Assessment of Genetic Diversity of Freshwater Mud Eel (Monopterus cuchia) Using RAPD and RFLP Markers

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Abstract

Background: Monopterus cuchia, an economically important eel of Bangladesh, is confronting the challenge of population reduction and germplasm degeneration since the vast majority of fingerlings are collected from natural habitats and artificial cultivation of the fish is yet to be established.

Materials and Methods: 30 individuals of M. cuchia from a natural population of Northern-East part of Bangladesh were analyzed using 8 decamer primers and glutamine synthetase gene was digested with 2 restriction enzymes.

Results: A total of 735 bands with 228 polymorphic loci were detected among the selected 30 individuals by using RAPD assay while 100% polymorphism was revealed by all the primers. The genetic distance among the individuals was calculated by using the data from pair-wise similarity index where 36 groups of genetic diversities were measured while the lowest and highest genetic distance were found 0.58 and 0.97 respectively with an average genetic diversity of 0.81. The Nei's genetic similarity values were found from 0.03 to 0.6 where average value was recorded 0.30 which was found significant regarding genetic distance. Phylogenetic relationships using UPGMA clustering revealed linkage distance ranged from 3.6 to 6.24 and generated 6 clusters by 11 clades with the involvement of 22 individuals while the rest of the samples were connected to those clades with specific linkage distances. Genetic diversity of glutamine synthetase gene was analyzed with two restriction enzymes, e.g. -CfrI and +Hpy17BIII while both the enzymes digested the gene fragment at a length of 541 bp and polymorphism was determined in terms of wild type homogoggles, polymorphic homozygote and heterozygosity. The genetic diversity was observed by using the RFLP band analysis and four different groups of individuals were identified with the p-values of 0, 0.033, 0.05 and 1 respectively whereas different distances were found among the groups indicating the polymorphism among experimental individuals of glutamine synthetase gene.

Conclusion: Herein, RAPD and RFLP analysis indicating the rich genetic diversity of this fish in the experimental ecological habitat and this would come helpful for the conservation of germplasm diversity and to support the sustainable breeding program of M. cuchia.

Keywords: RAPD; RFLP; Glutamine synthetase; Monopterus cuchia; Genetic diversity

Introduction

Monopterus cuchia [1], also known as Gangetic mud eel, freshwater mud eel, Kuicha, or Kunche, [2] of synbranchiformes family under the order of synbranchiformes [3,4] is one of the common freshwater eels found in Bangladesh. Although the fish has high nutritional as well as medicinal values, it is consumed only by peoples of some tribes and few other castes [5]. The trade of the fish offers a great export fishery to at least 15 foreign countries, including China, Malaysia, Singapore, Japan and Taiwan and almost all the harvested fish are exported [6]. The fish has the ability to survive in harsh conditions as it possesses systems for both aquatic and aerial respiration [7] and also the ability to survive 90 to 132 days without having foods. These offers the culture of a large number of fishes in small tanks, aquarium and other vessels as well as the transportation to distant places using simple earthenware or plastic jars [1,8].

Development of social fishery can help the socioeconomic welfare of an area in a unique way [9]. But due to overfishing as well as water obstruction, destruction of habitat, water pollution, emergence of diseases etc. the populations of M. cuchia in Bangladesh is falling-off at an alarming rate [10,11] and consequently, it is recorded as a least concerned species by IUCN [12]. Careful monitoring and research have become essential for its existence in the natural bodies. Being commercially important and presumed as a low-cost enterprise to the poor farmers, freshwater mud eel can be a good choice for artificial culture to meet the increasing demand of animal protein in Bangladesh as well as to earn foreign currencies [13] to help the development of our national economy.

As it has been mentioned earlier, M. cuchia can tolerate harsh conditions, its culture is easier and highly profitable than a number of other small size fish culture activities [14,15]. Several techniques of capturing the fish have been established, however, no cultural practice has been consecrated and only a few works have from different aspects been done on this so far [6,5,16-22]. The mud eel has also been identified for polyculture in seasonal and perennial ponds as well as in paddy fields, and in Bangladesh, some experiments were performed in captivity for observing growth, survival and diet condition [23].

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In order to develop a scientifically proved sustainable eel culture system, it is necessary to produce huge amounts of fry artificially while unsuccessful induced breeding was observed without spawning responses [24,25]. The unprosperous reports regarding induced breeding of the fish refer to the necessity of the studies to be conducted on the genetic characterization of the fish to obtain reliable seeds for aquaculture and to develop a sustainable selective breeding program. But, unfortunately, so far, no large-scale works have been found on the genetic characterization of M. cuchia barring some tiny DNA fingerprinting analysis using RAPD markers in China [4] and in Bangladesh [26-28].

Understanding the genetic diversity at molecular level is a prerequisite in developing a sustainable eel culture system along with effective conservation and utilization of M. cuchia genetic resources, and, in this circumstance, this current study represents a novel genetic survey investigating the genetic status of M. cuchia in Bangladesh. Genetic diversity is considered as a key tool for the development of breeding programs and conservation of species and in this study, we have assessed the genetic diversity of M. cuchia using RAPD (Random Amplified Polymorphic DNA) technique. Additionally, we have observed the genetic diversity of Glutamine Synthetase (GS) gene using PCR-RFLP technique, which is involved in the detoxification of ammonia and enabling the fish to survive in the extreme conditions and regarded as an important player for analysis of breeding biology of the fish [29,30]. We studied the genetic diversity and relationship among the M. cuchia individuals collected from the Hakaluki Haor (an aquatic ecosystem), a natural habitat of the fish in Bangladesh. Taking into account that, the fish possess high export demand and excellent nutritional values, the long-term goal of this work is to help the conservation of the species, sustainable commercial culture of it and also to tip an alternative livelihood for people involved in the fisheries sector [30].

Research Methodology

Collection of sample fishes and identification

Fishes were collected randomly by the help of the officials of regional office of Department of Fisheries, Government of Peoples Republic of Bangladesh and professional fish catchers from Hakaluki Haor (Haor=a natural water body; 23°35’-24°44’ N, 92°01’-92°09’ E) of Northern-East part of Bangladesh. A total of 30 individuals were selected for the study. Collected fishes were identified using morphometric characteristics as described by Shafi and Quddus, Rahman, Talwar and Jhingran [31-33]. Collected fishes were transferred to the Animal Keeping Laboratory of the department of Genetic Engineering and Biotechnology (GEB), Shahjalal University of Science and Technology (SUST), Sylhet 3114, Bangladesh and were kept in live fish preservation tanks until sacrificed humanely for tissues to be isolated. The study was undertaken after approval from the Animal Care and Ethics Committee of SUST.

Tissue isolation and DNA extraction

Each sacrificed fish was dissected and liver tissue was isolated from each individual. Isolated tissues were washed using distilled water and 70% alcohol and preserved separately in big Eppendorfs in 100% alcohol at -20°C temperature freezer cabin. The desired genomic DNA was extracted by using a commercially available kit, (Bioline) and DNA quality was checked on 0.8% agarose by gel electrophoresis using 100 bp ladder (GeneRuler™) and DNA quality was checked on 0.8% agarose by gel electrophoresis and using gel documentation system with digital camera (Panasonic DMC-fs20) image was captured. A good quality DNA was observed from each of the individual where 1 kb Plus DNA ladder (GeneRuler™) was used to compare the quality of the DNA. Finally, the extracted DNA of each sample was stored at -20°C. In order to perform PCR based examinations, a concentration of DNA around 200 ng/μl was required and hence, specimen with variable concentration of the DNA was adjusted to ~200 ng/μl before used in PCR.

PCR-RAPD assay

Eight decamer primers, e.g. B-03 (CATCCCCCTG), OPF 14 (TGCTGCAAGGT), C-04 (CCGCATCTAC), OPB 05 (TGCGCCCTTC), OPB 08 (GTCCACACGG), OPB 12 (CCTGACCGCA) OPB 19 (ACCCCCGAAG) and UBC 122 (GTAGACGAGC) (Operon Technologies Inc. USA) were adopted and utilized for RAPD assaying of freshwater mud eel. The PCR amplification was carried out in a total volume of 15 μl containing 8 μl of master mix (Promega Hot Start), 1 μl of 20 μM working solution of primer, 2 μl of template DNA and 4 μl deionized distilled water. After an initial 1-minute denaturation at 94°C, followed by 30 cycles of 94°C denaturation for 1 minute, annealing for a minute at 30°C for UBC122, 32°C for B-03, 33°C for OPF 14, C-04 and OPB05, and 34°C for OPB 08, OPB 12 and OPB 19 and 72°C extension for 2 minutes. A final 7-minute extension at 72°C completed the reaction.

RAPD data analysis

Different formulae, calculations and software were used for analyzing RAPD based genetic diversity of this experiment. The software AlphaEaseFC 4.0 was used for measuring molecular weight of bands. The proportion of polymorphic loci was calculated by the equation:

\[ P = \frac{n_{pol}}{n_{total}} \]

where, \( P \) = proportion of polymorphic loci, \( n_{pol} \) = number of polymorphic loci and \( n_{total} \) = total number of loci). Average number of alleles per locus was calculated by the following formula:

\[ N = \frac{1}{k} \sum_{i=1}^{k} n_{i} \]

Here, \( k \) is the number of loci and \( n_{i} \) is the number of alleles detected per locus. Genetic distance analysis by using pair wise similarity was calculated by the formula:

\[ D = 1 - \frac{N_{xy}}{N_{x} + N_{y} - N_{xy}} \]

where \( D \) = genetic distance between sample x and y, \( N_{xy} \) = number of bands shared by sample x and y, \( N_{x} \) = number of bands in sample x and \( N_{y} \) = number of bands in sample y). Using the equation,

\[ F = \frac{2N_{xy}}{N_{x} + N_{y}} \]

where \( F \) = Nei’s genetic similarity, \( N_{xy} \) = number of shared bands between x and y, \( N_{x} \) = number of bands in x and \( N_{y} \) is the number of bands in y). Nei’s genetic similarity was calculated. Linkage distance was calculated with squared Euclidean distances using the software new.stat and genetic relationships among individuals were observed based on linkage distance using the software “Statistica.”
PCR amplification for glutamine synthetase gene

Gene expression of the freshwater mud eel, *M. cuchia* was also analyzed using the Restriction Fragment Length Polymorphism (RFLP) method. A 541 bp long partial sequence of glutamine synthetase gene was allowed with two restriction enzymes. Gene specific primer of glutamine synthetase, accession no. Gbase 152041 (5'-GAGGGGCCTCAACAGCCTATGTA-3') and accession no. Gbase 152042 (5'-CTGAAGTTGTGATGGACGGCG-3') [34] were used for PCR amplification for the RFLP assay. The PCR amplification was carried out in a total volume of 15 μl containing 8 μl of master mix (Promega Hot Start), 1 μl of primers, 2 μl of template DNA and 4 μl deionized distilled water. The protocol for the process was as follows: preheating for 3 minutes at 94°C followed by denaturation at 94°C for 1 minute, annealing at 64°C for 1 minute and 2 minutes for elongation at 72°C and final step at 72°C for 7 minutes to allow complete extension of the amplified fragments. The efficiency of the PCR products of glutamine synthetase gene of *M. cuchia* was checked by agarose gel electrophoresis with 1.2% agarose. 1 kb plus DNA ladder (GeneRuler™) was used for checking the length of the DNA fragments at 541 bp length. The electrophoresis was run at 75 V for 40 minutes. The gel was then placed in gel documentation system and photograph was taken by digital camera (Panasonic DMC-fx20). After getting good bands of GS gene compare with ladders where ladder range was started from 250 bp lengths the PCR DNA products were kept in freezer at -20°C for further analysis by restriction enzymes.

PCR-RFLP analysis

The glutamine synthetase gene had an approximate size of 541 bp and it was digested with two restriction enzymes (+Hpy17II* [TCNNGA] and +CfrI [YGGCCR]) targeting a single cleavage site to detect polymorphism of freshwater mud eel in Bangladesh. Enzymes were identified using glutamine synthetase gene base pairs through blasting and the restriction site of the enzyme was searched out from the EBI nucleotide data base. The RFLP mix was prepared using milli-Q water, enzyme and enzyme's buffer and the mixture was run through PCR products at 37°C for 2 hours for the enzyme + Hpy17II* and at 55°C for two hours and 80°C for 20 minutes for the enzyme +CfrI. The efficiency of the PCR-RFLP products of freshwater mud eel was checked by 1.5% agarose at 100 V for 60 minutes. An image of the DNA fragments was taken by a digital camera through gel documentation system and photograph was compared with the 1 kb plus DNA ladder (GeneRuler™) while the ladder was started from 250 bp length.

Calculating the proportion of shared fragments and genetic distances

The restriction pattern of individuals of freshwater mud eel were compared to detect polymorphism, using matrices based on either presence or absence or relative abundance of bands. The similarity index or proportion of shared fragments (F) was calculated by comparing banding patterns between the two individuals. The formula used for this calculation is:

\[
F = \frac{2N_{xy}}{N_x + N_y}
\]

\(F\) = Nei's genetic similarity, \(N_{xy}\) = number of shared bands between \(x\) and \(y\), \(N_x\) = number of bands in \(x\) and \(N_y\) is the number of bands in \(y\).

A matrix of genetic distances (p-value) between all individuals based on dissimilarity indices were calculated and the formula used for this calculation is:

\[
P = 1 - \left[ \frac{(F^2 + 8F)^{\frac{1}{2}} - F^{\frac{1}{2}}} {2} \right]
\]

Here, \(r\) is the average number of enzyme sequences. Both calculations were done by using Microsoft Office Excel 2016.

Results

RFLP based genetic diversity

DNA profiling and data scoring: DNA profiling and data scoring were studied separately for each primer. Each amplified banding profile was defined by the presence (1) or absence (0) of bands at particular positions. The bands of different primers were seen in different levels of length of DNA. A total of 735 bands with 228 polymorphic loci were detected among the selected 30 individuals (Table 1). Polymorphic loci were revealed by all the primers with 100% polymorphism. The highest number of bands (120) was amplified by the primer C-04, while UBC122 produced the lowest. The highest number of bands (4.00) per individual was amplified from the primer C-04 and the lowest number of bands (1.63) per individual was amplified by the primer UBC122.

Genetic distances

Inter individual pair-wise similarity of *M. cuchia* was studied and based on the similarity data, 13 diverse groups of individuals (14, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2 and 1) were found. The genetic distance among individuals of *M. cuchia* was calculated by using the data from pair-wise similarity index where 36 groups of genetic diversities were measured. The lowest genetic distance was found between the individuals 7 and 16 (0.58) followed by 7 and 11 (0.59) and 11 and 20 (0.60), and the highest genetic distance was recorded 0.97 between the individual pairs of 17 and 22, 14 and 24 and 10 and 24 and second highest was found 0.96 between the individual pairs of 6 and 24 and 5 and 24 respectively. Relatively higher genetic distance was recorded in other individuals (Table 2). The average genetic diversity was found 0.8173716 and which indicates the good genetic status of this experimental fish in Bangladeshi nature especially in the experimental ecosystem. Considering the Nei's genetic similarity analysis, different values of similarities ranges from 0.06 to 0.6 were found by these 30 individuals where average value was recorded 0.301746 which was very relevant to genetic distance.

Genetic relationships among individuals

Based on Squared Euclidean Distances, the values of pair-wise comparisons of the linkage were computed from combined data of these experimental individuals. Different levels of values of linkage distance were found which was ranged from 3.6 to 6.24, and based on different linkage distances, cluster analysis using Unweighted Pair Group Method with Arithmetic Mean (UPGMA) was done to resolve the phylogenetic relationships among experimental individuals of *M. cuchia* (Figure 1). The UPGMA clustering system generated six clusters by 11 clades where 22 individuals/samples were involved to form 11 clades and the rest of the samples were connected to those clades with a specific linkage distance. Sample (S) 1 and Sample 4 (=S4) were found closely related as clade 1 which was created cluster 1 with clade 2 (S5, S6). Both of these clades were situated between the linkage distances of 3.5 to 4. Sample S3, S2, S13, S10 and S15 were individually connected to those clades with a specific linkage distance. Sample (S) 7 and Sample 8 (=S8) were observed approximately 4.9. Clade 3 (S7, S16) and clade 4 (S11, S20) were close enough between the linkage distance of 4.4.5 and formed...
<table>
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<th>Primers</th>
<th>Size of DNA (bp)</th>
<th>Total no. of Bands</th>
<th>Polymorphic loci (No.)</th>
<th>Monomorphic loci (No.)</th>
<th>Polymorphic loci (%)</th>
<th>Avg. no. of Bands per sample</th>
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<td>40-1473</td>
<td>120</td>
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Table 1: Summary of the bands revealed from eight RAPD primers.
Table 2: Genetic distance among individuals of M. cuchia.

Figure 1: Relationships among the 30 individuals of M. cuchia.
Genetic distance of 30 individuals of Monopterus cuchia.

Table 3: Genetic distance of 30 individuals of M. cuchia.
cluster 2 which was comparatively nearer to cluster 1. Clade 5 (S17, S28), clade 7 (S24, S26) and clade 8 (S29, S30) were very close to their respective pairs and their linkage distance was observed between 4-4.5 except clad 6 (S19, S27) which was around 4.6. All these clades were located under the cluster of 3 and formed cluster 4 when connected with cluster 2. S14 and S23 were found very similar to each other as well as S2, S18 and their linkage distances were seen 4.7 and 4.9 respectively. S23 was also found connected to the clade 9 (S9, S14) with a linkage distance of 5.2, clade 9 and S23 were placed under the cluster of 5 which was also connected to the cluster 3 and 4. Clade 10 was also linked to the cluster 5. The most distantly related samples were S8, S12 which produced a linkage distance of around 5.3 and they are connected through the most outer cluster, indicated cluster 6.

**RFLP based genetic diversity**

- Analysis of RFLP bands: PCR products of glutamine synthetase gene of freshwater mud eel, *M. cuchia* were digested with enzymes +CfrI and +Hpy17III and polymorphism was detected among the 30 individuals. These two enzymes have a single suitable restriction site to digest the glutamine synthetase gene of *M. cuchia* and a PCR amplified DNA was used without enzyme as a control to check proper enzyme activities, and this DNA did not cut and was found as a strong band on the gel indicating approximately 341 bp of the glutamine synthetase gene. 25 individuals were digested by the enzyme +CfrI and approximately 280 bp and 254 bp long fragments of DNA were found, comparing with 1 kb plus DNA ladder where the marker was started with 250 bp. These 2 fragments indicating the wild type homogygotes and rest of the 5 individuals were found uncut with single band by the +CfrI that is indicating the polymorphic homozygote. The same PCR products of glutamine synthetase gene were again digested by the enzyme +Hpy17III where 19 individuals were digested with two bands of 343 bp and 192 bp lengths indicating wild type homogygotes. The uncut DNA fragments with 6 individuals showed 541 bp lengths indicating polymorphic homogygote. Remaining 5 individuals with 3 bands e.g., 541 bp, 343 bp and 192 bp lengths were digested remark as heterozygocity.

**Observed genetic diversity**

Using the number of RFLP bands, proportions of shared fragments (F-value) were calculated from 30 individuals of freshwater mud eel where 7 different F-values such as 0.00, 0.4, 0.45, 0.5, 0.57, 0.67 and 1 were found. Using these F-values, genetic distances (p-value) were calculated and 7 different p-values such as 0.00, 0.02, 0.033, 0.04, 0.05, 0.06 and 1 were observed (Table 3). On the basis of genetic distances, four different groups of individuals were identified. 16 individuals were found in one group with zero (0) p-value. Second group with seven individuals were observed with p-value of 0.033. The third group with five individuals was recorded with p-value of 0.05 and finally, the fourth group with two individuals was recorded with p-value of 1. The genetic distance between individuals in group one and group two was found at p-value 0.02 which means they are closely related. Second group were also found at short genetic distance (p=0.04) with third group. The genetic distance between the individuals of third and fourth group was found 0.06, with large distance. Genetic distance between group one and three (p=0.04) as well as between group one and four (p=0.06) were found large distances and they are not closely related with each other.

**Discussion**

In this study, RAPD and RFLP based genotyping were performed to observe genetic status of freshwater mud eel, *M. cuchia* in Bangladesh. Using RAPD assay, in total 735 bands with 228 polymorphic loci was detected and polymorphisms were revealed by all the primers. Two previously conducted studies on the genetic diversity of freshwater mud eel, *M. cuchia* using three RAPD primers has reported higher genetic diversities alike the findings of this work [18,24]. Experimental data indicates that intra-specific polymorphism was observed 100% in this study; however, previous works has reported polymorphism with monomorphic loci as well [18,24,35]. Yin et al. [36] assessed the genetic differentiation and variation of the wild and raised swamp eels *M. albus* using RAPD technique and the results showed two wild polymorphic loci was 44.79% and 36.5%, while, in a different work, the percent polymorphic loci of wild samples was found to be 60.6% to 71% range and cultured samples was found to be in the 54% to 56.3% range by ISSR analysis [23]. The findings of this result disagree with the findings of [36], Wei et al. [35], Alam et al. [6], Li et al. [23], Miah et al. [18] and Miah et al. [24], while lowest polymorphism was observed there. Though little bit higher polymorphism was found by Alam et al. [6] in a population of freshwater mud eel in Bangladesh and Ruzainah et al. [35] also found higher genetic diversity with RAPD fingerprinting of two loaches, *Pangio pipera* and *Pangio filinaris*. But this work has come with the higher genetic diversity among all the previously conducted works. In the rice field eel (*Monopterus albus*), 30 microsatellites were analyzed by AFLP of sequences where 13 loci exhibited polymorphism and these loci should provide a sufficient level of genetic variation [37], while lots of polymorphic loci were recorded in the present study of *M. cuchia* by RAPD assay. The genetic diversity of Asian swamp eel *Monopterus albus* were analyzed with 16 polymorphic novel microsatellites with 11 loci [35] which was also lowest from the present study in *M. cuchia* by RAPD analysis. In this study, inter individual pair-wise similarity of freshwater mud eel were recorded; we found 13 different groups where highest and lowest similarity was 14 and 1 respectively, which are a bit different from the reports of Miah et al. [18] and Miah et al. [24]. Highest genetic diversity recorded in this study refers almost similar inference with the reports of Miah et al. [18], and Miah et al. [24]. Nei's genetic similarities were found identical to the report of Miah et al. [18] and Miah et al. [24]. Several values of linkage distances were found which ranges from 3.6 to 6.24, however, higher linkage distances were recorded in other populations by Miah et al. [18], Miah et al. [20]. A genetic relationship was observed through cluster analysis where 6 clusters were generated by 11 clades. Overall, higher genetic diversity has been recorded in this experimental using RAPD assay, which indicates the good genetic status of this fish in experimental nature. Miah et al. [18] found less genetic variation than the present study, but in another work, it was found higher [20].

In this study, partial sequence of glutamine synthetase gene of freshwater mud eel, *M. cuchia* was first time digested with two restriction enzymes and polymorphism was detected in an approximate size of 541 bp considering wild type homogygotes, polymorphic homogygotes and heterozygosity as well. The genetic diversity was observed with four different groups of individuals were identified and different genetic distances were recorded among the groups. The RFLP analysis of genetic polymorphism of glutamine synthetase gene was recorded higher than the findings of several previously reported studies [38]. Li et al. [15] analyzed hepcidin gene and they reported polymorphism in rice field *M. albus*. Tok et al. [37] analyzed glutamine accumulation and up-regulation of glutamine synthetase activity in the swamp eel, *M. albus*, however, no genetic diversity of glutamine synthetase gene of *M. albus* was studied. However, polymorphism was found among the population which is good for the experimental populations, though induced expression of downstream luc gene activities of *Monopterus albus* and *Channa straitus* were analyzed but no genetic diversity of this gene was analyzed [39].
Conclusion

Higher genetic diversity has been recorded in this study by RAPD and RFLP observation which means the good genetic status in experimental habitat. However, more studies are essential to reveal the most specified genetic status of this fish considering different genomic analysis in detail at population level.

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