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Mitochondrial DNA-based Analysis of Genetic Variation and Relatedness among Sri Lankan Indigenous Chickens and the Ceylon Junglefowl (*Gallus lafayetti*)

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Abstract

Indigenous chickens (IC) in developing countries provide a useful resource to detect novel genes in mitochondrial and nuclear genomes. Here, we investigated the nature and level of genetic diversity in IC from five distinct regions of Sri Lanka using a PCR-based resequencing method. Additionally, we investigated the relatedness of IC to different species of junglefowls including Ceylon (CJF; Gallus lafayetti), a subspecies that is endemic to Sri Lanka, green (Gallus varius), grey (Gallus sonneratii), and red (Gallus gallus) junglefowl. A total of 140 birds including eight CJF were used to screen 613 bp of IC and 675 bp of CJF control region of the mitochondrial DNA sequence for single nucleotide polymorphisms (SNPs) and other variants. We detected and validated 44 SNPs, which formed 42 haplotypes and six haplogroups in IC. The SNPs observed in the CJF were distinct and the D-loop appeared to be missing a 62 bp found in IC and the red junglefowl (RJF). Among the six haplogroups of IC, only one was region-specific. Estimates of haplotype and nucleotide diversities ranged from 0.901 to 0.965 and from 0.011 to 0.013, respectively. Estimates of genetic divergence were inconsistent but generally low in all the regions. Further, variation among individuals within regions accounted for 92% of the total molecular variation among birds. The Sri Lankan indigenous chickens were more closely related to red and grey junglefowls than to CJF, suggesting a multiple origin. The molecular information on genetic diversity revealed in our study may be useful in developing genetic improvement and conservation strategies to better utilize indigenous Sri Lankan chicken resources.

Keywords

Indigenous chicken; Genetic diversity; Sri Lanka

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Supplementary Material

The following supplementary material is available for the article online at

Table S1a Haplotypes and substitution sites of indigenous Chicken (between segment of 1-613 bp)

 $[\]label{eq:stable} \textbf{Table S1b} \ \text{Haplotypes and substitution sites of Ceylon Junglefowl (between segment of 1-675 bp)}$

Figure S1 The alignment of mitochondrial D-loop sequences of Sri Lankan Indigenous Chicken (IC), nine Ceylon Junglefowl, GenBank sequence of domestic chicken

Introduction

Diversity among farm animals within and among countries is of major interest to the scientific community as it is a significant resource for livestock development and for responding to changing needs and production requirements. With increasing world population, there is concern that the growing demands for animal products are eroding these genetic resources especially in developing countries, where most of the diversity is found. In recognition of this concern, many efforts have begun to characterize animals in developing countries to provide a foundation for developing sustainable genetic improvement approaches. Chief among these efforts is the program by the Food and Agricultural Organization (FAO) of the United Nations to develop a Global Strategy for the Management of Farm Animal Genetic Resources or FAnGR (Gibson et al., 2005; http://www.fao.org/ag/cgrfa/AnGR.htm).

Some efforts to understand diversity in native or indigenous chickens (IC) in developing countries have been described. Using microsatellites and DNA pools, Hillel et al. (2003) evaluated variation in 52 chicken populations that included selected and unselected chicken stocks. Though estimates varied, it was observed that unselected populations were more diverse. In another global diversity assessment in IC and commercial chickens, Granevitze et al. (2007) reported that populations from Asian countries that included China and Vietnam had high microsatellite-based heterozygosities that reflect their management histories. Mwacharo et al. (2007) also recently used microsatellites to assess genetic diversity among African IC from Ethiopia, Kenya, Sudan and Uganda. The populations appeared to cluster by region as could be expected from geographic (or reproductive) isolation. A similar analysis of relatedness within and among Chinese IC using microsatellites was reported by Qu et al. (2006). The indigenous breeds were also highly diverse and clustered by geographic regions.

Efforts to understand genetic diversity in commercial and non-commercial chickens have also involved characterization of variation in the mitochondrial genome. Though relatively few, the mitochondrial DNA (mtDNA) based studies have also provided insight into the maternal origins of chickens (Akishinonomiya et al., 1994; Liu et al. 2006; Niu et al., 2002). In a diversity study involving the D-loop of the mitochondria, Oka et al. (2007) evaluated genetic variation within and among Japanese IC and also assessed their routes of introduction into Japan. Using haplogroups, they showed that indigenous chickens in Japan have multiple origins that include both game and non-game chickens. Given the consensus on the Asian origins of domestic chickens (Akishinonomiya et al., 1994; 1996), characterization of IC in Asian countries thus continues to be of interest.

Sri Lankan IC are geographically isolated from the rest of the Indian sub-continent. This isolation may be responsible for the Ceylon Junglefowl (CJF, *Gallus lafayetti*) being endemic to Sri Lanka (Ceylon). Like with other Gallus species, the relationship between CJF and the domestic chicken as well as with other Galliformes remains of interest to scientists. There is a lack of consensus, for example, on whether the domestic chicken has a mono- or polyphyletic origin that could include the CJF. Recently, Krieg et al. (2007) used annotated retroposed element activity to find the evolutionary evidence of Galliformes. Using the fixation pattern of the transposed elements in different galliformes, they revealed that CJF and Red Junglefowl hold very close taxonomic positions in the phylogenetic tree of Galliformes. However, contribution of CJF in evolutionary process of domestic chicken is not yet been fully revealed. As a crossroads of ancient sea trade routes that connect Asia and the western world, Sri Lanka has been enriched with a variety of animal germplasms including chicken, which eventually developed into a distinct indigenous population. Diversity and relatedness among Sri Lankan IC have been little examined. Using randomly

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amplified polymorphic DNA analysis, Silva and Rajapaksha (2005) reported that the Sri Lankan IC, when considered as a relatively homogenous group, were more closely related to commercial chicken, the Rhode Island Red, than to CJF. To determine if chickens within Sri Lanka are a homogenous group and to further evaluate the relatedness of the IC with CJF, chickens from five different geographical locations were evaluated for genetic diversity in the mitochondrial D-loop. The data may provide some useful information in the ongoing debate about whether chickens have a single (Fumihito et al., 1996; Hillel et al., 2003) or multiple origins (Moiseyeva et al., 1998).

Materials and Methods

Samples

Blood samples were collected on FTA cards (Whatman, Inc.) from a total of 132 IC in five different geographical regions including north-central (NCP), north-western (NWP), southern (SP), uva (UP), and western (WP) provinces (Figure 1). Within each region, samples were collected from several birds from multiple households in different villages. To minimize the chances that the birds used from each village were related, a single bird was used from each household. The households within each village from which each bird was used were approximately 0.5 to one mile apart. Blood was also collected on FTA cards from 9 birds identified as CJF at the national Zoo (3 birds) and from the wild in central Sri Lanka (6 birds).

Molecular analysis

Extraction of DNA from the FTA cards was according to the recommended protocol of the manufacturer (Whatman, Inc.). The genomic DNA was used for PCR as previously described by Guan et al. (2007) with a minor modification. Though the forward primer, as described by Guan et al. (2007) was 5' AGGACTACGGCTTGAAAAGC 3', the reverse primer, 5' GCGATCACGGACTAAAGAGG 3', was developed for the present work using GenBank sequence with accession number NC_001323.. The amplicons, of expected size of 613 bp, were processed, sequenced, and the sequences analyzed using the approach of Guan et al. (2007).

Population genetic analyses

The sequences were initially analyzed using CLUSTALW (Higgins et al., 1994) as previously described (Guan et al., 2007). Using the DnaSP software (version 4.10.9, Rozas et al., 2003), the following statistics were estimated from the sequence comparisons: haplotype diversity (h), nucleotide diversity (π), genetic differentiation (F_{ST}) and mismatch distribution based on pairwise differences among all haplotypes (Rogers and Harpending, 1992). Parsimony network analysis of IC was done using TCS software version 1.21 (Clement et al., 2000). To further evaluate the partitioning of sequence variation in the five regions, analysis of molecular variance or AMOVA among the IC from the five regions was evaluated using ARLEQUIN (version 3.01, Excoffier, 2006). The D statistics (Tajima, 1989) were used to estimate whether the D-loop data in the IC, though relatively small in size, was consistent with the expectation of neutrality.

Phylogenetic analyses

Genetic relatedness among birds from the five regions as well as between the IC and CJF, green (*Gallus varius*), grey (*Gallus sonneratii*), and red (*Gallus gallus*) junglefowls were assessed using PAUP* version 4.0 (Swofford, 2002). Publicly available sequences in Genbank for the junglefowls, other than CJF were used in the analyses. Phylogenetic trees

were constructed using the neighbor-joining method. One thousand bootstrap replicates were used to assess confidence in the grouping (Felsenstein & Kishino,1993).

Results and Discussion

A total of 613 and 675 bp of the mitochondrial D-loop sequence from IC and CJF, respectively, were involved in the analyses. The difference in sequence length, considering that the amplicons sequenced were from the same primer pair, is due to an insertion of 62 bp in the CJF (Figure S1) at position 356 of the Genbank reference *Gallus gallus* mtDNA sequence (Accession number NC_001323). This insertion in the CJF has previously been reported, based on sequence from a single bird, to be about 61 bp and to also occur in the grey junglefowl (GrJF, Nishibori et al., 2005). The sequences flanking this insertion showed on average, the percentage similarity of 81.5% with the IC D-loop sequence. One of the CJF samples obtained from the Zoo, CJF141, lacked this insertion. Additionally, the sequence of the CJF141 also had a 99.5% sequence identity with the IC D-loop sequence as shown in Figure S1. Therefore, CJF141 was removed from further analyses as we believe it is most likely an IC.

Within the IC, a total of 42 haplotypes were detected from 44 polymorphic sites (Table 1). The sequences of all the haplotypes have been submitted to Genbank and assigned accession numbers. The haplotypes ranged in frequency from less than 1 to 12%. Only three haplotypes, SLvtHap1, 26 and 32, were observed in all five regions of Sri Lanka and 31 were detected in only one region. Sixty one percent of the haplotypes were unique to the SP region (Figure 1). Six haplogroups (A-F) based on shared SNPs as shown in Table S1 (Supplement) were identified from the 42 haplotypes. All but two, haplogroups E and F, were found in all the regions sampled (Figure 1). The haplogroups ranged in frequency from 0.02 to 0.33. In the CJF 21 SNPs formed six haplogroups (Table S2).

The diversity indices for IC ranged from 0.901 to 0.965 and from 0.011 to 0.13 for h and π , respectively (Table 2). Pairwise genetic (F_{st}) and nucleotide divergence (d_{xy}) estimates were significant for most of the comparisons (Table 3). Nucleotide divergence between CJF and each of the IC populations were also significant with F_{st} ranging from 0.921 to 0.932 (P<0.05). Both estimates of F_{st} and d_{xy} between the CJF and IC populations were several-fold higher than those between IC populations. The negative F_{st} values indicate negligible variation between the regions compared. Within the IC, birds from SP and WP were most divergent according to the estimates of inter-population nucleotide divergence. Further, the analysis of molecular variation revealed significantly high (92%) within region variation (P<0.05).

The consensus of an unrooted neighbor-joining (NJ) tree shows three distinct clusters for RJF, GrJF and IC, the green junglefowl, and the CJF (Figure 2). Within the RJF, GrJF and IC cluster, the RJF appears to be most closely related to haplogroup E and the GrJF to haplogroup A.

In both CJF and IC, Tajima's D statistics for neutrality test (data not presented) was not significantly different from zero (P>0.10). The average pairwise nucleotide differences were 10.63, 7.80 and 7.90 between IC and CJF, among IC, and among CJF, respectively. The distribution of observed mismatches of pairwise differences for IC and CJF populations are given in Figure 3. The IC population deviated from expected values and demonstrated a bimodal pattern of distribution (raggedness r=0.0129, calculated with parameters $\theta_{\text{final}} = \infty$, $\theta_{\text{initial}} = 2.878$, $\tau = 4.920$). The combined mismatch distribution analysis of IC+CJF (based on the parameters $\theta_{\text{final}} = \infty$, $\theta_{\text{initial}} = 9.432$, $\tau = 1.204$) showed two major peaks (raggedness r=0.0126) at around 1 and 9 differences and a smaller peak around 38. The

mismatch analysis among birds within each of the five geographic regions sampled also showed a bimodal pattern (data not shown).

The parsimony network analysis of IC haplotypes revealed five distinct groups of haplotypes with on average 7–8 bp difference (Figure 4). The five groups correspond to five of the six haplogroups identified in this study, and described in Table S1 and Figure 1. The root sequence (95% probability) is one of the most frequent haplotypes and it was included in haplogroup B. Haplogroup A was genetically diverse compared to other groups. The data further suggest that Haplogroups C, D, and E have diverged from haplogroup B and that haplogroup F, which is region specific, diverged from E. Haplogroups D and C appear to be more closely related. The network analysis also appears to support the wide geographic distribution in Sri Lanka of haplogroups A-D that was observed from the mismatch data.

The diversity estimates observed in our study were higher than the diversity indices reported for Japanese IC of 0.0016 (Oka et al., 2007), for Chinese IC of 0.045 (Niu et al., 2002), and for Indian IC of 0.66 (Pirany et al., 2007). The high genetic diversities revealed among Sri Lankan IC could be attributed to the evolutionary history of the population. The country's geographic location on the ancient trade routes, which connected East Asia to Europe and Middle East enabled the exchange of agriculturally important genetic materials including chicken. (Chandrasiri, 2002, McClellan & Dorn, 2006).

Since the CJF is endemic to Sri Lanka, the current comparison with the IC provides the first direct evidence that it is not the progenitor of the native domesticated chickens in the country. Our data shows that the IC are closely related to both RJF and GrJF, which appears to support reports of multiple origins of domestic chickens (Nishibori et al., 2005; Liu et al., 2006; and Oka et al, 2007). This, however, is inconsistent with earlier reports of a monophyletic origin of domestic chickens (Fumihito et al., 1994; 1996).

Given the free movement of birds and farming communities across Sri Lanka, it is not surprising that the current work detected no genetic subdivision of IC. However, the relatively high number of unique haplotypes in different geographical regions may be due to the breeding pattern practiced in rural areas where the birds are raised in free range and there is limited or no exchange of breeding birds among farms (Gunaratne et al., 1993). This tends to minimize inbreeding, and thus causing the relatively high level of haplotype diversity observed.

In summary, the current analyses suggest that in Sri Lanka, the IC have a relatively high level of genetic diversity that is consistent across a wide geographic area. The random geographic distribution of haplogroups indicates an extensive resource both for conservation and/or genetic improvement by breeding. This could be especially useful if divergent haplotypes and/or haplogroups are crossed to take advantage of heterosis. Our data, though based on only a few junglefowls and thus requiring further studies, appears to suggest that the Sri Lankan IC may have originated from either red or GrJF and not from the CJF that is endemic to the country.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Map adapted from www.lib.utexas.edu/maps/sri lanka.htm

Figure 1.

Sampling sites in Sri Lanka. NCP, NWP, SP, UP, and WP represent north central province, north western province, western province, uva province and southern province, respectively. Regional frequency distribution of the haplogroups is shown by each pie chart. The number of birds sampled from each region is presented in Table 2.



- 0.001 substitutions/site

Figure 2.

An unrooted neighbor-joining tree relating the mitochondrial D-loop haplotypes observed in the indigenous chickens of Sri Lanka (haplotypes 1–42) and the Ceylon (haplotypes CJF1-CJF6), Red (RJF, Accession number 71658078), Grey (GyJF, Accession number 71040179) and Green junglefowls (GrJF, Accession number 71040235). A - E refer to haplogroups, with D being that of the Genbank reference sequence, accession number NC_001323. Numbers in parentheses represent boostrap values from 1,000 replicates. Inset gives the enlargement of Haplgroup A



Figure 3.

Frequency distribution of the number of sequence differences observed in pairwise comparisons of 613 bp of mitochondrial D-loop of indigenous chickens (IC) of Sri Lanka and Ceylon Junglefowl (CJF). **a**. comparisons within the IC. **b**. Comparisons among IC and the CJF. **c** comparisons within the CJF. The within-indigenous chickens comparisons lags behind both the intra CJF and inter-CJF and indigenous chickens derivations.



Figure 4.

Parsimony network (minimum spanning tree) of 613-bp partial D-loop sequences from indigenous chickens in Sri Lanka. Circles represent individual haplotypes as described in Table 1. The line connecting two circles, independent of length, indicates a single base pair difference between the two haplotypes. The rectangle indicates the root haplotype based on 95% probability. Filled dots on the line represent intermediate haplotypes (theoretical) not found in the present analysis. The size of each circle is proportional to the frequency of the haplotype. The number in the circle indicate the haplotype and those in parenthesis correspond to the number of individuals with that haplotype (when number >1). Shading indicates haplogroups as described in Figure 1.

Table 1

Mitochondrial D-loop haplotype frequencies and distribution in Sri Lankan indigenous chicken.

Haplotype	Accession number		ΠN	nber ¹			Frequency ²
		NCP	NWP	WP	đ	SP	
SL vtHap1	EU199906	5	7	-	4	5	0.1061
SL vtHap2	EU199907	-	ı	ī		,	0.0076
SL vtHap3	EU199908	1	ī	ı.	,		0.0076
SL vtHap4	EU199909	ı	ı	ī		6	0.0152
SL vtHap5	EU199910	·	-	ī			0.0076
SL vtHap6	EU199911	ī	ī	ı.	,	-	0.0076
SL vtHap7	EU199912	ı	ı	ī		-	0.0076
SL vtHap8	EU199913	-	ı	ī	-	,	0.0152
SL vtHap9	EU199914	ī	ī	ī		1	0.0076
SL vtHap10	EU199915	ı	1	ī		'	0.0076
SL vtHap11	EU199916	ı	7	ī		,	0.0152
SL vtHap12	EU199917	ī	ī	ı.	,	-	0.0076
SL vtHap13	EU199918	-	2	-		-	0.0379
SL vtHap14	EU199919	-	·	ī			0.0076
SL vtHap15	EU199920	ī	ī	ı.	,	-	0.0076
SL vtHap16	EU199921	·	·	-	,	7	0.0227
SL vtHap17	EU199922	·	·	ī		-	0.0076
SL vtHap18	EU199923	'	'			7	0.0152
SL vtHap19	EU199924	,		·	-	,	0.0076
SL vtHap20	EU199925			·		7	0.0152
SL vtHap21	EU199926					-	0.0076
SL vtHap22	EU199927	,	'		-		0.0076
SL vtHap23	EU199928	·	·	ī	-		0.0076
SL vtHap24	EU199929		1	1	10	5	0.1288
SL vtHap25	EU199930	·	,	,		-	0.0076
SL vtHap26	EU199931	1	2	-	б	-	0.0606
SL vtHap27	EU199932	ï	ï		1	'	0.0076

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Haplotype	ACCESSION NUMBER		nZ	moer			F requency-
		NCP	AWN	WP	Ð	\mathbf{SP}	
SL vtHap28	EU199933	-					0.0076
SL vtHap29	EU199934	1	'	ı	·	'	0.0076
SL vtHap30	EU199935	1	,	1	4	4	0.0758
SL vtHap31	EU199936	'	'	,	7		0.0152
SL vtHap32	EU199937	3	4	5	З	7	0.1288
SL vtHap33	EU199938	-	-	-	ï	-	0.0303
SL vtHap34	EU199939	'	,	,		-	0.0076
SL vtHap35	EU199940	1	'	7	2	0	0.0530
SL vtHap36	EU199941	1	1	4	7		0.0606
SL vtHap37	EU199942	'	1	,			0.0076
SLvtHap38	EU199943	,	'	ı	·	-	0.0076
SLvtHap39	EU199944			,	ï	-	0.0076
SL vtHap40	EU199945	'	,	,	-		0.0076
SL vtHap41	EU199946	,	,	ı	-	'	0.0076
SL vtHap42	EU199947	,	,	7	,		0.0152

wince, western province, uva province and southern province, respectively.

 $^2\mathrm{The}$ cumulative frequency of the indigenous chicken population.

Table 2

Sampling sites, sample size (n), haplotype distribution (f), and haplotype (h) and nucleotide diversities (π), with standard deviations in parentheses, in the indigenous chickens (IC) of Sri Lanka and Ceylon Junglefowl (CJF) based on mitochondrial D-loop sequence comparisons.

Sampling sites	n	f	h	π
NCP	20	14	0.932 (0.044)	0.013 (0.001)
NWP	18	11	0.935 (0.038)	0.013 (0.001)
WP	20	11	0.905 (0.044)	0.013 (0.001)
UP	37	15	0.901 (0.033)	0.011 (0.001)
SP	37	23	0.965 (0.015)	0.012 (0.001)
CJF	8	6	0.929 (0.084)	0.012 (0.002)
Total (IC & CJF)	140	48	0.947 (0.008)	0.017 (0.001)
Total (only IC)	132	42	0.947 (0.009)	0.013 (0.000)

Table 3

Inter-region haplotype (above diagonal) and nucleotide divergence (below diagonal)

	NCP	SP	đ	NWP	WP	CJF
NCP		-0.007	0.107^{*}	-0.025	0.070	0.923
SP	0.012		0.066^*	0.014	0.130^{*}	0.926
UP	0.013	0.012		0.123^{*}	0.183^{*}	0.932
NWP	0.013	0.013	0.013		0.049^{*}	0.921
WP	0.014	0.014	0.014	0.014		0.923
CJF	0.062	0.061	0.060	0.061	0.063	

* Significantly different at P<0.05