Assessment of genetic diversity using mitochondrial DNA variation in Gir cattle of India

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Abstract: The present study is carried out with the aim to study the genetic diversity of 60 Gir cattle in the newly adopted herd of NDRI on the basis of mitochondrial D-loop hyper variable region. The animals were purchased from 4 major locations- IGFRI-Jhansi, Bhilwara, Ajmer and Kisangarh. After aligning the sequences a total of 53 haplotypes were identified. The Overall haplotype Diversity was observed to be 0.996±0.003 which shows that the population is diverse. Nucleotide diversity ranged from 0.693±0.35 to 0.707±0.35 with overall diversity of 0.704±0.34 across all 60 sequences and the average no. of nucleotide difference varied from 279.45±125.42 to 285.16±128.36. \( F_{ST} \) value is significantly different from 0 for all pair wise combinations representing significant amount of Genetic differentiation. \( F_{ST} \) estimates between the population shows that animals from Ajmer and Bhilwara are more genetically differentiated (0.02987) while animals from Kisangarh and Bhilwara are genetically more closer (0.00440). All the above results show substantial variability between populations. These findings can be used for designing proper breeding and management strategies for Gir cattle in NDRI Herd.

Keywords: Genetic Diversity, Gir cattle, Genetic Improvement, Mitochondrial DNA

The most important component of any breed improvement programme is genetic diversity which is the major cause of response to selection or genetic gain and plays a major role in improving livestock. Maintaining genetic diversity should be an important task in livestock breeding as it strengthens a population by increasing the likelihood that at least some individuals will be able to survive major disturbances and it enables the population less susceptible to inherited disorders and climate change. Animal genetic resources consist of species that are of agricultural and economic importance to man and food production systems depend heavily on utilization of locally adapted animal species. Genetic diversity and population structure need to be understood for guiding breed development programs en route for meeting the current production which allows sustained genetic improvement, to facilitate adaptation, and to device proper measures of utilization and conservation of livestock breeds (Notter, 1999; Dalvit et al. 2008). Thus, assessment of population structure and genetic diversity is an important basic tool for genetic improvement through modern breeding strategies (Toro et al. 2009; Sharma et al. 2013; Chung et al. 2017). Various technologies to study genetic diversity include PCR-RFLP, mini-satellites, micro-satellites, mitochondrial DNA analysis and SNP chips. The mtDNA has proved to be valuable in the study of genetic diversity as it shows maternal inheritance and changes rapidly than single copy nuclear DNA in mammals (Brown et al. 1982). The D-loop is the major control region for mitochondrial DNA (mtDNA) expression. The rate of nucleotide substitution in mtDNA is five to ten times higher than that of nuclear DNA (Brown et al. 1979). The mtDNA polymorphisms have been widely used to investigate the structure of populations, interspecies variability and identification of maternal lineages and postnatal growth (Bradley et al. 1998; Troy et al. 2001; Liu et al. 2004; Malau-Aduli et al. 2004; Yoon et al. 2005; Odahara et al. 2006; Lei et al. 2007). Considering the importance of cattle in Indian agriculture, few efforts have been made to evaluate the genetic diversity. Therefore, the present work was undertaken to quantify the genetic diversity of Gir breed of cattle in the National Dairy Research Institute, Karnal, Haryana. The objectives of this study was to use mitochondrial DNA hyper variable region polymorphisms to characterize the within-breed genetic diversity and to use the molecular information supplied to elucidate the
genetic diversity of this breed in order to establish adequate breeding strategy.

Blood samples were collected from the standing 60 female cattle which had completed their first lactation. Around 10 ml of venous blood was collected aseptically from the jugular vein of the animals in a 15 ml vacutainer tube under sterile condition using 0.5 ml of EDTA as an anticoagulant. Blood samples were stored in -20°C until DNA isolation. DNAs were extracted according to the manufacturer’s standard protocol of Wizard genomic DNA purification Kit (Promega, USA). Partial mitochondrial D-loop hyper variable region was amplified using the polymerase chain reaction. The reported primer with forward 5’-CCCAGGCAAGAGGTAATGTA-3’ and reverse primers 5’-TGTCCTGTGACCATTGACTG-3’ (Bhuiyan et al. 2007) was used to amplify 588 bp from the hyper variable region of D-loop. PCR amplification was carried out in a 25 μl reaction volume containing 100 ng of genomic DNA, 1 × PCR master mix buffer (50 mM KCl, 10 mM Tris-HCl, pH 8.3, 1.5 mM MgCl2, 200 μM dNTPs, 1 U Taq polymerase) (Promega, USA) and 0.5pM of each primer. Amplification was performed in a C1000 Thermal cycler (Applied Biosystems, USA) thermal cycler using a 10 min denaturation step followed by 30 cycles of 30 sec at 94°C, 30 sec at 62°C, 1 min at 72°C and a final extension at 72°C for 10 min. The size of amplification product was checked by loading 5 μL PCR product on to a 1.8 % agarose gel containing 0.5 μL/mL ethidium bromide. Sequencing was done directly by Sanger DNA Sequencing method (Apical Scientific Sdn Bhd, Malaysia). The sequences of the PCR product were analyzed using Chromas software. The sequences were edited and corrected by aligning forward and reverse sequences using BLASTN. Sites representing a gap in any of the aligned sequences were excluded from the analysis.

Alignment of 60 sequences of Gir cattle illustrated 53 different haplotypes in this investigation. Identical sequences were considered as the same haplotype. The populations from Bhilwara and Ajmer had highest no. of haplotypes with 18 and 17 respectively. The overall haplotype diversity was 0.98 throughout the population. Overall haplotype Diversity shows the population is diverse. The nucleotide diversity ranged from 0.693 to 0.707 (Table 1) with an overall nucleotide diversity of 0.704. The average no. of nucleotide differences (k) was 284.067 with a maximum of 285.169 in Kisangarh and minimum of 279.465 in population from Bhilwara. Tajima’s D value was found to be 0.37889, which showed that all populations evolved naturally and possesses greater genetic diversity.

In total, on an average 1092.5 variable substitutions were determined in the mtDNAs from 53 haplotypes of Gir cattle. Among them, 690 substitutions were transitions and 402.5 transversions (transitions/transversions rate = 1.72) indicating a heavy bias towards transition substitution that has previously been reported for bovine mtDNA (Loftus et al. 1994; Mannen et al. 1998; Henkes et al. 2005; Bhuiyan et al. 2007 and Sharma et al. 2015).

Table 2 represents Global Fixation values between the populations. FST value are significantly different from 0 for all pair wise combinations representing significant amount of genetic differentiation between population. The mean sequence divergence values ranged between 0.029 to 0.004 among different populations. FST estimates between the population shows that animals from Kisangarh and Bhilwara are more genetically differentiated while animals from Ajmer and Bhilwara are genetically closer values for pair wise FST was significant for populations of Ajmer and Bhilwara. Similarly, AMOVA revealed

| Locations | No. of Sequences | Nucleotide Diversity (pi) | Avg. Nucleotide Differences(k) | Haplotype Diversity(Hd) |
|-----------|------------------|---------------------------|-------------------------------|------------------------|
| Jhansi    | 13               | 0.704±0.36                 | 283.80±129.95                 | 1.000±0.030            |
| Ajmer     | 12               | 0.702±0.36                 | 283.24±130.49                 | 0.984±0.040            |
| Bhilwara  | 18               | 0.693±0.35                 | 279.45±125.42                 | 1.000±0.018            |
| Kisangarh | 17               | 0.707±0.35                 | 285.16±128.36                 | 1.000±0.020            |
| Overall   | 60               | 0.704±0.34                 | 284.06±123.15                 | 0.996±0.003            |

| Divergence Values between the populations | Jhansi | Ajmer | Bhilwara | Kisangarh |
|----------------------------------------|--------|-------|----------|-----------|
| Jhansi                                | 0.012  |       |          |           |
| Ajmer                                 | 0.020  | 0.029 |          |           |
| Bhilwara                              |        | 0.006 | 0.004    |           |
| Kisangarh                             |        |       | 0.006    |           |
that percent of variation among the populations was 0.61 % while within the population it was 99.39 %.

**Conclusions**

This study involves detailed analysis of the genetic diversity and indicates existence of genetic diversity and population structure in Gir cattle. It could generate the baseline information which will assist in formulating effective breeding strategies in future for overall genetic improvement of Gir cattle in the NDRI herd. However a well defined breeding plan is a must to maintain the existing genetic diversity which will help the future bull mother farm to assist in dissemination of high merit germplasm to the farmer’s herd.

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