The complete mitochondrial genome of *Trematomus loennbergii* (Perciformes, Nototheniidae)

Eunkyung Choi*a, ⁎, Tae-Eul Im*a, ⁎, Seung Jae Lee*b, Euna Jo*b, Jinmu Kim*c, Sun Hee Kim*a,b, Young Min Chi*a, Jeong-Hoon Kim*c and Hyun Park*a

aCollege of Life Sciences and Biotechnology, Korea University, Seoul, Korea; bGreenwitch Co, Chungcheongbuk-do, Korea; cDivision of Polar Life Science, Korea Polar Research Institute, Incheon, Korea

**ABSTRACT**

The complete mitochondrial genome of *Trematomus loennbergii* was studied using NGS technology with PacBio platform. The mitochondrial genome size was 19,374bp and it had 13 protein-coding genes, 22 tRNAs and 2 rRNAs. There were 4 types of stop codons which were TAA, TAG, AGG and T(AA) but start codon type was only one (ATG). The contents of GC were 44.09% and AT contents were 55.91%. To conduct phylogenetic analysis, 12 species in 3 families were used. The result suggested that *T. loennbergii* was close to *Pagothenia borchgrevinki* in Nototheniidae. This study would provide a fundamental data for molecular evolution of *T. loennbergii*.

**ARTICLE HISTORY**

Received 14 October 2020
Accepted 28 February 2021

**KEYWORDS**

Mitochondrial genome; *Trematomus loennbergii*; Scaly rockcod; PacBio

The Nototheniidae family of Antarctic fishes have various characters, for example, antifreeze glycoprotein (AFGP) and lack of hemoglobin and these are different with other teleost fishes (Clarke and Johnston 1996 and DeVries and Cheng 2005). The subfamily Trematominae has only 14 species and among these Trematomus is main genus having 11 species and their habit distribution is very wide (Lautredou et al. 2010). In addition, the morphology of Trematomus species is very similar, for example, *Trematomus loennbergii* (Regan, 1913) is known as Scaly rockcod and their distribution is Southern Ocean such as coasts of Antarctic Peninsula, Queen Mary, Adelie, South Victoria Coasts, Davis, Ross and Weddell seas according to fish base (https://www.fishbase.se/summary/7057). Their depth distribution is usually 65–832 m and common length is 20 cm. They feed on a wide range of prey and main food resources are epifaunal and tube-dwelling polychaetes (Mesa et al. 1993).

*Trematomus loennbergii* (Regan, 1913) was collected from Ross Sea (77° 05'S, 170° 30'E on CCAMLR Subarea 88.1), Antarctica and then transferred to −80°C freezer in laboratory. The specimen was deposited at the Earth Biocollection in the Division of Biotechnology, Korea University (http://genome.kusglab.org/collections.html, amy_choi@korea.ac.kr) under the voucher number KAN0003030. The DNA was extracted from frozen muscle tissue by the conventional phenol-chloroform method. For the quality and quantity check, Fragment analyzer (Agilent Technologies, CA, USA) and Qubit 2.0 Fluorometer (Life Technologies, CA, USA) were used respectively. To conduct sequencing step, the 20 kb fragmentation was done using Covaris G-tube (Covaris, MA, USA) and the SMRTbell library was constructed using SMRTbell™ Template Prep Kit 1.0 according to manufacturer’s protocol. Sequel Binding Kit 3.0 and SMRTbell Template Prep Kit 1.0 were used to construct SMRTbell-Polymerase and SMRTbell library Complex. After this step, the complex was load into the Sequel platform using Sequel Sequencing Kit 3.0 and SMRT cells 1 M v3. Total 4 SMRT cells were used to perform sequencing step in the Sequel sequencing platform (Pacific Biosciences, CA, USA). Subreads of PacBio for mitochondria genome assembly were filtered out using 16 s rRNA and COI gene sequences and then CANU assemble 1.8 (Koren et al. 2017) was used for de novo mitochondrial genome assembly. The assembled sequences were retrieved into MITOS (Bernt et al. 2013) web service for mitochondrial genome annotation.

The complete mitochondrial genome size of *T. loennbergii* (GenBank Number: MT447073) was 19,374bp and it consisted of 13 protein-coding genes, 2 rRNAs and 22 tRNAs. Start codon
was only ATG in 13 protein coding genes but there were 4 types of stop codon. 6 protein coding genes (ND3, ND2, COX1, ATP6, COX3 and ND4L) had TAA stop codon, 3 protein coding genes (ATP8, ND3 and ND5) had TAG stop codon, 3 protein coding genes (COX2, ND4 and Cytb) had T(AA) stop codon and only ND6 had AGG as a stop codon. The contents of GC were 44.09% and AT contents were 55.91%.

A phylogenetic tree (Figure 1) was analyzed to know the evolutionary position of T. loennbergii. The MEGA X (Kumar et al. 2018) by Maximum Likelihood method with 1,000 bootstrap replications and JTT matrix-based model (Jones et al. 1992) were used for this analysis. As a result, the data showed that T. loennbergii was clustered with Pagothenia borchgrevinki in Notototheniidae family. This study would provide the fundamental data to understand the evolutionary relationship with other fishes in Antarctic region.

**Disclosure statement**

The authors reported no potential conflict of interest.

**Funding**

This research was supported by the Ecosystem Structure and Function of Marine Protected Area (MPA) in Antarctica project (PM20060) funded by the Ministry of Oceans and Fisheries under Grant 20170336.

**ORCID**

Eunkyung Choi [http://orcid.org/0000-0003-4823-2532](http://orcid.org/0000-0003-4823-2532)
Seung Jae Lee [http://orcid.org/0000-0002-2404-3107](http://orcid.org/0000-0002-2404-3107)
Euna Jo [http://orcid.org/0000-0003-2666-6743](http://orcid.org/0000-0003-2666-6743)
Jinmu Kim [http://orcid.org/0000-0003-4045-7227](http://orcid.org/0000-0003-4045-7227)
Hyun Park [http://orcid.org/0000-0002-8055-2010](http://orcid.org/0000-0002-8055-2010)

**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. MT447073. The associated BioProject, SRA, and Bio-Sample numbers are PRJNA610666, SRS6351352, and SAMN14309518, respectively.

**References**

Bernt M, Donath A, Jühling F, Externbrink F, Florentz C, Fritzsch G, Pütz J, Middendorf M, Stadler PF. 2013. MITOS: improved de novo metazoan mitochondrial genome annotation. Mol Phylogenet Evol. 69(2):313–218.

Clarke A, Johnston IA. 1996. Evolution and adaptive radiation of Antarctic fishes. Trends Ecol Evol. 11(5):212–218.

DeVries AL, Cheng C-HC. 2005. Antifreeze proteins and organismal freezing avoidance in polar fishes. In: Farrell AP, Steffensen JF, editors. The physiology of polar fishes. San Diego: Elsevier Academic Press; p. 155–201.

DeWitt HH, Heemstra PC, Gon O. 1993. Notototheniidae. In: Gon O, Heemstra PC, editors. Fishes of the Southern Ocean. p. 279–399. Grahamstown, South Africa.

Jones DT, Taylor WR, Thornton JM. 1992. The rapid generation of mutation data matrices from protein sequences. Comput Appl Biosci. 8(3):275–282.

Koren S, Walenz BP, Berlin K, Miller JR, Bergman NH, Phillippy AM. 2017. Canu: scalable and accurate long-read assembly via adaptive k-mer weighting and repeat separation. Genome Res. 27(5):722–736.

Kumar S, Stecher G, Li M, Knyaz C, Tamura K. 2018. MEGA X: molecular evolutionary genetics analysis across computing platforms. Mol Biol Evol. 35(6):1547–1549.

Lautredou AC, Bonillo C, Denys G, Cruaud C, Ozouf-Costaz C, Lecointre G, Dettai A. 2010. Molecular taxonomy and identification within the Antarctic genus Trematomus (Nototothenioidei, Teleostei): How valuable is barcoding with COI? Polar Sci. 4(2):333–352.

Mesa ML, Vacchi M, Castelli A, Diviacco G. 1997. Feeding ecology of two nototheniid fishes, Trematomus hansoni and Trematomus loennbergii, from Terra Nova Bay, Ross Sea. Polar Biol. 17(2):62–68.