iDNA at sea

Recovery of whale shark (Rhincodon typus) mitochondrial DNA sequences from the whale shark copepod (Pandarus rhincodonicus) confirms global population structure

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The whale shark (Rhincodon typus) is an iconic and endangered species with a broad distribution spanning warm-temperate and tropical oceans. Effective conservation management of the species requires an understanding of the degree of genetic connectivity among populations, which is hampered by the need for sampling that involves invasive techniques. Here, the feasibility of minimally-invasive sampling was explored by isolating and sequencing whale shark DNA from a commensal or possibly parasitic copepod, Pandarus rhincodonicus that occurs on the skin of the host. We successfully recovered mitochondrial control region DNA sequences (~1,000 bp) of the host via DNA extraction and polymerase chain reaction from whole copepod specimens. DNA sequences obtained from multiple copepods collected from the same shark exhibited 100% sequence similarity, suggesting a persistent association of copepods with individual hosts. Newly-generated mitochondrial haplotypes of whale shark hosts derived from the copepods were included in an analysis of the genetic structure of the global population of whale sharks (644 sequences; 136 haplotypes). Our results supported those of previous studies and suggested limited genetic structuring across most of the species range, but the presence of a genetically unique and potentially isolated population in the Atlantic Ocean. Furthermore, we recovered the mitogenome and nuclear ribosomal genes of a whale shark using a shotgun sequencing approach on copepod tissue. The recovered mitogenome is the third mitogenome reported for
the species and the first from the Mozambique population. Our invertebrate DNA (iDNA) approach could be used to better understand the population structure of whale sharks, particularly in the Atlantic Ocean, and also for genetic analyses of other elasmobranchs parasitized by pandarid copepods.

**Keywords:** eDNA, sharks, minimally-invasive sampling, copepod, control region, population genetics, parasite, commensal

### INTRODUCTION

The term environmental DNA (eDNA) refers to DNA extracted from cells that are not collected directly from a target organism, but are obtained from the environments they inhabit such as oceans, river water, soil, and air (Ficetola et al., 2008; Fonseca et al., 2010; Andersen et al., 2012; Sigsgaard et al., 2016). Invertebrate-derived DNA (iDNA) is an offshoot of this approach that involves the extraction of genetic material of animals via the flesh-eating or haematophagous invertebrates that parasitise them (Schnell et al., 2015; Schubert et al., 2015; Lee et al., 2016). To date, most iDNA studies have focused on terrestrial vertebrates and have extracted host DNA from insects, ticks, or leeches. There have been no analogous studies in marine environments, despite the potential usefulness of the approach for sampling large marine megafauna such as cetaceans, sirenians, pinnipeds, marine reptiles (sea turtles), elasmobranchs, and teleosts. For such taxa, iDNA sampling offers both ethical and practical advantages as it is minimally-invasive and thus preferable to the direct collection of blood or tissue, particularly where target species are rare or endangered.

The whale shark (*Rhincodon typus*), similar to many large marine vertebrates, poses significant challenges for researchers trying to understand their biology and ecology and for managers attempting to develop conservation strategies (Graham and Roberts, 2007; Rowat et al., 2009). Tagging, genetic and modeling studies suggest that individuals can disperse widely (Sequeira et al., 2013), although there is evidence of isolation between populations in the Indo-Pacific and Atlantic oceans (Vignaud et al., 2014). However, the scale at which population structure can be discerned is dependent on sample sizes, with low numbers reducing analytical power to reject the null hypothesis of a panmictic population (Castro et al., 2007). For whale sharks, low sample size tends to be a result of the rarity of the species and the difficulties in obtaining biopsies (usually of skin tissue) for genetic analyses, which requires appropriate permits and ethical approvals for invasive sampling of a species that is categorized as “Endangered” by the International Union for the Conservation of Nature and the use of trained divers and technicians. In this situation there are many benefits to the use of an iDNA approach, provided that external invertebrate parasites or commensals that harbor the intact DNA of the whale shark host can be successfully identified. Although eDNA techniques have recently been used to obtain whale shark DNA (Sigsgaard et al., 2016), iDNA still offers a major advantage because it enables haplotypes to be linked to individual whale sharks for the analysis of the genetic structures of populations.

Copepods (Crustacea, Maxillopoda, Copepoda) are an excellent candidate species for iDNA studies on marine vertebrates. Free-living forms are a dominant element of marine zooplankton and may even exceed insects in terrestrial environments in terms of sheer abundance of individuals. Less recognized is that approximately half of the nearly 30,000 described species live in parasitic or commensal associations with a diverse range of taxa, including fish and mammals (Boxshall, 2005). The order Siphonostomatoida contains largely parasitic copepods that feed on the blood, epidermal tissue, or mucus of many marine teleost fishes, sharks, and rays. Approximately 550 genera, representing nearly 40 families, are placed in the order and include economically important species such as sea lice (Brachiura) that parasitise farmed fish (Gunn and Pitt, 2012). Eleven siphonostomatoid families have been reported as symbionts of a diverse range of elasmobranchs (Dippenaar, 2009) and one family, the Pandaridae, is composed of 23 genera that include species that are ectoparasites or commensals of large sharks (Cressey and Boyle, 1978; Walter and Boxshall, 2017). Within the Pandaridae, the genus *Pandarus* currently has 17 recognized species of which *P. rhincodonicus* (Norman et al., 2000) is noteworthy as it is appears to be associated exclusively with the whale shark, where it is predominately found on the leading and trailing edges of fins and on the lips. Although it is thought to be a commensal that feeds off bacteria and other microorganisms on the skin of the shark (Norman et al., 2000), dietary studies of the species are incomplete.

Here, we demonstrate that *P. rhincodonicus* sampled from sites across the Indian Ocean contain sufficient DNA from their whale shark host to routinely recover mitochondrial sequences that allow analyses of the genetic structure of host populations, and the recovery of the complete whale shark mitogenome and ribosomal nuclear genes. The implications of these findings are discussed in relation to the use of copepods for iDNA studies of marine teleosts and sharks and the ecological status of *Pandarus* spp. as commensals.
Mitochondrial Control Region Analysis

Authenticity of the DNA sequences of the mitochondrial control region of whale sharks were validated by BLAST searches, and sequences were aligned with previously generated sequences (Castro et al., 2007; Vignaud et al., 2014) using MAFFT version 7,310 with the option “--adjustdirection” activated (Yamada et al., 2016). The 5' and 3' ends of the alignment were manually trimmed using MEGA 6 (Tamura et al., 2013) so that each aligned control region had the same flanking sequence (Supplementary Figure 1). The trimmed alignment was subsequently de-gapped using Seqret (Rice et al., 2000) and clustered with cd-hit-est at a 100% sequence identity and 100% sequence length coverage cut-off as implemented using the –c 1.0 and -a 1.0 setting (Li and Godzik, 2006).

Analysis of Population Differentiation

The mitochondrial control regions of whale sharks generated using iDNA were combined with 613 sequences published by previous studies of whale shark populations (Castro et al., 2007; Ramirez-Macias et al., 2007; Schmidt et al., 2010; Vignaud et al., 2014). Identical iDNA sequences from the same whale shark host were removed prior to genetic analysis, to eliminate the chance of biasing estimates of population genetic differentiation (Supplementary Table 1). Arlequin suite version 3.5.2.2 (Excoffier and Lischer, 2010) was used to estimate global and pairwise FST. A median-joining haplotype network was constructed using PopArt version 1.7 (Leigh and Bryant, 2015) and to simplify network representation, only the common haplotypes (a haplotype with more than one sequence representation) were used to generate the network graph.

Recovery of Whale Shark Whole Mitochondrial Genome and Nuclear Ribosomal Genes

Assembly used a baiting and iterative mapping approach as implemented in MITObim version 1.8 (Hahn et al., 2013) from the complete mitogenome of a whale shark sampled off Taiwan (Accession Number: NC_023455) as the reference sequence (Alam et al., 2014). In-silico circularization and annotation of the assembled mitogenome followed (Iwasaki et al., 2013; Gan et al., 2014). To improve alignment accuracy and specificity, mitogenome coverage was calculated by mapping the raw paired-end reads to the assembled mitogenome sequence using Bowtie2 version 2.3.2 (Langmead and Salzberg, 2012) with the setting “—score-min L,0.2,—0.2.” BRIG was used to visualize the mitogenome annotation and read mapping coverage (Alikhan et al., 2011). Similar Bowtie2 mapping setting was used to map the raw reads to a whale shark assembly contig containing the nuclear ribosomal genes (18S rRNA and 28S rRNA) and the read alignment was subsequently visualized using Integrative Genomics Viewer (Thorvaldsdottir et al., 2013).

RESULTS

Mitochondrial Control Region Analysis

A ~1,000 bp DNA sequence fragment of the whale shark mitochondrial control region was successfully generated from the extracted gDNA from 44 copepods representing 31 sharks. Haplotype clustering confirmed a 100% sequence match for DNA sequences amplified from copepods collected from the same whale shark host (Supplementary Data 1). An initial alignment of the 31 newly-generated iDNA control region sequences with the 613 sequences provided by previous studies (Castro et al., 2007; Schmidt et al., 2010; Vignaud et al., 2014; Walter and Boxshall, 2017) yielded a 1,910 bp product (Supplementary Data 2). Trimming using 5'- and 3'-ends produced a final alignment length of 750 bp (Supplementary Data 3), from which a total of 163 unique haplotypes were evident (Supplementary Data 4). The Ningaloo Reef population had the highest number of sampled control region sequences (n = 163) and also had the greatest number of haplotypes (h = 36) (Figure 1A). Interestingly, the observed haplotype diversity for the Isla Holbox population was 50% lower (h = 11) than that of Djibouti population (Vignaud et al., 2014), despite similar sample sizes (Figures 1A,B).

Of the 31 deduplicated iDNA sequences reported in this study, five were found to represent replicate individuals from the Ningaloo population previously sequenced by Vignaud et al. (2014) using skin samples (Supplementary Table 1 and Supplementary Data 4). Despite the relatively low number of sequences reported by our study that represented sequences from new whale sharks (N = 27), our study contributed a 39 and 22% increase in the number of individuals sampled from the Mozambique and Seychelles populations, respectively (Figure 1A). In addition, four novel haplotypes for control region sequences of whale sharks were identified from copepod samples.
Population Structure of the Whale Shark

Global $\phi_{ST}$ was weak but significantly different from zero, indicating limited gene flow among some sampling locations. This pattern was driven by a single population, with all pairwise $\phi_{ST}$ calculations associated with the Isla Holbox population differing significantly from zero (average $\phi_{ST}$ of 0.2), whereas all estimates of $\phi_{ST}$ did not (Figure 2). This implied that the population at Isla Holbox was genetically isolated from an otherwise panmictic population spread across the Indo-Pacific Ocean. Three common haplotypes occurred at similar frequency across the sampling range (Figure 3). A few private haplotypes occurred in the Ningaloo Reef, Philippines and Seychelles populations. Consistent with $\phi_{ST}$ estimates, the population at Isla Holbox was the most genetically differentiated, as it lacked one of the three common ancestral haplotypes (H108, 110, 125, 150, and 152), and had a high frequency of others that were rare (H67, H68, and H76).

Recovery of the Whale Shark Mitogenome and Nuclear Genes from Shotgun Sequencing of Copepod Tissue

The complete mitochondrial genome of the whale shark was successfully assembled from partial genome sequencing of an iDNA sample from a copepod (isolate 366.1; Figure 4). Mapping of the 14 million paired-end reads to the reconstructed whale shark mitogenome gave a mapping rate of 0.006% (900 reads) representing approximately $5 \times$ mitogenome coverage (Figure 4). The mitogenomic composition of the whale shark was extracted was very similar (>99% identity) to those reconstructed from sharks from Taiwanese waters (Accession codes: KC633221 and NC_023445). A comparison of the 13 genes coding for mitochondrial proteins indicated one or two nucleotide mismatches, mostly associated with non-synonymous mutations found in the $\text{atp6}$, $\text{cox1}$, $\text{cytb}$, $\text{nad2}$, $\text{nad4}$, and $\text{nad5}$ genes. In addition, we also observed reads mapping to the whale shark 18S and 28S rRNA genes (Supplementary Figure 2 and Supplementary Data 5), indicating the presence of whale shark nuclear DNA in the copepod extracted gDNA, a phenomenon that will be useful for future population genetic studies of these sharks based on nuclear markers.

Data Deposition

Mitochondrial control regions were submitted to NCBI under the accession numbers MF872682-MF872725. Raw reads from the shotgun sequencing of the copepod were submitted to Sequence Read Archive under the run number SRR4111090 and the reconstructed whale shark mitogenome has been assigned the accession number MF872726.

DISCUSSION

Our study is the first demonstration of the use of iDNA sampling of a marine invertebrate in order to obtain mitochondrial genetic information from an elasmobranch host. We successfully amplified mitochondrial DNA fragments of whale sharks in the size range of $\sim$1,000 bp from a copepod, one of the largest host DNA fragments to be recovered by an iDNA study (Schnell et al., 2012, 2015; Lee et al., 2016; Pérez-Flores et al., 2016; Rodgers et al., 2017). This suggests the presence of largely intact whale shark DNA in or on copepods, although the exact source could not be identified as copepods were analyzed whole. Targeted efforts to amplify DNA fragments of whale sharks from the intestine and epidermal layers of $P$. rhincodon will be useful to resolve this issue and clarify the role of the copepod as a commensal or parasite. We found iDNA sequences from multiple copepods sampled from the same whale shark host to be identical, suggesting copepods have a persistent, long-term association with their host shark, which contrasts with the more generalist and mobile host associations of other invertebrate ectoparasites in terrestrial and aquatic ecosystems, such as leeches, blow flies, and mosquitoes (Calvignac-Spencer et al., 2013). However, these findings should be treated with some degree of caution and a multi-locus approach using microsatellite
markers (Olson et al., 2012) might offer a more powerful means to validate these findings. As reported by Vignaud et al. (2014), our estimates of genetic structure across the Indian, Pacific, and Atlantic oceans indicated the presence of two distinct populations, one in the Indo-Pacific and the other in the Atlantic Ocean. Isla Holbox was the only location sampled in the Atlantic Ocean and the minimally-invasive iDNA approach could potentially be used to improve the coverage of sampling in this region.

We also demonstrated that complete mitochondrial genome and nuclear ribosomal RNA sequences of whale sharks could be obtained using iDNA sampling. Although not the first for the species (Alam et al., 2014; Chen et al., 2016), the sequences generated by our study represent the first for the Mozambique population of whale sharks. The successful recovery of the mitogenome and nuclear ribosomal RNA from the shotgun sequencing of *P. rhincodon* was probably due to the high copy number of mitochondrial and nuclear ribosomal genes in somatic cells, coupled with high sequencing depth (~14 million paired-end reads). Efficiency of the recovery of whale shark mitogenomes from copepod tissue samples might be improved by sequencing gDNA extracted from the part of the copepod body that contains higher concentrations of whale shark tissue. In addition, given the intactness of whale shark gDNA (>1,000 bp) that was present in *P. rhincodon*, long range PCR of additional mitogenome regions followed by high throughput amplicon sequencing could also be explored to dramatically increase the coverage of the whale shark mitogenome for population genetic and phylogeographic applications (Deiner et al., 2017; Pavlova et al., 2017). Given that mtDNA only represents a fraction of the evolutionary history of a species due to its strict maternal inheritance and small genome size (16 kb), studies of the population genetics of whale sharks may benefit from the combination of both mtDNA and nuclear data.
FIGURE 4 | The mitogenome of *Rhincodon typus* (upper photo—note copepod parasites *Pandarus rhincodonicus* visible as black dots near the center of the upper lip) recovered from the shotgun sequencing of the parasitic copepod, *P. rhincodonicus* (lower photo). The purple ring indicates relative mapping coverage, and the transcriptional orientation of each protein coding gene is indicated by the red arrows. (Godinho et al., 2008; Schmidt et al., 2009; Vignaud et al., 2014). The presence of nuclear ribosomal RNA reads in the copepod shotgun sequencing library suggests that it is possible to amplify both nuclear DNA and mtDNA of whale sharks from the copepod and the recent publication of the first draft genome for the whale shark (Read et al., 2017) will greatly facilitate the design of a targeted next-generation sequencing panel to enhance future studies of whale shark population genetics.

The Pandaridae comprises 23 genera and more than 100 species, many of which are ectoparasites or symbionts of large sharks (Walter and Boxshall, 2017), and there are nearly 30 other families of copepods that include species that are parasites of fishes (Boxshall, 2005). These offer an opportunity to extend iDNA methods to other taxa of large marine vertebrates. Conversely, the presence of gDNA of whale sharks in the extracted DNA of *P. rhincodonicus* will need to be considered as a possible contaminant in future whole genome
sequencing or population genomic projects that target this copepod.

**AUTHOR CONTRIBUTIONS**


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**REFERENCES**

Alam, M. T., Petit, R. A. III., Read, T. D., and Dove, A. D. (2014). The complete mitochondrial genome sequence of the world's largest fish, the whale shark (Rhinodon typus), and its comparison with those of related shark species. *Gene* 539, 44–49. doi: 10.1016/j.gene.2014.01.064


**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2017.00420/full#supplementary-material

**Supplementary Table 1** | Sample collection information.

**Supplementary Figure 1** | Visualization of aligned control region sequences (one representative from each study). Arrows indicate regions upstream and downstream of the 5’ and 3’ ends, respectively, that will be trimmed off for subsequent haplotype clustering.

**Supplementary Figure 2** | Bowtie2 alignment of Illumina reads generated from copepod gDNA to the whale shark genomic region containing 18S and 28S nuclear rRNA genes. Red and blue arrow bars indicate forward and reverse pair-end reads, respectively.

**Supplementary Data 1** | Cd-hit clustering of trimmed whale shark control region sequences that were amplified from copepods.

**Supplementary Data 2** | Alignment of whale shark control region sequences.

**Supplementary Data 3** | Trimmed alignment of whale shark control region sequences based on the conserved region depicted in Supplementary Figure 1.

**Supplementary Data 4** | Cd-hit Clustering of trimmed whale shark control region sequences.

**Supplementary Data 5** | Pared-end reads mapped to the whale shark genomic region containing the 18S and 28S ribosomal RNA genes.


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

The reviewer GS declared a shared affiliation, with no collaboration, with several of the authors, CA, MT, AM, and HG, to the handling Editor.

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