Characterization of the complete mitochondrial genome of *Okenia hiroi* (Baba, 1938) (Nudibranchia, Goniodorididae)

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**ABSTRACT**

*Okenia* is a speciose genus of the family Goniodorididae with more than 50 valid species. The phylogenetic relationships within the genus are little known. The mitogenome is a good marker to understand the phylogenetic relationships of relative species. This study was performed to sequence the mitogenome of *O. hiroi*. The mitogenome of *O. hiroi* was 14,583 bp in size and was composed of 37 genes, including 13 protein-coding genes, two ribosomal *rRNA* genes, and 22 *tRNA* genes. The nucleotide composition was 30.5% A, 13.6% C, 16.5% G, and 39.4% T. The phylogenetic analysis showed that *O. hiroi* is sister to *Notodoris gardineri* (Aegiridae). This study recorded the first mitochondrial genome sequence of the family Goniodoridae.

The genus *Okenia* is composed of small to medium-sized nudibranchs (Pola et al. 2019). Because of its diversity, many new species of the genus *Okenia* have been recently described (Gosliner 2004; Pola et al. 2019). Currently, the genus includes more than 50 valid species that are distributed from the Pacific Ocean to the north and south Atlantic Ocean (Pola et al. 2019; Horton et al. 2021). In addition, there are limited studies on the phylogeny of the genus (Pola et al. 2019). *Okenia hiroi* is a species of the genus *Okenia*. This species was described from Japan and reported from Korea and Hong Kong (Jung et al. 2014). The analyses of the mitogenome genes of *O. hiroi* might be useful for understanding molecular identification and phylogeny of the genus. Therefore, this study aims at sequencing and analyzing structure of the mitogenome of *O. hiroi*.

The specimen of *O. hiroi* was collected from Munamjin-ri, Korea (38°18′14.75″N, 128°34′1.05″) in June 2020. Following the collection, the specimen (voucher number: SMU00087) was preserved in absolute ethanol and deposited at the Department of Biotechnology, Sangmyung University, Korea. Total DNA was extracted from the foot of the specimen. The extracted DNA was checked for puriyand concentration using a MaestroNano spectrophotometer (Maestrogen, Hsinchu, Taiwan). After library preparation, mitogenome sequencing was performed with MGISEQ-2000 (MGI, Shenzhen, China). MITObim (Hahn et al. 2013) was used for sequence assembly and MITOS (Bernt et al. 2013) was used for sequence annotation. Mitogenome sequences representing different families of the suborder Doridina and two species of the suborder Cladobranchia available in GenBank were obtained for phylogenetic analysis. All 37 genes of the mitogenomes were used for tree reconstruction. The partition schemes and best-fit model were searched in PartitionFinder 2 (Lanfear et al. 2017). MrBayes version 3.2.7a was applied to investigate the phylogenetic position of *O. hiroi* (Ronquist et al. 2012). For analysis, 10,000,000 generations were run and sampling was conducted every 1,000 generations.

The mitogenome of *O. hiroi* (GeneBank accession number: MW408699) was 14,583 bp long and contained 13 protein-coding genes, two *rRNA* genes, 22 *tRNA* genes. The nucleotide composition of the mitogenome was 30.5% A, 13.6% C, 16.5% G, and 39.4% T, with an A + T content of 69.9%. The size of 13 PCGs ranged from 156 bp (*atp8*) to 1,623 bp (*nd5*). Of 13 PCGs, there were nine genes (*cox1*, *cox2*, *cytb*, *nad1*, *nd2*, *nad4*, *nad4l*, *nd5*, and *nd6*) encoded on the H-strand and four remaining genes (*atp6*, *atp8*, *cox3*, and *nd3*) encoded on the L-strand. The mitogenome contained two *rRNA* genes with the size of 747 bp and 1,103 for 12S *rRNA* and 16S *rRNA*, respectively. Of the 22 *tRNA* genes, *tRNA*-Ser was the shortest gene (58 bp) and *tRNA*-Asn and *tRNA*-Asp were the longest genes (69 bp).

There were 27 intergenic regions in the mitogenome of *O. hiroi*, ranging from 1 to 108 bp. The longest intergenic region (108 bp) was located between *cox1* and *tRNA*-Val genes. Compared to the other available mitogenomes of nudibranchs, this is the longest intergenic region between *cox1* and *tRNA*-Val genes so far recorded. Also, there were five overlapping regions in the mitogenome of *O. hiroi*. The longest overlapping region (20 bp) was located over *nd1* and *nd5* genes.

Figure 1 shows the phylogenetic position of *O. hiroi* in the suborder Doridina. As seen from Figure 1, all species of Doridina were clustered together. The phylogenetic tree indicated that *O. hiroi* formed a clade with *Notodoris gardineri* (Aegiridae). In term of systematics, both Goniodoridae and Aegiridae belong to the super family Onchodoridae...
Previous study also revealed a close relationship between Goniodorididae and Aegiridae (Hallas et al. 2017). In the future study, additional mitogenomes of the family Goniodorididae should be generated to investigate the relationships within the family.

Disclosure statement
No potential conflict of interest was reported by the authors.

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Data availability statement
The genome sequence data that support the findings of this study is openly available in GenBank of NCBI at (https://www.ncbi.nlm.nih.gov/) under the accession no. MW408699. The associated BioProject, SRA, and Bio-Sample numbers are PRJNA693411, SRR13555242, and SAMN17386532, respectively.

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