Comparative Analysis on Mitochondrial Control Region DNA Diversity of Three Far Eastern Catfish (Silurus asotus) Populations

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Abstract
To investigate genetic diversity of the far eastern catfish (Silurus asotus) populations in the Yellow River, the Songhua River, and the Dayanghe River, mitochondrial control region (D-loop) DNA was analyzed for the samples collected in Zhengzhou (ZZ in the Yellow River), Haerbin (HEB in the Songhua River), and Donggang (DG in the Dayanghe River). There were totally twenty-one haplotypes identified in the three groups of samples. Haplotype networks by using neighbor-joining and median-joining clustering methods both indicated that the haplotypes 3, 8, and 19 constituted a single branch and the other 18 haplotypes constituted another branch. The highest genetic differentiation coefficients were observed between the ZZ and HEB samples (0.264), and the lowest was between the ZZ and DG (0.101). The average gene flow among the three populations was 3.063. The Fu's Fs and Tajima's D test values of the three populations were both negative, and indicated no significant difference between three populations (p=0.08 and 0.06> 0.05). Our results suggest that the far eastern catfish populations have not undergone significant population expansion and remained stable in the Yellow River, the Songhua River, and the Dayanghe River.

Introduction
The catfish (Silurus asotus), a member of the family Siluridae, is a freshwater species that is abundant and widely distributed throughout China, the Korean Peninsula, the Japanese islands, and the eastern regions of Russia. In the past several decades, catfish populations have been threatened by overfishing, water pollution, and water conservation projects (Hu, Pan, Hou, & Li, 2002; Yan, Jiang, & Liu, 2004). It is important to better understand the population dynamics of S. asotus, especially regarding the status of genetic diversity, and the conservation of endemic germplasm resources, with the purposes of sustaining the catfish populations.

To survey the extant natural resources of S. asotus is challenging, which requires acquisition of large amounts of data of genetic information from various catfish populations. The RAPD marker (Yoon, & Kim 2001), cytochrome B sequences of mitochondrial DNA (Xiao, Cui, Kang & Zhang, 2013; Xu, 2013), and microsatellite markers (Wu, Lian, Hou, Li, & Xiao, 2011) have previously been used to study the genetic diversity of catfish and related species. These methods have provided a wealth of information regarding the genetic fitness of these species, but the analysis of mitochondrial control region (D-Loop) DNA in catfish has not been reported. The D-loop region of mitochondrial DNA has been widely used in population genetic analysis, because it accumulates mutations more rapidly than other genomic regions and is not affected by selective pressure, as the D-loop region does not code for any protein (Xiao, Zhang, & Gao, 2009; Yuan, Zhang, Li, Zhu, & Luo, 2010).
The Songhua River, the Yellow River, and the Dayanghe River are chief rivers in the northern China, and contain rich catfish resources. In this study, the mitochondrial D-Loop was used to analyze the population genetic diversity and adaptive variation of three catfish populations in Zhengzhou (ZZ) of the Yellow River, Haerbin (HEB) of the Songhua River, and Donggang (DG) of the Dayanghe River. The results will be useful to inform catfish germplasm resource management, and better conserve the biodiversity and fitness of catfish populations.

Materials and Methods

Animals

The *S. asotus* samples were obtained from three catfish populations in ZZ, HEB and DG in 2017 (Figure 1). Thirty 1-2 age adults were selected from each of the population. The average body length is 254.83±41.07 mm, and the body weight is 158.73±90.86 g. The back-muscle tissues were collected from 15 catfishes in each of the populations and stored in 95% ethanol prior to DNA extraction. All surgeries were performed under sodium pentobarbital anesthesia, and every effort was taken to minimize suffering.

Extraction of Genomic DNA

Total genomic DNA was extracted from the back-muscle samples using the Animals DNA extraction kit (Tiangen biotech(Beijing)Co., LTD). Total DNA was adjusted to a concentration of 50 ng/ml and stored in ddH2O (Mascolo *et al.*, 2019). The DNA quantity was measured by nanodrop 2000 and agarose gel electrophoresis (Figure 2).

**Figure 1.** Sampling locations of the 3 *Silurus asotus* populations.

**Figure 2.** The electrophoretic pattern of *Silurus asotus* genomic DNA (portion). Back muscle tissue genomic DNA of *Silurus asotus* was extracted and electrophoresed on a 1.8% agarose gel. M: Marker (2000, 1000, 750, 500, 300, 100 bp).
Amplification of D-Loop Sequences

The mitochondrial D-loop was amplified by polymerase chain reaction (PCR) using the following primer set: forward primer 5’-AGGGGTATGTGGAGTAAA-3’ and reverse primer 5’-AGGGGTATGTGGAGTAA-3’. The primers were designed by the software of Primer 5.0 (Canada, Premier 5.0). Amplification was carried out in 50 μL reaction volume containing the following components: 1 μl genomic DNA, 25 μl Easy Taq® PCR Super Mix (Easy Taq® PCR System; TaKaRa, Dalian, China), 1.5 μl of each primer, and 21 μl ddH₂O (Table 1). The PCR conditions were set as follows: 94°C for 2 min followed by 33 cycles of denaturing at 94°C for 45 s, annealing at 56°C for 1 min, and elongating at 72°C for 1 min, then a final extension at 72°C for 8 min. Amplified products were purified with a Gel Extraction Kit (Generay Biotech Co., Ltd. Shanghai). Then, the PCR products were sent to the Generay Biotech Co. Ltd. for sequencing.

Sequence Alignments and Analyses

The sequences were proofread, edited, and aligned using BioEdit software version 7.0.5.3 (Ibis Biosciences, Carlsbad, CA). Mismatch distribution analysis of these polymorphisms was assessed using DnaSP v 5.0 (Rozas, Sánchez-Delbarrio, & Messeguer, 2003) (Universitat de Barcelona, Barcelona, Spain). Historical population expansion was examined using Tajima’s and Fu’s statistics with 10,000 demographic history records among the three populations. Parameters of genetic diversity within and between populations, such as the haplotype diversity (HD), nucleotide diversity (π), average number of nucleotide difference (K), and gene flow (Nm), were estimated with MEGA (Tamura, Peterson, & Peterson, 2011) (version 5.05). Phylogenetic reconstruction of the haplotypes was conducted using the maximum likelihood method, and a consensus of 10,000 bootstrap replicates. Arlequin (version 3.11; University of Berne, Berne, Switzerland) was used to estimate the genetic differentiation and Fst values with an analysis of molecular variance (Excoffier, Laval, & Schneider, 2005) (AMOVA). A haplotype network of the three catfish populations was constructed by Network (Bandelt, Forster, & Röhl, 1999). Statistical analyses were performed using SPSS19.0.

Results

Composition Analysis of the D-Loop Sequence

The D-Loop position for catfish between tRNA Pro and tRNA Phe. Approximately 535bp of the mtDNA control region, which shares extensive homology with other catfish populations (Mascolo et al., 2018), was analyzed from the ZZ (12), HEB (14), and DG (15) catfish populations. A total of 41 effective sequences were obtained and analyzed, and the estimated transition/transversion bias (R) was 6.13; A=29.6, C=27.8, G=14.9, and T=27.8; A+T (64.8%) was significantly higher than C+G (35.2%) (Table 2).

Analysis of Genetic Diversity and Population Genetics Structure

The estimates of haplotype diversity and nucleotide diversity are shown in Table 3. The DG population showed the highest HD (0.924±0.053), the

| Table 1. Reaction system and primer sequences of Silurus asotus D-loop PCR |
|-----------------------------|----------------|
| Component                    | Volume  |
| Genomic DNA(50ng/μl)         | 1μl     |
| F-Primer(5’-AGGGGTATGTGGAGTAA-3’) | 1.5μl |
| R-Primer(5’-AGGGGTATGTGGAGTAA-3’) | 1.5μl |
| 2xEasyTaq®PCRSuperMix        | 25μl    |
| ddH₂O                        | 21μl    |

| Table 2. Nucleotide composition of the D-LOOP sequences from three Silurus asotus groups (%) |
|--------------------------------------|--------|--------|--------|--------|--------|
| Population  | T     | C     | A     | G     | G+C    |
| ZZ          | 29.8  | 20.4  | 35.0  | 14.8  | 35.2  |
| DG          | 29.9  | 20.4  | 34.9  | 14.8  | 35.2  |
| HEB         | 29.7  | 20.5  | 35.0  | 14.8  | 35.3  |

| Table 3. Genetic diversity of the D-LOOP sequences from three Silurus asotus populations |
|--------------------------------------|--------|----------------|----------------|----------------|
| Population  | Haplotype(H) | Haplotype diversity (HD) | Polymorphic digit (P) | Nucleotide diversity (π) |
| ZZ          | 3        | 0.591±0.108       | 11              | 0.007±0.002     |
| DG          | 10       | 0.924±0.053       | 28              | 0.013±0.002     |
| HEB         | 8        | 0.890±0.060       | 19              | 0.012±0.001     |
highest polymorphic digit ($P=28$), and the highest nucleotide diversity ($\pi=0.013\pm0.002$). The genetic diversity of the three catfish populations was ranked as: DG, HEB, and ZZ.

The analysis resulted in 21 haplotypes from 41 individuals of $S. \textit{asotus}$. Haplotype Hap_20 was the most frequent one, and the distribution of haplotypes was unique within each population. No same haplotype was shared among all three catfish populations. Ten haplotypes were observed in the DG population, the most in any of the populations, and only three haplotypes were observed in the ZZ population, which is the least in any of the catfish populations (Table 4).

The genetic distance of the three populations was 0.011-0.015; the largest genetic distance was between the HEB and ZZ populations, and the smallest genetic distance was between the ZZ and DG populations (Table 5).

The UPGMA tree was constructed based on the genetic distance of the three populations of catfish, and showed that all the three populations were assigned to two clades, associated with a dispersion and expansion of the population, indicating the close relationship between the DG and ZZ populations (Figure 3).

Twenty-one kinds of haplotypes were found in three catfish populations, Hap_3 and Hap_8 were only found in DG population; Hap_19 is a haplotype unique to the ZZ population. The number of clustered haplotypes accounted for 86% of the total haplotypes. Cluster analysis of the haplotypes by neighbor-joining using the bootstrap method (repeat number 10000) test and it is shown in Figure 4.

| Table 4. Haplotype distribution in three $S. \textit{asotus}$ populations |
|---------------------------|---------------------------|---------------------------|
| Haplotype | ZZ | HG | HEB |
| Hap_1 | 0 | 4 | 0 |
| Hap_2 | 0 | 1 | 0 |
| Hap_3 | 0 | 2 | 0 |
| Hap_4 | 0 | 1 | 0 |
| Hap_5 | 0 | 1 | 0 |
| Hap_6 | 0 | 1 | 0 |
| Hap_7 | 0 | 1 | 0 |
| Hap_8 | 0 | 1 | 0 |
| Hap_9 | 0 | 1 | 0 |
| Hap_10 | 0 | 2 | 0 |
| Hap_11 | 0 | 1 | 1 |
| Hap_12 | 0 | 1 | 0 |
| Hap_13 | 0 | 1 | 0 |
| Hap_14 | 0 | 1 | 2 |
| Hap_15 | 0 | 1 | 1 |
| Hap_16 | 0 | 1 | 1 |
| Hap_17 | 0 | 1 | 4 |
| Hap_18 | 0 | 1 | 0 |
| Hap_19 | 4 | 0 | 0 |
| Hap_20 | 7 | 0 | 0 |
| Hap_21 | 1 | 0 | 0 |

| Table 5. Pairwise K2-P genetic distances (lower left) and Fst (upper right) between three $S. \textit{asotus}$ populations |
|---------------------------|---------------------------|---------------------------|
| Population | ZZ | DG | HEB |
| ZZ | - | 0.101 | 0.264 |
| DG | 0.011 | - | 0.130 |
| HEB | 0.015 | 0.013 | - |

Figure 3. The UPGMA clustering tree of three $S. \textit{asotus}$ populations based genetic distance.
Network analysis was used to identify the phylogenetic relationships among the 21 haplotypes within the three populations. The network analysis indicates that all haplotypes were divided into two groups, haplotype 3, 8 and 19 belongs to group A, and the other haplotypes belongs to group B (Figure 5). This is consistent with the phylogenetic analysis (Figure 4). The distribution frequency of haplotype 19 was 57% of the total A group, and haplotype 19 was the dominant haplotype. The distribution frequency of haplotype 1, 17, and 20 were 4% of the total B group; these three made up the dominant haplotypes. The dominant haplotypes within the DG, HEB, and ZZ populations were Hap_1, Hap_17, and Hap_20, respectively.

**Figure 4.** NJ phylogeny tree of D-loop haplotype of three *Silurus asotus* populations

**Figure 5.** Median joining network of three *Silurus asotus* populations
Analysis of Molecular Variance

The significance of the three population structures based on haplotype frequency: 16.32% molecular variance was attributed to the differentiation between the populations, whereas 83.68% molecular variance was observed within the populations (Table 6). Population structuring revealed that the Fst value was not significant (p=0.09>0.05) (Table 5). The genetic differentiation coefficient of the three populations ranged from 0.101-0.264, and the average value was 0.165. The genetic differentiation coefficient between the ZZ and HEB populations was the highest (Fst=0.264), and was lowest between the DG and ZZ populations (Fst=0.101).

The tests of neutral evolution (Tajima’s D and Fu’s F tests) indicated that there was no significant difference (p=0.08 and 0.06) among the three populations, indicating no significant population expansion. Mismatch distribution analysis of the mitochondrial D-loop regions sequences revealed no peaks (Figure 6).

Discussion

Genetic diversity of three *S. asotus* populations was assessed by analyzing the mtDNA D-loop sequences of 41 individuals from three populations. Within these three catfish populations the G+C content in the mtDNA D-loop region was significantly lower than the A+T content, which was consistent with the results of base composition of mitochondrial cytochrome B in a study of catfish from the Huaihe River (Xiao et al., 2013). The R of the three populations was larger than 2.0, indicating that the mitochondrial D-loop sequence can be used to assess genetic diversity between these populations (Kumar, Tamura, & Nei, 2004).

The haplotype diversity (HD) and nucleotide diversity in the DG and HEB populations was similar to previously reported (Xu et al., 2013), which identified haplotype diversity and nucleotide diversity of cytochrome B sequences from multiple Chinese catfish populations (HD=0.948±0.009, π=0.018±0.001). The HD and π described for the catfish populations in this study are higher than those reported in carp populations from the Yangtze and Pearl rivers (Fu, 2011). The genetic diversity of a species is closely related to the ability of that species to adaptation and survive to changing environmental conditions (Conrad, 1983; Féral, 2002; Frankham, Ballou, & Briscoe, 2002), and low diversity can indicate a decline in how well a species can adapt to environmental change (Nei, 1978; Gu, 2004). However, the values of HD and π of ZZ the population were significantly lower than those of the DG and HEB populations, indicating that the DG and HEB catfish populations may be better equipped to adapt to environmental changes than the ZZ population, due to higher genetic diversity within those populations. The low genetic diversity of the ZZ population may be related to environmental pollution and ecological damage in the Yellow River, and may also be caused by environmental deterioration of the Yellow River (Guan et al., 2016; Michel, Wolters, Sun, Huang, & Kuenzer, 2016).

The study of the genetic structure of a species can provide critical information regarding the genetic diversity within an individual population and between geographically diverse populations (Yan, Song, Wang, Lu, & Gao, 2015). Within these three catfish populations in the present study, the average individual genetic distance was 0.011, indicative of small differences between populations. In general, the genetic distance values of a genus, species, and population are 0.90, 0.30 and 0.05 (Shaklee, Tamaru, & Waples, 1982), respectively; the genetic distance between the three catfish populations was less than 0.05. Although the HEB and DG catfish populations are nearest in geographic proximity, the genetic distance between these populations was the greatest, indicating that the size of genetic distance between catfish populations does not necessarily correspond to the geographic distance between those populations. However, although analysis of molecular variation revealed that most variation arises from within an individual catfish population, and the overall variation between populations was small. Ultimately, no significant genetic differentiation was identified among the three catfish populations studied.

The value of gene flow is related to the exchange of genetic information between populations. Populations are isolated when the *Nm* is less than 1; an *Nm* greater than 1 indicates gene flow between populations; and an *Nm* greater than 4 indicates that each population exists as a random unit (Masatoshi, & Sudhir, 2000). In the current study, the *Nm* between ZZ and HEB was the smallest, but still greater than 1, indicating evidence of some gene flow between the two populations. The *Nm* between ZZ and DG was more than 4, which showed that two populations form a random unit. The gene flow between the three populations is closely related to the genetic differentiation of the

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Table 6. The analysis of molecular variance from three *Silurus asotus* populations

| Source of Variation | df | Sum of squares | Variance components | Percentage of variation(%) |
|---------------------|----|---------------|---------------------|---------------------------|
| Populations         | 2  | 23.634        | 0.631               | 16.32                     |
| Within populations  | 38 | 122.879       | 3.234               | 83.68                     |
| Total               | 40 | 146.512       | 3.864               |                           |

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Figure 6. The mismatch distribution of D-LOOP sequence of three *Silurus asotus* populations

Note: A. Total,  B. HEB,  C. DG,  D. ZZ.
populations, and may be the reason for the small genetic differentiation among the three catfish populations. In addition, the gene flows among the three populations do not correspond with geographical distance; factors such as commodity circulation, enhancement, and artificial breeding of catfish may cause this phenomenon.

The network method is commonly used to analyze molecular variation analysis of the species, and is especially suitable for the analysis of non-recombinant DNA sequences, such as mitochondrial DNA and Y chromosomal DNA (Bandelt et al., 1999). Phylogenetic relationships of 21 haplotypes were analyzed by adjacent tree and intermediary network diagramming in the present study. However, the intermediary network figure reveals that haplotype did not correspond to sampling sites. Hap_20, Hap_19, Hap_17, and Hap_1 have the highest frequency, suggesting these may be the original haplotypes.

The tests of neutral evolution (Tajima’s D and Fu’s Fs) were both negative in our study, which is consistent with Liu’ report (Liu, Yang, Wang, & Tang, 2009). Distribution of pairwise mismatches of S. asotos populations was multimodal, indicating no significant population expansion in the three rivers. Thus, it was concluded that the genetic differentiation among the three catfish populations was low.

The behavior of genetic variation within S. asotos populations provides important information in developing the conservation and management strategies for endangered fish populations. Despite the identification of the low levels of genetic differentiation among the ZZ, HEB, and DG catfish populations, the results indicate that geographic and environmental features of drainage regions may significantly impact the different catfish populations. Furthermore, these results suggest that conservation stations should be established on the three rivers to increase the number of catfish populations, and to improve the genetic diversity and overall fitness of this species. The results also improved the understanding about role of genetic variation in the catfish conservation. Genetic variation is a dimension of the biodiversity, to better monitor and compare the biodiversity of S. asotos populations the mechanism that driving the genetic variation should be researched in the future.

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