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First Mitogenome of Endangered *Enteromius thysi* (Actinopterygii: Cypriniformes: Cyprinidae) from Africa: Characterization and Phylogeny

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Abstract: The complete mitochondrial genome of endangered *Enteromius thysi* was determined from Cameroon in Western Africa. The genome was 16,688 bp in length, comprising 37 genes (13 PCGs, 2 rRNAs, 22 tRNAs, and an AT-rich control region). The heavy strand accommodates 28 genes (12 PCGs, 2 rRNAs, and 14 tRNAs), whereas the light strand holds 9 genes (NAD6 and 8 tRNAs). The *E. thysi* mitogenome is AT-biased (60.5%), as exhibited in other *Enteromius* species. Most of the PCGs start with the ATG initiation codon, except *COI*, with GTG, and seven PCGs end with the TAA termination codon, except some with an incomplete termination codon. Most of the tRNAs showed classical cloverleaf secondary structures, except tRNA-serine (*trnS1*). Bayesian phylogeny distinctly separated *E. thysi* from other congeners. The control regions of *Enteromius* species exhibited highly variable nucleotides, and parsimony-informative sites were found in the conserved sequence block-III (CSB-III) compared with other domains and a unique 9 bp tandem repeat (ATGCATGGT) in the variable-number tandem repeats (VNTRs) region of *E. thysi*. The present phylogeny with limited mitogenomes showed an uneven diversity and evolutionary patterns of *Enteromius* species distributed in the northwestern and southeastern riverine systems in Africa, which warrants further investigation. Based on the results of the present study, we recommend additional surveys, in-depth taxonomy, and the generation of more mitogenomes that could resolve the diversification pattern of *Enteromius* species in Africa.

Keywords: freshwater fish; threatened species; primer walking; mitogenome; evolution

1. Introduction

The family Cyprinidae (order Cypriniformes) is a species-rich group with 1781 valid species under 157 genera distributed in North America, Africa, and Eurasia [1]. African Cyprinids were broadly segregated into three groups: small diploid species, small-to-medium tetraploid species, and large hexaploid species [2]. Later, the generic status of small African diploid minnows (genus *Barbus*) was elevated to *Enteromius* [3–5]. To date, the genus *Enteromius* has been reported to include 224 valid species widely distributed in the African continent [6]. These *Enteromius* species are distinguished from other small minnows by their unique morphology [7]. However, the morphology and color pattern often overlapped within the *Enteromius* members poses difficulties for species-level

identification [4,8]. Consequently, the distribution patterns of many *Enteromius* are thought to overlap over a surprisingly wide geographic range across multiple river systems, avoiding dispersal limitations by terrestrial barriers [9]. Such a scenario evidences the lack of knowledge of the phenotypic and lineage diversification of African freshwater fishes in time and space. As a result, these taxonomic barriers are hampering biodiversity campaigns and conservation efforts.

Among extant African small barb, *Enteromius thysi* was discovered in the Kake River, Mungo River basin, Cameroon. The species has restricted distribution in Western Africa (Cameroon, Equatorial Guinea, and Bioco Island). Nevertheless, *E. thysi* faces several threats, such as habitat destruction, sedimentation, and pollution, as well as anthropogenic pressure [10]. Thus, the IUCN specialist group categorized *E. thysi* under 'endangered' status as per its extent of occurrence (EOO) and area of occupancy (AOO). To date, few studies have been conducted on *E. thysi* based on their morphology and molecular data. The nuclear (*RH*, *RAG1*, *IRBP*, and *EGR2B*) and mitochondrial (*NAD4*, *NAD5*, *CYTB*, *COI*, and *16S rRNA*) genes of *E. thysi* were generated to elucidate their phylogeny, resolving the classification of Cyprinids [3,11]. With advances in molecular studies of biodiversity and systematics, the evolutionary assessment of fishes through complete mitochondrial genomes has blossomed at a rapid pace worldwide [12,13]. Mitogenomic assessments often provide useful clues to understanding the architecture of mitochondrial genes that play important roles in adaptive divergence and speciation [14–16]. In addition, global communities are enriching future analysis pipelines by building mitochondrial genome databases to identify fish diversity from environmental DNA [17,18]. Prior to this study, the mitogenomes of a total of 13 *Enteromius* species had been generated around the world [19,20]. To improve in-depth phylogenetic knowledge of Cyprinids, the aim of the present study was to generate the complete mitogenome of *E. thysi* for the first time and characterize its genomic features. This genetic information can strengthen the mitogenomic library of Cyprinid fishes and would benefit future molecular examination and encourage scientific conservation of endangered *E. thysi* in Africa.

2. Materials and methods

2.1. Sampling and Species Identification

A single individual of *Enteromius* species was opportunistically collected from the Nyong River (3.7649 N 12.2468 E) in Cameroon (Figure 1). Experimental protocols were carried out in accordance with relevant ARRIVE 2.0 (<https://arriveguidelines.org>) guidelines and regulations. The ethics review board, IACUC (Institutional Animal Care and Use Committee), of the host institute confirmed that the use of muscle tissue from dead animals, which were not killed by the researcher, did not involve any animal ethical issue. The specimen was tentatively identified as *E. thysi* according to its morphological characteristics [7]. An adequate amount of tissue was collected and preserved in 70% ethanol in a 2 mL centrifuge tube at 4°C. To confirm the species identity, we amplified the partial *COI* sequences using published primers (FISH-BCL and FISH-BCH) [21]. The generated sequences showed 99.21% similarity with the published database sequence (KP712036) of *E. thysi* [3]. Hence, we confirmed the species' identity and proceed with further molecular investigation. The range distribution shape file of *E. thysi* was downloaded from the IUCN (<https://www.iucnredlist.org/>), and a map was prepared using ArcGIS 10.6 software (ESRI, CA, USA) and edited manually.

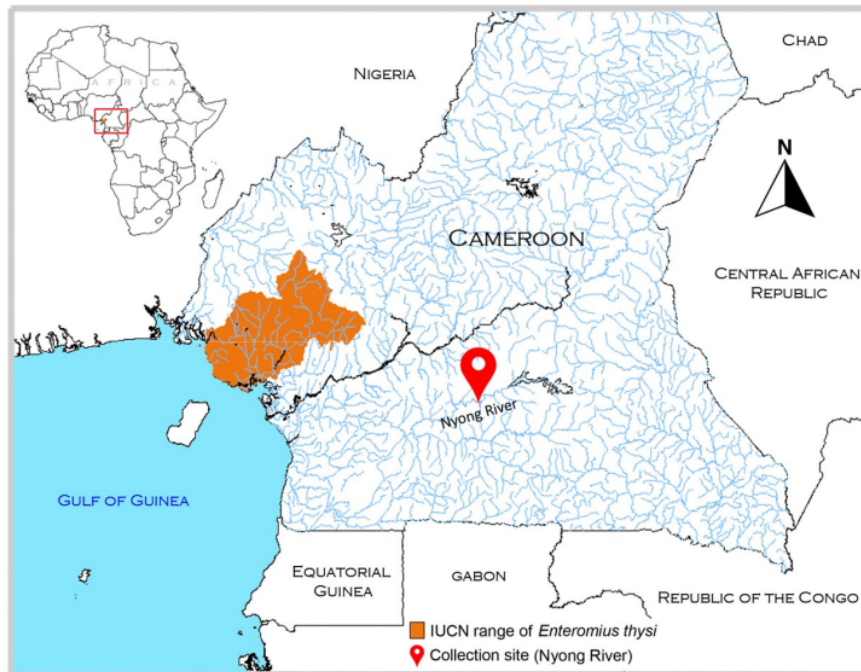


Figure 1. Map showing the IUCN range distribution and collection locality of *Enteromius thysi* from Cameroon.

2.2. DNA Extraction, Sequencing, and Assembly

Genomic DNA was extracted using an AccuPrep® genomic DNA extraction kit (Bioneer, Republic of Korea) according to the standard manufacturer protocol. The quality and quantity of genomic DNA were determined using a NanoDrop spectrophotometer (Thermo Fisher Scientific D1000). To amplify the long fragments of the mitogenome, primer walking with a total of six sets of primer pairs was used in the present study (Table S1). A total of three contigs (*NAD6* to *D-loop*, *D-loop* to *12S rRNA*, and *12S rRNA* to *16S rRNA*) were determined by typical amplification; however, to achieve the other three long contigs (*COI* to *COIII*, *COIII* to *NAD6*, and *16S rRNA* to *COI*) 2-step nested PCR was used. Each amplification reaction contained 1X PCR buffer, 10 pmol of each primer, 2.5 mM dNTPs, 1 U of Taq polymerase, and 1 µL of template DNA. The PCR amplification was performed by a TaKaRa PCR Thermal Cycler Dice® Gradient (Takara Korea Biomedical Inc. Seoul, Republic of Korea). The amplified products were purified using an AccuPrep® PCR/gel purification kit (Bioneer, Daejeon, Republic of Korea). The amplicons were further amplified with a BigDye® Terminator v3.1 cycle sequencing kit (Applied Biosystems) in DNA Engine Tetrad 2 Peltier Thermal Cycler (BIO-RAD) and sequenced bidirectionally in an automated sanger sequencer (96 capillaries; ABI PRISM 3730XL Analyzer) at Macrogen (<https://dna.macrogen.com/>) (Daejeon, Republic of Korea). The noisy parts were trimmed from each amplicon sequence using SeqScanner version 1.0 (Applied Biosystems Inc., CA, USA). The complete mitogenome was assembled by assuring the overlapping regions through MEGA v11 [22] and assured through nucleotide BLAST search (<https://blast.ncbi.nlm.nih.gov>). The contig, gene boundaries, and directions were confirmed by the MITOS v806 online webserver (<http://mitos.bioinf.uni-leipzig.de>) and MitoAnnotator (<http://mitofish.aori.u-tokyo.ac.jp/annotation/input/>) [18,23]. The protein-coding genes (PCGs) were confirmed after being translated into the putative amino acids as per the vertebrate mitochondrial

genetic code through the Open Reading Frame Finder web tool (<https://www.ncbi.nlm.nih.gov/orffinder/>). The generated mitogenome was submitted to the GenBank database using the Bankit submission tool.

2.3. Mitogenome Characterization and Phylogenetic Analyses

A circular illustration of the generated mitogenome of *E. thysi* was plotted through the MitoAnnotator webserver (<http://mitofish.aori.u-tokyo.ac.jp/annotation/input/>). The overlapping regions and intergenic spacers between the neighbor genes were verified manually through Microsoft Excel. The mitogenome size and nucleotide composition were calculated using MEGA v11. The base composition skew of the mitogenome was calculated as previously described: AT skew = $[A - T]/[A + T]$; GC skew = $[G - C]/[G + C]$ [24]. The boundaries of ribosomal RNA genes (rRNAs) and transfer RNA genes (tRNAs) were also confirmed by the MITOS online server, the tRNAscan-SE Search Server 2.0, and ARWEN 1.2 [25,26]. A total of eight *Enteromius* species (*E. thysi*, *E. guirali*, *E. callipterus*, *E. pobeguini*, *E. eburneensis*, *E. fasciolatus*, *E. hulstaerti*, and *E. trimaculatus*) were screened to determine the structural domains of control regions (CRs) by CLUSTAL X version 1.83 alignments [27]. The tandem repeats in the CRs were predicted by the online Tandem Repeats Finder web tool (<https://tandem.bu.edu/trf/trf.html>) [28]. To check the phylogenetic relationships, 13 *Enteromius* mitogenomes were downloaded from GenBank to form a dataset, and two sequences of *Barbus* species were incorporated as an outgroup (Table S2). The PCGs of all 16 mitogenomes were separately aligned and concatenated by the iTaxoTools 0.1 tool [29]. The best-fit model was calculated by partitioning each PCG using PartitionFinder 2 using CIPRES Science Gateway V. 3.3 [30,31]. A Bayesian (BA) tree was constructed using Mr. Bayes 3.1.2, and the MCMC was run for 1,000,000 generations with sampling at every 100th generation and 25% of samples rejected as burn-in [32]. The BA tree was further annotated by iTOL v4 (<https://itol.embl.de/login.cgi>) [33].

3. Results and Discussion

3.1. Mitogenome Structure and Organization

The mitogenome (16,688 bp) of endangered *E. thysi* was determined in the present study (GenBank accession no. OP819561) (Figure 2). Compared to the other *Enteromius* species, the mitogenome of *E. thysi* is longer than *E. eburneensis* (16,678 bp) and *E. fasciolatus* (16,566 bp); however, it is shorter than *E. hulstaerti* (16,775 bp), *E. guirali* (16,793 bp), and *E. callipterus* (16,859 bp). The longest mitogenome (16,933 bp) was observed in *E. pobeguini*, whereas the shortest genome (16,417 bp) was observed in *E. trimaculatus* (Table S2). The circular mitogenome of *E. thysi* constituted 13 protein-coding genes (PCGs), 22 transfer RNA genes (tRNAs), two ribosomal RNA genes (rRNAs), and a non-coding AT-rich control region (CR). A total of 28 genes (12 PCGs, 2 rRNAs, and 14 tRNAs) were located on the heavy strand, whereas *NAD6* and 8 tRNAs were located on the light strand (Table 1). The mitogenome of *E. thysi* is AT-biased (60.5%), with 32.8% A, 15.4% G, 24.1% C, and 27.7% T, as observed in the mitogenomes of other *Enteromius* species (Table S3). This AT-biased nucleotide composition was similar to that of the mitogenomes of other vertebrates [34,35]. A total of 22 bp overlapping regions and 42 bp intergenic spacer regions were detected in the *E. thysi* mitogenome. The AT skew and GC skew were 0.084 and −0.220, respectively, in the mitogenome of *E. thysi*. Comparative analysis showed that the AT skew ranged from 0.049 (*E. trimaculatus*) to 0.120 (*E. camptacanthus*), and the GC skew ranged from −0.187 (*E. hulstaerti*) to −0.262 (*E. camptacanthus*) (Table S3). The genetic variations observed in mitochondrial genes might be connected the mechanism of evolution and favor the energy metabolism of *Enteromius* species, as evidenced in other vertebrates [36].

Table 1. List of annotated mitochondrial genes of *Enteromius thysi*.

Gene	Start	End	Strand	Size (bp)	Intergenic	Anticodon	Start Codon	Stop Codon
<i>tRNA-Phe (F)</i>	1	69	+	69	.	GAA	.	.
<i>12S rRNA</i>	70	1026	+	957
<i>tRNA-Val (V)</i>	1027	1098	+	72	.	TAC	.	.
<i>16S rRNA</i>	1099	2792	+	1694
<i>tRNA-Leu (L2)</i>	2793	2868	+	76	1	TAA	.	.
<i>NAD1</i>	2870	3844	+	975	5	.	ATG	TAA
<i>tRNA-Ile (I)</i>	3850	3921	+	72	-2	GAT	.	.
<i>tRNA-Gln (Q)</i>	3920	3990	-	71	1	TTG	.	.
<i>tRNA-Met (M)</i>	3992	4061	+	70	.	CAT	.	.
<i>NAD2</i>	4062	5107	+	1046	.	.	ATG	TA-
<i>tRNA-Trp (W)</i>	5108	5178	+	71	2	TCA	.	.
<i>tRNA-Ala (A)</i>	5181	5249	-	69	1	TGC	.	.
<i>tRNA-Asn (N)</i>	5251	5323	-	73	33	GTT	.	.
<i>tRNA-Cys (C)</i>	5357	5423	-	67	-1	GCA	.	.
<i>tRNA-Tyr (Y)</i>	5423	5493	-	71	1	GTA	.	.
<i>COI</i>	5495	7045	+	1551	.	.	GTG	TAA
<i>tRNA-Ser (S2)</i>	7046	7120	-	75	1	TGA	.	.
<i>tRNA-Asp (D)</i>	7122	7193	+	72	6	GTC	.	.
<i>COII</i>	7200	7887	+	688	.	.	ATG	T--
<i>tRNA-Lys (K)</i>	7888	7963	+	76	1	TTT	.	.
<i>ATP8</i>	7965	8129	+	165	-7	.	ATG	TAA
<i>ATP6</i>	8123	8805	+	683	.	.	ATG	TA-
<i>COIII</i>	8806	9590	+	785	.	.	ATG	TA-
<i>tRNA-Gly (G)</i>	9591	9662	+	72	.	TCC	.	.
<i>NAD3</i>	9663	10,011	+	349	.	.	ATG	T--
<i>tRNA-Arg (R)</i>	10,012	10,081	+	70	.	TCG	.	.
<i>NAD4L</i>	10,082	10,378	+	297	-7	.	ATG	TAA
<i>NAD4</i>	10,372	11,752	+	1381	.	.	ATG	T--
<i>tRNA-His (H)</i>	11,753	11,821	+	69	10	GTG	.	.
<i>tRNA-Ser (S1)</i>	11,832	11,889	+	58	2	GCT	.	.
<i>tRNA-Leu (L1)</i>	11,892	11,964	+	73	3	TAG	.	.
<i>NAD5</i>	11,968	13,791	+	1824	-4	.	ATG	TAA
<i>NAD6</i>	13,788	14,309	-	522	.	.	ATG	TAA
<i>tRNA-Glu (E)</i>	14,310	14,378	-	69	5	TTC	.	.
<i>Cyt b</i>	14,384	15,520	+	1137	3	.	ATG	TAA
<i>tRNA-Thr (T)</i>	15,524	15,595	+	72	-1	TGT	.	.
<i>tRNA-Pro (P)</i>	15,595	15,665	-	71	.	TGG	.	.
Control region	15,666	16,688	+	1023

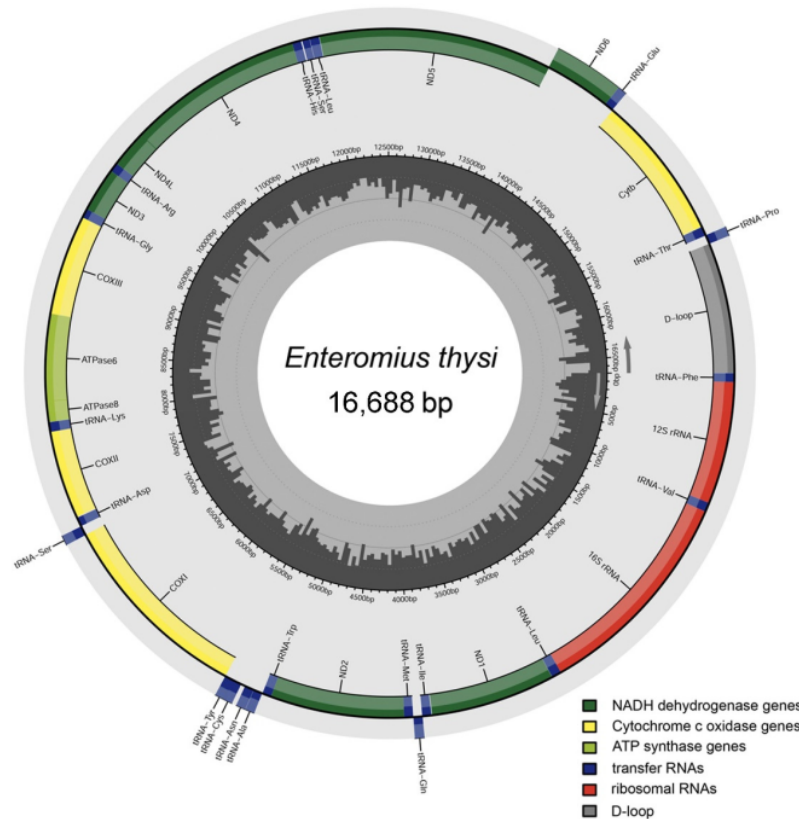


Figure 2. The mitochondrial genome of *E. thysi* drawn by the MitoAnnotator online server (<http://mitofish.aori.u-tokyo.ac.jp/annotation/input/>).

3.2. Protein-Coding Genes

A total of 13 canonical PCGs accounted for 68.33% of the mitogenome, measuring 11,403 bp in *E. thysi*. The shortest among the PCGs was *ATP8*, with a length of 165 bp, whereas the longest PCG was *NAD5*, with 1824 bp. The majority of the PCGs began with an **ATG** (Methionine) initiation codon, whereas *COI* begin with a **GTG** (valine) codon (Table 1). In terms of the termination codon, the typical TAA codon was observed among seven PCGs (*NAD1*, *COI*, *ATP8*, *NAD4L*, *NAD5*, *NAD6*, and *CYTB*), whereas the incomplete stop codons (TA-) and (T-) were exhibited among *NAD2*, *ATP6*, and *COIII* and among *COII*, *NAD3*, and *NAD4*, respectively, as maintained by other species. These incomplete termination codons may be completed to TAA by the addition of a poly A tail through RNA processing [37]. The PCGs (*COI*, *NAD3*, *NAD4L*, *NAD6*, and *CYTB*) were found to have negative GC skews, whereas *NAD6* was found to have a positive GC skew in the *E. thysi* mitogenome (Table S3). Such genetic variations in PCGs correlate with the missense mutations and their independent selection, as revealed in other fishes [38–40].

3.3. Ribosomal RNA and Transfer RNA Genes

In *E. thysi*, the length of the ribosomal RNA genes was 2651 bp (15.88% of the total mitogenome) comprising a small ribosomal RNA (*12S rRNA*) and a large ribosomal RNA (*16S rRNA*), with lengths of 957 bp and 1694 bp, respectively. The rRNAs of *E. thysi* are the third largest compared to the other *Enteromius* species (Table S3). Such structure and

variation, particularly the conserved multibranch loop of ribosomal subunits, allows us to gain substantial knowledge of catalytic reactions in protein synthesis [41]. Furthermore, the *E. thysi* mitogenome consists of 22 tRNA genes, which are characterized by a variation of a length of 67–76 bp and a total estimated length of 1558 bp (9.34% of the total mitogenome). Most of the tRNAs were predicted to fold in the typical cloverleaf secondary structure, except *trnS1*, which lacks d arms, as reported in other fish species [13]. These structural features are crucial for building the secondary structures of RNA structures and function in different biological systems [42]. The conventional base pairings (A=T and G=C) were detected in 14 tRNAs; however, wobble base pairing was observed in 14 other tRNA genes (*trnF*, *trnV*, *trnL2*, *trnQ*, *trnW*, *trnA*, *trnC*, *trnY*, *trnS2*, *trnD*, *trnK*, *trnG*, *trnE*, and *trnP*) (Figure S1).

3.4. Features of Control Region

The total length of *E. thysi* CR was 1023 bp, comprising 67.8% AT and 32.2% GC contents. The full-length sequence of the control region was obtained from the eight *Enteromius* species, ranging from 757 bp in *E. trimaculatus* to 1268 bp in *E. pobeguini* (Table S3). A total >13 copies of 9 bp (ATGCATGGT) tandem repeats were found in the VNTRs (Variable Number Tandem Repeats) region of *E. thysi* CR. Comparative analysis revealed variations in VNTRs regions in three species: *E. fasciolatus*, with >13 copies of 2 bp (TA); *E. hulstaerti*, with >15 copies of 17 bp (ACCCTTAATGGTATAGT); and *E. trimaculatus*, with >37 copies of 18 bp (TATGTACTTTGTACATAC) repeat regions. No repeats were found in the CR of *E. guirali*, *E. callipterus*, *E. pobeguini*, or *E. eburneensis*. Conserved sequence blocks (CSBs) were found to be similar in all the studied *Enteromius* species, as reported in other vertebrates [13,43]. The lengths of CSB-D and CSB-III were similar (19 bp); however, CSB-I was 22 bp, and CSB-II was 18 bp. The variable regions within the different CSB domains of the studied *Enteromius* species ranged from 12 to 15 sites (Figure 3). Comparative analysis revealed highly variable nucleotides (15), and parsimony-informative sites were detected in CSB-III. The variable structures of these CRs can be used to detect intraspecific variation and future phylogeographic studies, as suggested for teleost fishes [44]. The presence of conserved domains in *Enteromius* species may play an important role in the replication and transcription of the genome, as in other vertebrate species.

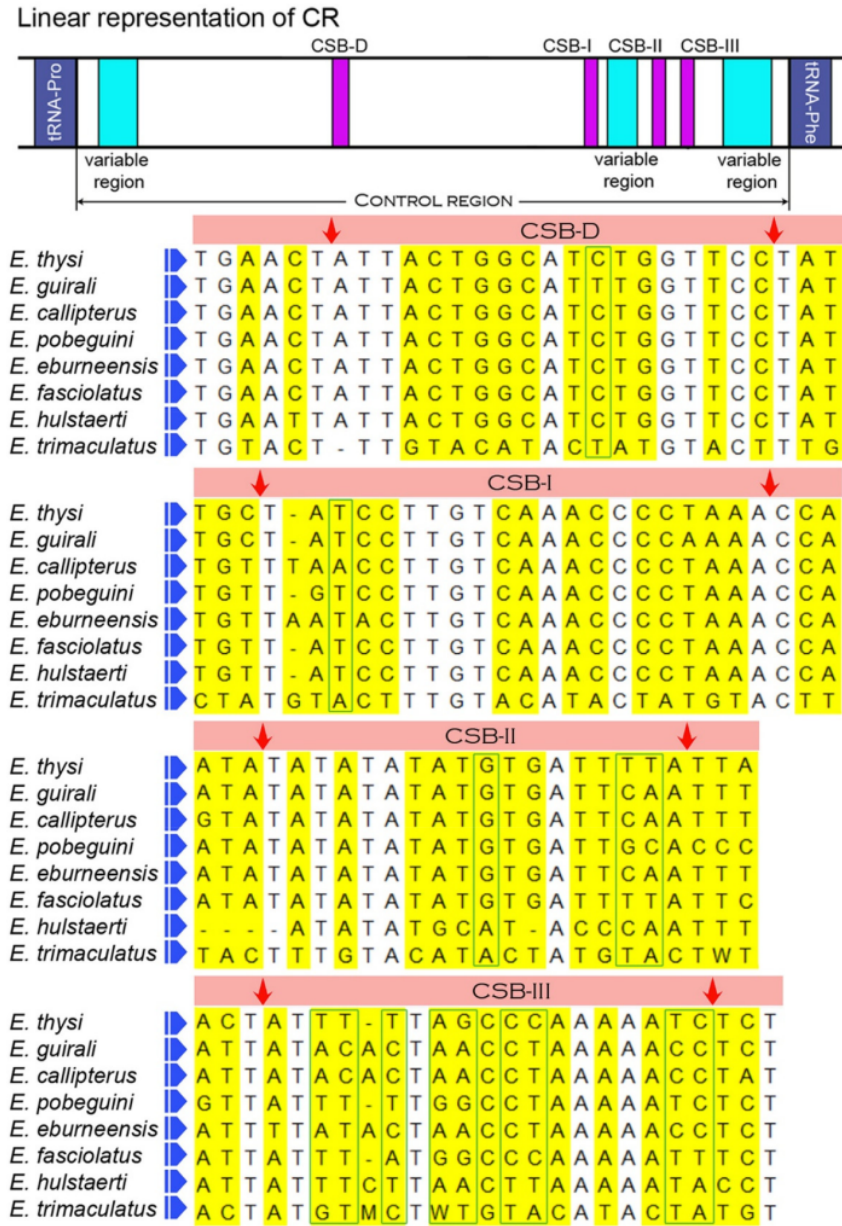


Figure 3. Structural variation of different domains of *Enteromius* control regions. The start and stop position of the different conserved domains are marked by red arrows. The variable regions are marked in yellow, and parsimony-informative sites are marked by green boxes.

3.5. Phylogenetic Relationship of *Enteromius*

The mitogenome dataset with 13 concatenated PCGs showed distinct clustering of all *Enteromius* species in the Bayesian (BA) topology. Among the studied species, *E. thysi* was clearly separated from all other congeners (Figure 4). The BA topology was further investigated based on the known distribution pattern of *Enteromius* species. The ranges of

Enteromius species were obtained from the IUCN (<https://www.iucnredlist.org/>) and mapped using the DIVA-GIS program (<https://www.diva-gis.org/>). Based on the known distribution pattern, the studied *Enteromius* species were categorized into two groups: (i) northwestern species distributed in three major rivers in Africa (Senegal River, Niger River, and Congo River) and (ii) southeastern species distributed in the Orange River, Limpopo River, Zambezi River, and Kalungwishi River. The *Enteromius* mitogenome generated from the Kalungwishi River is not named up to the species level and therefore not discussed in detail. Considering the BA phylogeny, *E. jae* and *E. hulstaerti* are distributed in northwestern riverine systems, separated from other congeners, and placed on the basal node of *Enteromius*. Furthermore, the four species (*E. camptacanthus*, *E. guirali*, *E. eburneensis*, and *E. callipterus*) distributed in the northwestern riverine systems closely clustered in the BA phylogeny with high posterior probability support. Three southeastern riverine species (*E. trimaculatus*, *E. cf. bifrenatus*, and *E. fasciolatus*) also revealed close assemblage in the BA phylogeny (Figure 4). However, *E. pobeguini* and the targeted taxa *E. thysi* distributed in the northwestern riverine systems were found to be surprisingly close to the southeastern riverine species in the present topology. Previous studies evidenced the non-monophyletic evolutionary pattern of *Enteromius* species (previously known as ‘*Barbus*’), with an unparallel diversification in African riverine systems. The present phylogenetic interpretation based on limited complete mitogenomic data was partially congruent with the earlier studies [3,7,11]. However, we suggest that more mitogenome sequences of other *Enteromius* are required to understand the clear evolutionary pattern of these charismatic fishes in Africa. This genetic information can help to elucidate the future population genetics of *E. thysi* in Africa.

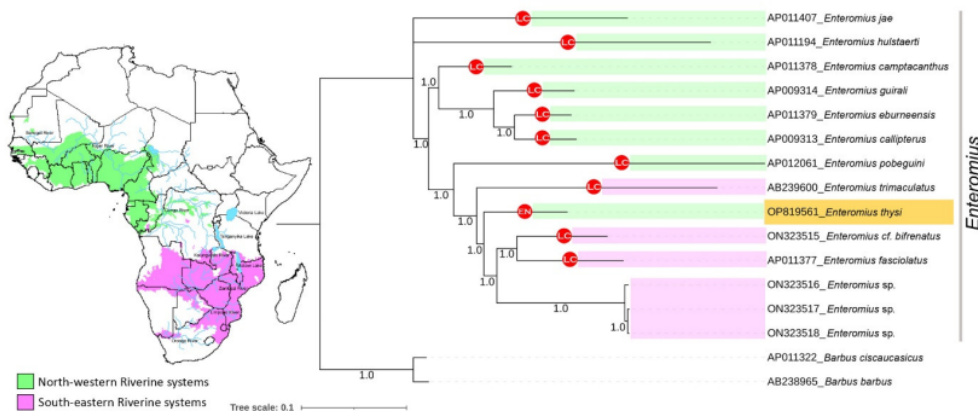


Figure 4. Bayesian phylogeny based on the concatenated nucleotide sequences of 13 PCGs shows the phylogenetic relationship of *E. thysi* with other *Enteromius* species. The posterior probabilities are superimposed with each node. The distribution pattern of *Enteromius* species in different riverine systems is marked by green and pink shades on the map, as well as in each node. LC = least concern; EN = endangered.

The freshwater ecoregions of Africa are home to rich biodiversity, with many endemic fishes, representing a unique entity for evolutionary studies [45–47]. However, this freshwater biodiversity is rapidly declining due to several anthropogenic threats and global climate change, including in Africa [48–50]. Owing to the threatened status of *E. thysi*, we further suggest that additional surveys are essential to attain a better understanding of the biogeography of this endemic fish and to mitigate the threats within and beyond its distribution range. We advocate the integration of non-invasive sampling and a mitochondriomics approach to assess the status and conservation management of this endangered fish species.

4. Conclusions

In the present study, we generated the first complete mitogenome of *E. thysi* and enriched the global nucleotide library. We elucidated the structural features of *E. thysi* and conducted a comparative analysis with other *Enteromius* species. Such empirical molecular data are significant for inferring the functions of the mitochondrial genome and its associated genes. Mitogenome-based phylogenomic analysis affirmed the classification of *Enteromius* species, in contrast to the earlier hypothesis. The results of the present study suggest that the generation of more mitogenomes of *Enteromius* species would be beneficial for systematics research, especially to confirm classifications. Similar genetic information could facilitate other studies on the population genetics and phylogeny of Cyprinid fishes and on the evolution of tandem repeats within control regions.

Supplementary Materials: The following supporting information can be downloaded at: <https://www.mdpi.com/article/10.3390/fishes8010025/s1>, Table S1. List of designed primer pairs used for the long PCR to assemble the *E. thysi* mitogenome. Table S2: Mitogenomes of *Enteromius* species used in phylogenetic analysis. Table S3. Nucleotide composition of the mitochondrial genome in different *Enteromius* species. Figure S1. Putative secondary structures for 22 tRNA genes in the mitochondrial genome of *E. thysi*. The last structure shows the nucleotide positions and details of the stem loop of tRNAs.

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