The complete mitogenome of *Hemiculter leucisculus* (Basilewsky, 1855) in Hainan Island and its phylogenetic status

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**ABSTRACT**

The sharpbelly, *Hemiculter leucisculus* (Basilewsky, 1855) is a small cyprinid fish that has a wide distribution in East Asia. In this study, we characterized the complete mitochondrial genome of *H. leucisculus* in Hainan Island using Illumina MiSeq platform. The mitogenome contained 16,621 bp with AT content of 56.2%. The mitogenome of *H. leucisculus* comprised 13 protein-coding genes, 22 transfer RNA genes, 2 ribosomal RNA genes, and one control region (D-loop). Phylogenetic analyses indicated that *H. leucisculus* in the Hainan Island formed independent lineage and the species of *H. leucisculus* might be a paraphyletic taxon.

**Key words**

Hemiculter leucisculus; mitogenome; phylogeny

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**Introduction**

The sharpbelly, *Hemiculter leucisculus* (Basilewsky, 1855), is a small cyprinid fish that has a wide distribution in the drainage basins of China, Korea island, Japan, Mongolia, and Russia (www.fishbase.org). In China, *H. leucisculus* widely occupies various freshwater habitats, such as rivers, lakes, reservoirs, and even pools (Luo and Chen 1998). Habitat generalist and wide distribution ranges of the *H. leucisculus* populations make this species possibly show complex phylogenetic status. Chen et al. (2017) demonstrated that *H. leucisculus* populations in southern China showed deep divergence with the Yangtze River populations and could be regarded as a cryptic species. Herein, we reported *H. leucisculus* mitogenome from Hainan Island and attempted to infer its phylogenetic status within genus *Hemiculter*.

Specimen of *H. leucisculus* was collected from the farmers market of Ledong County, Hainan Province, China (18.75219N, 109.17497E) on 27 October 2019 and was care-fully distinguished this species according to the ichthyography by Luo and Chen (1998). We extracted total genomic DNA from fin tissues using a Genomic DNA Isolation Kit (Qiagen, Germany). The sample and the total DNA was stored in the fish collection of Hunan University of Arts and Science (www.huas.edu.cn, Congqiang Luo and blackball@yeah.net) under the voucher numbers HL2019001. We sequenced the complete mitochondrial genome using the Illumina MiSeq platform (Illumina Inc, San Diego, CA, U.S.A.) and assembled the raw sequence reads into contigs using SPAdes 3.9.0. The complete mitochondrial genomes were finally obtained utilizing the contigs in SOAPdenovo (Luo et al. 2012) and were annotated using GeSeq (Tillich et al. 2017) with NCBI reference sequences.

Mitogenome length and AT content for the individual from Hainan Island are 16,621 bp and 56.2%, respectively (GenBank accession: MZ520999). This mitogenome showed similar AT contents with that in previous reported *H. leucisculus* mitogenome sampled from the Heilongjiang Province, China (GenBank accession: KF647872; Dong et al. 2015) and the Zhejiang Province, China (GenBank accession: KF956522; Xiang et al. 2016). The complete mitochondrial genome comprised of 13 protein-coding genes (PCGs), 2 ribosomal RNA (rRNA) genes (12S rRNA and 16S rRNA), 22 transfer RNA genes and a control region (D-loop). The 13 PCGs of the mitogenome had same initiating codon and stop codon with *H. leucisculus* mitogenome from the Heilongjiang Province (Xiang et al. 2016) but showed different stop codon with *H. leucisculus* mitogenome from the Zhejiang Province in ND5 gene (Dong et al. 2015).

To infer the phylogenetic status of *H. leucisculus* populations in the Hainan Island, we downloaded published *Hemiculter* mitogenomes and built Bayesian inference (BI) and maximum-likelihood (ML) trees using the combined DNA sequences of 13 PCGs and corresponding amino acids in RAXML-VI-HPC (Stamatakis 2006) and MRBAYES 3.2 (Ronquist and Huelsenbeck 2003), respectively. *Xenocypris davidi* (Genbank nos: KF039718) from Xenocyprinae was selected as an out-group. Optimal nucleotide substitution model of GTR+I+G for phylogenetic analyses were selected using the Akaake information criterion in MRMODELTEST version 2.3 (Nylander 2004). For the analyses with amino acids, we allowed MrBayes to estimate the best fixed-rate amino acid matrix (i.e. prset
aamodelpr = mixed) and used WAG model inferred from to construct ML tree. BI and ML trees based on 13 PCGs consistently obtained three lineages (Figure 1(a)), while BI and ML trees on the basis of amino acids congruously yielded two lineages (Figure 1(b)). The discordant result between the two molecular types was the placement of *H. bleekerii* (Figure 1). All trees showed that *H. leucisculus* in the Hainan Island formed an independent lineage and *H. leucisculus* was a paraphyletic taxon (Figure 1). However, further studies including morphology and genetics were needed to demonstrate this point.

**Ethical approval**

Experiments were performed in accordance with the recommendations of the Ethics Committee of Hunan University of Arts and Science. These policies were enacted according to the Chinese Association for the Laboratory Animal Sciences and the Institutional Animal Care and Use Committee protocols.

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**Author contributions**

Congqiang Luo: Conception, design, analysis, interpretation of the data and writing and revising the draft. Peng Chen and Jibin Zhu: Investigation. Zemin Ma: Conception and design. All authors agree to be accountable for all aspects of the work.

**Disclosure statement**

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

**Data availability statement**

The genome sequence data that support the findings of this study are openly available in GenBank of NCBI at [https://www.ncbi.nlm.nih.gov/](https://www.ncbi.nlm.nih.gov/) under the accession no. MZ520999. The associated BioProject, SRA, and Bio-Sample numbers of specimen are PRJNA745010, SRR15080965 and SAMN20130455, respectively.

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