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Complete mitochondrial genome of the gray reef shark, *Carcharhinus amblyrhynchos* (Carcharhiniformes: Carcharhinidae)

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**ABSTRACT**

We report the first mitochondrial genome sequences for the gray reef shark, *Carcharhinus amblyrhynchos*. Two specimens from the British Indian Ocean Territory were sequenced independently using two different next generation sequencing methods, namely short read sequencing on the Illumina HiSeq and long read sequencing on the Oxford Nanopore Technologies’ MinION sequencer. The two sequences are 99.9% identical and are 16,705 base pairs (bp) and 16,706 bp in length. The mitogenome contains 22 tRNA genes, two rRNA genes, 13 protein-coding genes and two non-coding regions; the control region and the origin of light-strand replication (OL).

**Main text**

The gray reef shark *Carcharhinus amblyrhynchos*, is a highly-social, reef-dependent species distributed widely in the tropical Indo-Pacific and currently listed as ‘Near Threatened’ in the IUCN Red List (Smale 2009). Populations have declined due to illegal fishing activities (Osgood and Baum 2015; Ferretti et al. 2018). Whilst there have been genetic studies conducted on the species (Holmes et al. 2009; Momigliano et al. 2015, 2017), its mitogenome has not yet been described.

We describe the complete mitochondrial genome of *C. amblyrhynchos*. Tissue was sampled as fin clips from two specimens in the British Indian Ocean Territory in March 2018. Specimen 1 (GenBank MT093205) was a female tagged at location 5.46386° 71.77841° and specimen 2 (GenBank MT104515) was a male tagged at −5.24956° 71.79906°. Samples were stored at Hopkins Marine Station before specimen 1’s tissue was transferred to Silwood Park, Imperial College London. The samples were then analyzed independently in two laboratories. The DNA from specimen 1 is available at Silwood Park DNA & Tissue Bank (CITES GB038) under accession VS8956-20002085971; DNA from specimen 2 is available at Hopkins Marine Station, Stanford University under accession 020002232485. For specimen 1, genomic DNA was extracted using Qiagen’s Blood & Tissue Kit and was sequenced using an Illumina HiSeq. The mitochondrial genome sequence was assembled using ABySS v2.0.2 (Jackman et al. 2017) and GapCloser v1.12 (Luo et al. 2012).

For specimen 2, the DNA was extracted and sequenced using the Oxford Nanopore Technologies’ MinION sequencer following Johri et al. (Johri et al. 2019). The MitoFish mitoannotator (Iwasaki et al. 2013) was used to annotate the sequences, and these were aligned against one another and mitogenomes from other Carcharhinid species using MUSCLE (Edgar 2004) within Geneious Prime (v2019.0.4). A phylogenetic tree was produced in Geneious Prime using MrBayes (Huelsenbeck and Ronquist 2001; Ronquist et al. 2012) plugin (v.3.2.6, substitution model: HKY85, burn-in length: 100,000) using the gray bamboo shark (*Chiloscyllium griseum*; NC_017882) and scalloped hammerhead shark (*Sphyrna lewini*; NC_022679) as outgroups.

The complete mitochondrial genomes are 16,705 bp (specimen 1) and 16,706 bp (specimen 2) in length. Each contains two rRNAs, 22 tRNAs, 13 protein-coding genes and a non-coding control region. The nucleotide base composition is identical with 31.5% A, 25.2% C, 13.2% G and 30.1% T, the overall GC content is 38.4%. The two sequences have 16,682 identical sites (99.9% pairwise identity). The differences include one base addition and 23 substitutions. Four substitutions result in a change to the amino acid sequence of COI. These differences could be due to the sequencing methods or represent evidence of population structure within the species in BIOT despite high spatial connectivity across the territory (Carlisle et al. 2019).

Whilst the fine-scale phylogenetic relationships within Carcharinidae remain unresolved (Naylor et al. 2012), the tree (Figure 1) supports the placement of *C. amblyrhynchos* in...
a clade with *C. albimarginatus*, *C. falciformis*, and *Prionace glauca*. The low posterior probability that supports the placement of *P. glauca* with *C. albimarginatus* and *C. falciformis* suggests that further work is required to fully resolve the tree. However, the high support for deeper clades within *Carcharhinus* adds to calls for a taxonomic revision of *P. glauca* (Naylor et al. 2012; Johri et al. 2019). The new mitochondrial genomes presented here will aid in conservation genetics, environmental DNA and population studies as researchers move toward assessing populations using genome sequences.

**Disclosure statement**

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

**Author contributions**

ND and SJ contributed to concept, bioinformatics and analysis, ND wrote the manuscript with comments from all authors, SJ contributed to sequencing and EAD provided laboratory support. BAB, VS and DC contributed to the concept of the manuscript and financial support of the project. BAB, TKC and DC contributed the samples.

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**Data availability statement**


**References**


