

Phylogeny and molecular delineation of leiognathids in the Southeast coastal waters of India

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Abstract

DNA barcoding is very useful in identification of animal species and resolving taxonomic ambiguity of morphologically similar animals. The aim of the present study was to investigate molecular identification of like Leiognathid fishes based on cytochrome c oxidase I (COI) gene sequence. Molecular level identification and comparison of six species from four genera of Leiognatidae were done. Results showed that A + T content of Leiognathids ranged from 55.27 to 53.32%. Mean genetic diversity within the species and overall species diversity within the family were calculated to be 0.004 and 0.178, respectively. Maximum K2P genetic distance (0.255) was observed between *Gazza minuta* and *Leiognathus equulus*. We obtained a monophyletic phylogenetic tree for Leiognathids with two major clades. One major clade contained only *Gazza*, and *Eubleekeria*, *Karalla* and *Leiognathus*, were grouped under another clade. Genus Leiognathus, which is not sexually dimorphic, get clustered as a separate sub-clade. Thus COI gene sequence could be used for species identification and delineation of complex groups like Leiognathids.

Keywords: DNA Barcoding, COI, Genetic variation, Leiognathidae, Phylogeny

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Introduction

Leiognathids commonly called ponyfishes, silverbellies or slipmouths and small sized. laterally compressed fishes inhabiting shallow water in marine and estuarine water with bioluminescent characteristic and schooling behaviour. They constitute nine genera that contains approximately 50 species (Sparks and Dunlap, 2004; Chakrabarty et al., 2010; Seah et al., 2012; Jawad et al., 2013; Eschmeyer et al., 2016) . Ponyfishes are mainly caught in bottom trawls, mostly treated as trash fish and are converted into fishmeal, poultry feed or may be marketed as fresh or dry - salted (Woodland et al., 2001). In wild, they are the crucial components of food chain and food web of marine ecosystem (Seth et al., 2016). They are internally and externally conservative in terms of morphology except at light organ system (LOS). The morphology based phylogenetic studies are less known in this family, and taxonomic comparison below family level is quiet difficult (Sparks et al., 2005). Even though, classification and identification of ponyfishes are mainly based on the phenotypic characters with detailed inspection of specimen (Mazlan and Seah, 2006) and molecular studies such as DNA barcoding makes it easy (Azaz et al., 2021).

DNA barcoding is an omnipresent system of identification of species and their products, based on sequence variation in mitochondrial gene cytochrome c oxidase I (COI) and new specimens can be identified by

analysing the gene sequence with barcoding reference library such as GenBank and **BOLD Systems** (Hajibabaei et al., 2007; Lakra et al., Molecular based recognition by utilizing DNA sequence diversity among different species aid to identify organisms and to reduce based morphology taxonomic ambiguity even in the identification of new or cryptic species and also to intensify taxonomic revisions (Chakrabarty and Sparks, 2007; Candek and Kuntner, 2015). Inaccuracy in identification morphological often happens and it had reported by many researchers in several groups like dinoflagellates organisms (Culverhouse et al., 2003) and some fish larvae (Ko et al., 2013). The Fish Barcode of Life campaign (FISH-BOL). an international research collaboration centre, is a DNA barcode library for reference and to monitor the DNA barcode project (Sachithanandam and 2020). Mohan. **Taxonomical** identification of different animals like fishes, birds and bats, through the COI gene sequence has been throughout the world and are used in the substantiation of species (Hebert et 2003. Ward etal.. 2005). al.. Trustability of DNA barcoding relies on barcode gap, which is the discontinuity in values of inter and intra-specific divergence, and more barcode gap reflects more accurate results (Hebert et al., 2004; Meyer and Paulay, 2005; Dasmahapatra and Mallet, 2006; Meier et al., 2008).

Molecular identification of several ponyfishes has been studied in the United Arab Emirates (Ludt et al., 2020) and Peninsular Malaysia (Seah et al., 2012) based on COI gene and in Perhent Islands of Malaysia based on 16S rRNA gene (Seah et al., 2008; Seth and Barik, 2021). In India, Banu et al. (2020) studied only three species of Leiognathids from Karaikal region including. This study is aimed at molecular studying analysis and phylogenetic characterisation of six Leiognathids species of including Eubleekeria splendens, Gazza achlamys, G. minuta, Karalla daura, K. dussumieri and Leiognathus equulus which are abundantly available in Tamil Nadu coast of India, using COI gene sequences.

Material and methods

Sample collection

Six morphotypes of Leiognathids were collected from Parangipettai landing centre, Tamil Nadu, India (11.4831° N, 79.7729° E) from January to June, 2018. The morphotypes were identified by FAO sheet (Bookstein *et al.*, 1985). After collection, the caudal fin of ten fishes from each morphotypes were dissected out and preserved in 95% ethanol for DNA isolation.

DNA isolation and PCR

The isolation of DNA was achieved by the standard phenol-chloroform ethanol method (Sambrook *et al.*, 1989). The absorbance ratio was taken at 260 nm and 280 nm (260/280) to estimate the quality of DNA. This DNA was diluted

with TAE buffer to a concentration of 100 ng/ml. PCR amplification of the COI gene was brought off in a 50 μ l volume PCR mix with 5 μ L of 10X Taq polymerase MgCl₂ (25 mM) buffer, 1 μ L of each dNTP (0.05 mM), 1 μ L of each primer (0.01 mM), 0.6 U of Taq polymerase, 2 μ L of genomic DNA and 36 μ L of double distilled water. *Fish F1*-

5'TCAACCAACCACAAAGACATTG GCAC3' FishR1and 5'TAGACTTCTGGGTGGCCAAAGA ATCA3' (Ward et al., 2005) were the universal primers employed in the amplification of the COI gene. The thermal cycle regime consisted of an initiation stage of 4 min at 95°C and variation of thermal subsequent exposure of 40 s at 94°C, 45 s at 54°C and 50 s at 72°C for 30 cycles and a final extension for 10 min at 72°C. The PCR product was run on 1.5% agarose after mixed with gel иL Ethidiumbrimode. The purified PCR product was sequenced by Sangers sequencing method.

Sequence analysis

The obtained sequences were edited with BioEdit v 7.2.6.1 and BLAST searched for similarity. All the sequences were submitted to NCBI through BankIt submission portal and got the accession numbers. Nucleotide composition, nucleotide diversity and genetic distance were calculated with the help of MEGA v 10.1.7 (Kumar, 2018) and the phylogenetic tree was based Neighbourconstructed on

Joining (NJ) method (Saitou and Nei, 1987).

Results and discussion

All the COI sequences were about 600 bp in length, without insertions, deletion or stop codon. In BLAST search, all the sequences showed more than 98% of similarity to specific species with more than 90% query coverage. Based on the BLAST results, the six morphotypes were identified as E. splendens, G. achlamys, G. minuta, K. daura, K. dussumieri and L. equulusand the accession numbers were MW843003, MW843014, MW843015, MW843018, MW843019, MW843034, MW843574, MW843598, MW844034, MW844035, MW844036, MW844041, MW844042, MW844046, MW846619, MW846620, MW850500 and MW856658.

Among six species, the the percentage of A+T content was ranged from 55.27 (L. equulus) to 53.32 (G. minuta) and the G+C content ranged from 46.68 to 44.73. The percentage values of nucleotides are displayed in These values Figure 1. accordance with the findings of Ward et al. (2005) where the G+C content in COI of fishes was ranged from 42.2 to 47.1%. The mean values of G+C content in this study resemble with previous observations of Lakra et al. (2011) and Seah et al. (2012) for Leiognathids and those values were 44.7% and 45.25% respectively. But a slight deviation from the observations of Banu et al. (2020), where the G+C content was 48.26%.

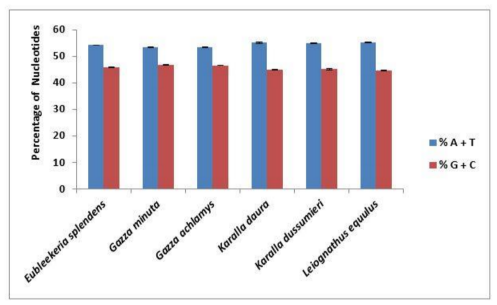


Figure 1: Percentage of nucleotide composition of COI gene in Leiognathids.

COI sequence divergence play a vital role in species identification. A fish species which shows a deep intraspecies divergence can be rejected by a taxonomic expert, from the same species (Bhattacharya *et al.*, 2016). According to Zhang and Hanner (2011), the mean intra-specific K2P distance for

marine fishes was 0.003 and the present study also displaysthe same intraspecific distance. Mean genetic distance within the species and the overall species distance were calculated to be 0.004 and 0.178, respectively, that show deviations from slight earlier observation by Banu et al. (2020). The mean intra-specific and intra-generic K2P distance of Leiognathids in this study is similar to the previous reports where it was 0.003 and 0.066 (Lakra et al., 2011) and 0.014 and 0.056 (Seah et al., (2017).

Among the six species of Leiognathids, the maximum interspecific K2P distance between *L*.

equulus and G. minuta was 0.255 and minimum between K. daura and K. dussumieri was 0.126 (Table 1). The observed low K2P distance between these two species is due to their taxonomic location in a same genus. Interestingly, a high K2P distance was observed between G. achlamys and G. minuta even though they are also coming under same genus. Maximum inter-generic K2P distance observed between Leiognathus and Gazza (0.228) and minimum was between Karalla and Eubleekeria (0.187). Maximum intrageneric distance was 0.118 in Gazza and minimum in both Eubleekeria and Leiognathus that was 0.007 (Table 2).

Table 1: K2P Genetic distance between and within species (bold).

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	E. splendens	G. minuta	G. achlamys	K. daura	K. dussumieri	L. equulus
E. splendens	0.007					
G. minuta	0.249	0.003				
G. achlamys	0.202	0.196	0.000			
K. daura	0.191	0.236	0.199	0.004		
K. dussumieri	0.183	0.227	0.189	0.126	0.001	
L. equulus	0.211	0.255	0.201	0.210	0.175	0.007

Table 2: K2P Genetic distance between and within genera.

	Eubleekeria	Gazza	Karalla	Leiognathus			
Eubleekeria	0.007						
Gazza	0.226	0.118					
Karalla	0.187	0.213	0.077				
Leiognathus	0.211	0.228	0.192	0.007			

The mean genetic distance within the genus is around 18 fold higher than the mean genetic distance within the species and this observation reflects the earlier reports on genetic variation studies of family Nemipteridae from Malasyian water by En *et al.* (2019). Increase in genetic diversity as we go from lower to higher taxonomic categories and it was supported by the

findings of Habert *et al.* (2004) and Lakra *et al.* (2011).Intra-specific variation ranged from 0.007 (*E. splendens*) to 0.00 (*G. achlamys*) and these K2P values may less and this should be confirmed by using more number of individuals. Similar low intra-specific genetic variation was observed by Banu *et al.* (2020) in Indian Leiognathids, Ward *et al.* (2005)

in Australian fishes and En *et al.* (2019) in fishes of Nemipteridae. Barcoding gap is the difference in the distribution of pairwise distance among intraspecific and inter-specific individuals, and the branch length among interspecies individuals serves to be much deeper than the intra-specific individuals (Meyer and Pauly, 2005).

Neighbour-Joining tree was constructed based on K2P boot strap method and *Caranx sexfasciatus* (Family: Carangidae) was selected as out group according to the postulates of Sparks *et al.* (2005) that Gerreids and Carangids are close relatives of ponyfishes. Figure 2 shows that family

Leiognathidae is monophyletic and the individuals belong to same genus and species are clustered as a definite clade high bootstrap Phylogenetic tree of Leiognathids consists mainly of two clades, first major clade comprises of genus Gazza which include G. achlamys and G. minuta. Second major clade includes genus Karalla, Leiognathus Eubleekeria. Genera Leiognathus do not exhibit sexual dimorphism even in the case of LOS (Sparks et al., 2005; Chakrabarty et al., 2011; Seah et al., 2012) and it gets clustered as a separate group in the second major clade.

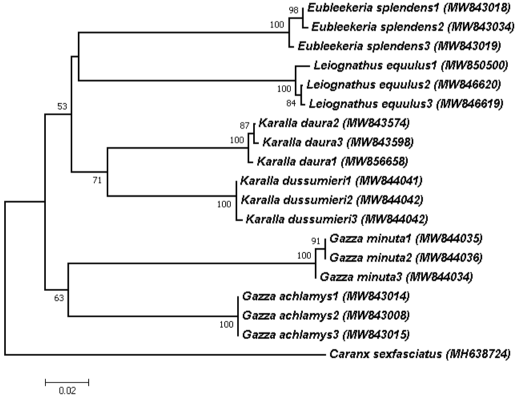


Figure 2: Neighbour joining tree of Leiognathidae based on COI sequences.

Anatomical changes of LOS in sexually dimorphic ponyfishes, may result in reproductive isolation and thus to sexual selection. Sexual selection can cause continuous diversification irrespective of habitat or environmental factors. Diversification rate ponyfishes are higher in sexually dimorphic taxa than in non-sexually dimorphic (Chakrabarty et al., 2011) and it reflects the clustering pattern of Leiognathus in this study. Karalla seems to be a valid genus in the NJ tree and it supports earlier findings of Seah et al. (2012) with the 16S rRNA sequences. Leiognathus equulus and E. splendens are found to be close relatives in the present study and which is not in accordance with the result obtained by Banu et al. (2020) by COI sequences as well as Seah et al. (2012) by 16S rRNA sequences and this may be due to the quality of sequences used in these studies. Monophyly of Gazza is very evident from the phylogenetic tree and which is similar to the previous findings of Spark and Dunalp (2004) and Ikeiima *et al*. (2004).paraphyly of Leiognathus was reported based on COI sequences (Spark and Dunalp, 2004; Ikejima et al., 2004) and 16S rRNA (Seah et al., 2008) but the same pattern is not observed in the present study. This may be due to the consideration of E. splendens as L. splendens in their study.

Conclusion

Misidentification exists in Leiognathids because of the morphological similarities among the species. Molecular identification is an inevitable technique and sufficient data that should be added in the BOLD system. Present study on Leiognathids from Parangipettai coast of Tamil Nadu provides taxonomists with new information the molecular to of identification Leiognathids. Grouping of non-sexually dimorphic species, mean genetic distance values within and between groups, G+C ratio of Leiognathids and monophylystatus of genera in this study are very relevant terms of their molecular identification and comparison. Even though the study is based on COI gene sequences, the findings are correlated with COI, and 16S rRNA sequences of previous reports. In short, it is understood that molecular phylogeny of Leiognathids is in compatible with their morphological delineation.

Conflict of interest

All the authors declare that, there is no conflict of interest.

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