The complete mitochondrial genome of *Chionoecetes japonicus* (Crustacea: Decapoda: Majoidea)

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**ABSTRACT**

The complete mitochondrial genome of *Chionoecetes japonicus* was sequenced using a specimen collected offshore in the East Sea. The genome includes 13 protein-coding genes (PCGs), 22 transfer RNA (tRNA) genes, two ribosomal RNA (rRNA) genes, and a control region (D-loop), with a total length of 16,060 bp. The overall nucleotide composition was 34.91% A, 17.29% C, 10.93% G, and 36.87% T, with 71.78% A + T. In the phylogenetic tree was constructed using maximum-likelihood and Bayesian inference analyses, *C. japonicus* and *C. pacificus* formed a genetic clade that was sister to *C. opilio*.

Red snow crab, *Chionoecetes japonicus* Rathbun, 1932, is found in the East Sea. Unlike *C. opilio*, which is found at 200–800 m, it can be observed at 500–2300 m (Park et al. 2003). The genus *Chionoecetes* is a very important fisheries resource, and its ecology and fishing are active research topics. However, for *Chionoecetes*, because only the mitochondrial genomes of *C. pacificus* and *C. opilio* are currently known, additional mitochondrial genomes need to be discovered to elucidate the systematic relationships among taxa in the genus.

The specimen used in this study was collected offshore of Ganggu, Yeongdeok-gun, the Republic of Korea on 1 June 2020 using a fish trap, fixed in 99.9% ethanol, and stored in the specimen storage facility of Soonchunhyang University (Voucher no. SUC19351). Genomic DNA was extracted from walking leg tissue using a HiGene™ Genomic DNA Prep Kit (Biofact, Daejeon, Republic of Korea), and a qualified library was constructed by sequencing 2 × 150 bp paired-end reads on an MGISEQ-2000 platform (MGI Tech Co. Ltd, Shenzhen, China) to generate raw reads with a total of 8,831,783,100 bp (SRA accession no. SRX12462354).

The mitochondrial genome sequence was assembled using Geneious R11 software (Kearse et al. 2012) and a reference sequence (GenBank accession no. AB735678), by mapping reads against contigs with the Geneious mapper tool (settings: no gaps allowed, 3% maximum mismatch per read, word length = 40). Annotations were performed with MITOS WebServer (Bernt et al. 2013) and corrected manually.

The mitochondrial genome of *C. japonicus* (GenBank accession no. MT750295) is 16,060 bp long and includes 13 protein-coding genes (PCGs), 22 transfer RNA (tRNA) genes, two ribosomal RNA (rRNA) genes, and a control region (D-loop). Four PCGs (nd5, nd4, nd4L, and nd1), eight tRNAs (tRNA^Met^, tRNA^Thr^, tRNA^Pro^, tRNA^Leu^, tRNA^Val^, tRNA^Glu^, tRNA^Gly^, and tRNA^Trp^), and two rRNAs (12S and 16S rRNA) are encoded on the light strand. The overall base composition is 34.91% A, 17.29% C, 10.93% G, and 36.87% T, with 71.78% A + T, which is similar to the base content and AT bias of mitochondrial genomes in other Crustacea (Xing et al. 2017; Lu et al. 2020; Wang et al. 2020).

Of the 13 PCGs, the start codon is ATG in six, ATT in three, and GTG in two; the start codons of the nd4L and nd6 genes are ATA and ATC, respectively. Three of the PCGs terminate with incomplete stop codons, T (nd5, nd4, and nd1), and the remaining 10 end with complete stop codons (TAA or TAG).

In addition, as in previous studies, all genes located in the β-strand in the *Maja* species compared here formed a single block located between tRNA^Glu^ and a control region. By contrast, in the genus *Chionoecetes*, the arrangements of the general mitochondrial genes and tRNAs were similar to those in Brachyura, such as in the genus *Damithrax* (Basso et al. 2017; Jeong et al. 2020).

All 13 PCGs on each mitochondrial genome used in the phylogenetic analysis were downloaded from the National Center for Biotechnology Information and subjected to an analysis based on the alignment results obtained with MAFFT 7.450 (Katoh et al. 2002; Katoh and Standley 2013). GTRGAMMA was found to be the optimal model based on the corrected Akaike information criterion (AICc) using jModelTest 2.1.10 (Guindon and Gascuel, 2003; Darriba et al., 2012). The maximum-likelihood (ML) tree was constructed with 1,000 bootstrap replications using PhyML 3.0.

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(Guindon et al. 2010), and the Bayesian inference (BI) tree was run for 1,000,000 generations using MrBayes 3.2.7 (Ronquist et al. 2012). In addition, two *Maja* species belonging to the superfamily Majoidea and one *Damithrax* species were used as the outgroup. The phylogenetic tree was constructed based on ML and BI analyses (Figure 1).

In the phylogenetic tree, two mitochondrial genomes (GenBank accession no. AB735378 and MT750295) from *C. japonicus* formed a clade sister to *C. opilio* (Figure 1). Similar results were observed for the co1 gene in Azuma et al. (2011). Altogether, these and previous findings support the current taxonomic system.

The basic data on the complete mitochondrial genome of *C. japonicus* provided in this study will be an important resource for population genetic analysis, and will also be helpful for molecular phylogenetic studies of other species within the genus *Chionoecetes* that have not yet been discovered.

**Disclosure statement**

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

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**Data availability statement**

The data that support the findings of this study are openly available in GenBank of NCBI at https://www.ncbi.nlm.nih.gov, reference number MT750295.

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