Complete mitochondrial genome sequences were determined for five species of Australian freshwater fishes, representing a diverse range of ecologies and life histories. Mitogenomes were sequenced and annotated for *Craterocephalus stramineus* (Atherinidae); *Hypseleotris klunzingeri* (Eleotridae); *Lovettia sealii* (Galaxiidae); *Leiopotherapon uniclor* (Terapontidae) and *Nematalosa erebi* (Clupeidae). The five new mitogenomes each share the typical vertebrate mitochondrial arrangement of 13 protein coding genes, 22 tRNA genes, two rRNA genes and a control region. These sequences will be a useful resource for studies of evolutionary relationships and for management applications.

**Introduction**

The mitochondrial genome (mitogenome) is a small molecule (16–17,000 bp) that is present in most eukaryotic organisms. Vertebrate mitogenomes feature highly conserved gene composition and gene order (Satoh et al. 2016). Complete mitochondrial sequences are now available for >2,000 species of ray-finned fishes, and this growing resource plays a key role in evolutionary studies and in environmental management applications such as environmental DNA (eDNA) and metabarcoding (Sato et al. 2018). The Australian freshwater fish fauna, with just over 250 species, is relatively depauperate in species-level diversity, but highly endemic (Unmack 2001). Relatively few Australian freshwater fishes have mitogenome sequences available, although recent studies have demonstrated the utility of fully characterized mitogenomes for analyses of selection and population structure in high profile Australian freshwater fishes (e.g. Harrisson et al. 2016; Bishop et al. 2018). The aim of this study was to assemble and annotate the first complete mitochondrial genomes for several ecologically significant Australian freshwater fish species. The selected taxa have wide distributions across multiple drainage basins, and exhibit a diverse range of ecological characteristics. Tasmanian whitebait (*Lovettia sealii*, Galaxiidae) has a temperate distribution and a semi-anadromous life history (Schmidt et al. 2014). Strawman (*Craterocephalus stramineus*, Atherinidae) occurs in tropical drainages of northern Australia and is a habitat specialist (Mondol 2016). Spangled perch (*Leiopotherapon uniclor*, Terapontidae) is Australia’s most widely distributed freshwater fish, capable of extreme long-distance dispersal (Schmidt et al. 2018b). The western carp gudgeon (*Hypseleotris klunzingeri*, Eleotridae) is an abundant and widely distributed species which does not engage in interspecific hybridisation unlike other taxa in the carp gudgeon species complex (Schmidt et al. 2011). The distribution of bony bream (*Nematalosa erebi*) includes dryland rivers of central Australia and this species has broad temperature tolerance and high dispersal capability (Hughes and Hillyer 2006).

**Methods**

Low-coverage, whole-genome shotgun libraries were prepared for each species using randomly-sheared genomic DNA. A TruSeq (Illumina, San Diego, CA) library preparation kit was used with a targeted insert size of 500 bp (see Schmidt et al. 2018a). Paired-end sequencing was performed on the Illumina MiSeq Sequencer at Australian Genomics Research Facility (AGRF), using a 600 cycle MiSeq reagent kit v3. Novoplasty v2.6.5 was used to assemble the pair-end reads generated for each species into a single circular contig (Dierckxsens et al. 2017). Mitogenome annotation was achieved using MitoFish (Iwasaki et al. 2013). Tissue samples used for sequencing were derived from previous genetic studies and relevant specimen voucher codes and associated publication details are provided below.

**Results and discussion**

Lovettia sealii (GenBank accession: MK029347). DNA was obtained from sample code 4021 collected from the Huon River, Tasmania (Lat. Long. –42.998483, 146.927883) (Schmidt et al. 2014). The seed sequence used to initiate assembly was *Galaxias maculatus* (NCBI RefSeq: NC_004594). A total of 3.8 × 10^6 reads were generated. Of these 4022 were incorporated into the new mitogenome assembly producing a 16,542 bp circular contig at 98% coverage.
**Craterocephalus stramineus** (GenBank accession: MK029350). DNA was obtained from specimen code DMC-CS8L-13 collected from the Daly River, Northern Territory (Lat. Long. /C0 13.8092666, 131.341133) (Mondol 2016). The seed sequence used to initiate assembly was **Craterocephalus eyresii** (NCBI RefSeq: NC_035148). A total of 4.4/10⁶ reads were generated. Of these 3,240 were incorporated into the new mitogenome assembly producing a 16,566 bp circular contig at 72x coverage.

**Leiopotherapon unicolor** (GenBank accession: MK024340). DNA was obtained from specimen code 11_04 collected from the Daly River, Northern Territory (Lat. Long. -14.3638, 131.557) (Schmidt et al. 2018a). The seed sequence used to initiate assembly was **Scortum barcoo** (NCBI RefSeq: NC_027171). A total of 3.4/10⁶ reads were generated. Of these 996 were incorporated into the new mitogenome assembly producing a 16,527 bp circular contig at 20x coverage.

**Hypseleotris klunzingeri** (GenBank accession: MK029348). DNA was obtained from specimen code BR28 collected from the Balonne River at St George, Queensland (Lat. Long. /C0 28.012917, 148.614753). The seed sequence used to initiate assembly was **Hypseleotris** sp. HAxB (GenBank: KT716513) (Schmidt 2016). A total of 4.6/10⁶ reads were generated. Of these 1710 were incorporated into the new mitogenome assembly producing a 16,508 bp circular contig at 37x coverage.

**Nematalosa erebi** (GenBank accession: MK029349). DNA was obtained from specimen code TM1 collected from the Bulloo River, Thargomindah, Queensland (Lat. Long. /C0 28.017222, 143.785556) (Hughes and Hillyer 2006). The seed sequence used to initiate assembly was **Nematalosa nasus** (NCBI RefSeq: NC_023824). A total of 3.8/10⁶ reads were generated. Of these 1,116 were incorporated into the new mitogenome assembly producing a 16,665 bp circular contig at 23x coverage.

![Phylogenetic relationship of five new mitogenomes along with the single closest match for each one derived from BLASTn search. Tip labels include GenBank accession and species name. New mitogenomes highlighted in bold font. Alignment of mitogenomes (excluding 16S, 12S and control region) was performed using MAFFT v7.017 (Katoh et al. 2002). A maximum-likelihood phylogenetic analysis was performed on the final alignment of 11,500 bp with RAxML v8.2.11 using the GTR + GAMMA substitution model (Stamatakis 2006).](image)

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The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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