Lack of Structural Variation but Extensive Length Polymorphisms and Heteroplasmic Length Variations in the Mitochondrial DNA Control Region of Highly Inbred Crested Ibis, *Nipponia nippon*

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Abstract

The animal mitochondrial DNA (mtDNA) length polymorphism and heteroplasmy are accepted to be universal. Here we report the lack of structural variation but the presence of length polymorphism as well as heteroplasmy in mtDNA control region of an endangered avian species – the Crested Ibis (*Nipponia nippon*). The complete control region was directly sequenced while the distribution pattern and inheritance of the length variations were examined using both direct sequencing and genotyping of the PCR fragments from captive birds with pedigrees, wild birds and a historical specimen. Our results demonstrated that there was no structural variation in the control region, however, different numbers of short tandem repeats with an identical motif of CA3CA2CA3 at the 3′-end of the control region determined the length polymorphisms among and heteroplasmic within individual birds. There were one to three predominant fragments in every bird; nevertheless multiple minor fragments coexist in all birds. These extremely high polymorphisms were suggested to have derived from the ‘replication slippage’ of a perfect microsatellite evolution following the step-wise mutational model. The patterns of heteroplasmy were found to be shifted between generations and among siblings but rather stable between blood and feather samples. This study provides the first evidence of a very extensive mtDNA length polymorphism and heteroplasmy in the highly inbred Crested Ibis which carries an mtDNA genome lack of structural genetic diversity. The analysis of pedigreed samples also sheds light on the transmission of mtDNA length heteroplasmy in birds following the genetic bottleneck theory. Further research focusing on the generation and transmission of particular mtDNA heteroplasmy patterns in single germ line of Crested Ibis is encouraged by this study.

Introduction

The Crested Ibis (*Nipponia nippon*) was widely distributed in Northeast Asia, however, the overexploitation and destruction of habitats led to the extinction of its Korean, Japanese and Russian populations in the wild in the late 20th century [1,2,3]. In May 1981, a remnant population consisting of two pairs of adults and three nestlings was re-discovered in Shaanxi Province, China [4]. Since then, intensive conservation efforts have been made to restore this species in the wild with up to 600 birds by 2007 [5] and to conserve it in captivity with 600 birds by 2009 [6] (Figure 1; also see Fig. 2 in [5] as well as Fig. 1 and 3 in [6] for more details). This species was once recognized as “Critically Endangered” but has been considered as “Endangered” since 2000 in the IUCN Red List of Threatened Species [7]. Understanding and identification of survived genetic variations are therefore critical for the long-term protection and management of all existing populations recovered from such limited founders which must have made this species to be highly inbred.

The polymorphic length heteroplasy (coexistence of two or more types of mtDNA genomes in different lengths within one individual) in mtDNA was reported in a wide range of animals, such as fruit flies [8,9], crickets [10,11,12], lizards [13], fishes [14,15] and elephant seals [16]. In avian species, mtDNA length heteroplasmy was detected in rails (*Rallus elegans* and *R. longirostris*) [17] and mourning dove (*Zenaida macroura*) [18] by restriction analysis of purified mtDNAs. Sequencing of PCR products revealed the length polymorphism and heteroplasy in mtDNA control region of 15 species of the order Charadriiformes [19] and two vulture species *Gyps alticeps* and *Neophron percnopterus* [20]. Intensive studies on the transmission of mtDNA genomes suggested a theory of ‘genetic narrow bottleneck’ following the proportion shift of mtDNA structural heteroplasmy between mother and offspring as well as between offspring in cattle, mice, human and salmon [21,22,23,24,25,26,27,28,29,30,31,32]. To our knowledge, the transmission of either length or structural heteroplasmy in mtDNA of birds remains to be examined.
Previous studies revealed an extremely low genetic diversity in the current populations of Crested Ibis both in the wild and captivity using mtDNA control region sequences and microsatellite DNA markers [33,34]. Sequences of domains II and III at 3'-end of the control region generated using PCR and cloning protocols were reported to have an intra-individual length heteroplasmy in the domain III. The heteroplasmy was defined by two different DNA structures of either a simple tandem repeat of (CA3CA2CA3)n or a complicated repeat (CA3CA2CA3)4CA3C

[34]. However, the distribution pattern and inheritance of the heteroplasmy were not fully investigated. In this study, both direct sequencing of PCR products and genotyping of PCR fragments were applied to investigate structural variation and length heteroplasmy in the mtDNA control region of the highly inbred Crested Ibis.

**Materials and Methods**

**Sampling and DNA Extraction**

Blood samples from 28 birds and both blood and feather samples from five individuals were collected in 2010 from Shaanxi Crested Ibis Breeding Center, Yangxian County, Shaanxi Province, China. These 33 captive adult Crested Ibis belonged to three families following pedigree records (Table 1). There were 23 individuals across three generations in family No. 1 and five individuals each across two generations in families No. 2 and No. 3. The current distribution range of the wild population, locations of the 43 nests banded in 2011 and 25 nests selected for this study in the wild as well as the Captive Breeding Center of Crested Ibis. Photo courtesy of Jiao Jingquan.

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**Figure 1.** The current distribution range of the wild population, locations of the 43 nests banded in 2011 and 25 nests selected for this study in the wild as well as the Captive Breeding Center of Crested Ibis. Photo courtesy of Jiao Jingquan.

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**Figure 2.** Structure of mtDNA control region and position of primers designed for this study. TAS = termination associated sequences, CSB = conserved sequence block. The primer positions are scored based on the complete mtDNA sequence of Crested Ibis deposited in GenBank (Accession no. NC_008132).

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No. 3. Another 27 feather samples were collected from nestlings with one each of 24 nests and three of one nest in the wild during banding season in 2011 in Shaanxi Hanzhong Crested Ibis National Nature Reserve (Figure 1). One foot pad sample was derived from a specimen, which was acquired from Xi’an of Shaanxi province in 1915 and is preserved in the State Zoological Museum of China in Beijing. This study was fully endorsed by both the Center and the Reserve and finally approved by the Institutional Animal Care and Use Committee (IACUC) of the College of Biological Sciences and Technology, Beijing Forestry University. All the sampling activities were conducted in collaboration with technical staff of the Center and the Reserve. Maximal care was taken during either bleeding for adults or banding for nestlings as part of the routine practices which have been applied to monitor the captive and wild flocks. The foot pad sample was donated by the Museum. These samples were strictly stored and used for research purpose only following specific collaborative agreements between the institutions.

The blood samples were collected from wing vein and mixed with same volume of anticoagulant buffer (0.1 M Tris-Cl, 0.1 M EDTA and 1% SDS, pH 8.0). Feather samples were kept in absolute alcohol. All the blood and feather samples were stored at −80°C. The foot pad sample was kept in a sealed envelope at room temperature until DNA extraction. After removal of the alcohol, the calamus (approximately 2 cm) of preserved feathers was washed twice with 1xosodium chloride-Tris-EDTA buffer (pH 8.0) and cut into pieces to get the pulp tissue inside the calamus. Total genomic DNA was extracted from all the blood, pulp tissue and foot pad samples using a conventional phenol/chloroform protocol [35]. After being qualified and quantified with one each of 24 nests and three of one nest in the wild during banding season in 2011 in Shaanxi Hanzhong Crested Ibis National Nature Reserve (Figure 1). One foot pad sample was derived from a specimen, which was acquired from Xi’an of Shaanxi province in 1915 and is preserved in the State Zoological Museum of China in Beijing. This study was fully endorsed by both the Center and the Reserve and finally approved by the Institutional Animal Care and Use Committee (IACUC) of the College of Biological Sciences and Technology, Beijing Forestry University. All the sampling activities were conducted in collaboration with technical staff of the Center and the Reserve. Maximal care was taken during either bleeding for adults or banding for nestlings as part of the routine practices which have been applied to monitor the captive and wild flocks. The foot pad sample was donated by the Museum. These samples were strictly stored and used for research purpose only following specific collaborative agreements between the institutions.

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Verification of Pedigree Records of the Three Families Using Microsatellite DNA Markers

Eight microsatellite DNA markers (NN01, NN03, NN04, NN12, NN16, NN21, NN25 [33] and NnNF5 [36]) were selected to genotype the 33 pedigreed Crested Ibis for verification of their genetic relationship. All forward primers were labeled by fluorescent dyes and similar PCR conditions were followed for genotyping using an ABI 3130xl Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) with detailed procedures described in later section.

Amplification of the Control Region and Direct Sequencing

Ten captive and 27 wild birds were chosen to sequence the complete control region. The 10 captive individuals included six parental birds of the first generation of the three families and four additional partners in the second generation of the family No. 1 (Table 1). The complete mtDNA sequence of Crested Ibis (GenBank accession number NG_008132 at http://www.ncbi.nlm.nih.gov/nuccore/NG_008132) was referred to design primers DL-F (5’- GTA AGT CAT AGC CAT TCC TGC T-3’) and DL-R (5’- TTTG TCC TTG GGT GCA AGA ATG-3’) (Figure 2) for the amplification of a fragment around 1290 base pairs (bp) covering the complete control region.

The amplification was performed in a 50 μl volume containing around 100 ng DNA, 3.0 units of EasyTaq DNA polymerase (TransGen Biotech, China), 5 μl of 10×EasyTaq Buffer, 4 μl of dNTPs (2.5 mM, Tiangen Biotech Co. Ltd., China) and 1 μl each of the forward and reverse primers (10 pmol/μl). Thermal cycling was performed using a GeneAmp PCR system 9700 thermal cycler (Applied Biosystems) started with 5 min at 94°C followed by 30 cycles of 40 s at 94°C, 30 s at 64°C and 90 s at 72°C, and finished by a final extension at 72°C for 7 min. Three-micro-liters each of the PCR products were run in a 1.5% agarose gel and visualized by GoldView™ Nucleic Acid Stain (SBS Genetech Co. Ltd., China) to confirm their quality and quantity. A total of 33 PCR products of captive and wild birds were purified and directly sequenced using the DL-F primer and two additional sequencing primers of DL-SF (5’- GGT TGG TCC TCA GGA ATT AA-3’) and DL-SR (5’- TAG GTG TAG TGG ATA CAC CA-3’) (Figure 2) for the complete control region including a repeat fragment. The remaining four captive samples were sequenced using the two PCR and two sequencing primers for complete amplicons.

PCR products amplified using the DL-F and DL-R primers from the remaining 23 captive individuals were only sequenced

| Table 1. Number of repeats in predominant fragments in 33 captive Crested Ibis. |
| --- |
| Family | Mating group | Parent | Offspring |
| No. 1 | 1 | Female A-05:11 Male A-04:11 | Female A-06:12 A-08:11 A-10:9 A-20:12 A-33:9 Male A-18:12 A-28:11 |
| No. 2 | 6 | Female A-26:9 Male A-25:10 | Female A-19:9 Male A-13:8 A-15:8 |
| No. 3 | 7 | Female A-01:11 Male A-03:11 | Female A-14:11 Male A-02:11 A-22:11 |

ID of individual is numbered as A-01 to A-33. doi:10.1371/journal.pone.0066324.t001
using the DL-SF primer for their last partial control region covering the repeat fragment. The single foot pad sample was amplified using the DL-SF and DL-R primers and directly sequenced using the DL-SF primer for the repeat fragment as well.

The BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems) was applied for sequencing reactions that were visualized on an ABI3730 Sequence Analyzer by the Beijing Sunbiotech Co. Ltd. (China). The raw data was manually edited using Chromas version 2.25 (http://www.technelysium.com.au/ chromas.html) and aligned along with all sequences retrieved from the GenBank or relevant publications for comparison with MEGAS software [37].

Genotyping of the Length Polymorphism and Heteroplasmy in Control Region

To detect the size variations in the repeat fragment, an additional forward primer DL-MF (5’-TCA CAT ACG GCC GCA AAA ACA-3’) (Figure 2) was designed according to the newly generated sequences from this study and labeled by a 6-FAM fluorescent dye at its 5’-end. It was applied together with the DL-R primer for the amplification of around 250 bp repeat fragment in the domain III among all 61 samples. The amplification was replicated in each DNA sample and performed in a 50 μl volume as above except 2.0 units of Taq DNA polymerase (Tiangen Biotech Co. Ltd., China) were used. Thermal cycling was started with 5 min at 94°C followed by 30 cycles of 40 s at 94°C, 30 s at 65°C and 30 s at 72°C, and completed by a final extension of 7 min at 72°C. Two micro-liters of the PCR product were mixed with 7.75 μl deionized formamide and 0.25 μl GeneScan™ 500 LIZ™ Size Standard (Applied Biosystems). This mixture was loaded on the ABI3130xl Genetic Analyzer and fluorescence signal data was collected using Data Collection v3.0 and then analyzed using GeneMapper v3.7 (Applied Biosystems).

Results

MtDNA Sequence Analysis

Four sequences of captive birds which were amplified using the DL-F and DL-R primers and sequenced using the DL-F, DL-R, DL-SF and DL-SR primers covered a repeat fragment ranging from the 1072nd nucleotide (scored according to JX853815) in the control region to the beginning of tRNA-Phe region. They all had 1179 nucleotides including 1071 bp remaining control region, 58 bp long segment covering the complete tRNA-Glu region, and 50 bp partial tRNA-Phe region. There was no polymorphism present within this 1179 bp long amplicon across these four samples, therefore a single representative sequence was deposited to the GenBank (accession no. JX853815). The control region of 1071 bp from the remaining 33 captive and wild samples shared this representative sequence.

Compared to the reference sequence of complete mtDNA of Crested Ibis (NC_008132), all 37 control region sequences in 1071 bp carried a 19 bp long motif of 10C+4T+8Gs starting from their 23rd nucleotide while the reference sequence carried only 9Gs, which were probably too difficult to be fully sequenced through. Our new data therefore filled this gap in the reference sequence and confirmed the presence of the interrupted poly(C) structure located at the 5’-end of mtDNA control region of Crested Ibis. A similar structure was identified across several bird groups such as Accipitriformes [20,38] (AF380303 for Buteo buteo; AV542900 for Gypaetus barbatus and AV542899 for Neoniphon percnopterus) according to the International Ornithology Committee World Bird List v 3.3 [39], Galliformes [40] (GU187969 for Tragopan caboti) and Sphenisciformes [41] (AF272143 for Pygoscelis adeliae).

Length Polymorphism and Heteroplasmy

The repeat fragment was detected in the four sequences amplified by the DL-F and DL-R primers and sequenced by the DL-F, DL-R, DL-SF and DL-SR primers. They all carried an identical repeat motif with 11 nucleotides in (CA9CA8CA7)n at the 3’-end of mtDNA control region. The remaining 33 captive and wild samples which were sequenced using the DL-F, DL-SF and DL-SR primers and the additional 23 captive and wild samples as well as the single historical foot pad sample which were sequenced only with the DL-SF primer shared the same repeat motif. This motif was also present in nine copies in the complete mtDNA sequence of Crested Ibis (NC_008132) and it was identical to the major motif reported by Zhang et al. [34]. Further examination of all 61 sequencing chromatograms generated using the DL-SF primer revealed different numbers of the repeat motifs among individuals (length polymorphisms) and also within individuals (length heteroplasmy) (Figure 3). Additional sequencing profiles of the DL-R primers confirmed our observation. Similar pattern was observed by Zhang et al. [34] and 6, 10 and 11 copies of the motif were identified among and within samples [42]. Clear heterogeneous sequencing profiles, resulted from intra-individual length heteroplasmy, were mostly detected after nine repeated motifs. Therefore it was not possible to determine the exact sizes of the repeat region that caused the variable length of the control region and/or mtDNA genome.

MtDNA length heteroplasmy due to long repeated sequences ranging from around 78 bp to 2 kb was first identified and genotyped using restriction analyses [8,9,13,44,45,46]. Following direct sequencing of PCR products of mtDNA control region, Ludwig et al. [14] confirmed the presence of such length heteroplasmy in 12 out of the 19 sturgeon species determined by one to seven central repeat units of 74 to 83 bp long fragments. After discovering a 75-bp tandem repeat segment, Munwes et al. [47] evaluated the length heteroplasmy in the eastern spadefoot toad (Pelobates syriacus) based on the number of distinct bands in PCR amplified repeat region on the agarose gel. Nesbo et al. [15] applied procedures of both cloning and direct sequencing of PCR products to assess the length variation and heteroplasmic length polymorphism of a tandem array of 10 bp repeats at the 5’-end of D-loop in two percid fish species. It is clear that the accuracy of these methods mostly depends on the degree and complexity of the heteroplasmic, especially for the low level heteroplasmy of short repeat motifs. Cloning and direct sequencing as well as fragment size analysis were used in interpreting the length heteroplasmy defined by several repeat sequences in human mtDNA control region [48,49,50].

To validate the length polymorphisms and heteroplasmonic length variations observed among the directly sequenced PCR products as described above, the size variations and their distribution patterns of the repeat fragment were further genotyped. It was evident that all genotyping profiles carried multiple peaks at least ranging from six to 14 repeats at different heights among and within all birds in this study. Comparison of sequencing and genotyping profiles across every sample confirmed the length polymorphisms among the samples and length heteroplasm among samples, both were attributed to highly variable numbers of the same repeat motif in (CA9CA8CA7)n. However, one to three predominant peaks at similar heights (fluorescent intensity) in genotyping profiles and relatively clear sequencing chromatograms carrying one to three motifs with respective number of repeats were observed within all samples (see Figure 3). On the other
hand, all the samples had relatively low sequencing signals and minor peaks at a regular interval of the 11 bp identical repeat. The same patterns were detected in both sequencing and genotyping results of duplicated blood and feather samples of five birds, demonstrating that the length polymorphisms and heteroplasmy may not be tissue specific in Crested Ibis mtDNA control region.

The museum foot pad specimen had a predominant fragment at 254 bp peak with 12 repeats. Among the 60 captive and wild birds, one captive bird of A-16 from No. 4 mating group carried three predominant fragments with 11, 12 and 13 repeats, one captive (A-30) and six wild birds had two predominant fragments carrying seven to 13 repeats while the rest 31 captive and 21 wild samples showed a single predominant fragment with number of repeats ranging from six to 13. The predominant fragment with nine repeats was the commonest among 21 wild birds across 19 nests while those with six, seven, eight, 10, 11 and 12 repeats were present at relatively low frequencies (Tables 1 and 2). A 'one sample Kolmogorov-Smirnov test' implemented in the SPSS 19.0 (IBM 2010) realized that the assignments of different predominant fragments among the 21 wild samples seems to have followed a pattern in normal distribution ($P=0.024$, Figure 4B). However, when the data were sorted out following the pedigree information of two major maternal lineages

Transmission of the Length Polymorphism and Heteroplasmy

A first examination of the distribution pattern of frequencies of different predominant fragments among the 31 captive birds revealed a significant deviation from the normal distribution ($P=0.024$, Figure 4B). However, when the data were sorted out following the pedigree information of two major maternal lineages

**Table 2. Number of repeats in predominant fragments in 27 wild Crested Ibis.**

| No. of repeats | ID of birds | No. of repeats | ID of birds |
|----------------|-------------|----------------|-------------|
| Single repeat  |             | Two repeats    |             |
| 6              | C-08        | 7, 8           | C-21        |
| 7              | C-13, C-22, C-33, C-59 | 7, 9           | C-26        |
| 8              | C-01, C-34, C-52 | 9, 10          | C-56, C-85  |
| 9              | C-18, C-31, C-36, C-38, C-62, C-71 | 9, 12          | C-37        |
| 10             | C-11, C-40, C-43 | 9, 13          | C-29        |
| 11             | C-54, C-55, C-78 |                |             |
| 12             | C-23        |                |             |

C-54, C-55 and C-56 are siblings.

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carrying predominant fragments with nine or 11 repeats in three mating groups each (Table 1), a match to the expected normal distribution was detected in the two lineages of either 10 progenies with their mother carrying a predominant fragment of nine repeats \( (P=0.162, \text{Figure 4C}) \) or 12 descendants with their mother having a predominant fragment of 11 repeats \( (P=0.079, \text{Figure 4D}) \).

To understand the inheritance mode of these intensive length polymorphisms and heteroplasmy in mtDNA control region, the pedigree records of 33 birds among three families were fully validated with the genotyping data of the eight microsatellite DNA markers (data not shown). Seventeen maternally-related birds in family No. 1 (Figure 5) across three generations showed that the types and/or combinations of predominant fragments, defined by different numbers of repeats, did not follow the rule of strict maternal inheritance across all mother-offspring pairs. For example, although the matriarch bird A-05 had a single predominant fragment with 11 repeats, only two out of her seven progenies shared the same predominant fragment but the remaining two and three descendants carried predominant fragments with nine and 12 repeats, respectively. Such irregular transmission was further detected in two out of the seven offspring in the third generation reproduced each by A-10 and A-33 shared the predominant fragment of nine repeats. Further assessment of the genotyping profiles showed that the secondarily predominant fragments with 11 repeats in A-10 and eight repeats in A-33 turned out to be the single dominant peaks in A-21 and A-32, respectively. In addition, four breeding males carrying different predominant genotypes from their mating females in families No. 1 and 2 allowed us to examine possible paternal leakage [53,54,55]. No shared predominant fragment was identified in nine out of the 11 father-offspring pairs (Table 1), thus a probable exclusion of paternal leakage in Crested Ibis mtDNA, though the evidence was not inclusive.

**Discussion**

**Lack of Structural Variation in the mtDNA Control Region of Crested Ibis**

As shown in Table 3, the single control region sequence obtained from this study was identical to five homologous sequences of the first part of control region (936 bp) obtained from the captive Crested Ibis population in China (accession no. AB104903–AB104907 at http://www.ncbi.nlm.nih.gov/nuccore/) but different by two nucleotides from the reference sequence (NC_008132) derived from ‘Midori’ which was one of the last survived birds in Japan. It is not clear how this complete mtDNA sequence of ‘Midori’ was generated. If the data was reliable, it may indicate the lost of a different maternal lineage once present in the Crested Ibis population in Japan.

This new sequence was also identical to the major control region haplotype 1 but different from the minor haplotype 2. Sequence fragments from both haplotypes, which represented the 36 birds of captive and wild Crested Ibis populations in China, were cloned from a 563 bp long fragment adjacent to the repeat region [34]. A surprisingly high control region diversity with four haplotypes (298 bp) defined by two transversions was identified in 42 captive birds [56]. However, the haplotype 1 identical to the sequence obtained in this study was reported to be present in nine out of the 42 samples in Zhang [56] and in all seven birds in Lei.
et al. [57]. In fact, all the samples for these three studies were
derived from the same genetic origin of the remnant ancestral
population in China [4]. Thus a maximum of two different
mtDNA lineages, assuming one each from the two maternal
ancestors, are expected in the restored populations. The single
transition reported by Zhang et al. [34] could be true for 26 birds
in captivity and 10 individuals in the wild sampled from 1990 to
2000. The four haplotypes I–IV identified using the primers L438
and H1248 [56], which were claimed from Genovart et al. [58]
but cited with different sequences, clearly had some errors. The
same dataset was further mistakenly quoted as to have five
haplotypes defined by three polymorphic nucleotides [59]. Our
samples in this study had the largest number of birds from the wild
(27 from 25 nests) while the 10 parental birds in captivity fully
represented all maternal lineages of all 33 birds across three
families. The single haplotype of mtDNA sequence covering the

Figure 5. Pedigree chart represents three generations with 17 maternally-related individuals in family No. 1. G-1, G-2 and G-3 indicate
the generations. Squares stand for males and circles for females. Numbers in brackets right to the squares or circles are the number of repeats in
every sample.
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Table 3. Nucleotide variation in mtDNA D-loop sequences of Crested Ibis.

| Sequence/haplotype (samples) | Sequencing method | Nucleotide position | Reference |
|-----------------------------|-------------------|---------------------|-----------|
| JX853815 (27 wild and 10 captive birds) | PCR and direct sequencing | A C C T C | This study |
| NC_008132 (Midori, the last wild bird in Japan) | n/a | G T . . . | http://www.ncbi.nlm.nih.gov/nuccore/NC_008132 |
| AB104903–AB104907 (the five birds were descendants of Chinese captive population) | n/a | . . . . . | http://www.ncbi.nlm.nih.gov/nuccore/ |
| Haplotype I (9 captive birds) | PCR and direct sequencing* | – – – C A | Zhang (2004) [56] |
| Haplotype II (8 captive birds) | PCR and direct sequencing* | – – A . G | Zhang (2004) [56] |
| Haplotype III (2 captive birds) | PCR and direct sequencing* | – – A . . | Zhang (2004) [56] |
| Haplotype IV (23 captive birds) | PCR and direct sequencing* | – – . G | Zhang (2004) [56] |
| One haplotype (7 captive birds) | PCR and direct sequencing | – – . . . | Lei et al. (2004) [57] |

n/a means no information on how these sequences were generated. * represents no information on how the PCR products were sequenced. A dash means that the
nucleotide was not available. A dot represents the nucleotide identical to JX853815.
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hyper-variable segment in the control region clearly demonstrated the absence of structural genetic diversity in mtDNA genome among the current Crested Ibis populations in China.

Function, Origin and Inheritance of the mtDNA Length Polymorphism and Heteroplasmy in Crested Ibis

Similar to the repeat motif in (CA\textsubscript{3}CA\textsubscript{2}CA\textsubscript{3})\textsubscript{n} identified among all the captive and wild Crested Ibis, such short CA-type tandem repeats located in the control region close to the rRNA-Phe region have been observed in several bird species of Alcidae, Laridae, Scolopacidae, Spheniscidae and Threskiornithidae families [19,41,60,61,62,63], which were all thought to be related to other groups in the order Ciconiformes following the Sibley-Ahlquist taxonomy [64]. However, the former three families are now placed in the order Charadriiformes, the forth in Sphenisciformes and the fifth in Pelecaniformes [39]. These changes seem to suggest a deep common ancestry of this type of the tandem repeats. The control region contains the replication origin for heavy strand and the promoters for transcription of both heavy and light strands [65,66]. In human, certain mutations in the control region were accumulated with aging in specific tissues and considered to have been subject to climatic selection [67,68]. Munwes et al. [47] studied the length heteroplasmy of a 75 bp long tandem repeat in the control region of eastern spadefoot toads (Pelobates syriacus) distributing from edge to core with different environmental conditions. They found that the number of repeats and level of heteroplasmy increased from edge to core, suggesting that the control region may not be neutral but under selection. Length heteroplasmy spanning the functional regions such as termination associated sequences (TAS) and light and heavy strand promoter sites (LSP and HSP, respectively) was detected in several species [14,41,43]. The putative TAS and conserved sequence blocks were mapped to the 5’-end to middle range of the complete control region sequence of Crested Ibis; therefore this repeat motif was believed to be an evolutionary outcome of a simple perfect microsatellite DNA structure without immediate inheritance but the effect of possible genetic hitch-hiking on its inheritance could not be ruled out as well [69].

It is believed that the Taq polymerase slippage rate is inversely related to repeat motif length in microsatellites [52,70,71]. No stutter band was observed at chicken LEI0258 locus carrying intensive length variations in this case) repopulate the offspring of next generation from the germ line [21,22,25,26,27,75]. No reduction of mtDNA content in early primordial germ cells [76,77] but the relaxed amplification of a subpopulation of mtDNA genomes in early oogenesis [23] or during postnatal folliculogenensis [78] in mice or by the end of oogenesis in fish [32] is suggested to have contributed to the random mtDNA segregation [31]. However, the differences in the properties of heteroplasmic mtDNA structures (point mutation v.s. length variation) and in the cleavage patterns of embryos between fishes (discoïdal meroblastic), birds (discoïdal meroblastic) and mammals (rotational holoblastic) require further investigation on how the bottleneck influences the transmission of mtDNA mutations in birds. The rapid segregation of intensive mtDNA length polymorphism and heteroplasmy firstly detected in the highly inbred Crested Ibis populations carrying no structural mtDNA diversity offers an opportunity for further investigation of the mechanism of such bottleneck responsible for random mtDNA segregation in birds.

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Author Contributions

Conceived and designed the experiments: JLH CQD XLH. Performed the experiments: XLH. Analyzed the data: XLH JLH CQD. Contributed reagents/materials/analysis tools: CQD JLH. Wrote the paper: XLH JLH CQD.

References

1. Archibald GW, Lantis SDH (1978) Conservation of the Japanese Crested Ibis. Proceedings of the 1976 Conference of the Colonial Waterbird Group: 1–15.
2. Ding CQ (2004) Research on the Crested Ibis. Shanghai: Shanghai Scientific and Technological Publishing House. (in Chinese).
3. Yamashina Y, Nakanishi G (1983) Proceedings of the 1978 Conference of the Colonial Waterbird Group: 1–15. (in Chinese).
4. Liu YZ (1981) The re-discovery of the Crested Ibis in Qinling Mountain. Acta Zoologica Sinica 27: 275. (in Chinese).
5. Zhang Z, Ding C (2008) The in-situ conservation of Crested Ibis and relate research progress. Science and Technology Review, 26: 48–53. (in Chinese with English abstract).
6. Ding C (2010) Crested Ibis. Chinese Birds 1: 156–162.
7. BirdLife International (2012) Nipponia nippon. In: IUCN 2012. IUCN Red List of Threatened Species. Version 2012 2. <www.iucnredlist.org>. Downloaded on 09 January 2013.

8. Hale LR and Singh RS (1986) Extensive variation and heteroplasmy in size of mitochondrial DNA among geographic populations of Drosophila melanogaster. Proc Natl Acad Sci USA 83: 8813–8817.
9. Solignac M, Monnerot M, Mounouda JC (1983) Mitochondrial DNA heteroplasmy in Drosophila melanogaster. Proc Natl Acad Sci USA 80: 6941–6946.
10. Harrison RG, Rand DM, Wheeler WC (1985) Mitochondrial DNA size variation within individual crickets. Science 228: 1446–1448.
11. Rand DM, Harrison RG (1986) Mitochondrial DNA transmission genetics in crickets. Genetics 114: 935–950.
12. Rand DM, Harrison RG (1989) Molecular population genetics of mtDNA size variation in crickets. Genetics 121: 551–569.
13. Denison LD, Wright JW, Brown WM (1983) Length variation and heteroplasmy are frequent in mitochondrial DNA from parthenogenetic and bisexual lizards (genus Cnemidophorus). Genetics 110: 689–707.
42. Zhang B (2005) Studies on the mitochondrial DNA diversity and MHC class II B genes diversity in Crested Ibis (Nipponia nippon). [Dissertation]. Zhejiang University. (in Chinese with English abstract).

43. Buroker NE, Brown JR, Gilbert TA, O'Hara PJ, Beckenbach AT, et al. (1990) Length heteroplasmy of sturgeon mitochondrial DNA: an illegitimate elongation model. Genetics 124: 157-163.

44. Boyce TM, Zwick ME, Aquadro CF (1989) Mitochondrial DNA in the bark snake: Sinea, structure and heteroplasmy. Genetics 123: 825-836.

45. Brown JR, Beckenbach AT, Smith MJ (1992) Mitochondrial DNA length variation and heteroplasmy in populations of white sturgeon (Acipenser transmontanus). Genetics 132: 525-535.

46. Munews I, Geffen E, Friedmann A, Tikuomski Y, Gafni S (2011) Variation in repeat length and heteroplasmy of the mitochondrial DNA control region along a core-edge gradient in the eastern spadefoot toad (Pelobates fuscus). Mol Ecol 20: 2878-2887.

47. Berger C, Hatzer-Gruitwieser P, Hohoff C, Parson W (2011) Evaluating sequence-determined mtDNA heteroplasmy by allele size analysis. Forensic Sci Int Gen 5: 142-145.

48. Forster I, Forster P, Gurney SM, Spencer M, Huang C, et al. (2010) Evaluating length heteroplasmy in the human mitochondrial DNA control region. Int J Legal Med 124: 133-142.

49. Latz-Bonengel S, Sanger T, Pollak S, Szöbor R (2004) Different methods to determine length heteroplasmy within the mitochondrial control region. Croat Med J 45: 274-275.

50. Konko F, Satta Y, Matsuru ET, Ishiwa H, Takahata N, et al. (1999) Incomplete maternal transmission of mitochondrial DNA in Drosophila. Genetics 126: 657-663.

51. Kua K, Martens J, Nazareno AA, Orell M (2003) Paternal leakage of mitochondrial DNA in the great tit (Parus major). Mol Biol Evol 20: 243-247.

52. Schwartz M, Vising J (2002) Paternal inheritance of mitochondrial DNA. N Engl J Med 347: 576-580.

53. Zhang YD (2004) Study on DNA polymorphism in artificially reared Nipponia nippon. [Dissertation]. Shanxi: Northwest Sci-Tech University of Agriculture and Forestry. (in Chinese with English abstract).

54. Lei CZ, Zhang YD, Cheng H, Fan GL (2004) Analysis of mtDNA D-loop sequence polymorphism in Nipponia nippon. Animal Biotechnology Bulletin 9: 104-110.

55. Shinde D, Lai Y, Sun F, Arnheim N (2003) Tandem duplication of mitochondrial DNA in the black-faced spoonbill, Platalea minor. Genes Syst Evol 84: 297-305.

56. Sibley CG, Ahlquist JE (1990) Phylogeny and classification of birds. New Haven: Yale University Press.
74. White DJ, Gemmell NJ (2009) Can indirect tests detect a known recombination event in human mtDNA? Mol Biol Evol 26: 1435–1439.
75. Bergstrom CT, Pritchard J (1998) Germline bottlenecks and the evolutionary maintenance of mitochondrial genomes. Genetics 149: 2135–2146.
76. Cao L, Shitara H, Horii T, Nagao Y, Imai H, et al. (2007) The mitochondrial bottleneck occurs without reduction of mtDNA content in female mouse germ cells. Nat Genet 39: 386–390.
77. Cao L, Shitara H, Sugimoto M, Hayashi J, Abe K, et al. (2009) New evidence confirms that the mitochondrial bottleneck is generated without reduction of mitochondrial DNA content in early primordial germ cells of mice. PloS Genet 5: e1000756.
78. Wai T, Teoli D, Shoubridge EA (2008) The mitochondrial DNA genetic bottleneck results from replication of a subpopulation of genomes. Nat Genet 40: 1484–1488.