Association of polymorphism at Exon-1 of the alpha 1-antitrypsin gene with milk production traits in Sahiwal and Karan Fries Cattle

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Abstract: PCR-RFLP analysis of PCR products was carried out using SphI/PaeI enzyme for 100 Sahiwal cattle and 115 Karan Fries cattle. In Sahiwal cattle, 448 bp exhibited three genotypes viz., AA (448), AB (448+313+135bp), BB (313+135 bp) with corresponding genotype frequencies of 0.20, 0.45, 0.35 and gene frequencies 0.42 (A) and 0.58 (B). In Karan Fries cattle, genotype frequencies were 0.29, 0.21, 0.50 respectively with corresponding gene frequencies 0.39 (A) and 0.61 (B). These genotypes of 448 bp are highly significant for FL305DMY, FLTMY, FL305DSNFY and non significant for FL305 DFY and FL305DPY in Sahiwal. In Karan Fries cattle these genotypes are highly significant for FL305DMY, FLTMY, FL305DSNFY and FL305DPY and non significant for FL305DFY. AB genotype was superior for FL305DMY, FLTMY, FL305DSNFY traits and AA genotype was superior for FL305DFY and BB genotype was superior for FL305DPY trait in Sahiwal cattle and In Karan Fries Cattle, AB genotype was superior for all traits.

Keywords: Sahiwal, Karan Fries, Cattle, Alpha 1-antitrypsin gene, Milk constituents traits

Introduction

India is a rich reservoir of genetic diversity in cattle with 43 recognized cattle breeds. The total number of cattle population was 190.90 million (37.28% of total livestock population) out of which indigenous cattle population comprised of 151.17 million (19th Livestock Census, 2012). Milk is an important source of essential nutrients for lactating calves and a key raw material for human food preparations (Reinhardt et al. 2012). The total milk production in India was 132.43 million tons (cattle contributes 43.11% with 59.92 million tons milk) in 2012-13 (BAHS, 2014) and in terms of milk production, India ranks first in the world. As far as milk production is concerned Sahiwal is the best dairy breed of the Indian subcontinent. Sahiwal breed had been termed due to its habitat in Montgomery district of Pakistan which is now named as Sahiwal and in India, number of herds of this breed is maintained. This is a comparatively heavy breed with a symmetrical body and loose skin. This breed is therefore, also known as Lola (loose skin), Lambi Bar, Montgomery, Multani, Teli (Nivsarkar et al. 2000). The Karan Fries breed has been evolved from crossbreeding between Tharparkar and Holstein Friesian at the ICAR-National Dairy Research Institute, Karnal, Haryana. The breed has 50% inheritance from Friesian. The breed carries black patches and sometime is completely dark with white patches on the forehead and the switch of the poll. The udder is also dark with white patches in teats. The animals are extremely docile and very good yielders.

Alpha 1-antitrypsin (AAT), a strong protease inhibitor, also known as α1-protease inhibitor (α1PI), belongs to the super family of serpins or serine proteinase inhibitors in addition to others C1 esterase, antithrombin and α1-antichymotrypsin. AAT is a glycoprotein which forms a sodium dodecyl sulfate (SDS) staple complex with elastase. The molecular mass of AAT is about 52 kDa, and carbohydrates account for 15% of its mass (Carrell et al. 1982). The bovine AAT gene consists of five exons, spanning about 9 kb of genomic DNA and encoding a 416-AA protein. Alpha 1-antitrypsin (AAT) can protect vulnerable elastic tissues from degradation by neutrophil elastase, this is an important issue as protein degradation in bovine milk affects the quality of dairy products.
Materials and methods

Experimental animals and genomic DNA isolation

The data for present study pertained to various milk production and milk composition traits were collected from history sheets and milk composition constituents registers, data on milk production and milk constituent records of 100 Sahiwal and 115 Karan Fries cattle spread over a period of 13 years period 2004 to 2016 were collected from Animal Genetics and Breeding division of ICAR-National Dairy Research Institute, Karnal, Haryana. Blood samples were collected from the targeted population. About 10 ml of venous blood was collected aseptically from the jugular vein of the animals in a 15ml polypropylene centrifuge tube under sterile condition using 0.5 ml of EDTA as an anticoagulant. The tube was shaken gently to facilitate thorough mixing of blood with the anticoagulant. The tubes containing blood samples were transported to the laboratory as soon as possible in an icebox containing ice packs and were stored in the refrigerator at –20°C temperature until the isolation of DNA was done. Phenol extraction method as described by Sambrook and Russell, (2001) was used for isolation of genomic DNA. Horizontal submarine agarose gel electrophoresis was used to check the quality of genomic DNA. The purity of genomic DNA was checked by spectrophotometer. The 6 µl of genomic DNA of each sample was dissolved in 294 µl of triple distilled water and spectrophotometer readings at OD$_{260}$ and OD$_{280}$ was recorded against 300µl double distilled water as a blank. Genomic DNA samples showing the OD ratio in the range of 1.7 to 1.9 was used further in the study.

Amplification of targeted region of AAT genes

Only good quality genomic DNA was used for amplification of Exon 1 of 448 bp of AAT by polymerase chain reaction under optimized conditions.

F: 5'-ATGGCACCCTCCATCAAGCG-3'
R: 5'-CCACTAGCTTTGCACCTCCAT-3'

Forward and reverse primers were used to amplify targeted region of AAT gene, which was reported by Li et al. (2010) the best amplification of the desired fragment was taken for further analysis. The standard programme is given as under (Table 01):

Polymerase chain reaction amplification of targeted region of AAT genes

Various annealing temperatures were tried for PCR amplification of AAT genes. Total volume of 25 µl for each sample was used to set up the PCR reactions. The set of primer was used to amplify target regions of AAT genes in Sahiwal and Karan Fries cattle. The best results were obtained when amplification was performed in PCR thermal cycler (Eppendorf Germany) programmed for 32 cycles with an initial denaturation at 94°C for 05 minutes, denaturation at 94°C for 30 second, annealing at 62°C for 30 second and extension at 72°C for 30 second with a final extension at 72°C for 10 minutes (Plate 01 and 02).

Agarose gel electrophoresis

The amplified product was checked for quality and quantity by agarose gel electrophoresis as described by Sambrook and Russell (2001).

PCR amplification and restriction digestion

This is a useful technique for screening of sequence variations that give rise to RE sites. Such analysis involves amplification of a specific region of DNA encompassing the polymorphic RE sites and digestion of the amplified DNA fragment with the respective RE. This technique allowed the identification of only two alleles per locus and is slow to be used with large genome size in mammals, where about 3 x10$^9$ individual nucleotides are present in the total DNA content.

Restriction digestion of the PCR product was performed in a total volume of 30 µl, having 10X buffer Tango 2 µl, PCR reaction mixture 10 µl, restriction enzyme fast digest (Sph I /PaeI) (10 units/µl) 1 µl and nuclease free water 17 µl. The reaction mixture was spinned for few seconds for uniform mixing and then incubated at 37°C for 30 minutes in the water bath (Table 2).

Agarose gel electrophoresis of digested product

The restriction fragments of different lengths were separated on 2% agarose gel electrophoresis run at 80V for 1hour. The lengths of all fragments generated by restriction enzyme ((Sph I /PaeI) digestion were compared with O’ GeneRuler™ (Fermentas Life Science) 100bp DNA ladder (range 100-1000bp) as a molecular size marker. The PCR-RFLP DNA bands were visualized under UV light and documented by gel documentation system (BioRad, USA). The band size of each sample was judged by comparing with molecular size marker and recorded (Plate 03, 04).

Statistical analysis

The analysis was carried out with appropriate Statistical method using software’s in the computer centre of the institute under the following headings:

Restricted Maximum Likelihood Method (REML)

Estimation of breeding value

The single trait animal model was considered for estimation of breeding value using WOMBAT software (Meyer, 2010). The following animal model was considered:

$$Y_{jk} = X b_i + Z u_j + e_{jk}$$
\[ Y_{ijk} = k^{th} \text{ observation of } j^{th} \text{ random effect of } i^{th} \text{ fixed effect} \]

\[ b_i = \text{ Vector of observation of fixed effect} \]

\[ X = \text{ Incidence matrix of fixed effect} \]

\[ u_j = \text{ Vector of additive genetic effect (animal effect)} \]

\[ Z = \text{ Incidence matrix of random effect} \]

\[ e_{ijk} = \text{ Vector of residual errors} \]

**Association estimation**

Based on the adjusted records, pertaining to milk yield and its constituents on Sahiwal and Karan Fries cattle maintained at ICAR-NDRI, Karnal, regression analysis was carried out to identify SNPs contributing significantly to the variation in milk and its constituents.

\[ Y_{ijk} = a + b_i \text{SNP}_1 + b_j \text{SNP}_2 + \ldots + b_n \text{SNP}_n + e_{ijk} \]

where,

\[ Y_{ijk} = \text{ Adjusted observation on } k^{th} \text{ animal of } i^{th}, j^{th}, \ldots, n^{th} \text{ SNPs} \]

\[ a = \text{ Intercept} \]

\[ b_i = \text{ Partial regression coefficient for SNPs considered} \]

\[ \text{SNP}_{ij...n} = \text{ Effect of SNPs taken as independent variable} \]

\[ e_{ijk} = \text{ Random error NID} (0, \sigma^2_e) \]

**Effect of genotypes on breeding value**

The relative contribution of Genotypes to breeding value of the animal for milk yield and milk constituents was assessed using the following model:

\[ Y_i = \mu + G_i + e_{ij} \]

where,

\[ Y_i = \text{ Breeding value of } j^{th} \text{ animal of } i^{th} \text{ genotype} \]

\[ m = \text{ Overall mean} \]

\[ G_i = \text{ Effect of } i^{th} \text{ genotypes (SNPs/haplotypes)} \]

\[ e_{ij} = \text{ Residual error NID} (0, \sigma^2_e) \]

**Results and discussion**

PCR-Restriction fragment patterns and genotyping at Exon-1 of AAT gene (448 bp) in Sahiwal

PCR-RFLP analysis of PCR products were carried out using reported Sph I/Pae I enzyme for 100 Sahiwal cattle. Exon 1 with 448 bp exhibited three genotype AA (448), AB (448+313+135 bp), BB (313+135 bp) having corresponding genotype frequencies 0.20, 0.45, 0.35 and gene frequencies of 0.42 (A) and 0.58 (B). These genotypes of 448 bp are highly significant for FL305DMY, FLTMY, FL305DSNFY and non significant for FL305 DFY and FL305DPY. The mean ± SE of AA genotype for FL305DMY, FLTMY, FL305DFY, FL305DSNFY, FL305DPY were 1711.95 ± 8.06 kg, 1923.30 ± 11.5 kg, 100.5 ± 0.66 kg, 154.03 ± 0.13 kg and 43.99 ± 0.15 kg respectively and for AB genotype were 1869.67 ± 5.37 kg, 2057.16 ± 7.65 kg, 100.33 ± 0.44 kg, 155.35 ± 0.08 kg, 43.77 ± 0.10 kg respectively and for BB genotype were 1789.01 ± 6.09 kg, 2001.15 ± 8.67 kg, 100.25 ± 0.50 kg, 154.79 ± 0.10 kg, 44.02 ± 0.11 kg respectively. AB genotype was found to be superior for FL305DMY, FLTMY, FL305DSNFY traits and AA genotype was superior for FL305DFY and BB genotype was superior for FL305DPY trait (Table 03 and 04). The result was in agreement with Li et al. (2010) in Chinese Holstein and reported significant for milk fat percentage, milk protein percentage and 305-day milk yield. They concluded that AAT is a potential candidate gene influencing milk production traits and could be implemented in breeding programmes to improve the production performance of Chinese Holstein cattle.

### Table 1 The standard programme for PCR Reaction mix

| PCR component                      | Volume (µl) | Final concentration |
|------------------------------------|-------------|---------------------|
| 2X PCR Master Mix (Fermentas)      | 12.5        | 02X                 |
| Deionised water (DNAase free water)| 7.5         |                     |
| Forward primer                     | 1.0         | 10 pmole            |
| Reverse primer                     | 1.0         | 10 pmole            |
| Genomic DNA (30ng/µl)             | 3.0         | 90 ng/µl            |
| Total                              | 25.0        |                     |

### Table 2 Compositions of restriction enzyme reaction mixture

| Reagent                            | Quantity (µl) |
|------------------------------------|---------------|
| Restriction enzyme (10 units/µl)   | 1.0           |
| 10X Buffer Tango                   | 2.0           |
| PCR reaction mixture               | 10.0          |
| Nuclease free water                | 17.0          |
| Total                              | 30.0          |
The result is also in agreement with Kheiripour et al. (2014) in Holstein dairy cows and reported that the cows of AB genotype had higher milk fat percentage than those of genotype AA. Cows with AB genotype showed 0.07% higher fat % and 0.02% higher protein % as compared to AA genotype. It was concluded that the association value could be implemented as a marker in breeding programmes for these traits.

Regression equation
The significance of association of SNP with different performance traits were estimated by constructing the regression equation and the best fit equation for each of them is given below:

\[
\text{FL305DMY} = 1790.21 - 78.26 \text{ SNP}_{AA} + 79.46 \text{ SNP}_{AB} - 1.20 \text{ SNP}_{BB} \quad (R^2=74.51)
\]

\[
\text{FL305TMY} = 1993.86 - 70.58 \text{ SNP}_{AA} + 63.29 \text{ SNP}_{AB} + 7.29 \text{ SNP}_{BB} \quad (R^2=49.80)
\]

\[
\text{FL305DFY} = 100.36 + 0.14 \text{ SNP}_{AA} - 0.03 \text{ SNP}_{AB} - 0.10 \text{ SNP}_{BB} \quad (R^2=0.09)
\]

\[
\text{FL305DSNFY} = 154.72 - 0.69 \text{ SNP}_{AA} + 0.62 \text{ SNP}_{AB} + 0.06 \text{ SNP}_{BB} \quad (R^2=41.85)
\]

\[
\text{FL305DPY} = 43.93 + 0.06 \text{ SNP}_{AA} - 0.15 \text{ SNP}_{AB} + 0.08 \text{ SNP}_{BB} \quad (R^2=3.04)
\]

PCR-Restriction fragment patterns and genotyping at exon 1 of AAT gene (448 bp) in Karan Fries Cattle

PCR-RFLP analysis of PCR products were carried out using reported Sph I/PaeI enzyme for 115 Karan Fries animals. The 448 bp has three genotypes AA (448), AB (448+313+135bp), BB (313+135 bp) having genotype frequencies 0.29, 0.21, 0.50
respectively and gene frequencies are 0.39 (A) and 0.61 (B). These
genotypes are highly significant for FL305DMY, FLTMY,
FL305DSNFY and FL305DPY and non significant for FL305 DFY.
The mean ± SE of AA genotype for FL305DMY, FLTMY,
FL305DFY, FL305DSNFY, FL305DPY were 3443.20 ± 6.32 kg,
4462.96 ± 6.32 kg, 130.70 ± 6.93 kg, 277.36 ± 0.07 kg and 112.57 ±
0.06 kg respectively and for AB genotypes were 3621.18 ± 7.42
kg, 4640.94 ± 7.42 kg, 139.89 ± 8.13 kg, 279.00 ± 0.08 kg, 114.35 ±

Table 4  Least Square Mean and Standard Error for milk production traits

| Genotype   | N  | Mean (Kg) ± SE          |
|------------|----|-------------------------|
| FL305 DMY  |    |                         |
| AA         | 20 | 1711.95 ± 8.06<sup>a</sup> |
| AB         | 45 | 1869.67 ± 5.37<sup>b</sup> |
| BB         | 35 | 1789.01 ± 6.09<sup>c</sup> |
| FLTMY      |    |                         |
| AA         | 20 | 1923.3 ± 11.50<sup>a</sup> |
| AB         | 45 | 2057.16 ± 7.65<sup>b</sup> |
| BB         | 35 | 2001.15 ± 8.67<sup>c</sup> |
| FL305DFY   |    |                         |
| AA         | 20 | 100.51 ± 6.64            |
| AB         | 45 | 100.33 ± 4.43            |
| BB         | 35 | 100.25 ± 5.02            |
| FL305DSNFY |    |                         |
| AA         | 20 | 154.03 ± 13.30<sup>a</sup> |
| AB         | 45 | 155.35 ± 8.90<sup>b</sup> |
| BB         | 35 | 154.79 ± 10.00<sup>c</sup> |
| FL305DPY   |    |                         |
| AA         | 20 | 43.99 ± 15.30<sup>a</sup> |
| AB         | 45 | 43.77 ± 10.20<sup>b</sup> |
| BB         | 35 | 44.02 ± 11.60<sup>c</sup> |

The mean values with different superscript alphabet indicate highly significant difference (p<0.05) among themselves

Table 5  ANOVA for Exon 1 of AAT gene in Karan Fries

| Source     | df  | SS     | MSS   | F-value |
|------------|-----|--------|-------|---------|
| FL305 DMY  |     |        |       |         |
| Genotype   | 2   | 443214 | 221607| 167.93**|
| Error      | 112 | 147797 | 1320  |         |
| Total      | 114 | 591011 |       |         |
| FLTMY      |     |        |       |         |
| Genotype   | 2   | 443214 | 221607| 167.93**|
| Error      | 112 | 147797 | 1320  |         |
| Total      | 114 | 591011 |       |         |
| FL305DFY   |     |        |       |         |
| Genotype   | 2   | 1270.83| 635.41| 0.40    |
| Error      | 112 | 177744.97| 1587.00|         |
| Total      | 114 | 179015.81|       |         |
| FL305DSNFY |     |        |       |         |
| Genotype   | 2   | 38.35  | 19.17 | 115.47**|
| Error      | 112 | 18.60  | 0.16  |         |
| Total      | 114 | 56.95  |       |         |
| FL305DPY   |     |        |       |         |
| Genotype   | 2   | 44.29  | 22.14 | 140.28**|
| Error      | 112 | 17.68  | 0.15  |         |
| Total      | 114 | 61.97  |       |         |

**Highly significant (p<0.01)
The result is in agreement with Li et al. (2010) in Chinese Holstein and reported significant for milk fat percentage, milk protein percentage and 305-day milk yield. They concluded that AAT is a potential candidate gene influencing milk production traits and could be implemented in breeding programmes to improve the production performance of Chinese Holstein cattle.

The result is also in agreement with Kheiripour et al. (2014) in Holstein dairy cows and reported that the cows of AB genotype had higher milk fat percentage than those of genotype AA. Cows with AB genotype showed 0.07% higher fat % and 0.02% higher protein % as compared to AA genotype. It was concluded that the association value could be implemented as a marker in breeding programmes for these traits.

### Regression equation

The significance of association of SNP with different performance traits were estimated by constructing the regression equation and the best fit equation for each of them is given below:

\[
FLW305DMY = 3530.96 - 87.76 \text{SNP}_{-AA} + 90.22 \text{SNP}_{-AB} - 2.47 \text{SNP}_{-BB} (R^2=74.99)
\]

\[
FLTMY = 4550.72 - 87.76 \text{SNP}_{-AA} + 90.22 \text{SNP}_{-AB} - 2.47 \text{SNP}_{-BB} (R^2=74.99)
\]

\[
FL305DFY = 135.66 - 4.97 \text{SNP}_{-AA} + 4.22 \text{SNP}_{-AB} + 0.74 \text{SNP}_{-BB} (R^2=0.71)
\]

\[
FL305DSNFY = 278.19 - 0.83 \text{SNP}_{-AA} + 0.81 \text{SNP}_{-AB} + 0.01 \text{SNP}_{-BB} (R^2=67.34)
\]

\[
FL305DPY = 113.46 - 0.88 \text{SNP}_{-AA} + 0.88 \text{SNP}_{-AB} - 0.33 \text{SNP}_{-BB} (R^2=71.47)
\]

## Conclusions

PCR-RFLP analysis of PCR products was carried out using Sph I/PaeI for 100 Sahiwal cattle. These genotypes of exon 1 with 448 bp are highly significant for FL305DMY, FLTMY, FL305DSNFY and non significant for FL305 DFY and FL305DPY. AB genotype was superior for FL305DMY, FLTDMY, FL305DSNFY traits and AA genotype was superior for FL305DFY and BB genotype was superior for FL305DPY trait. PCR-RFLP analysis of PCR products were carried out using, Sph I/PaeI for 115 Karan Fries animals. These genotypes of exon 1 with 448 bp are highly significant for FL305DMY, FLTMY, FL305DSNFY and FL305DPY and non significant for FL305 DFY. AB genotype was observed superior for all traits under present investigation.
References

BAHS (2014) Basic Animal Husbandry Statistics. Department of Animal Husbandry, Dairying & Fisheries. Ministry of Agriculture, Govt. of India
Carrell RW, Jeppsson JO, Laurell CB, Brennan SO, Owen MC, Vaughan L, Boswell DR (1982) Structure and variation of human alpha-1-antitrypsin. Nature 298: 329-334
Kheiripour MH, Mahdavi AH, Rahmani HR, Soltani-Ghombavani M, Edriss MA (2014) Association of polymorphism in the alpha-1-antitrypsin gene with milk production traits in Holstein dairy cows. South Afr J Anim Sci 44 (2): 155-160
Li QL, Zhang ZF, Wang CF, Yang H, Wang HM, Li JB, Huang JM, Zhong JF (2010) Association of polymorphism of the alpha 1-antitrypsin gene with milk production traits in Chinese Holstein. South Afr J Anim Sci 40: 113-120
Livestock Census, (2012) 19th All India Livestock Census. Department of Animal husbandry, Dairying and Fisheries. Ministry of agriculture. Government of India
Meyer K (2010) WOMBAT—A Tool for Mixed Model Analyses in Quantitative Genetics by Restricted Maximum Likelihood (REML). J Zhejiang Univ Sci 8(11): 815-821
Nivsarkar AE, Vij PK, Tantia MS (2000) Animal Genetic Resources of India: Cattle and Buffalo. ICAR publication, New Delhi
Reinhardt TA, Lippolis JD, Nonnecke BJ, Sacco RE (2012) Bovine milk exosome proteome. J Proteomics 75(5): 1486-92
Sambrook J, Russell DW (2001) Molecular cloning. A laboratory manual 3rd Edn. Cold Pring Harbor Laboratory Press, New York