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FROM THE COVER

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Comparative population genomics of manta rays has global implications for management

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Abstract

Understanding population connectivity and genetic diversity is of fundamental importance to conservation. However, in globally threatened marine megafauna, challenges remain due to their elusive nature and wide-ranging distributions. As overexploitation continues to threaten biodiversity across the globe, such knowledge gaps compromise both the suitability and effectiveness of management actions. Here, we use a comparative framework to investigate genetic differentiation and diversity of manta rays, one of the most iconic yet vulnerable groups of elasmobranchs on the planet. Despite their recent divergence, we show how oceanic manta rays (*Mobula birostris*) display significantly higher heterozygosity than reef manta rays (*Mobula alfredi*) and that *M. birostris* populations display higher connectivity worldwide. Through inferring modes of colonization, we reveal how both contemporary and historical forces have likely influenced these patterns, with important implications for population management. Our findings highlight the potential for fisheries to disrupt population dynamics at both local and global scales and therefore have direct relevance for international conservation of marine species.

KEYWORDS

conservation genomics, fisheries management, genetic diversity, population structure, ddRAD

1 | INTRODUCTION

Understanding the extent to which populations are connected is key to exploring population dynamics, predicting extinction risk and informing conservation management (Hanski & Gilpin, 1991; Lowe & Allendorf, 2010; Mills & Allendorf, 1996). In species with isolated populations characterized by limited dispersal, the risk of extirpation from local depletion is high (Reed, 2004). In such cases,

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local and regional scale management will be most appropriate for preventing and reversing population declines (Palumbi, 2003). In contrast, species with high rates of gene flow are potentially demographically and genetically more resilient to extrinsic factors (Lowe & Allendorf, 2010; Pascual et al., 2017). However, in order to maintain connectivity and mitigate genetic diversity loss in these taxa, management measures must be coordinated and encompass migratory corridors. As overexploitation and habitat destruction threaten to disrupt population dynamics at a global scale, characterizing genetic variation and connectivity has become more important than ever before (Funk et al., 2012; Kardos et al., 2021; Palsbøll et al., 2007).

In widely distributed marine species with high dispersal potential, genetic differentiation is often found to be subtle or non-existent (Palumbi, 2003; Waples, 1998; Ward et al., 1994). Such patterns can arise from a range of mechanisms - from high contemporary gene flow to recent divergence of historically large populations (Palumbi, 2003; Waples et al., 2008; Waples & Gaggiotti, 2006) - and can therefore be difficult to interpret. The latter scenario reflects a disconnect between demographic and genetic connectivity and has important implications for species resilience (Bailleul et al., 2018; Lowe & Allendorf, 2010; Waples, 1998). This is because populations that appear genetically connected may not operate as a single demographic unit, making them more vulnerable to overexploitation. High-resolution SNP datasets go some way to addressing this problem by providing greater power to detect subtle differences at both neutral and adaptive loci (Gagnaire et al., 2015; Hauser & Carvalho, 2008). However, since population genetic differentiation can be affected by past, as well as contemporary patterns, parallel inference of historical relationships and genetic variation can allow the relative contribution of historical processes to be explicitly evaluated (Foote & Morin, 2016; Liu et al., 2022; Louis et al., 2021). Furthermore, when carried out within a comparative framework, such an approach can provide powerful insights into the drivers of population divergence and therefore improve recommendations for conservation management (Gagnaire, 2020).

Manta rays are large, mobile elasmobranchs inhabiting tropical and sub-tropical oceans (Couturier et al., 2012) (Figure 1a,c) and provide an excellent opportunity to evaluate the genomic consequences of historical and contemporary population processes within a comparative framework. They comprise two described species estimated to have diverged less than 0.5 Mya as a result of distinct habitat preferences (Kashiwagi et al., 2012). The reef manta ray (Mobula alfredi) frequents near-shore tropical reef environments, such as coral atolls and barrier reefs (Kashiwagi et al., 2011), with a high degree of residency (Braun et al., 2015; Deakos et al., 2011; Germanov et al., 2022; Jaine et al., 2014; Knochel, Hussey, et al., 2022; Peel et al., 2019; Setyawan et al., 2018). In contrast, while the oceanic manta ray (Mobula birostris) also inhabits near-shore environments, it is often found ranging into sub-tropical habitats along continental coastlines and at oceanic islands, usually adjacent to productive deep-water

upwellings (Andrzejaczek et al., 2021; Kashiwagi et al., 2011). As a result of these differences in habitat use, M. alfredi and M. birostris have long been considered to display marked differences in their migratory abilities and levels of gene flow. Yet, only a handful of long-distance movements have ever been recorded in M. birostris (Andrzejaczek et al., 2021; Knochel, Cochran, et al., 2022) alongside observations of site fidelity (Cabral et al., 2023; Garzon et al., 2023; Gordon & Vierus, 2022), raising questions about the extent to which population structure and genetic variation may differ across species. To date, assessments of genetic differentiation in M. alfredi have focussed on local and regional patterns (Lassauce et al., 2022; Venables et al., 2021; Whitney et al., 2023), and we have little understanding of how genetic variation is distributed across the species' range. In, M. birostris, the situation is even less clear, with studies reporting both widespread connectivity and population differentiation (Hosegood et al., 2020; López et al., 2022; Stewart et al., 2016). Critically, these differences and uncertainties exist against a background of ongoing global exploitation and uncertain implications for management.

Targeted and incidental fisheries, driven in part by increasing demand for mobulid gill plates (Couturier et al., 2012; O'Malley et al., 2017), have led to widespread population declines in manta rays (Carpenter et al., 2023; Croll et al., 2016; Moazzam, 2018; Rohner et al., 2017; Ward-Paige et al., 2013). Currently, both species are managed through a patchwork of local, regional and international measures with varying levels of implementation and enforcement (Fernando & Stewart, 2021; Lawson et al., 2017; Lawson & Fordham, 2018). To determine the appropriateness of management measures and assess population vulnerability, a global assessment of management units is urgently required (Lawson et al., 2017: Stewart, Jaine, et al., 2018). Here, we undertake a comparative genomic analysis of manta ray populations from across their global distribution to investigate connectivity, genetic variation and historical relationships with an aim to guide effective fisheries management and emphasize the value of genomic research for advancing knowledge of understudied elasmobranchs.

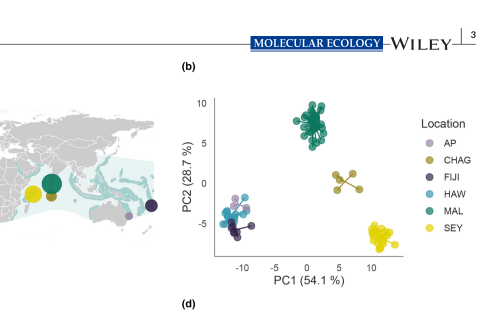
2 MATERIALS AND METHODS

2.1 Sample collection

Tissue samples were opportunistically collected from 12 geographic locations to represent the global distribution of each species (Figure 1a,c). For M. alfredi (total n = 120), these originated from the Chagos Archipelago (n=5), the Maldives (n=48), Seychelles (n = 24), Australia Pacific (n = 4), Fiji (n = 9) and Hawaii (n=30). For M. birostris (total n=112), these originated from Sri Lanka (n=43), the Philippines (n=37), South Africa (n=3), Mexico Caribbean (n = 4), Mexico Pacific (n = 13) and Peru (n = 12). Samples from Mexico Caribbean, where a third putative manta ray species occurs in sympatry (Mobula cf. birostris, Hinojosa-Alvarez et al., 2016; Hosegood et al., 2020), were visually and genetically

(a)

(c)



Oceanic manta ray (Mobula birostris)

Reef manta ray (Mobula alfredi)

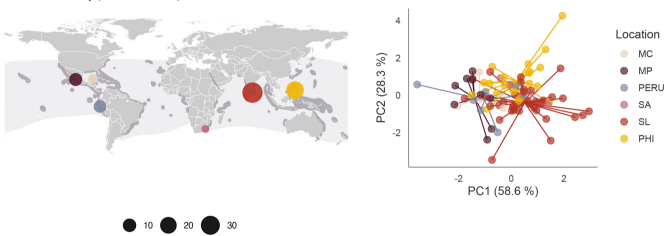


FIGURE 1 Contrasting patterns of population structure in manta rays. (a, c) Geographic distributions of (a) *Mobula alfredi* and (c) *Mobula birostris* visualized together with the locations of samples used in this study. Dark-shaded distributions denote the confirmed species range, and light-shaded distributions denote the expected species range. Sampling location points are distinguished by colour and scaled by the number of samples. Further details are provided in the Appendix S1. (b, d) Scatterplots showing individual variation in principal components (PC) one and two derived from discriminant analysis of principal components analysis for (b) *M. alfredi* and (d) *M. birostris* individuals. The amount of variance explained by each PC is shown in parentheses. Population abbreviations: AP, Australia Pacific; CHAG, Chagos; FIJI, Fiji; HAW, Hawaii; MAL, Maldives; MC, Mexico Caribbean; MP, Mexico Pacific; PERU, Peru; PHI, the Philippines; SA, South Africa; SEY, Seychelles; SL, Sri Lanka.

confirmed as *M. birostris*. For both species, samples were collected from a combination of live animals and fisheries specimens (see Appendix S1 for further information).

2.2 | DNA extraction and ddRAD sequencing

Genomic DNA was extracted using the Qiagen DNeasy Blood and Tissue Kit and quantified using a Qubit 3.0 Broad Range Assay. Double digest restriction-site associated DNA (ddRAD) libraries were prepared following the Peterson et al. (2012) protocol with modifications described in Palaiokostas et al. (2015) and were 125 bp paired-end sequenced on an Illumina HiSeq. Sequencing reads from both species were assessed for quality using FastQC and processed together using the Stacks v2.54 de novo assembly pipeline (Catchen et al., 2013). The three main assembly parameters were chosen following the optimization procedure outlined in Rochette and Catchen (2017) (Figures S1 and S2). Initial quality filters were applied to the resulting genotypes before generating three high-quality datasets for use in downstream analysis: two speciesspecific datasets, and one dataset comprising both species. For the species-specific datasets, we extracted either *M. birostris* or *M. alfredi* individuals, removed individuals with high relatedness coefficients (Korneliussen & Moltke, 2015; Waples et al., 2019, Figure S3) and filtered out SNPs with a minor allele count of less than 3, a genotyping rate less than 90% and that were in linkage disequilibrium using PLINK. For the dataset comprising both species, we first removed closely related individuals and then filtered out SNPs with a minor allele count of less than 90%. Population genetic summary statistics were calculated using the R package diversity (Keenan et al., 2013, Table S1). See Appendix S1 for further information on library preparation, read processing and SNP and individual filtering.

2.3 | Population structure

To investigate population structure, we used the species-specific datasets and three complementary approaches. First, we carried out a discriminant analysis of principal components (DAPC) using the R package adegenet (Jombart, 2008). This approach initially transforms the SNP data using a principal components analysis (PCA) and then performs a discriminant analysis on the retained PCs. This serves to maximize discrimination of individuals between groups while minimizing variation within (Jombart et al., 2010). Following the recommendations outlined in Thia (2023), the number of PCs retained as predictors was determined based on the K-1 criterion, where K is equal to the number of effective populations. For M. alfredi, this was set to 5, under the assumption that each sample site reflects a separate population. For M. birostris, this was set to 4 under the assumption that Mexico Pacific and Peru may represent a single population given their close geographic proximity. Second, we estimated admixture proportions for the individuals in each dataset using ADMIXTURE. Admixture runs were performed for ancestry clusters ranging from K=1-8, with 10 runs for each K. The optimal K was identified based on the lowest cross-validation error. The runs with the highest likelihood were visualized. Third. we estimated pairwise genetic differentiation between populations within each species using the Weir and Cockerham F_{ST} value (Weir & Cockerham, 1984) calculated in the R package dartR (Gruber et al., 2018). Confidence intervals and p-values were estimated based on bootstrap resampling of individuals within each population 1000 times. Mobula alfredi samples from Australia Pacific and M.birostris samples from South Africa were excluded from this analysis due to low sample sizes.

2.4 | Isolation by distance

To investigate patterns of isolation by distance, we examined the relationship between genetic and geographic distance between all pairs of populations in each species. Genetic distances were based on the pairwise $F_{\rm ST}$ estimates calculated above. Geographic distances were determined based on a least-cost path analysis implemented using the R package marmap (Pante & Simon-Bouhet, 2013) with a minimum depth constraint of -10m in order to prevent paths overland. The significance of associations between genetic and geographic distance matrices was inferred using distance-based Moran's eigenvector maps (dbMEM) by redundancy analysis (RDA, Legendre et al., 2015). For this, geographical distances were transformed into dbMEMs using the R package adespatial, and genetic distances were decomposed into principal components using the R function prcomp. RDA was then

performed using the R package vegan, with significance tested using 1000 permutations.

2.5 | Contemporary gene flow

To infer the strength and directionality of contemporary gene flow between populations, we used the programme BA3-SNPs BayesAss v1.1 (Mussmann et al., 2019) which estimates the proportion of immigrants in a given population using Bayesian inference. This analysis was restricted to M. alfredi as it assumes low levels of connectivity and imposes an upper bound on the proportion of non-migrants in a population. We first performed initial runs of BayesAss to determine optimal mixing parameters (dM=migration rate, dA=allele frequency and dF=inbreeding coefficient) using the autotune function in BA3-SNPs. We then ran BayesAss-3 with 10,000,000 iterations, a burnin of 1,000,000 and a sampling interval of 1000. Mixing parameters were set to dM = 0.21, dA = 0.44 and dF = 0.08. Results were averaged across five replicate runs, and migration rates were considered significant if 95% credible sets (mean migration rate \pm 1.96 \times mean standard deviation) did not overlap zero. Chain convergence was assessed, and migration rates were visualized using R (Figure S4).

2.6 | Historical relationships among populations

To explore historical relationships among populations of M. alfredi and M. birostris, we used the programme TreeMix (Pritchard et al., 2000). TreeMix uses population allele frequencies to estimate a bifurcating maximum likelihood tree with which to infer historical population splits, admixture events and the degree of genetic drift. We first supplemented the M. alfredi dataset with one randomly selected M. birostris individual and the M. birostris dataset with one randomly selected M. alfredi to act as outgroups when rooting the trees. Both datasets were then filtered for linkage, a minor allele count of less than 3, a genotyping rate less than 90% and related individuals using PLINK v1.9 (Purcell et al., 2007). Allele frequencies for each population were then calculated using the -freq and -within arguments in PLINK. For both the M. birostris and M. alfredi datasets, we then performed 10 initial runs of TreeMix for each migration event (M) ranging from 0 to 10. The number of migration edges that explained 99.8% of the variance was selected as the best model for each species (M. birostris: M=0; M. alfredi: M=2, Figure S5). We then re-ran TreeMix 100 times using the optimal number of migration edges. Consensus trees and bootstrap values were estimated and visualized using code modified from the BITE R package (Milanesi et al., 2017).

2.7 | Genome-wide heterozygosity

To assess levels of genetic variation both within and between species, we used the high-quality SNP dataset comprising both species.

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We then calculated multi-locus heterozygosity for each individual using the R package inbreedR (Stoffel et al., 2016).

3 | RESULTS

High-quality SNPs were genotyped in 173 individuals from 12 locations representing the global distribution of each species (Figure 1a,c). The species-specific datasets contained a total of 1553 SNPs in 91*M. alfredi* individuals, and 6278 SNPs in 82*M. birostris* individuals, while the full dataset contained a total of 15,312 SNPs called across both species. See Materials and Methods and Appendix S1 for details.

3.1 | Contrasting patterns of population structure at a global scale

To investigate population differentiation within each species, we used four complementary approaches: discriminant analysis of principal components (DAPC), admixture, pairwise F_{ST} and isolation by distance analysis. In *M. alfredi*, all methods supported the presence of strong population structure at both global and regional scales. Populations inhabiting different ocean basins displayed the highest degree of differentiation in the DAPC, with Pacific and Indian

Ocean populations forming distinct clusters along PC1 (Figure 1b). Regional differentiation was also detected, with Seychelles, Chagos and the Maldives clustering apart along PC2, and Hawaii separating from Australia Pacific and Fiji along PC3 (Figure 1b and Figure S6A). These patterns were reinforced in the admixture analysis which highlighted two major ancestral source populations, inferred an optimal value of K=4 and resolved hierarchical structure up to K=7(Figures S7 and S8A). Interestingly, only weak separation was observed between Australia Pacific and Fiji; however, this pattern may be confounded by the small sample size of the former, which can lead to spurious merging of distinct populations (Puechmaille, 2016). Pairwise F_{sT} estimates between ocean basins were on average over two times higher than those within (mean pairwise F_{ST} between ocean basins = 0.30, mean pairwise F_{ST} within ocean basins = 0.13, Figure 2a); yet all population comparisons were found to be significant (Figure S9A, mean=0.23, min=0.08, max=0.43). Finally, we detected a significant relationship between pairwise F_{ST} and geographic distance (adjusted R^2 =.65, p=.03) indicating an effect of isolation by distance (Figure 2b).

In stark contrast, *M. birostris* displayed little evidence of strong population structure across all methods. Individuals from different ocean basins clustered closely together along each axis in the DAPC (Figure 1d and Figure S6B). Admixture identified K=1 as the optimal number of clusters, with increasing values of *K* merely introducing additional mixing (Figures S7 and S8B). Pairwise F_{ST} estimates

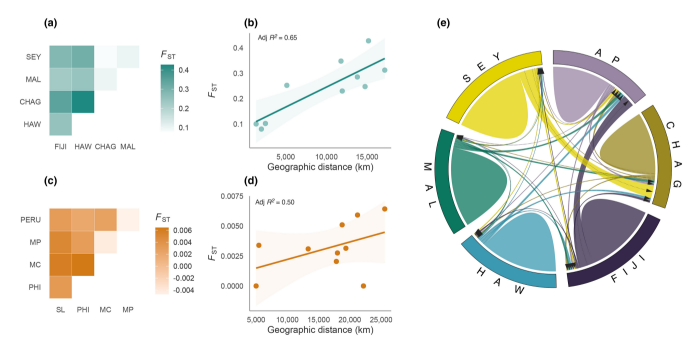


FIGURE 2 Population genetic differentiation, isolation by distance and contemporary migration in manta rays. (a, c) Pairwise F_{ST} estimates between sampling locations for (a) *M. alfredi* and (c) *M. birostris*. Samples from Australia Pacific and South Africa were excluded from this analysis due to low sample sizes. (b, d) Relationship between genetic (F_{ST}) and geographic distance as calculated by least-cost path analysis for all pairwise population comparisons in (b) *M. alfredi* and (d) *M. birostris*. Solid lines and shaded areas reflect the regression slopes and standard errors, respectively, based on a linear model. Samples from Australia Pacific and South Africa were excluded from this analysis due to low sample sizes. Note that F_{ST} scales differ between (a) and (c), and axis scales differ between (b) and (d). (e) Contemporary gene flow estimates between populations of *M. alfredi*. The direction of each arrow represents the direction of gene flow, and the width of each ribbon reflects the relative amount of gene flow. Population abbreviations: AP, Australia Pacific; CHAG, Chagos; FIJI, Fiji; HAW, Hawaii; MAL, Maldives; MC, Mexico Caribbean; MP, Mexico Pacific; PERU, Peru; PHI, the Philippines; SEY, Seychelles; SL, Sri Lanka.

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were two-fold lower than in M.alfredi, with no pairwise comparison falling above 0.007 (mean=0.002, min=-0.005, max=0.006, Figure 2c). Nevertheless, despite these broad patterns, several lines of evidence indicate the presence of subtle geographic differentiation in this species. First, individuals from Mexico Pacific, Peru and Mexico Caribbean clustered separately from those sampled in South Africa, Sri Lanka and the Philippines along PC1 (Figure 1d). Second, despite pairwise F_{sT} estimates being low, comparisons between Eastern-Pacific and Indo-Pacific populations, and between Sri Lanka and the Philippines were statistically significant (Figure S9B). Small F_{sT} values are expected when minor allele frequencies are low and therefore do not necessarily reflect an absence of differentiation (Jakobsson et al., 2013). Furthermore, it is possible that overall levels of population structure were underestimated due to the small sample size of two of our six populations (Puechmaille, 2016). Finally, while no significant relationship was observed between pairwise F_{sT} and geographic distance (adjusted $R^2 = .50$, p = .10), there was a tendency for populations separated by greater distances to display higher differentiation (Figure 2d).

3.2 Contemporary gene flow

To characterize the strength and direction of gene flow between populations, we used the programme BA3-SNPs (Mussmann et al., 2019) to estimate recent migration. As this method assumes low levels of connectivity and imposes an upper bound on the proportion of non-migrants in a population, this analysis was restricted to M.alfredi. As expected, contemporary gene flow was low (Figure 2e); the average migration rate between populations, measured as the estimated number of migrants per generation, was 0.029 (min=0.008, max = 0.15), with this figure falling to 0.018 (min = 0.008, max = 0.03) when considering gene flow between populations in different ocean basins. Migration into both Hawaii and the Maldives was lowest, indicating these populations are the most isolated of those sampled (Table S2). Migration rates were only deemed significant between Seychelles and Chagos (0.15) and between Fiji and Australia Pacific (0.15), in line with these populations being last to separate in the admixture analysis. These patterns highlight that while M. alfredi may have the propensity to travel over large distances, restricted movement likely dominates.

3.3 Historical relationships among populations

To place patterns of genetic differentiation into a historical context, we investigated population origins and colonization patterns using TreeMix (Pritchard et al., 2000). This programme uses allele frequency data to infer patterns of population splits and admixture events through the construction of a maximum likelihood tree. In M. alfredi, internal branch lengths were relatively long, with an initial split clearly separating populations in the Indian and Pacific Oceans (Figure 3a). The Maldives and Australia Pacific were the first to separate within

each locality and displayed the lowest levels of genetic drift overall. Hawaii was among the last populations to split and displayed the highest amount of drift, in line with its geographic isolation. The best supported model inferred two migration events (Figure S5A): one from the M. alfredi population in Seychelles into M. birostris and one from Chagos into Hawaii. However, because not all geographic regions are represented in our dataset, the true sources and sinks of these admixture events may originate from related ghost populations. In contrast to M. alfredi, the addition of migration events led to no substantial improvement in the model for M. birostris (Figure S5B), and therefore the tree without migration is presented here. Interestingly, internal branch lengths were considerably shorter in M. birostris, indicating rapid radiation from a shared ancestral source population (Figure 3b). External branch lengths were also short, consistent with larger populations displaying marginal drift and low divergence. Nevertheless, despite these patterns, some geographic signal was detected in the M. birostris tree, with individuals originating from the Eastern Pacific and the Caribbean (Peru, Mexico Caribbean and Mexico Pacific) grouping separately from those originating from the Atlantic and Indo-Pacific (South Africa, the Philippines and Sri Lanka), although bootstrap support was overall low.

3.4 | Heterozygosity landscape across species and populations

To explore how patterns of population structure and colonization are associated with genome-wide variation, we compared individual multi-locus heterozygosity between species and among populations. Strikingly, heterozygosity was on average over three times higher in M. birostris (mean=0.10, min=0.053, max=0.12) than in M. alfredi (mean = 0.03, min = 0.01, max = 0.051), with every individual displaying a higher value than any *M*. alfredi (β = .07, 95% CI = 0.06 - 0.07, $p = <2.2 \times 10^{-16}$, Figure 3c). This finding is in line with the patterns of population structure and historical splits we observed in each species. Variation in heterozygosity was also observed at a population level (Figure 3d,e). In M. alfredi, the Maldives and Australia Pacific had the highest levels of heterozygosity within each ocean basin, in line with these populations being first to split in the TreeMix analysis. Indian Ocean populations displayed higher overall heterozygosity than Pacific populations and had mean values that were overall similar. Interestingly, heterozygosity within Pacific populations declined steeply from west to east, with heterozygosity in Hawaiian individuals being around half that of the Australian animals (Figure 3d), in line with this population being last to split in the TreeMix analysis and displaying the highest amount of drift. In contrast, M. birostris populations displayed less extreme variation in heterozygosity, with mean values differing by less than 0.02 and having no clear geographical pattern (Figure 3e). Furthermore, variance within populations was an order of magnitude greater in M. birostris than in M. alfredi, and was particularly large in Sri Lanka, the Philippines and Mexico Pacific populations.

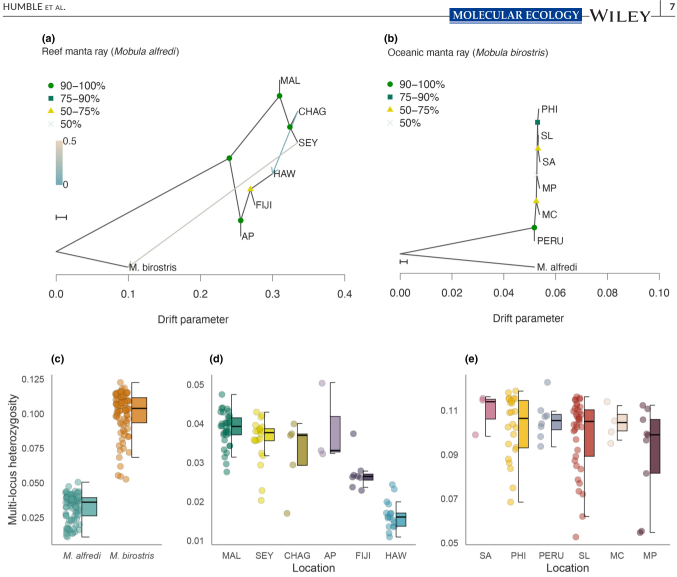


FIGURE 3 Historical relationships and heterozygosity in manta rays. (a, b) TreeMix maximum likelihood consensus tree displaying the historical relationships among (a) M. alfredi and (b) M. birostris populations. Horizontal branch lengths reflect the amount of genetic drift that has occurred along each branch. Bootstrap support values for each node are indicated. Migration edges inferred using TreeMix are represented as arrows and coloured according to their migration weight. The scale bar reflects 10 times the average standard error of the entries in the sample covariance matrix. (c-e) Variation in individual multi-locus heterozygosity between (c) species and among populations of (d) M. alfredi and (e) M. birostris. Note that y-axis scales differ for (d) and (e). Centre lines of boxplots reflect the median, bounds of the boxes extend from the first to the third quartiles, and upper and lower whiskers reflect the largest and smallest values but no further than 1.5×the interquartile range from the hinge. Population abbreviations: AP, Australia Pacific; CHAG, Chagos; FIJI, Fiji; HAW, Hawaii; MAL, Maldives; MC, Mexico Caribbean; MP, Mexico Pacific; PERU, Peru; PHI, the Philippines; SA, South Africa; SEY, Seychelles; SL, Sri Lanka.

DISCUSSION 4

Manta rays are iconic megafauna with cultural, socio-economic and ecological significance. Due to targeted and bycatch fisheries operating across their broad-ranging distributions, populations are declining worldwide. Elucidating levels of connectivity and genetic variation remain a crucial priority for conservation management. We use reduced-representation sequencing on a global set of samples and reveal striking differences in the population genetic landscape of two recently diverged manta ray species. By considering the influence of both contemporary and historical processes, our study

provides a precautionary framework for assessing conservation units in widely distributed marine species.

We first demonstrate the presence of strong genetic differentiation in M. alfredi at a global and regional scale. From a total of six sampling locations, we found evidence for at least five genetically distinct, and by extension, demographically independent populations. Two of these were separated by a distance of ~1200km, which is close to the maximum recorded movement in the species (Armstrong et al., 2019; Jaine et al., 2014), indicating that long-distance migrations are likely rare. Indeed, contemporary gene flow was low - especially between geographically

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distant locations - with only a small proportion of individuals in any population being identified as first- or second-generation migrants. Furthermore, when gene flow was observed, it tended to be unidirectional. These results are in line with recent studies demonstrating population differentiation between Western Australia and Mozambique (Venables et al., 2021) and between Eastern Australia and New Caledonia (Lassauce et al., 2022), together highlighting how large ocean basins form significant barriers to dispersal in coastal elasmobranchs (Hirschfeld et al., 2021). High site fidelity has been widely reported in M. alfredi based on tagging and mark-recapture studies (Braun et al., 2015; Deakos et al., 2011; Germanov et al., 2022; Jaine et al., 2014; Knochel, Hussey, et al., 2022; Peel et al., 2019; Setyawan et al., 2018). However, the degree of residency has been shown to vary, with movements rarely exceeding a few hundred kilometres in some locations (Braun et al., 2015; Deakos et al., 2011; Kessel et al., 2017; Knochel, Hussey, et al., 2022; Setyawan et al., 2018) yet reaching over 1000 km in others (Armstrong et al., 2019; Germanov & Marshall, 2014). Our study presents a comparatively broad-scale analysis relevant to regional and global management planning. Further work on local patterns of population structure will shed light on the nuances and drivers of fine-scale movement patterns in this species (Whitney et al., 2023).

To explore the mechanism by which manta rays colonized their distribution, we reconstructed historical relationships and assessed levels of heterozygosity. In M. alfredi, we found strong evidence for an initial split between the Indian and Pacific Oceans followed by further separation within each locality. Furthermore, populations in the Pacific displayed a signal of declining heterozygosity from west to east, together suggesting that M. alfredi underwent a stepping-stone pattern of range expansion from an Indo-Pacific Ocean origin, involving opportunistic long-range movements and associated founder events. This is consistent with a recent observation of a pregnant M. alfredi individual at Cocos Island, Costa Rica (Arauz et al., 2019), almost 6000km east of the nearest confirmed sighting, and the first record of M. alfredi in the Eastern Pacific. Range expansion inherently impacts genetic variation, with a stepping-stone model of colonization predicted to result in the strongest cumulative effect of founder events (Le Corre & Kremer, 1998). Among our sampled populations, Hawaii is the most geographically isolated, situated at the edge of the *M. alfredi* distribution. Interestingly, not only was Hawaii the most genetically differentiated from all populations in our study, but it displayed the longest external branch lengths in the TreeMix analysis and the lowest levels of heterozygosity. This could be suggestive of a single founder event by a small population. Genetic variation is fundamental for enabling populations to adapt in response to selection (Bonnet et al., 2022; Kardos et al., 2021; Lai et al., 2019). Our findings therefore expose how isolated M. alfredi populations at the periphery of their distribution may be intrinsically more vulnerable to changing environmental conditions and the genetic impacts of population decline.

In stark contrast to the patterns observed in M. alfredi, M. birostris displayed markedly higher levels of heterozygosity and only [365294x, 0, Downloaded from https://onlinelibrary.wiley.com/doi/10.1111/mec.17220 by Bangor University,

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subtle genetic differentiation across ocean basins. Weak population structure is common in highly mobile marine species (Leslie & Morin, 2018; Nikolic et al., 2023; Vignaud et al., 2014), and yet warrants careful interpretation, particularly considering management recommendations (Younger et al., 2017). On the one hand, these findings may be an indication of high contemporary gene flow and low natal philopatry, in line with the species' occurrence at remote oceanic islands, tendency to range into sub-tropical habitats and lower overall re-sight rates than M. alfredi (Couturier et al., 2014; Harty et al., 2022; Rambahiniarison et al., 2023). To date, our understanding of the movement behaviour in M. birostris has largely been based on coastal aggregations of adult individuals over relatively short timeframes (Beale et al., 2019; Harty et al., 2022; Rohner et al., 2013; Stewart et al., 2016). Such studies have a tendency to capture seasonal migrations as opposed to dispersal events and may explain why only a handful of long-distance (~1000km) movements have been recorded in the species (Andrzejaczek et al., 2021; Knochel, Cochran, et al., 2022). In an infinite island model, only a few migrants per generation are required to obscure strong population structure when N_{a} is large (Wright, 1931), and therefore it is possible the patterns we observe translate to infrequent dispersal events. Furthermore, dispersal could be segregated by age and/or sex (McClain et al., 2022; Phillips et al., 2021) and may vary among individuals (Papastamatiou et al., 2013; Perryman et al., 2022; Thorburn et al., 2019). While challenging, there is benefit in extending future tagging efforts to transient individuals away from known aggregation sites (Garzon et al., 2023), as well as previously underrepresented age classes - such as juveniles - to capture what may be infrequent yet evolutionarily relevant movements.

An alternative explanation for the patterns we observe in M. birostris is that insufficient time has elapsed to reliably identify recent genetic divergence among localities. In contrast to M. alfredi, our TreeMix analysis indicated that M. birostris rapidly radiated from a large ancestral source, with only marginal genetic drift occurring between regions. This was further evidenced by substantially higher levels of genetic variation that differed little across sampling locations. In addition, little differentiation was observed between Mexico Pacific and Mexico Caribbean, two regions that have been geographically separated since the emergence of the Isthmus of Panama. These findings are consistent with a recent mark-re-sight analysis that estimated the population of M. birostris in coastal Ecuador to number at least 22,000 individuals (Harty et al., 2022). Large effective population sizes and high genetic variation increase the time taken for populations to diverge due to genetic drift (Bailleul et al., 2018; Taylor & Dizon, 1996; Wright, 1931). This is further compounded in species with long and overlapping generations (Hoffman et al., 2017) as is the case for manta rays (Dulvy et al., 2014). Taken together, genetic similarities among M. birostris localities may be partially confounded by recent shared ancestry and large effective population size.

On the basis of these considerations, we propose that a combination of large historical population size and contemporary gene flow has contributed to the comparatively high levels of heterozygosity and genetic homogeneity in M. birostris. The subtle population differentiation we observe between the Indian Ocean, South-East Asia and the Eastern Pacific is likely best explained by the geographic limits of dispersal as opposed to complete geographic isolation. Yet, unlike in M. alfredi where genetic clusters almost certainly reflect discrete demographic units relevant for conservation management, the extent to which genetic connectivity in M. birostris reflects demographic connectivity is less clear. For example, in extreme cases, the number of migrants required to eliminate signals of population structure will not be enough to demographically link populations, and more importantly, replenish those that have been depleted (Waples, 1998). Interestingly, while re-sight rates are typically lower in M. birostris than M. alfredi, demographic independence has been implicated in several mark-recapture studies where re-sightings follow predictable patterns (Beale et al., 2019; Cabral et al., 2023). Furthermore, a population genetic analysis based on F_{sT} outliers uncovered allele frequency differences between two Mexican locations and Sri Lanka (Stewart et al., 2016), suggesting recent divergence against a background of ongoing gene flow. Taken together, we highlight the potential for further work investigating adaptive divergence between M. birostris populations and emphasize the need to combine molecular measures of connectivity with empirical demographic data in this species (Cayuela et al., 2018; Lowe & Allendorf, 2010; Younger et al., 2017).

4.1 | Conservation implications

The remarkable differences we observe in the population genetics of manta rays directly inform likely response to continued exploitation and respective conservation measures. At present, M. alfredi is among the most protected mobulid species worldwide, with some management frameworks in place at local, national and international levels (Lawson et al., 2017; Stevens et al., 2018). Our findings of global population structure underline how local initiatives recognizing populations as distinct management units will be most appropriate for this species. However, we also demonstrate the consequence of geographic isolation on genetic variation and reveal how M. alfredi likely faces a greater risk from local depletion. This is especially true for populations at the edge of the species range and in regions with high coastal fishing pressure. Prioritizing these populations in conservation action plans and maintaining local connectivity will therefore be crucial for boosting resilience and preventing local extinction in this vulnerable species.

The implications of our findings for *M. birostris* are more nuanced. Despite detecting only subtle population genetic differentiation, we cannot rule out the possibility that historical processes and large effective population size are obscuring a higher degree of contemporary demographic separation. Together with studies reporting high site fidelity and restricted movement patterns, our findings strongly suggest that local and national management actions should be considered essential for protecting resident aggregations of *M. birostris*. Nevertheless, we expect that weak population structure and high - MOLECULAR ECOLOGY – WILEY

genetic variation are simultaneously being driven by some degree of contemporary dispersal. Consequently, any fishing activity taking place along migratory corridors threatens to disrupt a mode of gene flow that may be fundamental for long-term resilience of the species. Similarly, although we have limited understanding of the number and distribution of breeding and nursery grounds (Knochel, Cochran, et al., 2022; Pate & Marshall, 2020; Stewart, Nuttall, et al., 2018), significant reduction in local stocks may impact long-term recruitment at oceanic and even global scales. We therefore emphasize the escalating need to improve the implementation of regional and international measures that seek to protect taxa in the high seas. Together with local scale management, appropriate evidence-based actions will contribute to maintaining large, connected and genetically diverse populations of manta rays into the future.

AUTHOR CONTRIBUTIONS

EH, JH, GC, MdB, SC, GMWS and RO conceived and designed the study. GMWS, AA, RB, MD, DF, NF, LRP, SP, AP, JDS and SW provided samples. JH carried out laboratory work. EH analysed the data with input from JH. EH wrote the paper with input from all other co-authors.

AFFILIATIONS

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CONFLICT OF INTEREST STATEMENT

The authors declare no competing interests.

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BENEFIT-SHARING STATEMENT

A research collaboration was developed with scientists from regions providing genetic samples to advance the conservation of manta rays and their relatives through evidence-based research, international collaboration and institutional capacity building. Data and analysis pipelines associated with this work have been shared on public databases for the benefit of this wider cause.

DATA AVAILABILITY STATEMENT

Sequencing data have been deposited to the European Nucleotide Archive under study accession number PRJEB66437. Analysis code is available at https://github.com/elhumble/manta_pop_gen_2022.

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