Complete mitochondrial genomes of two water mite species: Hygrobates (H.) longiporus and Hygrobates (rivobates) taniguchii (Acari, Trombidiformes, Hygrobatoidea)

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ABSTRACT

We determined the complete mitochondrial genome sequences of two water mites, Hygrobates (Hygrobates) longiporus and H. (Rivobates) taniguchii. The length of the entire mitogenome of these two species is 13,721 bp and 13,770 bp long, respectively. Both of them contain 13 proteins, two rRNAs, and 22 tRNAs for a total of 37 gene products. The gene order of Hygrobates is almost identical to the two species of Unionicola included in the analysis, with some difference in the position of transfer RNA genes. Phylogenetic analyses highly support these Hygrobates species form a clade with other species of the Hygrobatoidea.

Diverse assemblages of water mites are present in almost all freshwater environments around the world except Antarctica. The diversity of the water mites is now rapidly becoming apparent, with over 10,000 species estimated to be present (Di Sabatino et al. 2008), while approximately 7,500 species have been described so far (Smit in press). Recently as well as molecular research on water mites greatly developed; molecular studies showed the monophyly of the Hydrachnidia as well as all but one of the traditional superfamilies (Dabert et al. 2016). In several studies, cryptic species, as well as species complexes, have been detected (Martin et al. 2010; Pešić & Smit 2016, 2017, 2018; Pešić et al. 2017, 2019; Blattner et al. 2019). Water mites reach great abundance and diversity in several freshwater environments and are associated with a variety of macrofauna—adults and deutonymphs of most species are predators mainly on microcrustaceans and aquatic insect eggs and larvae (one group has been found to parasitize amphibians (Goldschmidt et al. 2020)) but the larvae are parasites of aquatic insects. Therefore, it has been noted that they are excellent bioindicators (Goldschmidt 2016). On the contrary, their diversity is greatly underestimated, suggesting the presence of multiple cryptic species within many morphological species (Pešić et al. 2017; Blattner et al. 2019).

Thus, more genetic information is urgent to elucidate phylogenetic relationships and genetic structure for revising taxonomy and species diversity in water mites. However, there was no mitogenome record for the family Hygrobatidae, one of the major families in water mites. So, we choose two Hygrobates species for the representative of the family and determined the whole mitogenome sequences by shotgun sequencing for both species.

Total DNA was extracted using DNeasy Blood & Tissue Kit (QIAGEN, Hilden, Germany) and processed by Qiaseq FX DNA Library kit (QIAGEN). Paired-end sequencing (300 cycles) was conducted by National Museum of Nature and Science, Tokyo on MiSeq, with inserts of ca. 50–200 bp, for a total of ca. four million reads. Assembly was performed using CLC Genomics Workbench ver. 12 (QIAGEN) with default setting. Ambiguous part of the contig was proofread by 3500 xL Genetic Analyzer (Thermo Fisher Scientific, Waltham, MA, USA). Gene identification was done using MITOS web server (Bernt et al. 2013). Voucher specimens with extracted DNA were deposited to National Museum of Nature and Science, Tokyo (NSMT-DNA 49380 and 49381).

The entire mitogenome of Hygrobates (Hygrobates) longiporus Thor, 1898 (GenBank/DDBJ/EMBL accession number LC552026) and Hygrobates (Rivobates) taniguchii Imamura, 1954 (LC552027) is 13,721 bp and 13,770 bp long, respectively. Both of them contain 13 proteins, two rRNAs, and 22 tRNAs for a total of 37 gene products. The overall A+T content of the H. (H.) longiporus and H. (R.) taniguchii
mitochondrial genome is 72.5 and 67.9%, respectively, which is in ordinal range among Parasitengona species (67.5–73.6%). In the mitogenomes of both species, ND5 starts with CTT codon and stops with incomplete termination codon T. The gene order of *Hygrobates* is almost identical to the species of genus *Unionicola*, with some difference in the position of transfer RNA genes.

The maximum-likelihood phylogenetic analysis (ML) based on 13 protein coding genes was conducted by RAxML-NG ver.0.9.0 (Kozlov et al. 2019) with bootstrap analyses of 1,000 replicates. The phylogenetic tree also with posterior probability from Bayesian analyses (BA) conducted by MrBayes 3.2.6 (Ronquist et al. 2012). *H. (H.) longiporus* and *H. (R.) taniguchii* made a sister clade with species belonging to superfamly *Hydrobatidea* (*Unionicola parkeri* and *U. foili*) with high nodal support value (Figure 1). These mitogenomes would be useful for reconstructing higher systematics of Hydrachnid mites.

**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 the National Center for Biotechnology Information database (NCBI/GenBank) at https://www.ncbi.nlm.nih.gov/, accession numbers LC552026 and LC552027.

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