Phylogenetic analyses to uncover the evolutionary relationship of a newly sequenced mitochondrial genome from an Eastern spinebill (Acanthorhynchus tenuirostris)

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\section*{ABSTRACT}

The Eastern spinebill (Acanthorhynchus tenuirostris), a passerine bird in the family Meliphagidae (honeyeaters), a dominant group of birds in Australia and New Guinea. The aim of this study was to sequence the complete mitochondrial genome of the Eastern spinebill and use its sequence to better define the phylogeny of this species. The complete mitogenome sequence of A. tenuirostris was circular and 16,614 bp in length, and its architecture was conserved in comparison to other mitogenome sequences under the family Meliphagidae. The Eastern spinebill mitogenome had the highest sequence identity with mitogenome sequences of two other honeyeaters, the white eared honeyeater, Nesoptilotis leucotis, (84.9\%) and the white-plumed honeyeater, Ptilotula penicillata (85.5\%). The maximum-likelihood topology distinctly discriminated the Eastern spinebill sequence against all other species of the Meliphagidae with significant bootstrap supports. We suggest the widespread sampling and complete mitogenome sequencing would be valuable in establishing the most accurate phylogenetic taxonomy of the family Meliphagidae.

\section*{1. Introduction}

The family Meliphagidae (honeyeaters) is species-rich (38 genera and 180 species), morphologically diverse, and widely distributed group across Australia, New Guinea, and oceanic islands across Wallacea and the Pacific (Driskell and Christidis 2004; Andersen et al. 2019). In certain habitats, more than 12 species of honeyeater can co-occur seasonally (Keast 1985). They can be either nectarivorous, insectivorous, frugivorous, or more commonly, a combination of nectar- and insect-eating (Lea and Gray 1936).

Despite ecological and evolutionarily significance of the Meliphagidae the phylogenetic relationships for some species within this large family remain unclear (Andersen et al. 2019). The Eastern spinebill (Acanthorhynchus tenuirostris) is a honeyeater found in the woodlands of Eastern, Southeast and Tasmania Australia. Using a nuclear gene and selected mitochondrial genes, the Eastern spinebill and the Western spinebill (Acanthorhynchus superciliosus) have been found to form a sister clade to the remainder of the Meliphagidae (Driskell and Christidis 2004; Gardner et al. 2010; Andersen et al. 2014). A recent study on genus-level phylogeny of the honeyeaters using 4397 ultraconserved elements from 57 species demonstrated that Acanthorhynchus spinebill was sister to all remaining clades (A–H), but a completely resolved phylogeny of honeyeaters based on genome-wide data remains elusive (Andersen et al. 2019). Therefore, we believe that the complete mitogenome could play a significant role to provide more clear evolutionary relationships, the divergence time of speciation and influencing management decisions of species.

\section*{2. Materials and methods}

\subsection*{2.1. Source of sampling and extraction of DNA}

Droppings were collected from a holding bag of an Eastern spinebill caught in a mist net as part of a fauna survey in the Windsor Downs Reserve Nature (33\(^\circ\)39\'0.425", 150\(^\circ\)48\'31.97"E), New South Wales (Australian Bird and Bat Banding Scheme (ABBBS) Banding Authority Number 1893, ABBBS project approval—cooperative project 8529, New South Wales National Parks and Wildlife Service—Scientific License No. SL101929). The total genomic DNA was extracted using a commercial kit (Purelink\textsuperscript{\textsuperscript{TM}} Genomic DNA Mini Kit, Invitrogen, California, CA, USA) according to the manufacturer's instructions.
2.2. Library construction and sequencing

The library preparation and sequencing were performed as previously described (Sarker et al. 2017; Sarker et al. 2019a; Sarker et al. 2019b). Briefly, the paired-end library with an insert size of 150 bp was prepared using the QIAseq FX DNA Library Kit (Qiagen) starting with ten ng of total genomic DNA (gDNA). The amplified library was cleaned to remove PCR-generated adaptor-dimers using JetSeq™ Clean beads (Bionicle) according to the protocol described in QIAseq FX DNA Library Kits. The quality and quantity of the prepared library was assessed using an Agilent Tape Station (Agilent Technologies) by the Genomic Platform, La Trobe University. The prepared library was normalized and pooled in equimolar quantities. Cluster generation and sequencing of the pooled DNA-library were sequenced on Illumina® NextSeq500 platform according to the manufacturer’s instructions (Australian Genome Research Facility, Melbourne).

2.3. Assembly protocol

Sequencing data used in this study were analyzed according to a previously established pipeline (Sarker et al. 2017; Sarker et al. 2019a; Sarker et al. 2019b) using Geneious (version 10.2.2, Biomatters, New Zealand) and CLC Genomics Workbench (version 9.5.4). Briefly, a total of 11.7 million reads with a read length of 150 bp was used to obtain the complete mitochondrial genome of Eastern spinebill. Preliminary quality evaluation for all raw reads was generated, pre-processed to remove ambiguous base calls and poor-quality reads, and trimmed to remove the Qiagen Universal adapter sequences. The trimmed reads that passed the quality control based on PHRED score were used as input data for de novo assembly using SPAdes assembler (version 3.10.0) (Bankevich et al. 2012) in Geneious. Annotation was performed using default parameter under the genetic code of vertebrate mitochondrial (transl_table 2) in Geneious (version 10.2.2).

2.4. Phylogenetic analysis

Phylogenetic analyses were performed using the mitogenome sequence of Eastern spinebill determined in this study with other selected mitogenome sequences belong to the order Passeriformes in the GenBank database. Mitogenome sequences that showed more than 82% similarity (based on Blastn) with newly sequenced mitogenome of the Eastern spinebill were included to construct genome level phylogenetic tree. Nucleotide sequences of partial cytochrome B gene were selected from the family Meliphagidae and used to align with MAFFT L-INS-I algorithm implemented in Geneious (version 7.388) (Katoh and Standley 2013). Phylogenetic analyses were performed under the GTR substitution model with 1000 bootstrap support in Geneious (version 10.2.2) and CLC Genomics Workbench (version 9.5.4). The calculations of sequence similarity percentages were performed using tools available in Geneious (version 10.2.2).

3. Results

3.1. Eastern spinebill mitogenome structure

The overall genome architecture of the mitogenome sequence of Eastern spinebill was mostly conserved in comparison to other mitogenome sequences under the family Meliphagidae.

It had a circular genome of 16,614 bp, containing 13 methionine-initiated protein-coding genes (PCGs), two rRNA genes, 22 tRNA genes, and a control region (D-loop). The percentage of A, T, C, and G were 29.2%, 23.9%, 31.9%, and 15.1%, respectively. The AT and GC content of this complete mitogenome was 53.0% and 47.0%, respectively. The proportion of coding sequences with a total length of 11,405 bp (68.65%), which encodes more than 3801 amino acids, and all protein-coding genes started with Met. The lengths of 12S and 16S ribosomal RNA were 982 bp and 1605 bp, respectively.

3.2. Evolutionary relationship of mitogenome of the Eastern spinebill to other passerine species

To understand the evolutionary relationship of the newly sequenced mitogenome of the Eastern spinebill to other passerine species, phylogenetic analysis was conducted with the inclusion of other selected complete mitogenome sequences under the order Passeriformes. In the resulting maximum-likelihood (ML) tree, mitogenome of the Eastern spinebill was placed in a well-separated monophyletic clade between other five representative species of the order Passeriformes (bootstrap support 100%) (Figure 1). Based on this analysis the Eastern spinebill is the sister taxon of white-eared honeyeater (Nesoptilotis leucotis) (GenBank accession numbers KY994583.1 and KY994594.1) found in Australia. In agreement with phylogenetic relationships, the mitogenome of the Eastern spinebill sequence demonstrated much higher sequence identities with mitogenome sequences of white-eared honeyeater (Nesoptilotis leucotis) and white-plumed honeyeater (Ptilotula penicillate) (84.9% and 85.5%, respectively).

By also building phylogenetic trees using selected partial nucleotide sequences of the cytochrome B gene (Figure 2) from Meliphagidae family, we observed that the Eastern spinebill was placed in a sister clade with olive straightbill (Timelipsis fulvigula) (GenBank accession no. AY488387) and western spinebill (Acanthorhynchus superciliosus) (GenBank accession no. AY488331), which is consistent with the previous studies that examined the molecular phylogenetic relationship among Australian passerines birds species using partial mitochondrial and nuclear genome sequences (Dirskell and Christidis 2004; Gardner et al. 2010; Andersen et al. 2014).

4. Discussion

This paper describes the characterization and phylogenetic relationship of a novel mitochondrial genome sequenced from an Eastern spinebill. Phylogenetic analysis using the
mitogenome of the Eastern spinebill positioned it among
the other five representative species of the order
Passeriformes, which has shown strong bootstrap
support (100%) (Figure 1), but the Eastern spinebill
remains sister to the representative
clade. As shown in Figure 1, it is evident that the mitoge-
nome of *A. tenuirostris* is most closely related to *Nesoptilotis
leucotis* (84.9%), inferring that these species likely share
the common evolutionary history. A well-supported phylogenetic

**Figure 1.** Maximum-likelihood phylogenetic tree to infer host-phylogeny relationship using complete mitochondrial genome sequenced from *A. tenuirostris* along with other selected species under the order of *Passeriformes*. The new complete mitogenome of *A. tenuirostris* was highlighted by bold font.
tree was also constructed using partial nucleotide sequences of cytochrome B gene, and they showed that the Eastern spinebill was more closely related with other two species, olive straightbill and western spinebill within the family Meliphagidae and was placed in the same subclade (Figure 2), which is consistent with the previous findings (Driskell and Christidis 2004; Gardner et al. 2010; Andersen et al. 2014).

The spinebills, Acanthorhynchus, which are perhaps the most obviously specialized nectarivores in the family Meliphagidae and inhabit dry heathland and woodlands (Driskell and Christidis 2004), have no complete mitochondrial genome from their close relatives. Therefore, the well-defined phylogenetic relationship of the species is equivocal. Our findings demonstrated that A. tenuirostris species appear to be sister to remaining Meliphagidae species such as white-eared honeyeater (Nesoptilosis leucotis), white-fronted chat (Epthianura albifrons), tui (Prosthemadera novaeseelandiae), noisy miner (Manorina melanocephala) and white-plumed honeyeater (Ptilotula penicillate) (Figure 1). There were many key nodes resolved in the recent phylogenetic trees using unconserved elements of about 30% of the honeyeater species (Andersen et al. 2019); however, ample work remains to be done before achieving a solid

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**Figure 2.** Maximum-likelihood phylogenetic tree to infer host-phylogeny relationship using partial cytochrome B gene sequenced from A. tenuirostris along with other selected species under the family of Meliphagidae. The A. tenuirostris species was highlighted by black box. The tree is displayed as circular phylogram and the labels are aligned. Bootstraps support higher 80% are highlighted by line width.
phylogenetic framework of the honeyeaters. Particularly, it will be important to make availability of the genome-level data sets for the comparative studies of the morphological, behavioral and biogeographical diversification of the honeyeaters.

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Data availability statement
Raw sequencing datasets that support the findings of this study are accessible through Mendeley Data repository via the following Link (http://dx.doi.org/10.17632/57zhllhs7xz.1) (Sarker 2021).

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