

DNA Barcoding of Stone Fish *Uranoscopus Oligolepis*: Intra Species Delineation and Hypothetical Protein Analysis

Chandan Tiwary¹, M.A.Badhul Haq*¹, S.Vaitheeswari¹, M.Kalaiselvi¹, Mohammad Nurul Azim Sikder² and Wah Wah Min³

¹Faculty of Marine Sciences, CAS in Marine Biology, Annamalai University, Parangipettai – 608502, India

²Institute of Marine Sciences and Fisheries, University Of Chittagong, Chittagong-4331, Bangladesh

³Department of Zoology, University of Yangon, Myanmar.

*Corresponding Author: drhaqmarinevirology@gmail.com

Abstract: The present study addresses this issue by examining the patterning of Cytochrome Oxidase I diversity in the stone fish *Uranoscopus oligolepis* the structurally diverse group of Family *Uranoscopidae*. The sequences were analyzed for their species identification using BOLD's identification engine. The COI sequences of *U. oligolepis* from different geographical regions were extracted from NCBI for intra species variation analysis. All sequences were aligned using Clustal W. The sequences were trimmed using software and phylogenetic tree was constructed with bootstrap test. The results showed that the cytosine content was high (31%). The least molar concentration was observed in guanine (19.5%) and Adenine (19.6%). Thymine was the second predominant in molar concentration next to thymine which is followed by adenine. The G+C content was found to be 49.6% and A+T content was 50.4%. Leucine and Alanine content was high in the amino acid composition. From the study it is assumed that the mitochondrial gene COI can be the potential barcoding region to identify an organism up to the species level.

Keywords: COI, intra species, *Uranoscopus oligolepis*, barcoding, phylogenetic

INTRODUCTION

DNA barcoding—the sequencing of a short standardized region of DNA—has been proposed as a new tool for animal species identification [1]. The DNA barcode itself consists of a 648 bp region of the cytochrome c oxidase 1 (COI) gene. Additionally to the mitochondrial COI gene, nuclear loci are sometimes also considered to improve assignment performance [2, 3]. This has been shown to provide species level resolution of the vast bulk of species in a wide range of animal taxa, including ants, bats, birds, butterflies, crustaceans, fish, and spiders [4, 5,6,7,8 and 9]. It is based on the postulate that every species will most likely have a unique DNA barcode (indeed there are 4⁶⁵⁰ possible ATGC-combinations compared to an estimated 10 million species remaining to be discovered, Wilson [10].

DNA barcoding has been effectively tested on diverse taxa, from invertebrates [11, 12, 13] to vertebrates [14] allowing the discrimination of different species, often consistently with traditional morphological approaches [15]. The escalating use of DNA barcoding approach in the identification of fish species [16] insisted in raising a new research project with the support from Consortium for the Barcoding of Life (<http://www.barcoding.si.edu/>): the Fish Barcode of Life initiative (FISHBOL; <http://www.fishbol.org>), in which the sequence datas are included into a main unique database called BOLD (Barcode of Life Data System, <http://www.barcodinglife.org/views/login.php>; [17, 18]. The cost and time-effectiveness of DNA barcoding enables automated species identification, which is particularly useful in large sampling campaigns. In this way, DNA barcoding could also improve large surveys aiming at unknown species detection and identification of pathogenic species with medical, ecological and agronomical significance [19].

MATERIALS AND METHODS

2.1. Wet lab methodologies

2.1.1. Sample preservation

The *U. oligolepis* fish samples were collected from Parangipettai (South east coast of India) fish landing centre and the tissue samples was excised and cut into small pieces (< 5-7 mm) and preserved in fresh 95% ethanol using 1.5 ml labeled tubes.

2.1.2. DNA extraction

Salting out procedure was adapted to extract DNA from *U. oligolepis* tissues. The preserved tissue in ethanol was washed four to five times with sterile distilled water to get clear of the ethanol content. The ethanol free tissues were transferred in to 1.5 ml tube and grounded in micro pestle with 500µl of solution 1 (500mM Tris-HCL, 20mM EDTA and 2% SDS). After homogenizing the tissues were added with 5µl of Proteinase K (20mg/ml). The tubes were incubated at 55°C in water bath for 2 hours with occasional mixing by inverting the tubes. Following incubation the samples were chilled on ice for 10 minutes and about 250µl of solution 2 (6M NaCl) was added to it and mixed well by inverting the tubes several times. Tubes were then chilled on ice for 5 minutes. Then the tubes were centrifuged at 8000 rpm for 15 minutes and following centrifugation, 500 µl of clear supernatant was collected in a 1.5 ml tube. Equal

volume of (1ml) of 100% analytical grade ethanol was added to precipitate the DNA. A thin hair like precipitate was observed after addition of ethanol. After 30 minutes the tubes were allowed to spin at 11,000rpm for 5 minutes. The supernatant was removed and partially dried in room temperature. The DNA pellets were washed thrice with 70% cold ethanol. The pellets were suspended in 100 μ l of sterile distilled H₂O.

Quantitation of DNA by Spectrophotometric method [20]

- 10 μ l of DNA solution was diluted with 990 μ l of TE.
 - Mixed well and absorbance at 260nm and 280nm was measured.
- The absorbance at 260nm can be used to calculate the concentration of DNA as follows:

Calculations

OD₂₆₀ of 1 = 50 μ g/ml DNA

7Dilution factor = 100

Concentration of DNA in a given solution = $\frac{50 \times \text{OD} \times \text{Dilution factor } \mu\text{g}/\mu\text{l}}{1 \times 1000}$

2.1.3. PCR amplification

The primer set MAB Fw and MAB Rw designed in the conserved region was used for the amplification of the COI region of the test organisms and the primer sequences are;

MABFw: (5'-TCAACCAACCACAAAGACATTGGCAC-3') and MABRw: (5'-TAGACTTCTGGGTGGCCAAAGAATCA-3'). A 1.0 μ l of Sample DNA (approximately 100 ng/ μ l) was added to PCR Mixture containing 100mM Tris HCl (pH 8.3), 500mM KCl (pH 8.3), 2.5 μ l MgCl₂ (25mM), 2.0 μ l dNTP's (2.5mM), 1.0 μ l Primer Forward & Reverse (each of 10pm/ μ l) and 1u / μ l of Taq Polymerase (Bioserve Make) & the final volume made to 25 μ l with nuclease free water. The thermal profile consisted of 35 cycles at 94 °C for 50 s, 54 °C 50 s and 72 °C for 1 min. The amplicons obtained were about 629 bp long.

2.1.4. DNA sequencing

QIAGEN QIAquick™ kit was used for sequencing reaction. PCRs products were gel purified and directly sequenced using MegaBase sequencer- Bioserve India, Hyderabad. Sequences were checked by eye with Bioedit sequence alignment editor using GenBank sequences as reference sequences and unambiguously aligned using Clustal W.

2.2. Dry lab methodologies

2.2.1. BOLD's identification engine

BOLD (Barcoding of life database) is an online workbench that aids in collection, management, analysis, and use of DNA barcodes. Identification engine is the one of the important components of BOLD database which consists of large volume of barcode sequences for both plants (intranuclear spacer gene) and animals (cytochrome c oxidase subunit gene). BOLD-IDS provide a species identification tool that accepts DNA sequences from the barcode region and returns a taxonomic assignment to the species level when possible. The BOLD identification system (IDS) accepts sequences from the 5' region of the mitochondrial gene cytochrome oxidase subunit I and returns species-level identification when one is possible. Further validation with independent genetic markers will be desirable in some forensic applications. This identification engine was accessible online through <http://www.barcodinglife.org/views/idrequest.php>. The sequences were given in FASTA file format in the query box and results were obtained similar to that of BLAST search.

2.2.2. Profiling the barcode region of *Uranoscopus oligolepis*

The molecular weight of the single stranded barcode DNA was calculated as the sum of the monophosphate forms of each deoxyribonucleotide minus one water molecule each. One water (18 Da) was added at the end to represent the 3' hydroxyl at the end of the chain and one more hydrogen atom at the 5' phosphate end. Nucleotide composition summaries and plots were obtained by choosing "Nucleotide Composition" form the "Nucleic Acid" submenu of the "Sequence" menu. Bar plots showed the Molar percent of each residue in the sequence. The degenerate nucleotide designations were added to the plot wherever they are encountered. Any DNA sequence has only A, G, C and T and these were represented by four bars on the graph.

2.2.3. Barcode protein profiling

DNA to Protein: The online software at www.insilico.ehu.es was used to extract hypothesized amino acid sequences from the COI region of *U. oligolepis*. This software allowed modeling and modifications of already existing techniques, as well as new theoretical approaches. Standard genetic code translation was used. DNA sequences were fed in to the query box in FASTA format. Minimum size of protein sequence for Open Reading Frames (ORF) is customizable and they were trimmed to MET-to-Stop.

CLUSTAL W: ClustalW is a general purpose global multiple sequence alignment program for DNA or proteins. It produces biologically meaningful multiple sequence alignments of divergent sequences. It calculates the best match for the selected sequences,

and lines them up so that the identities, similarities and differences can be seen. Evolutionary relationships can be seen via viewing Cladograms or Phylograms.

Phylogenetic tree construction using MEGA: Neighborhood joining (NJ) method of phylogenetic tree construction was preferred for accurate establishment of phylogenetic relationship and to trace out the presence of phylogenetic signals in the DNA sequences [21]. The distance was calculated between every pair of sequences and these were used to construct the phylogenetic tree which guided the final multiple alignment. The scores were calculated from separate pair wise alignments.

MEGA (Molecular Evolutionary Genetic Analysis) version 5: MEGA is an integrated tool for conducting automatic and manual sequence alignment, inferring phylogenetic trees, mining the web base data bases, estimating the rates of molecular evolution, and testing evolutionary hypothesis [22].

Bootstrapping: One of the most commonly used tests of the reliability of an inferred tree is Felsenstein's [23] bootstrap test which is evaluated using [24] bootstrap resampling technique. If there are m sequences, each with n nucleotides (or codons or amino acids) a phylogenetic tree can be reconstructed using the same tree building method. From each sequence n nucleotides were randomly chosen with replacements, giving rise to m rows of n columns each. These now constitute a new set of sequences.

A tree is then reconstructed with these new sequences using the same tree building method as before. Next the topology of this tree was compared to that of the original tree. Each interior branch of the original tree that was different from the bootstrap tree the sequence it partitions is given a score of 0 all other interior branches was given the value 1 was noted. This procedure of re-sampling the sites and the subsequent tree reconstruction was repeated several hundred times and the percentage of times each interior branch was given a value of 1 was noted. This is known as the bootstrap value.

The multiple aligned sequences from Clustal X were loaded into MEGA through Create New Alignment option in Alignment menu. The sequences were trimmed for their conserved regions and saved in MEGA format for phylogram construction. Bootstrap test for phylogeny was preferred to detect the reliability of each branch in phylogram. As a general rule if the bootstrap value for a given interior branch is 95% or higher than the topology of that branch then the value is considered "correct" [25].

RESULTS

3.1. Quantitation of DNA by electrophoresis

A thick band was seen above the 300kb band of λ Hind III marker (marker not shown here). This indicates high molecular nature of genomic DNA. In the electropherogram the bands of the size ~700bp (for sample MAB03) was observed against 100bp DNA ladder. There was no overlapping of the bands in the case of test organisms and that way the bands were clear.

3.2. Top 10 Sequences Producing Significant Alignments from NCBI

The sequences were checked for considerable alignments from NCBI. About 10 sequences showed significant alignments of which the maximum identity ranged from 86% to 100%. The maximum score ranged from 1135 to 742. The query coverage was found to be 100%. The summary of the results is depicted in Fig 1. The distance tree comparison of the study organism showed similar evolutionary similarity with *U. oligolepis* (Fig 2).

3.3. BOLD's search

Identification summary (Fig 3) showed the probability of placement (100%) along with taxonomic level and taxon assignment. The distance summary is illustrated in Fig 4. A species level match has been made. This identification is solid unless there is a very closely allied congeneric species that has not yet been analyzed. The bolds search showed top 20 specimen similarity with 88.14% to 100%. (Fig 5). The COI species database tree confirmed that the study organism belongs to the order Carangidae which resembled much similarity with *U. oligolepis* (Fig. 6).

3.4. Accession numbers of sequences closely related to the Test organism used in the analysis & their locations.

The test organisms were reviewed for close relations to the test organism (MAB03 -JX120607) for which the accession numbers were cross checked from the database. The accession numbers were HM422426, GU804916, GU805062 (Ontario) and FJ237963 (China).

3.5. Profiling the barcode region of *Uranoscopus oligolepis*

Nucleotide composition summaries were obtained and shown in Table 1. The table represents the Molar concentration of DNA nucleotides in the COI region *U. oligolepis* sample (MAB03) from Parangipettai waters versus closely related organisms. The results showed that the Cytosine content was high (31%). The least molar concentration was observed in Guanine (19.5%) and Adenine (19.6%). Thymine was the second predominant in molar concentration next to thymine which is followed by adenine. The G+C content was found to be 49.6% and A+T content was 50.4%. Upon comparison with other samples the GC content ranged from 49.6% to 52.3% and the AT content ranged from 47.5 to 50.4%.

3.6. Barcode protein profiling

3.6.1. DNA to Protein

The translation alignment was optional, and amino acids were displayed as a 1-letter amino acids code. Amino acid composition summaries and plots were obtained by choosing “Amino Acid Composition” from the “Protein” submenu of the “Sequence” menu. Bar plots showed the Molar percent of each residue in the sequence (Fig 7). Amino Acid plots and summaries were similar, though residues other than the standard 20 amino acids were ignored. Leucine and Alanine content was high in the amino acid composition. A helical wheel is a type of plot or visual representation used to illustrate the properties of alpha helices in proteins. The sequence of amino acids that make up a helical region of the protein's secondary structure are plotted in a rotating manner where the angle of rotation between consecutive amino acids is 100°, so that the final representation looks down the helical axis. The plot reveals whether hydrophobic amino acids are concentrated on one side of the helix, usually with polar or hydrophilic amino acids on the other (Fig 8)

3.6.2. CLUSTAL W

The similarities between two or more DNA sequences were compared using multiple sequence alignments. The query sequences were posted on the query box in Clustal W from the tools option of EMBL. The results page displays the similarities between the sequences. The similarities in sequences of the study animal with intra species is shown in fig 11. Minimum evolutionary distance of 0.1 in the scale was observed (Fig 9).

DISCUSSION

The efficiency of DNA barcoding has been reported in the detection and description of new cryptic species [26, 27, and 28]. This identification tool can clearly give support to improve classifications and to critically examine the precision of morphological traits commonly used in taxonomy [29]. For a barcoding approach to succeed, within species DNA sequences need to be more similar to one another than those between species and recent studies confirmed that the majority of species examined are well delineated by a tight cluster of very similar sequences [30, 31, 32, 33 and 34]. The methodology requires that intra-species DNA barcode variation is substantially less than interspecies variation, allowing accurate identification of individuals [35]. In the present study *U. oligolepis* from China origin showed more similarity with *U. oligolepis* of Parangipettai waters than with the same species from Ontario. Phylogeographical signals and the arrangement of *U. oligolepis* from China in a separate branch indicate that environmental parameters influence genetic diversity among the same species of organisms.

Freshwater fishes show more population differentiation than marine species, although marine species can show significant differentiation [36] Indeed, several studies have already illustrated the advances provided by the iterative processes between morphological- and DNA barcode-based studies in [37,38 and 39]. Exploring the microscopic eukaryotic life diversity can be achieved by the COI-based barcode [40, 41 and 42]. The profiling study on the barcode regions of *U. oligolepis* revealed that barcode region was rich in cytosine and least in adenine content. The molar concentration of cytosine was found higher when compared to other nucleotides in barcode region of *U. oligolepis* from Parangipettai waters where as the molar concentration of adenine was in the lower side. From the analysis it is assumed that the mitochondrial gene COI can be the potential barcoding region to identify an organism up to the species level. This study clearly revealed that COI could be a barcode sequence distinguishing *U. oligolepis* to its species level both through the phylogram and by search result of barcode of life database.

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Sequences producing significant alignments:

Accession	Description	Max score	Total score	Query coverage	E value	Max ident
FJ237963.1	<i>Uranoscopus oligolepis</i> voucher MBCSC:HN SY08334 cytochrome ox	1135	1135	100%	0.0	100%
FJ237961.1	<i>Uranoscopus oligolepis</i> voucher MBCSC:HN SY08506 cytochrome ox	1130	1130	100%	0.0	99%
FJ237962.1	<i>Uranoscopus oligolepis</i> voucher MBCSC:HN SY08335 cytochrome ox	1130	1130	100%	0.0	99%
HM422426.1	Perciformes sp. BOLD:AAJ8254 voucher BW-A7438 cytochrome oxi	908	908	99%	0.0	92%
GU805069.1	<i>Uranoscopus archionema</i> voucher ADC09_230.2#2 cytochrome oxid	848	848	99%	0.0	90%
GU804916.1	<i>Uranoscopus archionema</i> voucher ADC09_230.2#5 cytochrome oxid	848	848	99%	0.0	90%
GU805062.1	<i>Uranoscopus archionema</i> voucher ADC09_230.2#1 cytochrome oxid	843	843	99%	0.0	90%
GU804944.1	<i>Uranoscopus archionema</i> voucher ADC09_230.2 #4 cytochrome oxi	843	843	99%	0.0	90%
HQ920475.1	Perciformes sp. BOLD:AAG7346 voucher I.44773-019 cytochrome o	758	758	99%	0.0	87%
GU674178.1	Perciformes sp. BOLD:AAD7191 voucher BW-A6917 cytochrome oxi	742	742	99%	0.0	86%

Figure 1. Top 10 Sequences producing significant alignments from NCBI

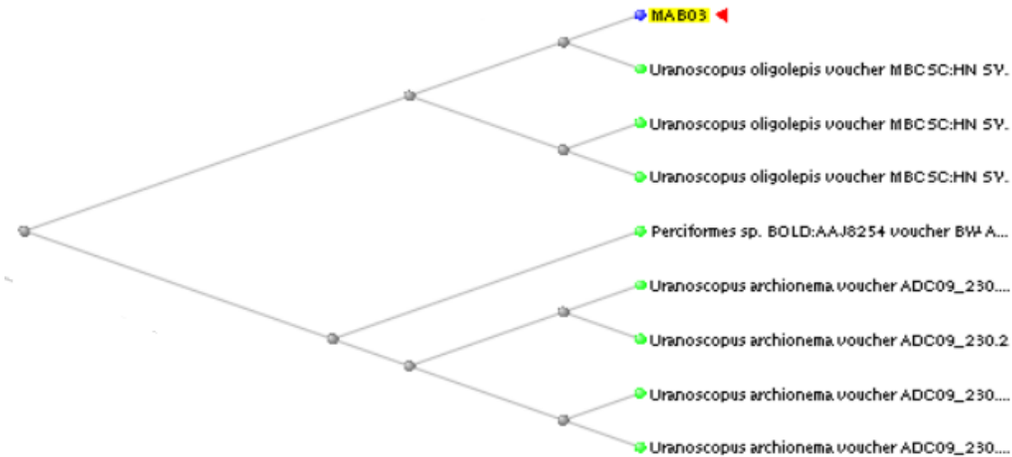


Figure 2. Distance tree of the study animal extracted from NCBI

Search Request:

Type : COI SPECIES DATABASE

Search Result:

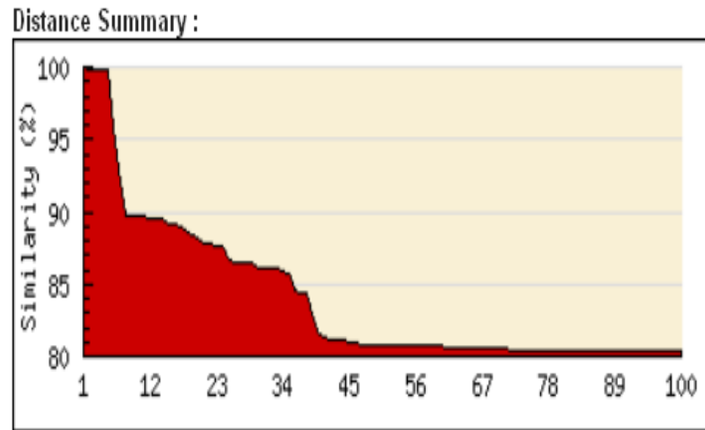
Identification Summary :

Taxonomic Level	Taxon Assignment	Probability of Placement (%)
phylum	Chordata	100
class	Actinopterygii	100
order	Perciformes	100
family	Uranoscopidae	100
genus	Uranoscopus	100

A species level match could not be made, the queried specimen is likely to be one of the following :

- Uranoscopus oligolepis*
- Uranoscopus* sp.

Figure 3. Results of the identification summary from BOLD search



Similarity scores of the top 100 matches

Figure 4. Results of the distance summary

TOP 20 Matches :

Phylum	Class	Order	Family	Genus	Species	Specimen Similarity (%)
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>oligolepis</i>	100
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>oligolepis</i>	99.84
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>oligolepis</i>	99.84
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	sp.	99.84
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>oligolepis</i>	99.83
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	sp.	95.83
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>kaianus</i>	91.99
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>n.spscalymape</i>	89.74
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>cognatus</i>	89.74
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>archionema</i>	89.74
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>archionema</i>	89.74
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>archionema</i>	89.58
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>archionema</i>	89.58
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>archionema</i>	89.58
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>japonicus</i>	89.08
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>japonicus</i>	89.05
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>japonicus</i>	88.94
Chordata	Actinopterygii	Pleuronectiformes	Cynoglossidae	Cynoglossus	<i>arel</i>	88.7
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>bicinctus</i>	88.32
Chordata	Actinopterygii	Perciformes	Uranoscopidae	Uranoscopus	<i>bicinctus</i>	88.14

Figure 5. Results of top 20 specimen similarity with 88.14% to 100%

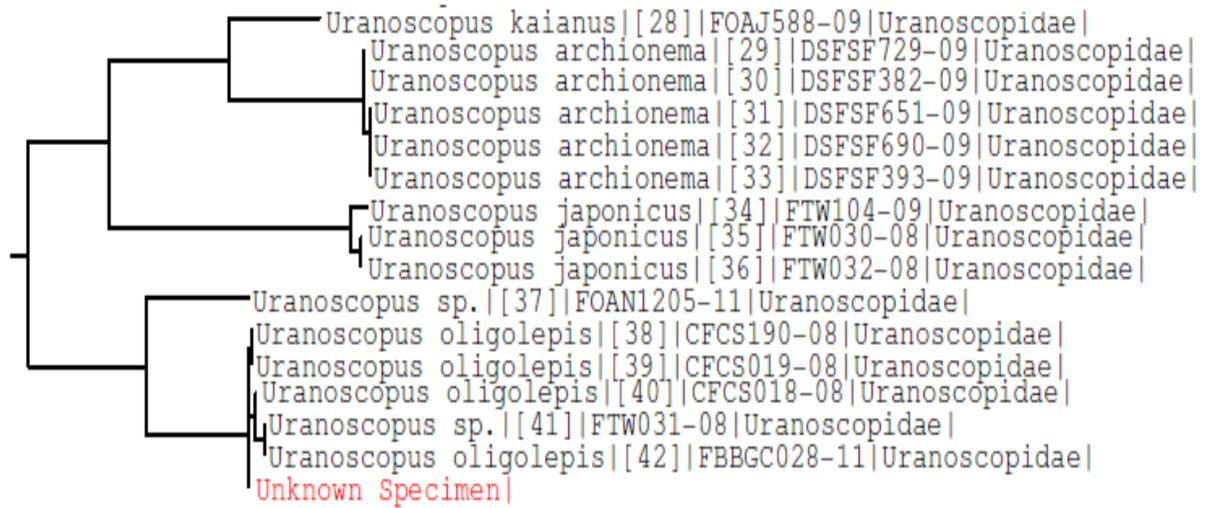
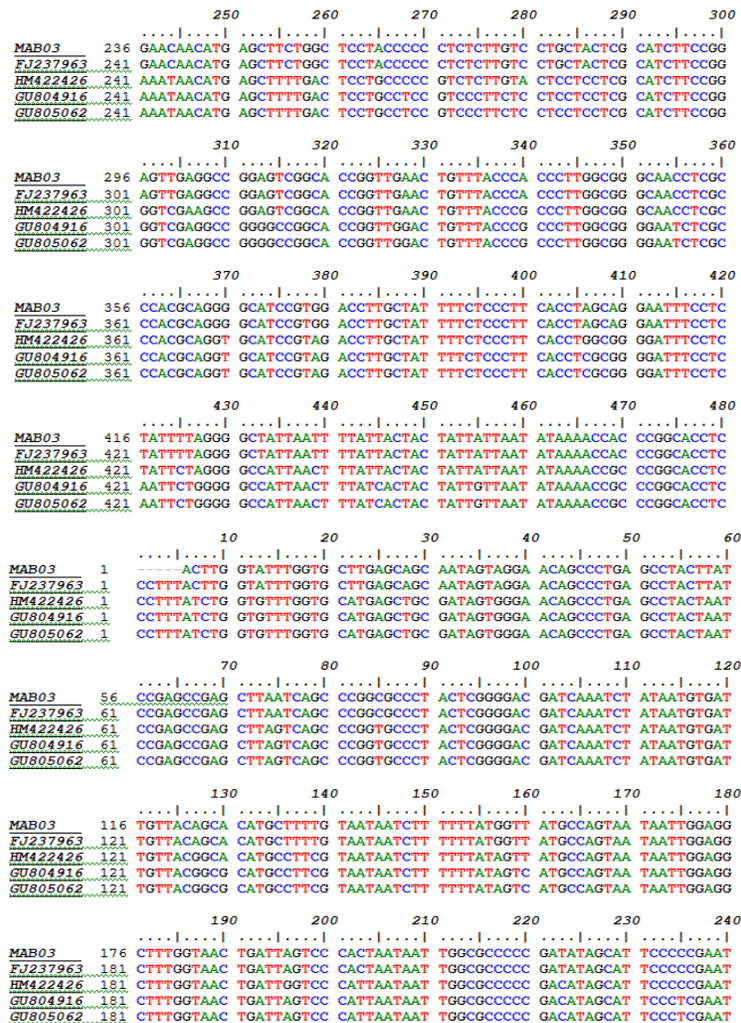


Figure 6. COI species database tree



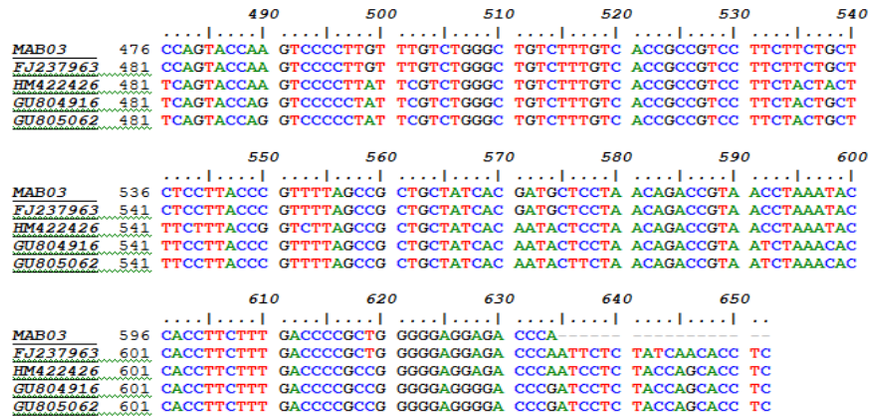


Figure 7. Clustal W alignments for the test organism versus closely related specimens

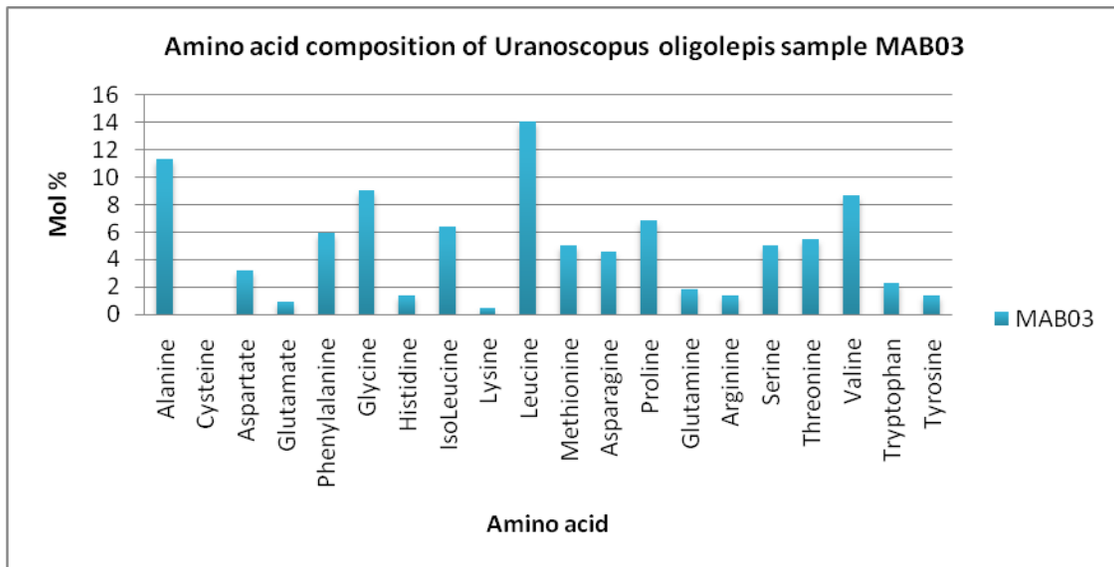


Figure 8. Graph showing hypothetical barcode profiling of *T. blochi* of Parangipettai waters

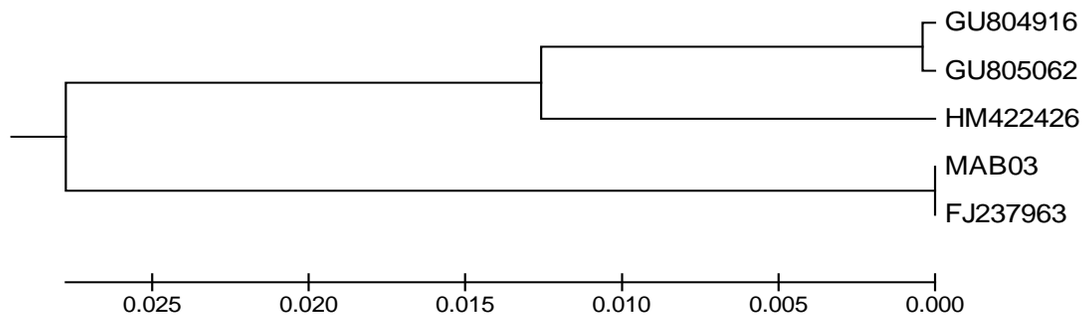


Figure 9. Evolutionary relationships of taxa of study animal (MAB03 - JX120607) by UPGMA Method

Table 1. The Molar concentration of DNA nucleotides in the COI region *U. oligolepis* sample from Parangipettai waters versus closely related specimens

Accession ID	Base pair length	G+C content (%)	A+T content (%)	Nucleotide Number and Mol%			
				A	T	G	C
JX120607 (MAB03)	629	49.6%	50.4%	134 21.3%	183 29.1%	128 20.3%	184 29.3%
FJ237963	652	49.2%	50.8%	139 21.3%	192 29.4%	128 19.6%	193 29.6%
HM422426	652	50.6%	49.4%	136 20.9%	186 28.5%	133 20.4%	197 30.2%
GU804916	652	52.5%	47.5%	127 19.5%	183 28.1%	140 21.5%	202 31.0%
GU805062	652	52.3%	47.7%	127 19.5%	184 28.2%	140 21.5%	201 30.8%