Structural characteristics of the control region of the Beaufortia kweichowensis mitochondrial genome

Abstract

In present study, we identified the structural characteristics of the Beaufortia kweichowensis mtDNA control region using the next-generation sequencing method. Result showed that the control region could be further divided into three parts, including the extended termination associated sequence domain (ETAS), the central conserved domains (CSB-F, CSB-E, CSB-D) and the conserved sequence block domains (CSB-1, CSB-2, CSB-3), and their conserved sequences were identified. Additionally, we found the ETAS domain could be folded and form a 25 bp loop, which are usually considered to play an important role in mtDNA transcript termination. Finally, we reconstructed the phylogenetic relationship among the family of Balitoridae based on the nucleotide sequence of the control region, result showed that the Balitoridae can be divided into two clades, one clade is Gastromyzoninae and the other is Homalopterinae, and the B. kweichowensis was grouped into clade Gastromyzoninae and shared a close relationship with B. szechuanensis.

Introduction

Mitochondrial DNA (mtDNA) is a circular double-strand DNA with a 15–20 kb length in animals, and it usually encodes 37 genes, including 13 protein-coding genes (PCGs), 22 transport RNA (tRNA), and 2 ribosomal RNAs (rRNAs) [1,2]. To date, mtDNA has been widely used for genetic research, taxon classification, phylogenetic evolution research and population studies attribute to its fast variation, maternal inheritance, rapid evolution and lack of recombination [3,4]. Recent years, the mitochondrial genome database growth rapidly following the fast development of high throughput sequencing technology [5], which might provide more chance for scientists to solve the biological mystery in the future.

The control region, also called displacement–loop region (D–loop), is a DNA fragment with fastest evolution rate in mtDNA because it composes of the single strand nucleotide [6]. It can be divided into three domains, including the extended termination associated sequence domain (ETAS), the central conserved domains (CSB–F, CSB–E, CSB–D) and the conserved sequence block domains (CSB–1, CSB–2, CSB–3) [7,8], and it also has been extensively used for phylogenetic analyses and populational genetics researches [9,10].

Fish is one of the most amount groups in vertebrate, and now more than 30000 species has been discovered [11]. Thus far, more than 200 fish genome and 1000 fish mitochondrial genome has been sequenced and deposited into NCBI database, and these data are available for all researchers, thus then some important scientific issue such as the evolutionary relationship of fish has been well investigated [12,13], which also will be helpful for species classification and phylogenetic analyses in these groups.

B. kweichowensis (Fang) is an endemic fish of the upper reaches of the Beipan River of China, belonging to the family of Balitoridae (Cypriniformes), and has not yet been well studied about its phylogenetic evolutionary status [2]. In present study, the structure of control region of B. kweichowensis mtDNA was investigated, and the conserved sequences also was identified. Subsequently, the phylogenetic tree of the family Balitoridae was reconstructed based on the sequence information of mtDNA control region, which could be helpful to establish the evolutionary status of this species.

Material and Methods

Sample collection and DNA isolation

One adult B. kweichowensis used in this study were obtained from Conservation and Utilization of Fishes resources in the Upper Reaches of the Yangtze River Key Laboratory of Sichuan Province, Neijiang Normal University. A 20–30 mg fine clip was collected and preserved in 95% ethanol at 4 °C. Total genomic DNA was extracted with a Tissue DNA Kit (OMEGA E.Z.N.A.) following the manufacturer’s protocol.
Sequencing and structure analysis of control region

The genomic DNA was sequenced using the next-generation sequencing, and then the mitogenome was assembled using *B. szechuanensis* (GenBank accession number: KP716708.1) as reference. The non-coding control region structure was analyzed and drawn according to the control region structure of other Botiinae fish.

Phylogenetic analysis

The phylogenetic tree was constructed with nucleotide sequence of control region using MEGA 6.0 software [14]. The HKY+G model was selected as the best evolutionary model after testing, and the tree was performed using maximum likelihood and bootstrapped with 1000 replications. The species *Cycleptus elongatus* and *Catostomus commersonii* were selected as outgroups.

Results and Discussion

The full length of the control region of *B. szechuanensis* mtDNA is 930 bp in length, and it can be divided into three domains including the extended termination associated sequence domain (ETAS), the central conserved domains (CSB-F, CSB-E, CSB-D) and the conserved sequence block domains (CSB-1, CSB-2, CSB-3), result is shown in figure 1. Simultaneously, the conserved sequences of these domains were identified, the conserved sequence of ETAS is CATATATGCATAATATTACATAATGTAATAG-TACATATATGTA, and it can be folded into a 25 bp long loop structure (Figure 2), which is usually considered to be connection with mitochondria transcript termination [2], and this phenomenon also can be found in other fishes [15]. The conserved sequence of the central conserved domains are ATGTAGTA-AGAAACCAGACCCAGTCCA, TCAAGGACTATGTGTGAGG and TGAATTATTACTGGCATCTGGTTCCTATTTCAGG, which represent the conserved sequence of CSB-F, CSB-E, CSB-D domains, respectively. Moreover, the conserved sequence of the conserved sequence block domains is AGGTTAATGATTAAATGACATACTCAAGA, CGCGAGAGGCCCCCTTACCCCTTACAC and CCTGTG-CAAAACCGCAAAAACAGGA, which represent the conserved sequence of CSB-1, CSB-2, CSB-3 domains, respectively. Furthermore, the TATA element also was identified, which is considered to play an important role in transcript origination [9].

The Balitoridae is regarded as a distinct family in order Cypriniformes. Hora divided this family into two subfamilies Gastromyzoninae and Homalopterinae, and he considered the former as derivative of the Cobitidae and the latter as a descendant of the family Cyprinidae [16]. In order to better understand the relationship of this family, maximum likelihood phylogenetic tree was reconstructed based on the nucleotide sequence of mtDNA. Results showed that the family Balitoridae exactly can be divided into two clades, including subfamily Gastromyzoninae and Homalopterinae, and the *B. kweichowensis* was grouped into clade Gastromyzoninae and shared a close relationship with *B. szechuanensis* (Figure 3). These findings are consistent with our previous study based on whole mtDNA sequence, suggesting the *B. kweichowensis* should be origin from the subfamily Gastromyzoninae [17].
Conclusion

The control region of *B. kweichowensis* could be divided into two three domains, and their conserved sequence were identified. Furthermore, a 25 bp long loop domain was found in the ETAS. Finally, the phylogenetic relationship of the family Balitoridae was analyzed, result highly supported the *B. kweichowensis* belongs to the subfamily Gastromyzoninae.

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