Data in Brief 5 (2015) 281-284

Contents lists available at ScienceDirect

Data in Brief

journal homepage: www.elsevier.com/locate/dib



Data Article

Mitochondrial and nuclear phylogenetic trees and divergence time estimations of Sulawesi endemic Adrianichthyidae



Daniel F. Mokodongan, Kazunori Yamahira

Tropical Biosphere Research Center, University of the Ryukyus, Okinawa 903-0213, Japan

ARTICLE INFO

Article history: Received 29 July 2015 Received in revised form 15 August 2015 Accepted 25 August 2015 Available online 3 September 2015

ABSTRACT

This data article is related to the research article entitled "Origin and intra-island diversification of Sulawesi endemic Adrianichthyidae" by Mokodongan and Yamahira [1]. In this data article, we present phylogenetic trees of Sulawesi adrianichthyids separately reconstructed using mitochondrial (cytochrome *b*: cyt *b* and NADH dehydrogenase subunit 2: ND2) and nuclear (*tyrosinase*) sequences. We also present Bayesian chronograms of Sulawesi adrianichthyids separately estimated using a substitution rate for cyt *b* and for ND2. © 2015 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY license

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Specifications table

Subject areaBiology, Genetics and GenomicsMore specificPhylogenetics and phylogenomicssubject areaType of dataType of dataPhylogenetic trees, chronogramsHow data wasPhylogenies were acquired using maximum likelihood and Bayesian inference methods.acquiredChronograms were acquired using Bayesian inference methodsData formatAnalyzed

DOI of original article: http://dx.doi.org/10.1016/j.ympev.2015.07.024 *E-mail address:* yamahira@lab.u-ryukyu.ac.jp (K. Yamahira).

http://dx.doi.org/10.1016/j.dib.2015.08.032

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	Experimental factors	Phylogenies were estimated by raxMLGUI and MrBayes using mitochondrial (cyt <i>b</i> and ND2) and nuclear (<i>tyr</i>) sequences. Bayesian chronograms were estimated by BEAST using two different molecular clocks.
	Experimental features	Codon-specific $GTR+I+G$ models and a rapid bootstrap analysis of 1000 bootstrap replicates were used in raxmlGUI. MrBayes was run using ngen=3000000, samplefreq=100, and burnin=30000. BEAST analyses were performed using ngen=50000000, samplefreq=1000, burnin=12500000, and a substitution rate of 2.65% and 2.8% per My for cyt <i>b</i> and ND2, respectively.
	Data source location	n/a
	Data accessibility	With this article

Value of the data

- Data on phylogenies separately estimated using mitochondrial and nuclear sequences enable researchers to examine how the topologies differ from each other.
- Data on phylogenies of Sulawesi adrianichthyids enable researchers to compare their diversification history with those of other Sulawesi taxa.
- Data on two alternative chronograms enable researchers to infer the possible ranges of time frames in the divergence events of Sulawesi taxa.

1. Data, experimental design, materials and methods

1.1. Phylogenetic trees of Sulawesi adrianichthyids

The data presented here represent the phylogenetic trees of Sulawesi adrianichthyids separately reconstructed using mitochondrial (cytochrome b: cyt b and NADH dehydrogenase subunit 2: ND2) and nuclear (tyrosinase: tyr) sequences (Fig. 1A and B). The phylogenetic tree using concatenated sequences of the mitochondrial and nuclear sequences was presented in [1]. The data of the mitochondrial, nuclear, and combined phylogenetic trees in the Newick format are available as Supplementary data 1–3, respectively.

All of the sequence data used in this data article have been deposited into DNA Data Bank of Japan (DDBJ) under accession numbers LC51634-LC51792 for adrianichthyids and AP002932, AB569470, AB188743, and AB188744 for the two beloniform outgroups. They were aligned using ClustalW version 1.4 [2], and the alignment was later corrected by eye. The aligned sequences of the mitochondrial genes (cyt b: 1141 bp and ND2: 1046 bp), were concatenated into a single sequence (2187 bp). For the aligned sequence set of the nuclear gene (tyr), any site in intron 1 where a gap was found in any of the populations/species was removed, which resulted in a 1253-bp sequence (exon 1 = 809 bp, intron 1 = 269 bp, and exon 2 = 175 bp). Heterozygous sites in the tyr sequences (R, Y, M, K, W, and S) were replaced with N. The aligned mitochondrial and nuclear sequences were separately analyzed. Thereafter, the sequences were combined into a single 3440-bp concatenated alignment, and analyses were performed on the combined sequences.

Phylogenies were estimated by maximum likelihood (ML) and Bayesian inference (BI) methods. ML analyses were performed by raxmIGUI version 1.31 [3] using codon-specific GTR+I+G models for each sequence set (however, codon positions were not partitioned for tyr intron 1), where a rapid bootstrap analysis of 1000 bootstrap replicates was conducted. BI analyses were conducted with MrBayes version 3.2.4 [4]. Appropriate substitution models were separately determined for cyt b, ND2, tyr exon 1, and tyr exon 2 codon positions using jModelTest version 2.1.7 [5] based on the Akaike Information Criterion (AIC) (Table 2 in [1]). Similarly, the GTR+G model was chosen for tyr intron 1 without separating codon positions. Analyses were run using the following settings: ngen=3000000, samplefreq=100, and burnin=30000.



Fig. 1. Bayesian inference phylogenies of Sulawesi adrianichthyids based on (A) mitochondrial sequences (cyt *b*: 1141 bp and ND2: 1046 bp) and (B) nuclear sequences (*tyr*: 1253 bp). Numbers on branches represent Bayesian posterior probabilities (top) and maximum likelihood bootstrap values (bottom). The scale bar indicates the number of substitutions per site. The major lineages of Sulawesi adrianichthyids are color coded.



Fig. 2. Bayesian chronograms of Sulawesi adrianichthyids based on the mitochondrial sequences (cyt *b*: 1141 bp and ND2: 1046 bp) using a substitution rate of (A) 2.65% per My for cyt *b* and (B) 2.8% per My for ND2. Numbers on branches are Bayesian posterior probabilities. Bars represent 95% high posterior density. The major lineages of Sulawesi adrianichthyids are color coded.

1.2. Bayesian chronograms of Sulawesi adrianichthyids

The data presented here represent the Bayesian chronograms of Sulawesi adrianichthyids separately estimated using a substitution rate for cyt b and for ND2 (Fig. 2A and B). The data of the two chronograms in the Newick format are available as Supplementary data 4 and 5.

Lognormal relaxed clock analyses were performed in BEAST version 1.8.1 [6] on the 2187-bp mitochondrial sequence dataset. Appropriate substitution models were determined for each gene (GTR+I+G for both cyt *b* and ND2) using the jModelTest. Analysis was run by partitioning codon positions and using the following settings: birth-death process, ngen=50000000, samplefreq=1000, and burnin=12500000. Analysis was run separately specifying substitution rates as (1) 2.65% per My for cyt *b* and "estimate" for ND2 and as (2) 2.8% per My for ND2 and "estimate" for cyt *b*. Both of the clocks were based on previous studies on divergence time estimations of Adrinichthyidae [7,8].

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi. org/10.1016/j.dib.2015.08.032.

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