

Comparison of Mitochondrial Genome Arrangement of *Labeorohita* with Some Selected Vertebrae Species.

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ABSTRACT

The mitochondrial genome of *Labeorohita* serves as a microcosm for exploring genetic diversity and evolutionary relationships within vertebrates. Through comprehensive analysis of the mitochondrial genome, encompassing 37 genes including protein-coding genes (PCGs), transfer RNA (tRNA), and ribosomal RNA (rRNA), we elucidated its organization and compared it with other vertebrate taxa.

Our investigation revealed a compact mitochondrial genome of approximately 16 kb, comprising 13 PCGs, 22 tRNA genes, and 2 rRNA genes, with a distinctive dispersed loop region (D-loop). Notably, the arrangement of PCGs formed three distinct clusters, shared across *Labeo* species, suggesting evolutionary conservation within the genus. Comparative analyses with other osteichthyes species corroborated this conservation, underscoring *L. rohita*'s representative status within the class.

Furthermore, comparisons with chondrichthyes, lungfish, amphibians, reptiles, birds, and mammals unveiled evolutionary trends in mitochondrial genome organization. Noteworthy findings include the unique replication origin in lungfish and amphibians, and transposition events in birds. Phylogenetic reconstructions based on mitochondrial gene sequences affirmed the close relationship between lungfish and amphibians, supporting lungfish as the closest relatives of class Amphibia.

In summary, our study provides comprehensive insights into the mitochondrial genome organization of *L. rohita* and its implications for vertebrate evolution. The findings contribute to a deeper understanding of genetic diversity and evolutionary relationships within vertebrates, with potential applications in taxonomy, conservation, and evolutionary biology.

Keywords: *Labeorohita*, mitochondrial genome, genetic diversity, evolutionary relationships, vertebrates, phylogenetic analysis, protein-coding genes, transfer RNA, ribosomal RNA, conservation.

INTRODUCTION

The genus *Labeo* contains 103 species, with four species found in Bangladesh (Rahman, 2005). *Labeo* species constitute an important group of fish with intense diversity and potential for commercial aquaculture in many Southeast Asian nations, including the Indian subcontinent.

There are only two types of DNA in animals: nuclear DNA and mitochondrial DNA (mtDNA).

Mitochondria are cellular organelles that have the function of oxidative phosphorylation and the formation of ATP. In humans, mtDNA is a double-stranded, circular, covalently closed molecule of 16.5 kb. MtDNA

is inherited as a haploid from the mother, and heteroplasmy has been rarely found. From a population perspective, it could be considered as a system of small, sexually isolated demes or clonal lineages, with an evolutionary rate 5 to 10 times faster than the nuclear genome (José A. Castro, Antònia Picornell, & Misericòrdia Ramon, 1998).

Complete mitochondrial DNA analysis has proven useful in clarifying relationships among closely related species (Mayer, 1993). The 16S, 12S, Co1, and Cytb genes of the mitochondrial genome have been used as useful molecular markers for species-specific identification (Sperling & March, 1994; Girish et al., 2005).

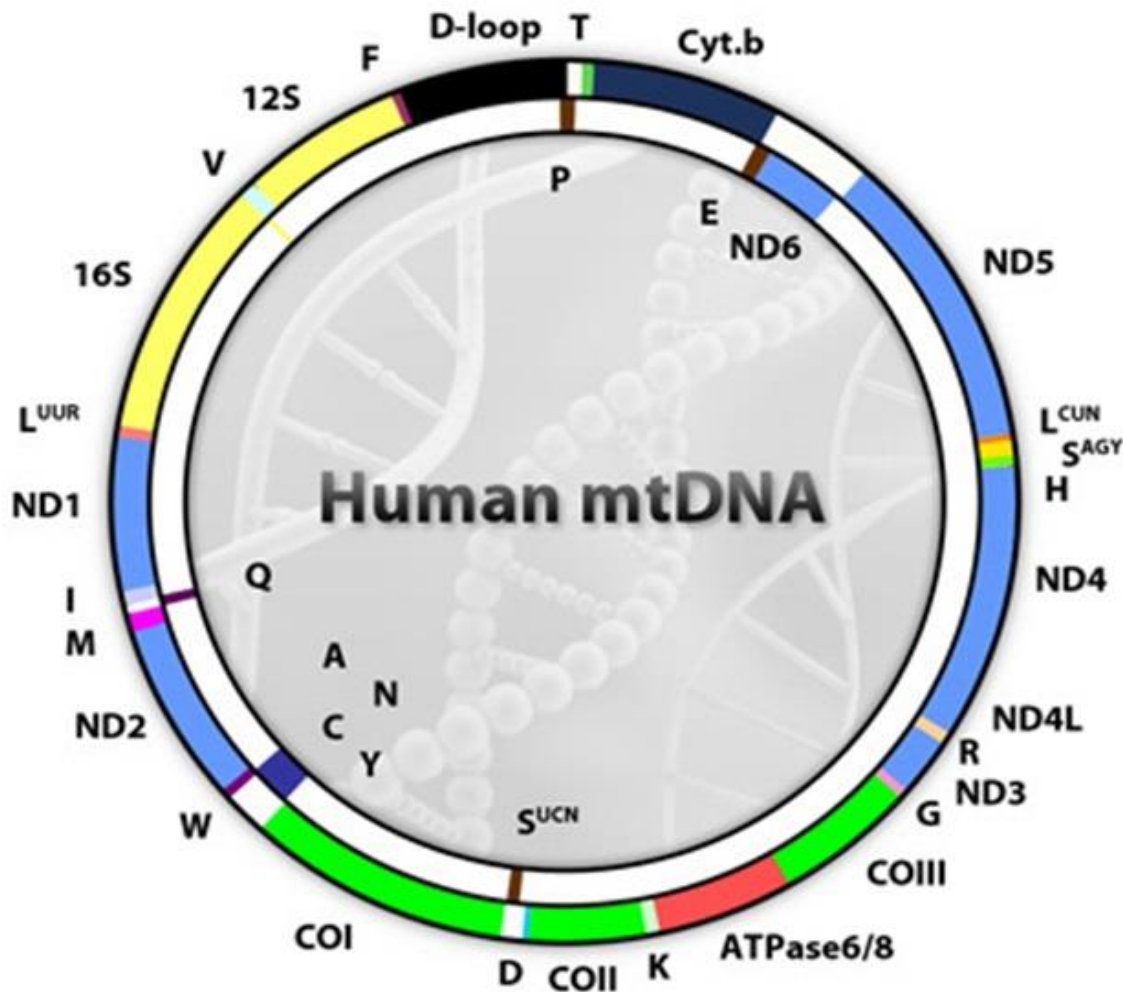


Figure 1: Animal mitochondrial DNA showing position of all the genes. Figure collected from Wikipedia (Mitochondrial DNA; <http://commons.wikimedia.org>).

The animal mitochondrial DNA (mtDNA) is a closed circular molecule (see Figure 1). Only the cnidarian classes Cubozoa, Scyphozoa, and Hydrozoa have been found to have linear mtDNA chromosomes (Bridge et al., 1992). In general, the vertebrate mt genome is highly compact, around 16 kb in length, and contains only 37 genes: 13 protein-coding genes, two ribosomal RNA genes (rRNAs), and 22 transfer RNA genes (tRNAs), along with one long noncoding region known as the control region containing the signals for regulation and initiation of mtDNA replication and transcription (e.g., Wolstenholme, 1992). Mt gene arrangement tends to be conserved among vertebrates, with all 37 genes and the control region organized in relatively the same order from teleost fish to eutherian mammals (Boore, 1994a).

Complete mitochondrial DNA (mtDNA) sequences have been determined for 177 vertebrate species and 75 invertebrate species, with partial sequences for hundreds of others (Boore et al., 1999). All are circular DNA

molecules containing 37 genes: 13 for proteins (COI-III, ND1-6, ND4L, Cytb, A6, A8); two for rRNAs (srRNA and lrRNA); and 22 for tRNAs (designated by the one-letter amino acid code, with the two S and two L tRNAs differentiated by the codons recognized [AGN/UCN and CUN/UUR, respectively]). The genes are arranged very compactly, with no introns and few intergenic nucleotides.

Studies on the genome rearrangement of *Labeorohita* and its comparison with other species have not been conducted yet. Genome arrangement of *Labeorohita* has not been compared with other chondrichthyes and osteichthyes species. Furthermore, lower vertebrate species have not been compared with genome arrangement with that of *Labeorohita*. Similarly, higher vertebrate species have not been compared with the genome arrangement of *Labeorohita*. Despite the extensive work on the complete mitochondrial genome or partial sequencing of mtDNA and identification of species based on 16S, CO1, and Cyt b genes, there has been no research on rearrangement. Therefore, I attempted to study this topic.

METHODOLOGY

Sample Collection and DNA Extraction:

- i. Specimens of *Labeorohita* and other vertebrate species were collected from diverse geographical locations.
- ii. Genomic DNA was extracted from tissue samples using the CTAB method.

PCR Amplification and Sequencing:

- i. A short region of the mitochondrial genome, such as the 16S rRNA gene, was amplified using polymerase chain reaction (PCR) with species-specific primers.
- ii. PCR products were purified and sequenced using Sanger sequencing technology.

Data Retrieval and Assembly:

- i. Mitochondrial DNA sequences of 17 species, namely *L. rohita*, *L. bata*, *L. calbasu*, *Channa marulius*, *Sicamugilcascasia*, *Tenualosailisha*, *Heteropneustesfossilis*, *Macragnathus aculeatus*, *Mastacembelusarmatus*, *Neoceratodusforsteri*, *Chimaera monstrosa*, *Dasyatiszugei*, *Scoliodonlaticaudus*, *Bufo melanostictus*, *Gecko gekko*, *Acridotheres tristis*, and *Cavia porcellus*, were collected from the GenBank Database (NCBI).
- ii. The genes of the collected sequences were manually arranged according to their position in the collected sequence using Microsoft Office. To construct a phylogenetic tree from the nucleotide sequences of the 16S and COI genes of the 17 species mentioned above, different software programs were used, including Notepad, Serial Cloner, and SeaView.
- iii. Sequences were assembled and annotated using bioinformatics tools to identify protein-coding genes, tRNA, rRNA, and the D-loop region.

Genome Arrangement Analysis:

- i. The arrangement of protein-coding genes, tRNA, rRNA, and the D-loop region in the mitochondrial genome of *rohita* was analyzed and compared with other vertebrate species.
- ii. Manual arrangement was performed using software like Microsoft Office or specialized bioinformatics tools.

Phylogenetic Analysis:

- i. Nucleotide sequences of conserved mitochondrial genes (e.g., CoI, 16S rRNA, cytochrome b) from

rohita and representative vertebrate species were aligned.

- ii. Phylogenetic trees were constructed using maximum likelihood (ML), neighbor-joining (NJ), or maximum parsimony (MP) methods.
- iii. Bootstrap analysis was conducted to assess the robustness of tree topologies.

Comparative Genomics:

- i. Comparative analyses were performed to identify conserved regions, gene clusters, and unique features in the mitochondrial genomes of different vertebrate taxa.
- ii. Differences in gene arrangements, sequence lengths, and structural elements like replication origins were examined.

Data Interpretation and Conclusion:

- i. Results from genome arrangement analysis, phylogenetic reconstruction, and comparative genomics were interpreted to infer evolutionary relationships and genetic diversity.
- ii. Conclusions regarding the representative nature of *rohita* mitochondrial genome, evolutionary trends, and relationships among vertebrates were drawn.

Validation and Review:

- i. Results were validated through rigorous statistical analyses and peer review processes.
- ii. Feedback from experts in the field was incorporated to refine interpretations and ensure the robustness of conclusions.

RESULT & DISCUSSION

Comparison of mtDNA arrangement in different vertebrate

Figure 2: Schematic diagram of the mitochondrial genome of *Labeorohita*(AP 011201). Different color showing the position of 13 protein coding, 22 tRNA and 2 rRNA genes with D-loop region.

Mitochondrial Genome Arrangement:

From the NCBI, it was revealed that the complete mitochondrial genome of *Labeorohita* is 16 kb long. It consists of 37 genes with one dispersed loop region or D-loop region. Among them, 13 protein-coding genes, 22 tRNAs, and 2 rRNA genes are found.

1. Protein coding genes:

There are 13 protein-coding genes in the mitochondrial genome of *Labeorohita*, namely NADH1, NADH2, Cytochrome Oxidase I, Cytochrome Oxidase II, ATPase 8, ATPase 6, Cytochrome Oxidase III, NADH 3, NADH 4L, NADH 4, NADH 5, NADH 6, Cytochrome b, and the D-loop or control region. These 13 protein-coding genes are interrupted by 22 tRNAs except for three clusters, namely ATPase 6 – ATPase 8 – COIII, ND4L – ND4, ND5 – ND6. The arrangement of these three clusters of protein-coding genes is commonly observed in vertebrate mitochondrial genomes.

Comparing the mitochondrial genome of three *Labeo* species, namely *Labeorohita*, *L. calbasu*, and *L. bata*, it was observed that these three clusters of protein-coding genes are the same in all three species of *Labeo*. However, differences were observed only in gene length, which varied from 1 to 10 bp. From this observation, it was concluded that the genome arrangement of *Labeorohita* is a representative or typical

genome arrangement for *Labeo* species in the case of protein-coding genes.

Typical *Labeo* species were compared with the genome arrangement of *Tenualosailisha*, *Channa marulius*, and *Heteropneustes fossilis*, and with *Mastacembelus armatus*, *Macrognathus aculeatus*, and *Sicamugilcascasia*. The same result was found as before, that is, all three clusters are similar in these species. From those comparisons, it was eventually concluded that the mitochondrial genome arrangement of *L. rohita* can represent the class Osteichthyes for protein-coding genes. Thus, the arrangement of the mitochondrial genome of *L. rohita* can be described as the typical mitochondrial genome arrangement for class Osteichthyes.

<i>Labeorohita</i>	<i>Labeocalbasu</i>	<i>Labeo bata</i>	Typical <i>Labeosp</i>
Phe, 69	Phe, 69	Phe, 69	Phe,
12S,954	12S, 954	12S,954	12S
Val,71	Val, 71	Val,71	Val
16S,1686	16S, 1689	16S,1686	16S
Leu,75	Leu, 75	Leu,75	Leu
ND1,974	ND1, 974	ND1,974	ND1
Ile,71	Ile, 71	Ile,71	Ile
Gln,70	Gln, 70	Gln,70	Gln
Met,68	Met, 68	Met, 68	Met
ND2,1044	ND2, 1044	ND2,1044	ND2
Trp,70	Trp, 70	Trp, 70	Trp,
Ala,68	Ala, 68	Ala, 68	Ala
Asn,72	Asn, 72	Asn, 72	Asn
Cys,66	Cys, 66	Cys, 66	Cys
Tyr,70	Tyr, 70	Tyr, 70	Tyr
COI,1550	COI, 1550	COI,1550	COI
Ser,70	Ser, 70	Ser, 70	Ser
Asp,71	Asp, 71	Asp,71	Asp
COII,680	COII, 690	COII,680	COII
Lys,75	Lys, 73	Lys, 75	Lys
ATP8,164	ATP8, 164	ATP8, 164	ATP8
ATP6, 682	ATP6, 682	ATP6, 682	ATP6
COIII, 784	COIII, 785	COIII, 784	COIII
Gly,71	Gly, 71	Gly,71	Gly
ND3, 348	ND3, 348	ND3, 348	ND3
Arg, 69	Arg, 69	Arg, 69	Arg,
ND4L,296	ND4L, 296	ND4L, 296	ND4L
ND4, 1380	ND4, 1380	ND4, 1380	ND4
His, 68	His, 68	His, 68	His
Ser, 68	Ser, 68	Ser, 68	Ser
Leu, 72	Leu, 72	Leu, 72	Leu
ND5, 1823	ND5, 1823	ND5, 1823	ND5

ND6, 521	ND6, 521	ND6, 521	ND6
Glu, 68	Glu, 68	Glu, 68	Glu
Cyt b, 1140	Cyt b, 1140	Cyt b, 1140	Cyt b
Thr,71	Thr, 71	Thr,71	Thr
Pro,69	Pro, 69	Pro, 69	Pro
D-loop, 940	D-loop, 937	D-loop, 940	D-loop

Figure 3: Mitochondrial genome arrangement of three species of *Labeo* namely *Labeorohita*(AB 011201), *L. calbasu* (AP012143) and *L. bata* (AP011198). Number showing the total length of particular genes in the boxes. Protein-coding genes are represented bold. tRNA genes are represented as three letter code.

- When the typical mitochondrial genome arrangement of fish is compared with chondrichthyes such as *Chimaera monstrosa*, *Dasyatis zugei*, and *Scoliodon laticaudus* (see Figure 6), it was found that the arrangement of protein-coding genes is the same with the three clusters. So, it can be concluded that in the case of protein-coding genes, the genome arrangement of *L. rohita* is typical for chondrichthyes.
- When the typical mitochondrial genome arrangement of fish is compared with the lungfish *Neoceratodus forsteri* (see Figure 6), it was found that the arrangement of protein-coding genes is the same, and the three clusters are also the same. So, it can be concluded that in the case of protein-coding genes, the genome arrangement of *L. rohita* is typical for *Neoceratodus forsteri*.
- When the typical mitochondrial genome arrangement of fish is compared with amphibian *Bufo melanostictus* (see Figure 7), it was revealed that the protein-coding genes are the same, but the positions are different, and the number of base pairs also varies from that of the fish mitochondrial genome. Additionally, it was observed that in fish, between the clusters ND4L-ND4 and ND5-ND6, there are three tRNAs present (tRNA^H, tRNA^S, tRNA^{L2}), but in the case of Amphibia (*Bufo*), only tRNA^H and tRNA^S are present, and tRNA^{L2} is absent. So, it can be concluded that amphibians are distant from the fish mitochondrial genome.
- When the typical mitochondrial genome arrangement of fish is compared with reptiles (*G. gekko*), birds (*A. tristis*), and mammals (*C. porcellus*), it was revealed that the protein-coding genes are the same in all cases. Additionally, the three clusters ATPase 6 – ATPase 8 – COIII, ND4L – ND4, ND5 – ND6 are similar to that of reptiles and mammals genome arrangement, but in the case of birds, there is an ND5-Cyt b cluster present instead of the ND5 – ND6 cluster. This is due to the transposition of the ND6 and tRNA^{Glu} gene.
- The typical mitochondrial genome arrangement of fish was also compared with *Drosophila melanogaster*, and it was revealed that tRNA^H, tRNA^T, and tRNA^P were present between the ND5 and ND6 genes (see Figure 9). The same result was found in the sessile barnacle *Tetraclita japonica* (Rowshan et al., 2004). Therefore, it can be concluded that the mitochondrial genome arrangement between fish and invertebrates is significantly different when protein-coding genes are considered.

Typical <i>Labeo</i> sp	<i>Tenualosa</i>	<i>Channa</i>	<i>Heteropneustes</i>
Phe,	Phe, 69	Phe, 70	Phe, 69
12S	12S, 953	12S, 944	12S, 954
Val	Val, 71	Val, 71	Val, 71
16S	16S, 1679	16S, 1685	16S, 1663
Leu	Leu, 75	Leu, 75	Leu, 74
ND1	ND1, 974	ND1, 974	ND1, 974
Ile	Ile, 71	Ile, 69	Ile, 71

Gln	Gln, 70	Gln, 70	Gln,70
Met	Met, 68	Met, 69	Met,69
ND2	ND2, 1049	ND2, 1045	ND2, 1044
Trp,	Trp, 71	Trp, 70	Trp, 70
Ala	Ala, 68	Ala, 68	Ala, 68
Asn	Asn, 73	Asn, 72	Asn, 72
Cys	Cys, 65	Cys, 66	Cys, 65
Tyr	Tyr, 69	Tyr, 69	Tyr, 69
COI	COI, 1550	COI, 1550	COI, 1550
Ser	Ser, 70	Ser, 70	Ser,70
Asp	Asp, 68	Asp, 71	Asp, 70
COII	COII, 690	COII, 690	COII, 690
Lys	Lys, 73	Lys, 74	Lys, 73
ATP8	ATP8, 167	ATP8, 167	ATP8, 167
ATP6	ATP6, 683	ATP6, 682	ATP6, 682
COIII	COIII, 785	COIII, 784	COIII, 783
Gly	Gly, 70	Gly, 68	Gly, 72
ND3	ND3, 350	ND3, 348	ND3, 348
Arg,	Arg, 68	Arg,69	Arg, 68
ND4L	ND4L, 296	ND4L,296	ND4L, 296
ND4	ND4, 1380	ND4, 1380	ND4, 1380
His	His, 68	His, 69	His,69
Ser	Ser, 66	Ser, 69	Ser, 66
Leu	Leu, 70	Leu, 72	Leu, 72
ND5	ND5, 1847	ND5, 1838	ND5, 1826
ND6	ND6, 521	ND6, 521	ND6, 518
Glu	Glu, 68	Glu, 67	Glu, 68
Cyt b	Cyt b, 1140	Cyt b, 1140	Cyt b, 1137
Thr	Thr, 71	Thr,71	Thr, 72
Pro	Pro, 69	Pro,69	Pro, 69
D-loop	D-loop, 1168	D-loop, 911	D-loop, 858

Figure 4: Compare the mitochondrial genome arrangement of *Labeo* with those of *Tenualosailisha* [Clupeiformes, (AP 011610)], *Channa marulius* [Channiformes(AB 968638)], *Heteropneustesfossilis* [Siluriformes (AP 012013)].

2. tRNA:

There are 22 tRNA genes observed in *L. rohita*, consistent with the observation of Wolfe (1993). The tRNA genes of *L. rohita* range in size from 66 to 76 bp, and they form 5 clusters, namely WANCY, IQM, SD, HSL2, and TP (see Figure 3).

- a. The arrangement of the 22 tRNA genes of three *Labeo* species, *L. rohita*, *L. calbasu*, and *L. bata*, was

studied, and the aforementioned 5 clusters are the same in all three species. However, the lengths differ from each other by only 1-2 bp (see Figure 3). From this observation, we can conclude that the genome arrangement of *Labeorohita* is a representative or typical genome arrangement for *Labeo* species for tRNA genes.

- b. Subsequently, the typical *Labeo* species' tRNA gene arrangement was compared with *Tenualosailisha*, *Channa marulius*, and *Heteropneustes fossilis* (see Figure 4), and with *Mastacembelus armatus*, *Macragnathus aculeatus*, and *Sicamugil cascasia* (see Figure 5). The same result was found as before; all 5 clusters are similar in these species. From these comparisons, it was concluded that the mitochondrial genome arrangement of *L. rohita* can represent the class Osteichthyes for tRNA gene arrangement. Thus, the arrangement of the mitochondrial genome of *L. rohita* can be described as the typical mitochondrial genome arrangement for class Osteichthyes or the typical mitochondrial genome arrangement for fish.

Typical <i>Labeosp</i>	<i>Mastacembelus</i>	<i>Macragnathus</i>	<i>Sicamugil</i>
Phe,	Phe, 69	Phe, 69	Phe, 69
12S	12S, 948	12S, 944	12S, 950
Val	Val, 71	Val, 72	Val, 70
16S	16S, 1670	16S, 1669	16S, 1696
Leu	Leu, 73	Leu, 74	Leu, 73
ND1	ND1, 971	ND1, 971	ND1, 974
Ile	Ile, 70	Ile, 70	Ile, 69
Gln	Gln, 70	Gln, 70	Gln, 70
Met	Met, 68	Met, 68	Met, 69
ND2	ND2, 1045	ND2, 1045	ND2, 1044
Trp,	Trp, 71	Trp, 71	Trp, 71
Ala	Ala, 68	Ala, 68	Ala, 68
Asn	Asn, 72	Asn, 72	Asn, 72
Cys	Cys, 66	Cys, 66	Cys, 65
Tyr	Tyr, 66	Tyr, 66	Tyr, 67
COI	COI, 1550	COI, 1550	COI, 1592
Ser	Ser, 70	Ser, 70	Ser, 70
Asp	Asp, 71	Asp, 71	Asp, 72
COII	COII, 690	COII, 690	COII, 690
Lys	Lys, 74	Lys, 74	Lys, 74
ATP8	ATP8, 167	ATP8, 167	ATP8, 167
ATP6	ATP6, 682	ATP6, 683	ATP6, 683
COIII	COIII, 784	COIII, 784	COIII, 783
Gly	Gly,70	Gly, 69	Gly, 72
ND3	ND3, 345	ND3, 345	ND3, 348
Arg,	Arg, 68	Arg, 69	Arg, 68
ND4L	ND4L, 297	ND4L, 296	ND4L, 296

ND4	ND4, 1380	ND4, 1380	ND4, 1380
His	His, 64	His, 69	His, 68
Ser	Ser, 66	Ser, 66	Ser, 67
Leu	Leu, 72	Leu, 72	Leu, 72
ND5	ND5, 1832	ND5, 1838	ND5, 1841
ND6	ND6, 515	ND6, 521	ND6, 521
Glu	Glu, 68	Glu, 68	Glu, 68
Cyt b	Cyt b, 1137	Cyt b, 1137	Cyt b, 1134
Thr	Thr, 71	Thr, 71	Thr, 76
Pro	Pro, 69	Pro, 69	Pro, 69
D-loop	D-loop, 871	D-loop, 917	D-loop, 1389

Figure 5: Compare the mitochondrial genome arrangement of *Labeo* with *Mastacembelusarmatus* [synbranchiformes (NC_023977)], *Macrogathus aculeatus* [synbranchiformes (NC_027435)] , *Sicamugilcascasia* [mugiliformes (NC_017898)].

- When the typical mitochondrial genome arrangement of osteichthyes is compared with that of chondrichthyes such as *Chimaera monstrosa*, *Dasyatis zugei*, and *Scoliodon laticaudus* (see Figure 27), it was found that the arrangement of tRNA was almost the same. However, a replication origin containing 37 base pairs was observed within the cluster WANCY, more specifically, between tRNAN and tRNAC. So, it can be concluded that in the case of tRNA genes, chondrichthyes differ from osteichthyes.
- When the typical mitochondrial genome arrangement of fish is compared with lungfish *Neoceratodus forsteri* (see Figure 6), it was found that the arrangement of tRNA is the same, but the difference is that in the cluster WANCY, there lies a replication origin of 35 base pairs. So, it can be concluded that in the case of tRNA, lungfish is slightly different from osteichthyes and resembles that of chondrichthyes.
- When the typical mitochondrial genome arrangement of fish is compared with amphibian *Bufo melanostictus* (see Figure 7), it was revealed that the tRNA clusters are different. In amphibians, the tRNA clusters are LITPF, IQM, WANCY, SD, and HS. A replication origin of 10 base pairs is present between tRNAN and tRNAC. Lungfish also has a replication origin in the same clusters between tRNAN and tRNAC (see Figure 7). The position of the origin of replication is a unique identifying characteristic of lungfish, which is similar to that of amphibian mtDNA and dissimilar to that of other osteichthyes mtDNA.
- When the typical mitochondrial genome arrangement of fish is compared with reptiles (*G. gekko*), birds (*A. tristis*), and mammals (*C. porcellus*), it was revealed that the 5 clusters (WANCY, IQM, SD, HSL2, TP) are similar in all cases. However, the position of the TP cluster is different in the case of birds.
- By comparing the typical mitochondrial genome arrangement of osteichthyes with that of *Drosophila melanogaster*, it was revealed that the tRNA clusters of *Drosophila* are different. In *Drosophila*, the tRNA clusters ARNS1EP, IQP, WCY, KD, and TP were found (see Figure 9). The same result was found in the sessile barnacle *Tetraclita japonica* (Begum, R.A. et al., 2004). So, it can be concluded that the mitochondrial genome arrangement of osteichthyes, more commonly known as fish, and invertebrates are far different from each other whenever vertebrate and invertebrate mtDNA is

compared.

Typical Osteichthyes	<i>Neoceratodus</i>	<i>Chimaera</i>	<i>Dasyatis</i>	<i>Scoliodon</i>
Phe,	Phe, 68	Phe, 70	Phe, 71	Phe, 70
12S	12S, 951	12S, 948	12S, 963	12S, 951
Val	Val, 70	Val, 70	Val,71	Val, 71
16S	16S, 1679	16S, 1664	16S, 1704	16S, 1669
Leu	Leu, 74	Leu, 74	Leu, 74	Leu, 74
ND1	ND1, 971	ND1, 971	ND1, 977	ND1, 974
Ile	Ile, 76	Ile, 72	Ile, 68	Ile, 69
Gln	Gln, 70	Gln, 69	Gln, 72	Gln, 71
Met	Met, 68	Met, 68	Met, 70	Met, 68
ND2	ND2, 1043	ND2, 1041	ND2, 1046	ND2, 1044
Trp,	Trp, 71	Trp, 68	Trp, 69	Trp, 70
Ala	Ala, 68	Ala, 68	Ala, 68	Ala, 68
Asn	Asn, 72	Asn, 72	Asn, 72	Asn, 72
Cys	Rep org, 35	Rep org, 52	Rep org, 35	Rep org, 37
Tyr	Cys, 70	Cys, 73	Cys, 66	Cys, 66
COI	Tyr, 69	Tyr, 67	Tyr, 69	Tyr, 68
Ser	COI, 1556	COI, 1559	COI, 1556	COI, 1556
Asp	Ser, 70	Ser, 71	Ser, 71	Ser, 70
COII	Asp, 68	Asp, 71	Asp, 70	Asp, 68
Lys	COII, 712	COII, 690	COII, 69 0	COII, 690
ATP8	Lys, 71	Lys, 72	Lys, 73	Lys, 73
ATP6	ATP8, 167	ATP8, 167	ATP8, 167	ATP8, 167
COIII	ATP6, 683	ATP6, 683	ATP6, 683	ATP6, 683
Gly	COIII, 785	COIII, 785	COIII, 785	COIII, 785
ND3	Gly, 70	Gly, 68	Gly, 70	Gly, 69
Arg,	ND3, 348	ND3, 348	ND3, 348	ND3, 348
ND4L	Arg, 69	Arg, 69	Arg, 68	Arg, 69
ND4	ND4L, 296	ND4L, 296	ND4L, 296	ND4L, 296
His	ND4, 1380	ND4, 1374	ND4, 1380	ND4, 1380
Ser	His, 68	His, 68	His, 68	His, 68
Leu	Ser, 66	Ser, 68	Ser, 66	Ser, 66
ND5	Leu, 68	Leu, 71	Leu, 71	Leu, 71
ND6	ND5, 1829	ND5, 1838	ND5, 1844	ND5, 1829
Glu	ND6, 515	ND6, 521	ND6, 515	ND6, 521
Cyt b	Glu, 68	Glu, 69	Glu, 68	Glu, 69
Thr	Cyt b, 1143	Cyt b, 1143	Cyt b, 1142	Cyt b, 1144
Pro	Thr, 72	Thr, 70	Thr, 72	Thr, 69

D-loop	Pro, 69	Misc_feature,	Pro, 69	Pro, 68
	D-loop, 925	1533(CRI)	D-loop, 2522	D-loop, 1062
		Pro, 68		
		Misc_feature,		
		1434(CRI)		

Figure 6: Compare the mitochondrial genome arrangement of *Labeo* with other chondrichthyes namely *Neoceratodusforsteri* (AJ 584642), *Chimaera monstrosa* (AJ 310140), *DasyatisZugei* (NC_019643), and *ScoliodonLaticaudus*(KP 336547).

3. rRNA:

Two ribosomal RNA genes were found in the *Labeo* mitochondrial DNA, one encoding the large subunit, 16S rRNA (1686 bp), and the other encoding the small subunit, 12S rRNA (954 bp). The tRNA^{Val} is present between the 12S rRNA and 16S rRNA genes (see Figure 24), representing a unique arrangement among vertebrate and invertebrate mitochondrial genomes (Begum, R.A. et al., 2004).

- Comparing *L. rohita* with the other two *Labeo* species, it was observed that the number of base pairs is almost similar, differing only by 1-3 bp (see Figure 3).
- Comparing with other osteichthyes, it was found that the number of base pairs differs by up to 22 bp (see Figure 4 & 5).
- Similar results were found when comparing with chondrichthyes and lungfish.
- When compared with amphibians, it was revealed that the number of base pairs differs by 82 bp (see Figure 7).
- When the typical mitochondrial genome arrangement of fish was compared with reptiles (*G. gekko*), birds (*A. tristis*), and mammals (*C. porcellus*), it was found that tRNA^{Val} is present between the small rRNA (12S) and large rRNA (16S) like in other vertebrates and invertebrates. However, the number of base pairs differs by 3 to 25 bp.
- Comparing with *Drosophila* (invertebrate), it was found that there is a large difference between vertebrate and invertebrate rRNA base pair numbers. In *Drosophila*, 12S rRNA has 785 bp, whereas in vertebrates, it is around 954 bp, and 16S rRNA has 1324 bp in *Drosophila*, whereas in vertebrates, it is around 1686 bp.

Typical chondrichthyes	Typical osteichthyes (<i>Labeo</i>)	<i>Neoceratodus</i> (Lung Fish)	<i>Bufo melanostictus</i> (Amphibia)
Phe, 70	Phe,	Phe, 68	Phe, 67
12S, 951	12S	12S, 951	12S, 936
Val, 71	Val	Val, 70	Val, 68
16S, 1669	16S	16S, 1679	16S, 1597
Leu, 74	Leu	Leu, 74	Leu, 72
ND1, 974	ND1	ND1, 971	ND1, 960
Ile, 69	Ile	Ile, 76	Ile, 70
Gln, 71	Gln	Gln, 70	Gln, 68
Met, 68	Met	Met, 68	Met, 72
ND2, 1044	ND2	ND2, 1043	ND2, 1032

Trp, 70	Trp,	Trp, 71	Trp, 69
Ala, 68	Ala	Ala, 68	Ala, 68
Asn, 72	Asn	Asn, 72	Asn, 72
Rep org, 37	Cys	Rep org, 35	Rep org, 10
Cys, 66	Tyr	Cys, 70	Cys, 65
Tyr, 68	COI	Tyr, 69	Tyr, 69
COI, 1556	Ser	COI, 1556	COI, 1541
Ser, 70	Asp	Ser, 70	Ser, 67
Asp, 68	COII	Asp, 68	Asp, 68
COII, 690	Lys	COII, 712	COII, 687
Lys, 73	ATP8	Lys, 71	Lys, 70
ATP8, 167	ATP6	ATP8, 167	ATP8, 164
ATP6, 683	COIII	ATP6, 683	ATP6, 683
COIII, 785	Gly	COIII, 785	COIII, 783
Gly, 69	ND3	Gly, 70	Gly, 67
ND3, 348	Arg,	ND3, 348	ND3, 339
Arg, 69	ND4L	Arg, 69	Arg, 70
ND4L, 296	ND4	ND4L, 296	ND4L, 299
ND4, 1380	His	ND4, 1380	ND4, 1364
His, 68	Ser	His, 68	His, 68
Ser, 66	Leu	Ser, 66	Ser, 66
Leu, 71	ND5	Leu, 68	ND5, 1788
ND5, 1829	ND6	ND5, 1829	ND6, 494
ND6, 521	Glu	ND6, 515	Glu, 67
Glu, 69	Cyt b	Glu, 68	Cyt b, 1145
Cyt b, 1144	Thr	Cyt b, 1143	D-loop, 1967
Thr, 69	Pro	Thr, 72	Leu, 71
Pro, 68	D-loop	Pro, 69	Thr, 71
D-loop, 1062		D-loop, 925	Pro, 68

Figure 7: Mitochondrial genome arrangement showing the evolutionary relation among fishes and amphibian. Rearrangement shown in arrow marks.

4. D-loop region:

The D-loop region or control region is observed between two tRNA genes, namely tRNAP and tRNAF, in the mitochondrial genome of *L. rohita* (Accession Number: AB011201).

- a. Comparing the mitochondrial genome of three *Labeo* species – *Labeorohita*, *L. calbasu*, and *L. bata* (see Figure 3) – it was observed that the position of the D-loop region is the same. Its length differs by only (1-3) bp. Therefore, it can be said that in the case of the D-loop region, the mitochondrial genome arrangement of *L. rohita* can represent the *Labeo* species.

- b. When the mitochondrial genome arrangement of *L. rohita* is compared with that of *Tenualosailisha*, *Channa marulius*, and *Heteropneustes fossilis* (see Figure 4), and with *Mastacembelus armatus*, *Macrognathus aculeatus*, and *Sicamugil cascasia* (see Figure 5), it was found that although the position is the same, the length of the D-loop region differs from 82 bp to 228 bp, which suggests placing these osteichthyes in different orders. However, from the arrangement comparison, it can be concluded that the arrangement is typical.
- c. When the typical mitochondrial genome arrangement of fish is compared with chondrichthyes fish such as *Chimaera monstrosa*, *Dasyatis zugei*, and *Scoliodon laticaudus* (see Figure 6), it was found that the arrangement and position of the D-loop region are the same. However, the length differs from (122-1582) bp, which indicates that these fish cannot be placed in the same class. It was observed that they belong to the class Chondrichthyes.

Typical Fish	Amphibia <i>Bufo melanostictus</i>	Reptile <i>Gekko gekko</i>	Bird <i>Acridotheres tristis</i>	Mammal <i>Cavia porcellus</i>
Phe,	Leu, 71	Phe, 73	Phe, 69	Phe, 65
12S	Thr, 71	12S, 957	12S, 979	12S, 942
Val	Pro, 68	Val, 69	Val, 70	Val, 69
16S	Phe, 67	16S, 1585	16S, 1598	16S, 1565
Leu	12S, 936	Leu, 75	Leu, 75	Leu, 74
ND1	Val, 68	ND1, 962	ND1, 978	ND1, 960
Ile	16S, 1597	Ile, 72	Ile, 72	Ile, 65
Gln	Leu, 72	Gln, 72	Gln, 71	Gln, 71
Met	ND1, 960	Met, 69	Met, 69	Met, 69
ND2	Ile, 70	ND2, 1035	ND2, 1040	ND2, 1044
Trp,	Gln, 68	Trp, 71	Trp, 70	Trp, 70
Ala	Met, 72	Ala, 68	Ala, 69	Ala, 69
Asn	ND2, 1032	Asn, 72	Asn, 73	Asn, 73
Cys	Trp, 69	Cys, 62	Cys, 67	Rep_origin, 37
Tyr	Ala, 68	Tyr, 65	Tyr, 71	Cys, 67
COI	Asn, 72	COI, 1544	COI, 1551	Tyr, 67
Ser	Rep org, 10	Ser, 74	Ser, 73	COI, 1542
Asp	Cys, 65	Asp, 67	Asp, 69	Ser, 69
COII	Tyr, 69	COII, 692	COII, 684	Asp, 69
Lys	COI, 1541	Lys, 68	Lys, 68	COII, 684
ATP8	Ser, 67	ATP8, 169	ATP8, 168	Lys, 67
ATP6	Asp, 68	ATP6, 686	ATP6, 684	ATP8, 204
COIII	COII, 687	COIII, 783	COIII, 784	ATP6, 681
Gly	Lys, 70	Gly, 69	Gly, 69	COIII, 784
ND3	ATP8, 164	ND3, 339	ND3, 351	Gly, 68

Arg,	ATP6, 683	Arg, 68	Arg, 70	ND3, 347
ND4L	COIII, 783	ND4L, 296	ND4L, 297	Arg, 68
ND4	Gly, 67	ND4, 1380	ND4, 1378	ND4L, 297
His	ND3, 339	His, 71	His, 70	ND4, 1378
Ser	Arg, 70	Ser, 64	Ser, 64	His, 69
Leu	ND4L, 299	Leu, 70	Leu, 71	Ser, 57
ND5	ND4, 1364	ND5, 1829	ND5, 1818	Leu, 70
ND6	His, 68	ND6, 509	Cyt b, 1143	ND5, 1812
Glu	Ser, 66	Glu, 65	Thr, 70	ND6, 531
Cyt b	ND5, 1788	Cyt b, 1134	Pro, 70	Glu, 69
Thr	ND6, 494	Thr, 70	ND6, 519	Cyt b, 1140
Pro	Glu, 67	Pro, 67	Glu, 72	Thr, 69
D-loop	Cyt b, 1145	D-loop, 1172	Misc_feature, 1250	Pro, 68
	D-loop, 1967			D-loop, 1357

Figure 8: Comparison of mitochondrial genome arrangement of typical fish with other vertebrate such as Amphibian (*Bufo melanostictus*, A/N: NC_005794), Reptile (*Gekko gekko*, A/N: HM 370130), Bird (*Acridotheres tristis*, A/N: NC_015195), Mammal (*Cavia porcellus*, A/N: AJ222767).number showing the total length of particular gene in the boxes, Protein coding genes are represent bold. tRNA gene are represent as three-letter code.

1. When the typical mitochondrial genome arrangement of fish is compared with lung fish *Neoceratodus forsteri* (see Figure 6), it was found that the arrangement and position of the D-loop region are the same. The length differs from that of typical fish by only about 15 bp. This finding suggests that this lungfish can be placed with typical fish in the class Osteichthyes.
2. When the typical mitochondrial genome arrangement of fish is compared with amphibian *Bufo melanostictus* (see Figure 7), it was revealed that the D-loop region is 1967 bp long and positioned between the cytochrome b gene and tRNA^{Leu}, which differs from the typical mitochondrial genome of fish. Specifically, it can be said that this position of the D-loop region in amphibian *Bufo melanostictus* is different from that in fish. Therefore, it can be concluded that amphibians belong to a different class.
3. When the typical mitochondrial genome arrangement of fish is compared with reptiles (*G. gekko*), birds (*A. tristis*), and mammals (*C. porcellus*), it was revealed that the position of the D-loop region of reptiles and mammals is similar to that of typical fish genome. However, the position of the D-loop region in birds is between tRNA^{Glu} and tRNA^{Phe}. This is due to the transposition of the *cyt b* gene.
4. When the typical mitochondrial genome arrangement of fish is compared with *Drosophila melanogaster* (Invertebrate) (see Figure 9), it was found that the D-loop region of *Drosophila* is located between the small ribosomal RNA (12S rRNA) and tRNA^{Ile}, with a length ranging from 14917 bp to 19517 bp. As the position and length vary significantly from vertebrates, it can be concluded that *Drosophila* should be placed in another subkingdom. The same result was found in the case of barnacles *Tetraclita japonica* (Rowshan et al., 2004).

Thus, we can conclude that the vertebrate order is first Osteichthyes, then Chondrichthyes, followed by lungfish, and immediately after lungfish is the position of amphibians. Therefore, it can be said that lungfishes are the closest relatives of the class Amphibia.

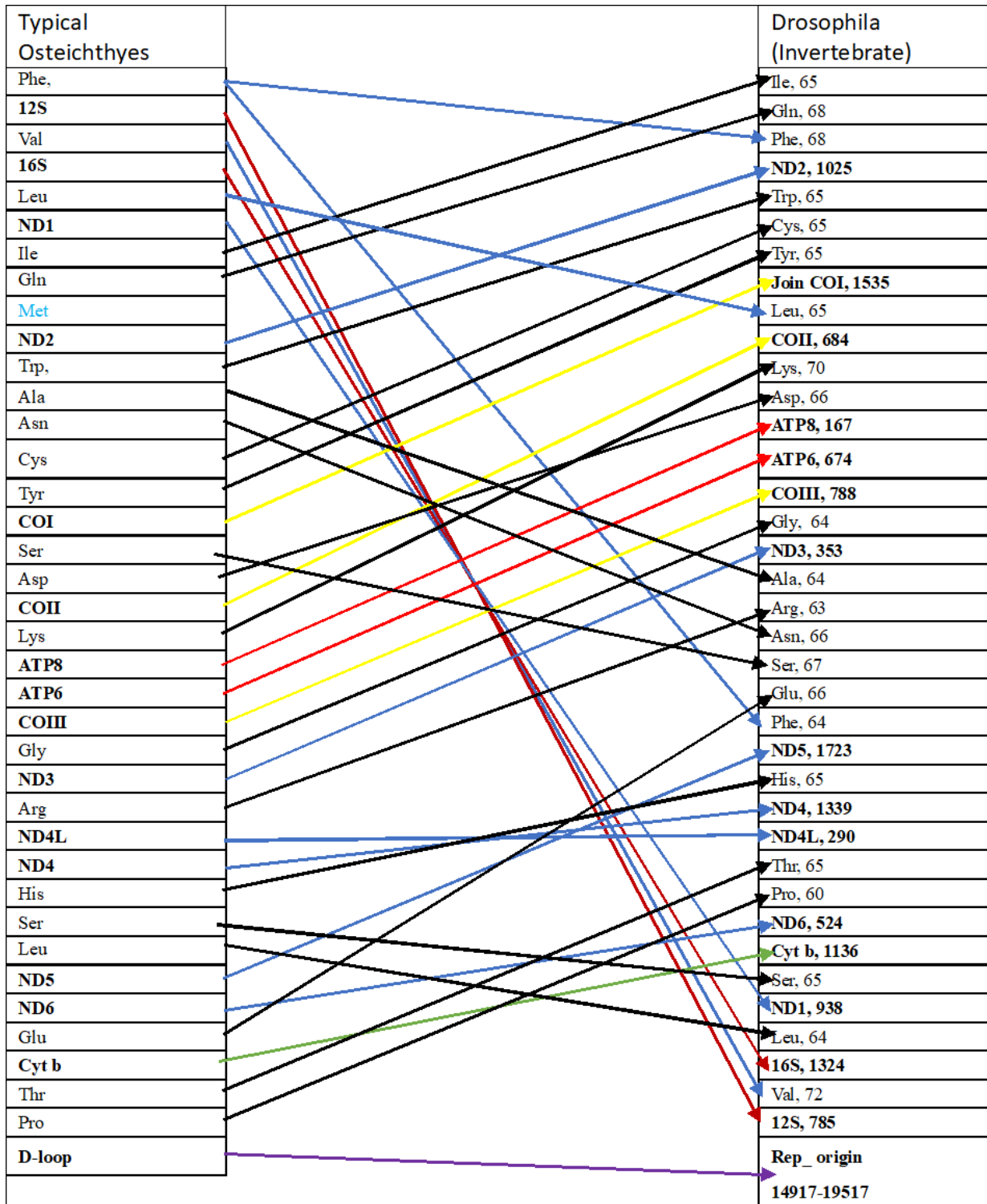


Figure 9: The comparison of mitochondrial genome arrangement between typical Osteichthyes (*Labeo*) and *Drosophila*. Rearrangement are shown in arrow marks (black= a.a; yellow=CO gene; blue=ND gene).

Phylogenetic Tree:

The relationships among *Labeo* species, *Labeo* species with other Osteichthyes, Chondrichthyes, lungfish, and amphibians were investigated using a phylogenetic tree based on the nucleotide sequences of CoI, 16S, and cytochrome b genes. The molecular phylogenetic tree with bootstrap values was constructed with particular attention to different groups of animals.

1. Phylogenetic Tree based on Nucleotide Sequence of CoI Gene:

- a. For the nucleotide dataset of the concatenated CoI, the Maximum Likelihood (ML) method produced distinct branching orders. Three species of Labeo formed a monophyletic group with the highest bootstrap value of 100% in the ML tree. This relationship is consistent with the genome arrangement and the phylogeny based on the CoI gene (see Figure 10).
- b. Osteichthyes such as Tenualosailisha, Channa marulius, Heteropneustesfossilis, Mastacembelusarmatus, Macrogna thus aculeatus, and Sicamugilcascasia form a monophyletic group according to ML trees, consistent with the phylogeny based on taxonomy.
- c. Neoceratodusforsteri (lungfish) is placed just after the Osteichthyes group in the Maximum Likelihood tree.
- d. The phylogenetic tree reveals that Chondrichthyes fish such as Chimaera monstrosa, Dasyatiszugei, and Scoliodonlaticaudus form a monophyletic group, positioned after Amphibia (Bufo melanostictus).
- e. ML shows that Bufo melanostictus (amphibian) is positioned after the lungfish. This relationship is consistent with the genome arrangement and the phylogeny based on the CoI gene (see Figure 10).
- f. The Maximum Likelihood tree shows that birds are situated just before mammals, with a bootstrap value of 0.93, indicating the relevance of this position. This relationship also supports the taxonomic hierarchy of vertebrates.
- g. In the case of Drosophila, the ML tree shows Drosophila as an outgroup. This finding is consistent with the comparison of mitochondrial genome arrangements.

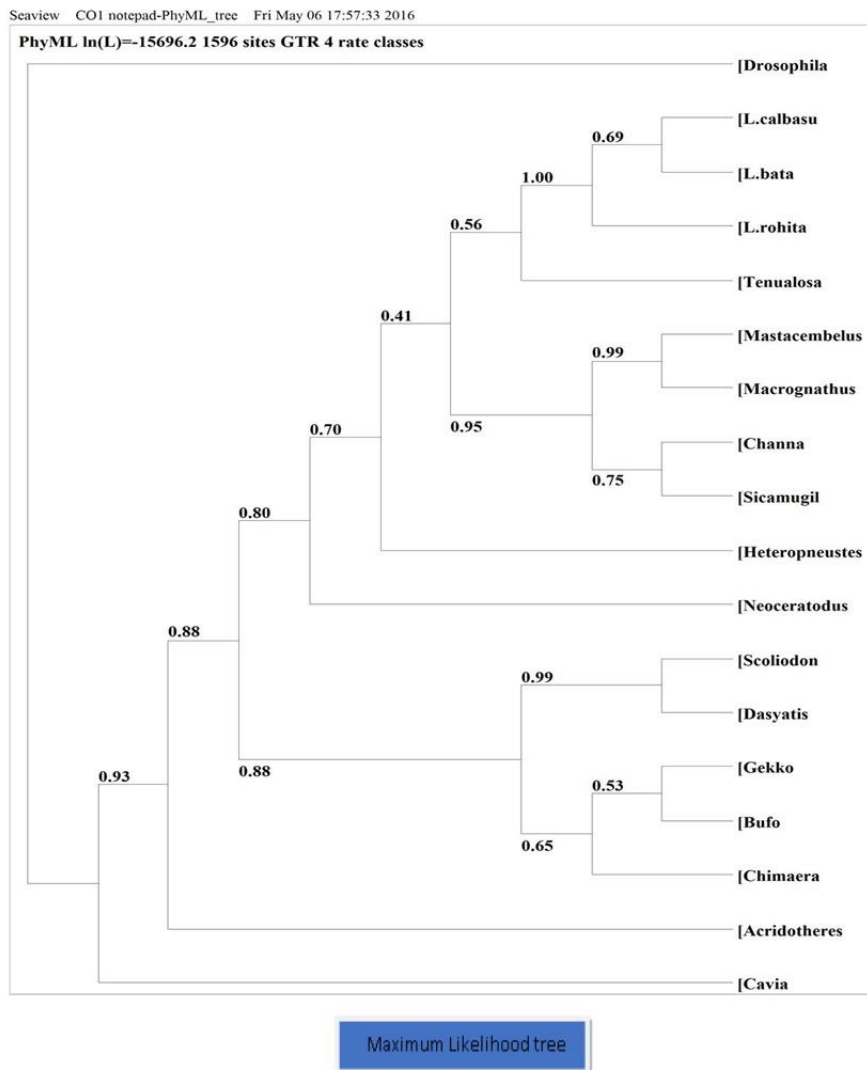


Figure 10: Phylogenetic tree (Maximum Likelihood) based on nucleotide sequence of CO1 gene, number

showing the bootstrap values. Nucleotide sequence of *Drosophila* was used as outgroup.

2. Phylogenetic Tree based on Nucleotide Sequence of 16S rRNA Gene:

- Analysis of the phylogenetic tree based on the 16S rRNA gene of 15 vertebrate species revealed that the Maximum Parsimony and Neighbor Joining methods produce different branching orders. However, a common finding in both trees is that *Labeo* species form a monophyletic group with a bootstrap value of 100.
- According to both Maximum Parsimony and Neighbor Joining trees, *Tenualosailisha*, *Channa marulius*, *Heteropneustesfossilis*, *Mastacembelusarmatus*, *Macrognathus aculeatus*, *Sicamugilcascasia*, including *Labeo* species, form a group, although there are some sub-groups within this larger group. This grouping can be referred to as the Osteichthyes fish group.
- The phylogenetic tree reveals that Chondrichthyes fish such as *Chimaera monstrosa*, *Dasyatiszugei*, and *Scoliodonlaticaudus* form a monophyletic group, positioned with the lungfish *Neoceratodusforsteri*.
- Neoceratodusforsteri* (lungfish) is placed just after the Chondrichthyes group in the Maximum Likelihood tree.
- When a phylogenetic tree (maximum likelihood) was constructed from the 16S nucleotide sequence of different selected species, it was observed that higher vertebrates form a monophyletic group.
- In the case of *Drosophila*, the Neighbor Joining tree shows *Drosophila* as an outgroup, while the Maximum Parsimony tree shows both *Drosophila* and *Bufo* as monophyletic. *Bufo*'s position is after the fish group (as Osteichthyes, *Neoceratodus*, and Chondrichthyes are all fish and sequentially situated in the phylogenetic tree, they can be referred to as the fish group).

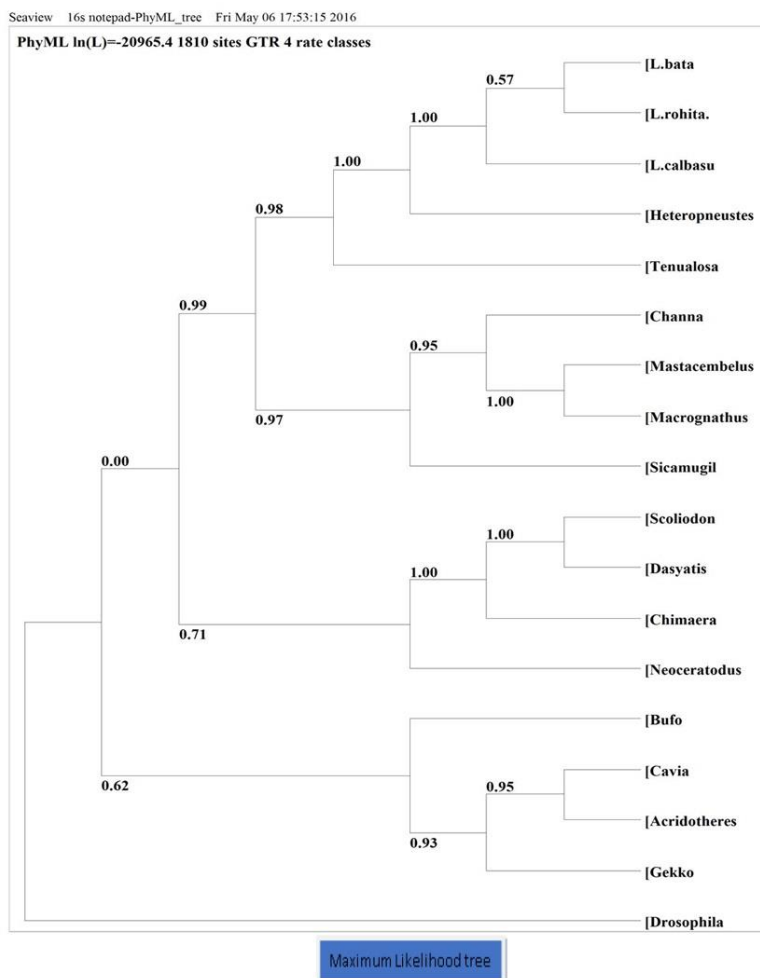


Figure 11: Phylogenetic tree (Maximum Likelihood) based on nucleotide sequence of 16S gene, number

showing the bootstrap values. Nucleotide sequence of *Drosophila* was used as outgroup. CONCLUSION

1. The genome arrangement of *Labeorohita* is conserved in most osteichthyes species, indicating a common evolutionary history among these species.
2. The phylogenetic tree constructed from the nucleotide sequence of 16S rRNA supports the conclusion that lungfish is the closest relative of amphibians.
3. The arrangement of the 37 genes in *L. rohita*'s genome could serve as representative data for vertebrates, excluding birds due to a distinct arrangement.
4. The genome arrangement of invertebrates differs significantly from that of vertebrates, highlighting a clear distinction in evolutionary paths between these two groups.

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