DNA barcoding of *Schizothorax* species from the Neelum and Jhelum Rivers of Azad Jammu and Kashmir

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Abstract

The mitochondrial *Cytochrome Oxidase 1* gene is used as a standardized, authenticated, and reliable genetic marker for a global species-level bio-identification system. The present study was conducted to determine whether barcoding can help accurate species identification in fishes. The overall base composition of *Schizothorax* species was 29.6% of T, 25.5% of C, 26.5% of A, and 18.4% of G, A+T content 56.1% and G+C content 43.9%. The Ts/Tv bias (R) was 2.51. Complete COI gene was amplified using PCR and sequenced from 17 samples collected from river Neelum and Jhelum, and identification of species were done by following Mirza (1991), Jhingran (1991) classification and also through BOLD (99.3 to 99.9%) and NCBI (99.6 to 99.9%) reference sequences of those species. Multiple alignments of COI mtDNA gene resulted in a range of 1535–1551 base pairs. Out of 1535 consensus sites, 1490 were constant, 61 characters were variable, in which 54 were parsimony informative, and 7 variables were parsimony uninformative. This is the very first study reported from a reservoir of cold water bodies of Azad Kashmir which have a great potential for conservation of cold water fish species. We emphasized that, DNA barcoding is an accurate, reliable and has the great potential for identification of freshwater fish species.

Accurate identification of fishes is very important in many areas and would improve the fish conservation and ecosystem research and contribute for long-term fish management and its sustainability. For this purpose, a large variety of DNA and protein-based methods has been used. A short fragment of *COI* gene (655 bp) has been used for identification of species from 30 years, but in different laboratories, different DNA sequences have been also used for species identification (Hebert et al. 2003). The family Cyprinidae containing the genus *Schizothorax*, are locally known as snow trout, containing 20 genera and more than 150 species throughout the world (Mirza 1991). The genus *Schizothorax* contains the remarkably similar morphology, and difficult to distinguish based on the external morphological characters. For the current study, *Schizothorax* samples were collected from the river Neelum and Jhelum Azad Kashmir, Pakistan (34°23′03.0″N and 73°27′53.8″E). Total DNA was isolated by standard phenol–chloroform extraction by Sambrook et al. (1989). Sequencing primers of complete *COI* gene of *Schizothorax* species were designed by using program Primer-3. After sequencing and alignment, these sequences were deposited in Genbank for accession numbers (Table 1). The nucleotide composition, nucleotide and haplotype diversity and neutrality test were examined by MEGA6 (Tamura et al. 2013) and DnaSP 5.0 program.

A total of 17 COI barcodes were obtained from four species of Schizothorax. The absence of stop codons and well defined peaks indicated that co-amplification of nuclear pseudo-genes did not occur (Zhang & Hewitt 1996). The total 17 sequences of the same species were downloaded from Genbank (NCBI). Multiple alignments of COI gene resulted in a range of 1535–1551 base pairs. Out of 1535 consensus sites, 1490 were constant, 61 characters were variable, in which 54 were parsimony informative, and 7 variables were singleton. The nucleotides of *COI* gene sequenced were globally G-deficient (18.4%), whereas (A, 26.5%; C, 25.5%; T, 29.6%). Such type of nucleotide composition pattern has been widely stated in many other fish species with the smaller variations (Khan et al. 2015). The A+T content 56.1% and G+C content 43.9% showing an obvious anti-G bias as appear commonly in teleost fishes (Zhu et al. 2012). The average percentage divergence (K2P) distance of individual’s species of *S. plagiostomus* is 0.003% and 0.002% for *S. esocinus* and 0.001% for *S. niger* and *S. progastus*. There is high inter-specific sequence divergence for studied *Schizothorax* species i.e. 0.009% as compared to intra-specific sequence divergence. The possibility of inter-specific sequence divergence is due to hybridization and ancestral polymorphisms (Hajibabaei et al. 2006). The lack of difference in the mitochondrial sequence data of some *Schizothorax* species may be explained in terms of introgressive hybridization, incomplete lineage sorting.
rapid radiation in lineages, and multiple hits (homoplasy) (Qi et al. 2007).

The neighbour-joining (Figure 1) analysis of the COI barcode region placed *S. plagiostomus* and *S. esocinus* as sisters to *S. progastus* while, *S. niger* form a separate cluster. As the *S. niger* inhabiting cold streams and rivers is distributed in the inland waters of occupied Kashmir (Kullander et al. 1999), but in present study *S. niger* was first time collected from river Jhelum near Muzaffarabad city. This fish is restricted to upper part of river Jhelum due to damming of waters. However, fortunately due to floods, the eggs and larvae of *S. niger* migrated from upper (Eastern) side of river Jhelum (India) toward its lower side (Pakistan).

The COI gene of *Schizothorax* species was sequenced and aligned with the 17 global sequences available in BOLD and Genbank database using the Clustal W program. This alignment allowed grouping of species into definite clusters. These *Schizothorax* species showed maximum sequence homology with the BOLD (99.3 to 99.9%) and NCBI (99.6 to 99.9%) reference sequences of those species. These four species have 14 haplotypes with haplotype diversity ($H_d$) $0.9779 \pm 0.027$. The average value of nucleotide diversity ($P_i$) was $P_i: 0.00878 \pm 0.003$.

The transitional substitutions are outnumbered the transversional substitutions. The estimated $T_s/T_v$ bias ($R$) was $=2.51$. According to the Tajima D test and the Fu and Li $D^*$ and $F^*$ tests, the genetic variation between populations were

### Table 1. List of samples used in this study, including species name, code, sample locality and Genbank Accession numbers.

| No | Species name | Code | Sex | Genbank accession no. | Collection locality          |
|----|--------------|------|-----|------------------------|-----------------------------|
| 1  | *S. plagiostomus* | SP-AM | Male | KU317684 | River Jhelum, Air Port    |
| 2  | *S. plagiostomus* | SP-AX | Female | KU317689 | River Jhelum, Kohala      |
| 3  | *S. plagiostomus* | SP-BH | Male | KU317694 | River Jhelum, Domel       |
| 4  | *S. plagiostomus* | SP-AT | Male | KU317687 | River Jhelum, Domel       |
| 5  | *S. plagiostomus* | SP-AV | Male | KU317688 | River Jhelum, Chatter Kalass |
| 6  | *S. plagiostomus* | SP-AZ | Female | KU317690 | River Jhelum, Chatter Kalass |
| 7  | *S. plagiostomus* | SP-BG | Male | KU317693 | River Jhelum, Chatter      |
| 8  | *S. esocinus* | SE-AL | Male | KU317698 | River Jhelum, Ambor        |
| 9  | *S. esocinus* | SE-BG | Female | KU317699 | River Jhelum, Ambor        |
| 10 | *S. esocinus* | SE-BB | Female | KU317700 | River Jhelum, Ambor        |
| 11 | *S. esocinus* | SE-BE | Male | KU317701 | River Neelum, Chella Bandi |
| 12 | *S. esocinus* | SE-H  | Male | KU317702 | River Neelum, Chella Bandi |
| 13 | *S. niger* | SN-AS | Female | KU317703 | River Jhelum, Domel       |
| 14 | *S. niger* | SN-AW | Female | KU317704 | River Jhelum, Domel       |
| 15 | *S. progastus* | SPR-AU | Female | KU317705 | River Jhelum, Garhi Dupatta |
| 16 | *S. progastus* | SPR-AW | Male | KU317706 | River Jhelum, Garhi Dupatta |
| 17 | *S. progastus* | SPR-BC | Female | KU317707 | River Jhelum, Garhi Dupatta |

Figure 1. The phylogenetic analysis of *Schizothorax* species of present study and international sequences (with accession number) by neighbour-joining method using MegAlign program (DNASTAR).
not neutral \( (\text{Tajima } D = -0.957, \ p > .10; \ \text{Fu and Li's } D^* \ \text{test statistic } = 1.05942, \ p > .10; \ \text{Fu and Li's } F^* \ \text{test statistic } = 0.55331, \ p > .10) \). The negative values of Tajima's D indicated that the genetic variations between populations were not neutral under the random effects of genetic drift and mutation which reflect the excess of external mutation.

**Disclosure statement**

The authors alone are responsible for the content and writing of the paper. The authors report no conflicts of interest.

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