shallow differentiation is a result of sufficient gene flow occurring between hemispheres to prevent accumulation of larger genetic differences, it is also possible that fine scale demographic independence exists between roundscale spearfish from the WNA and WSA. This latter assertion is based on the concordance of significant differentiation from both nuclear and mitochondrial markers between populations. Furthermore, it is also possible that the observed shallow differentiation between hemispheres was a result of a sampling artifact caused by assessing individuals from distinct populations, captured as a mixed assemblage of migratory adults. These equivocal results, especially placed in context of known overfishing of other billfish species - which is very likely also occurring for roundscale spearfish [18] - and the need for precautionary management principles for billfish in general [77], leads us to recommend that roundscale spearfish be recognized for future assessments and conservation on a two MU basis comprising northern and southern hemisphere stocks.

Comparative mitochondrial DNA-based genetic diversity and demographic histories of the roundscale spearfish and white marlin

With one exception (see below), all statistical tests (Table 2 and Suppl. online Additional file 1) and Bayesian coalescent-based methods for inferring historical population trends were concordant in supporting strong signals of population expansion for both billfish species. However, the mismatch distributions for the roundscale spearfish and white marlin differed, being smooth and unimodal for the roundscale spearfish but ragged and multimodal for the white marlin (Suppl. online Additional file 2).

The distribution curve for the roundscale spearfish was consistent with a demographic history of sudden expansion (or exponential growth [54]), but the distribution for white marlin was inconsistent with the expansion model. We note, however, that the white marlin mismatch distribution failed to statistically deviate from model expectations of expansion (see *Hri* and SSD in Suppl. Online Additional file 1). The reason for this discrepancy between the observed multimodal mismatch distribution curve and statistical fit is unclear. Collectively, however, the majority of the demographic results overwhelmingly support the scenario that both species have experienced substantial historical growth throughout the Pleistocene, consistent with findings for a number of other large pelagic species (e.g., [3,78,79]). Interestingly, the roundscale spearfish and white marlin mismatch distributions suggest that a population expansion began between ~200 000 - 600 000 ybp. This temporal window overlaps several Pleistocene interglacial periods, including one of the warmest and longest interglacials (the M11) which occurred approximately 400 000 ybp [80], which would have provided billfish with the opportunity for population expansion. However, we recognize that these estimates are based entirely on the assumed mutation rate, and may not accurately reflect the appropriate temporal window of population growth.

Estimates of roundscale spearfish nucleotide and haplotype diversity fell within those reported for the mtCR of other billfishes [23,43,81,82]. Interestingly, however, comparison of the genetic diversity indices of the roundscale spearfish and white marlin revealed estimates (nucleotide and Θ) to be consistently higher for white marlin, although substantial overlap of confidence intervals was present (Table 2). Overall, estimates of white marlin diversity (nucleotide and Θ) were 1.5-1.9 times those of the pooled collections of roundscale spearfish. Assuming equal mutation rates, generation times, and the selective neutrality of mtCR, these results suggests that the historical Nf_E of white marlin may be larger (1.5 to 1.9 times) than roundscale spearfish. However, it is important to note additional caveats related to this inference.

To date, no information is available on the generation time and growth of the roundscale spearfish [83], and small differences in generation time between the two species may lead to notable differences in estimates of their effective population size. Both species, however, occupy similar habitats and likely possess many similar life history characters, supporting the hypothesis of similar generation times.

Coalescent-based estimates of the female effective population size also suggested a higher effective size for white marlin. Final median estimates of Nf_E derived from the BSPs (effective female population size x generation time) for white marlin were approximately 1.5 to 1.9 times greater than for roundscale spearfish (Figure 3), although

again substantial overlap of credibility intervals was present. Furthermore, LAMARC-derived [59] coalescent-based estimates of white marlin and roundscale spearfish N_{fE} similarly showed a higher population size of white marlin relative to roundscale spearfish. These N_{fE} estimates represent the long-term female effective population size of these two species (weighted harmonic mean over $2N_E$ generations), suggesting that historically white marlin population size has been larger than that of the roundscale spearfish.

As stated above, because the estimates of female effective population size generated herein are based on a single mitochondrial region, and the assumption of equal generation times and mutation rates for the two species, the absolute and even relative N_{fE} s should be considered highly provisional. Moreover, it is important to note that coalescent-based analyses incorporate numerous sources of error when estimating both the genealogy and population history from a small sample of individuals [84], and that similar population histories may generate highly variable BSPs (including variability in coalescent times) [85], suggesting that our BSPs and our relative estimates of population size should be interpreted with caution. Interestingly, however, although the higher N_{fE} s for white marlin compared to roundscale are preliminary, they are concordant with recent species observations of catch proportion data from fisheries which suggest that white marlin census population size is higher than that of roundscale spearfish, at least in the WNA and Caribbean waters [18,86,87]. Assuming an equal susceptibility of both billfish species to the fishery and a 1:1 sex ratio, the comparative N_{fE} estimates suggest that white marlin census size may also be close to two times higher than that of roundscale spearfish. What are the implications of a roundscale spearfish population size that is much lower than the white marlin population size? It is well established that white marlin have been severely overfished [10] and that its population biomass may be as low as 12% of the biomass required to produce maximum sustainable yield. There are no data on roundscale spearfish landings to assess the status of its stocks, but it is very likely that it has historically been landed as white marlin because its taxonomic existence in fisheries data has been recognized only recently [11]. Since its recognition, three studies conducted in commercial and recreational fisheries have confirmed that the roundscale spearfish can make up a substantial proportion (~22-27%) of the putative white marlin catch [18,86,87].

To examine the effects of misidentification of roundscale spearfish on population trends of both roundscale spearfish and white marlin, Beerkircher et al. [18] conducted population assessment simulations under various demographic scenarios. The majority of simulation outcomes showed steeply declining population trends for both species, but even greater declines for roundscale spearfish relative to

white marlin. With a much smaller apparent population size for roundscale spearfish, as suggested by the genetic analyses and fisheries data, continued exploitation of this species at current levels raises considerable concern about the long-term population health of this recently recognized species.

Conclusions

The population structure and comparative demographics results presented here underscore the importance of changing from the current international management model for white marlin as a two species complex to management of the roundscale spearfish and white marlin as distinct evolutionary lineages. We recognize that a shift to species-specific management will be challenging due to misidentifications. However, there are now morphological identification tools available to distinguish the two species [12,17], and the increasing use of genetic tools can assist in this process.

Finally, we recognize that our conclusions pertaining to roundscale spearfish population structure and demographics are based on analyses of samples from only the western Atlantic, which may represent just part of this species potential Atlantic-wide distribution. However, our findings on these parameters placed in context of general billfish overfishing, almost no information on the life history of the roundscale spearfish, and the difficulty of enforcing pelagic fishery regulations on the international level, call for an aggressive precautionary management policy for this enigmatic species. Individual species management and conservation attention, including management on a two stock precautionary basis, are prudent to avoid inadvertent, drastic reductions in what appears to be a lower abundance species, and unrecognized population collapse of either potential roundscale spearfish hemispheric stock.

Additional files

Additional file 1: Mitochondrial control region parameter estimates for mismatch distributions. RS, roundscale spearfish; WM, white marlin; WNA, western North Atlantic; WSA, western South Atlantic; τ , tau derived from mismatch distribution; Θ_0 , theta at time 0 derived from mismatch distribution; Θ_1 , theta at time 1 derived from mismatch distribution; H_r , Harpending's (1994) Raggedness index derived from mismatch distribution; SSD, sum of squared differences derived from mismatch distribution; P , probability.

Additional file 2: Mismatch distributions for pooled samples of roundscale spearfish (RS: *Tetrapturus georgii*) and white marlin (WM: *Kajikia albida*).

Additional file 3: Bayesian skyline plots (BSPs) for western North Atlantic (WNA) and western South Atlantic (WSA) roundscale spearfish (*Tetrapturus georgii*) populations. BSPs derived using a mitochondrial control region mutation rate of (a) 1.8% per site per million years, and (b) 4.5% per site per million years.

Competing interests

The authors declare that they have no competing interests.

Authors contributions

AMB generated all data and performed the data analyses. The study was designed and manuscript written by AMB and MSS. KAF provided analytical support and provided manuscript edits. Samples and general manuscript comments were provided by EDP, FHVH, FA, and AD. All authors read and approved the final manuscript.

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