Chromosomal assembly of the Antarctic toothfish (Dissostichus mawsoni) genome using third-generation DNA sequencing and Hi-C technology

DEAR EDITOR,

The Antarctic toothfish, Dissostichus mawsoni, belongs to the Nototheniidae family and is distributed in sub-zero temperatures below 60° latitude in the Southern Ocean. Therefore, it is an attractive model species to study the stenothermal cold-adapted character state. In this study, we successfully generated highly contiguous genome sequences of D. mawsoni, which contained 1,062 scaffolds with a N50 length of 36.98 Mb and longest scaffold length of 46.92 Mb. Repetitive elements accounted for 40.87% of the genome. We also inferred 32,914 protein-coding genes using in silico gene prediction and transcriptome sequencing and detected splicing variants using Isoform-Sequencing (Iso-Seq), which will be invaluable resource for further exploration of the adaptation mechanisms of Antarctic toothfish. This new high-quality reference genome of D. mawsoni provides a fundamental resource for a deeper understanding of cold adaptation and conservation of species.

Many unique fish live in the Southern Ocean surrounding Antarctica within the coldest waters on Earth. It has been isolated by the Antarctic circumpolar current (Eastman, 2005; Livermore, et al., 2005) where sea temperatures range around the ice point (−1.9 °C) for most of the year. Antarctic fish, which include cold-adapted teleosts, are dominated by a single lineage belonging to the Perciformes suborder Notothenioidei. This suborder consists of eight families and >100 species and accounts for ~90% of total fish biomass in the Antarctic Ocean (Eastman & De Vries, 1981). From a common ancestor, a variety of closely related species with distinct ecological status as well as size, shape, and color have emerged in the Southern Ocean. Therefore, genomic research is essential to understand the environmental adaptation and evolution of these fish.

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The Antarctic toothfish, Dissostichus mawsoni, belongs to the family Nototheniidae of the order Perciformes and is native to the Southern Ocean. It is distributed below 60° latitude and is the largest of all Antarctic fish (2.0 m in length and 140 kg in mass) (Eastman & De Vries, 1982). Its stenothermal cold-adapted state makes the species an attractive model for evolutionary and genomic studies among Antarctic fish. The Antarctic toothfish is an economically important fishery species, with a commercial catch in Subarea 88.1 of 2.680 tons in 2018 (Maschette et al., 2019).

Recently, de novo assembly of the Antarctic toothfish genome and extensive transcriptomic characterization using short-read Illumina data have been reported (Chen et al., 2019), though the genome was fragmented into many scaffolds due to sequencing by synthesis technology. The development of third-generation single-molecule sequencing technology has enabled the production of long-read sequences and the discovery of the features of previously unavailable DNA regions. Here, we report on a re-assembled whole-genome of D. mawsoni using long-read sequencing and Hi-C technology, which should help provide comprehensive insight into its adaptive mechanisms.

Antarctic toothfish (length ~50 cm, sex not determined) were collected using a vertical setline in the eastern Ross Sea (Subarea 88.1), Antarctica (http://www.fao.org/fishery/area/Area88/en), during the austral summer season (December 2018). Specimens were killed for tissue sampling and then rapidly frozen for further analysis. All sample collection and experimental protocols were in compliance with the laws regarding activities and environmental protection in Antarctic and were approved by the Ministry of Foreign Affairs and Trade of the Republic of Korea.

To obtain sufficient high-quality DNA molecules for the
PacBio Sequel platform (Pacific Biosciences, USA), one D. mawsoni fish was dissected and muscle tissue was used for DNA extraction using the phenol/chloroform extraction method. DNA quality was checked using a fragment analyzer system (Agilent Technologies, USA) and Qubit 2.0 fluorometer (Invitrogen, Life Technologies, USA). The single-molecule real-time (SMRT)bell library was sequenced using eight SMRT cells (Pacific Biosciences, Sequel™ SMRT Cell 1M v2) with a Sequel Sequencing Kit 2.1 (Pacific Biosciences, USA) and 1×600 min movies were captured for each SMRT cell using the Sequel sequencing platform (Menlo Park, USA). The average coverage of the SMRT sequences was ~81-fold (Supplementary Table S1).

Muscle tissue from the same sample was used to construct a Hi-C chromatin contact map for chromosome-level assembly. Tissue fixation, chromatin isolation, and library construction were performed according to the manufacturer’s instructions (Dovetail Genomics, USA) (Belton et al., 2012). After checking the insert size, concentration, and effective concentration of the constructed libraries, the final libraries were sequenced using the Illumina NovaSeq 6000 platform (San Diego, USA) with a 150-bp paired-end strategy. A total of 874 million raw reads were generated from the Hi-C libraries (Supplementary Table S1). The Iso-Seq sequencing data were analyzed using Sequel Sequencing Kit 2.1 (Pacific Biosciences, USA) and SMRTbell Template Prep Kit 1.0-TM (Invitrogen, Life Technologies, USA). The single-molecule real-time (SMRT)bell library was sequenced using eight SMRT cells (Pacific Biosciences, Sequel™ SMRT Cell 1M v2) with a Sequel Sequencing Kit 2.1 (Pacific Biosciences, USA) and 1×600 min movies were captured for each SMRT cell using the Sequel sequencing platform (Menlo Park, USA). The average coverage of the SMRT sequences was ~81-fold (Supplementary Table S1).

For transcriptome sequencing, we prepared 1 μg of pooled total RNA from the muscle and skin of D. mawsoni. Using a SMARTer PCR cDNA Synthesis Kit (Clontech, USA), RNA was synthesized to cDNA. The SMRTbell library was constructed using the SMRTbell™ Template Prep Kit 1.0-SPv3. The SMRTbell library was sequenced using SMRT cells (Pacific Biosciences, Sequel™ SMRT Cell 1M v2) and the Sequel Sequencing Kit 2.1. For each SMRT cell, 1×600 min movies were captured with the use of the Sequel sequencing platform; the pre-extension time was 240 min (Supplementary Table S1). The Iso-Seq sequencing data were analyzed using SMRT Link (v6.0.0).

For de novo genome assembly, the FALCON-Unzip assembler (v0.4, Falcon, RRID:SCR 016089) was used (Chin et al., 2013) with parameters of length_cutoff=12 000 and length_cutoff_pr=10 000 and with filtered subreads from SMRT Link (v4.0.0) (minimum subread length=50). To improve the quality of the genome assembly, the FALCON-Unzip assembler was polished using the Arrow algorithm with unaligned BAM files as raw data.

A draft D. mawsoni genome was previously generated using Illumina short-read sequencing (Chen et al., 2019). However, since several gaps prevailed in the draft genome set and there was no information about the linkage group, it was difficult to compare the structure of the Antarctic toothfish genome at the chromosomal scale. To improve this genome resource, long-read SMRT sequencing from Pacific Biosciences and Hi-C scaffolding were implemented. First, we performed de novo assembly of the long PacBio sequence reads using the FALCON-Unzip tool and obtained a genome assembly with a size of 924.75 Mb, an N50 contig size of 3.23 Mb, and longest contig size of 24.49 Mb (Supplementary Table S2). To construct the reference genome at the chromosome level, we constructed a Hi-C library and anchored the scaffolds into chromosomes after quality control using the HiC-Pro, Juicer (v1.5) (Durand et al., 2016) and 3D-DNA (v170123) pipeline (Dudchenko et al., 2017) based on the draft genome assembly (Figure 1A). The assembled genome was 926.3 Mb (GC content: 41.57%) in length with a scaffold N50 of 36.98 Mb and longest scaffold of 46.82 Mb. In total, there were 1 062 scaffolds in the D. mawsoni genome assembly, with 24 chromosome-scale scaffolds occupying 91.3% of the assembly (Supplementary Tables S2, S3 and Figure S1).

Benchmarking Universal Single-Copy Orthologs (BUSCO) v3.0 (Simão et al., 2015) (RRID:SCR 015008) was used along with the actinopterygiiodb9 database to assess the completeness of the new D. mawsoni genome assembly. Among 4 584 BUSCO groups searched, 4 197 and 194 BUSCO core genes were completely and partially identified, respectively, contributing to a total of 95.7% BUSCO genes in the D. mawsoni genome (Supplementary Table S4).

The diploid chromosome number (2n) of D. mawsoni is 48 (Ghigliotti et al., 2007). Comparison of its chromosome-scale assemblies with those of the Gasterosteus aculeatus genome (2n=42) showed a highly similar synteny (Figure 1B). However, each of three chromosomes (from Groups 1, 4, and 7) of G. aculeatus matched with two chromosomes in D. mawsoni (scaffolds 23 and 35, scaffolds 13 and 22, scaffolds 11 and 24, respectively) (Supplementary Figure S2).

A de novo repeat library was constructed using RepeatModeler (v1.0.3) (Bao & Eddy, 2002), which included RECON (RRID:SCR 006345) and RepeatScout v1.0.5 (RRID:SCR 014653) (Price et al., 2005) software with default parameters. The Tandem Repeats Finder (Benson, 1999) program was used to predict the consensus sequences and classification information for each repeat. We analyzed the repetitive sequences in the D. mawsoni genome, including those in the tandem repeats and transposable elements (TEs). TEs play an important role in shaping genome architecture and are a source of regulatory mutations in evolution. A difficulty arises in representing TEs in genome assemblies based on short Illumina sequence reads. Therefore, our long-read sequences greatly improved both the length and quantity of the TE repeats in the D. mawsoni genome assembly compared to the published short-read assembly. The D. mawsoni genome contained 40.87% of repeat sequences, including 36.27% (336 Mb) of TEs such as long terminal repeats (LTRs, 4.21%), short interspersed nuclear elements (SINES, 0.50%), long interspersed nuclear elements (LINEs, 5.35%), and DNA transposons (15.51%) (Supplementary Table S5 and Figure S3). Divergence of TEs was examined using RepeatMasker software, where Kimura distances (Kimura, 1980) estimated for aligned TEs (calcDivergenceFromAlign.pl) were used to draw repeat landscapes (createRepeatLandscpae.pl). The D. mawsoni genome had a higher number of recent TE insertions (Kimura, 2021).
divergence K-values(5) that were strongly shaped by DNA transposons (Supplementary Figure S4). Because K-values calculated for TEs can reflect age and transposition history (Chalopín et al., 2015), we concluded that there have been recent transposable element bursts in the Antarctic toothfish.

Genome annotation was conducted using MAKER v2.28 (RRID:SCR_005309) (Holt & Yandell, 2011), which is a portable and easily configurable genome annotation pipeline. Subsequently, repeat masked genomes were used for ab initio gene prediction with SNAP v2006–07-28 (SNAP, RRID:SCR 002127) (Korf, 2004) and Augustus (Augustus: Gene Prediction, RRID:SCR_008417) software. MAKER was initially run in the est2genome mode based on the Iso-Seq data, including 57,406 full-length transcripts. Additionally, protein evidence was obtained from the genomes of 19 teleosts, including three Antarctic fish (Supplementary Table S6). Exonerate software, which provides integrated information for the SNAP program, was used to polish MAKER alignments. MAKER was then used to select and revise the final gene model considering all available information. Other non-coding RNAs in the Antarctic toothfish assembly were identified using Infernal (v1.1) (Nawrocki & Eddy, 2013) and covariance models (CMs) from the Rfam database v12.1 (Rfam, RRID:SCR_007891) (Griffiths-Jones et al., 2005). Putative tRNA genes were identified using tRNAscan-SE v1.3.1 (IRNAScan-SE, RRID:SCR 010835) (Lowe & Eddy, 1997), which uses a CM that scores candidates based on their sequences and predicted secondary structures.

We identified orthologous gene clusters using the OrthoMCL (Li et al., 2003) pipeline, which applied the Markov Clustering Algorithm (MCL) with default options in all steps for the genome sequences of the 20 species (Supplementary Table S6). It was critical for analysis to include representative species of diverse phylogenetic clades and the 20 species were selected among those with well-annotated and well-assembled genomes.

Phylogenetic tree construction was performed based on single-copy orthologous genes. The sequences of protein-coding genes were aligned using a Probabilistic Alignment Kit (PRANK) (Löytynoja & Goldman, 2005) with the codon alignment option. The maximum-likelihood method was applied to construct a phylogenetic tree using RAxML with 1,000 bootstraps, and divergence times were calibrated with TimeTree (median estimates of pairwise divergence time for D. rerio and G. morhua: 230.4 million years ago) (Hedges et al., 2006).

Ortholog gene families from each species were identified: 7,731 orthologous gene families were commonly identified among the 20 teleosts, including 434 (1,431 genes) paralogous gene families that were D. mawsoni-specific (Supplementary Table S8). The phylogenetic tree of D. mawsoni and the 19 teleost species was constructed using 1,422 single-copy orthologs (Figure 1C). Among the 20 fish species, D. mawsoni and three other Antarctic fish were clustered together on the branch of a non-Antarctic fish, G. aculeatus, with a divergence time of around 105 million years ago. Furthermore, D. mawsoni diverged approximately 28 million years ago from the Antarctic fish Chaenocephalus aceratus. Analysis of gene gain-and-loss among genomes enables the reconciliation of a species tree with the gene tree for each family. Here, D. mawsoni had 659 (including 214 genes) significantly expanded and 116 (including 136 genes) significantly contracted gene families (Figures 1C, D). The vast majority of the expanded biological pathways belonged to two functional categories: (i) involved in nervous system functions (neuron projection development, GO:0031175; neuron development, GO:0048666: cell morphogenesis involved in neuron differentiation, GO:0048667; generation of neurones, GO:0048689; neuron projection morphogenesis, GO:0048812; axon development, GO:0061564) and (ii) cellular component morphogenesis (cell projection organization, GO:0030030; cell part morphogenesis, GO:0032990; cell projection morphogenesis, GO:0048858). In the molecular function category, peptidase regulator activity (endopeptidase regulator activity, GO:0061135; peptidase inhibitor activity, GO:0030414; endopeptidase inhibitor activity, GO:0004866), and signaling receptor binding (endopeptidase inhibitor activity, GO:0004866) were the major expanded pathways (Supplementary Tables S9, S10). In addition, 14,055 orthologous gene families containing 16,162 genes in D. mawsoni were commonly identified in the four Antarctic fish. Moreover, 621 gene families were D. mawsoni species-specific paralogs (Figure 1E) involved in DNA metabolic processes (DNA biosynthetic process, GO:0071897; DNA integration, GO:0015074; RNA-dependent DNA biosynthetic process, GO:0006278) (Supplementary Table S11).

Splicing variants were analyzed using SQANTI2 (Tardaguila et al., 2018) with the Iso-Seq data as full-length transcript
In the current study, we presented a high-quality chromosome-level genome assembly of the Antarctic toothfish, *D. mawsoni*, using PacBio Sequel sequencing and Hi-C chromatin contact maps. The *D. mawsoni* genome assembly (926 Mb) included 24 chromosomes that accounted for 91% (840 Mb) of all genome sequences. The *D. mawsoni* genome contained 32,914 protein-coding genes and 434 paralogous *D. mawsoni*-specific gene families among 20 teleost fish and 621 paralogous *D. mawsoni*-specific gene families among the four Antarctic teleost fish. This chromosome-length genome assembly will not only provide insights into the molecular and ecological adaptations of Antarctic fish to extreme environments but will also facilitate exploration of genomic adaptations to a wide range of evolutionary, ecological, metabolic, developmental, and biochemical features of Antarctic fish.

**DATA AVAILABILITY**

The *Dissostichus mawsoni* genome project was deposited in NCBI under BioProject No. PRJNA574770 and the Whole-Genome Shotgun project was deposited at DDBJ/ENA/GenBank under accession No. JAAKYF0100000000. The version described in this paper is JAAKYF0100000000. The genome browser, assembly, and annotation data are accessible on http://genome.kusglab.org/.

**SUPPLEMENTARY DATA**

Supplementary data to this article can be found online.

**COMPETING INTERESTS**

The authors declare that they have no competing interests.

**AUTHORS’ CONTRIBUTIONS**

H.P. and H.-W.K. conceived the study. S.J.L., E.J., S.-G.C., S.C. E.C., J.K., and H.P. performed genome sequencing, assembly, and annotation. E.J. and S.J.L. performed experiments. S.J.L., J.-H.K., H.-W.K., and H.P. mainly wrote the manuscript. All authors contributed to writing and editing the manuscript as well as collating the supplementary information and creating the figures. All authors read and approved the final version of the manuscript.

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