Association of *IGF-I* gene polymorphisms with milk yield and body size in Chinese dairy goats

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Abstract

The association of *IGF-I* gene polymorphisms with certain traits in 708 individuals of two Chinese dairy-goat breeds (Guanzhong and Xinong Saanen) was investigated. Polymerase chain reaction-single strand conformation polymorphism (PCR-SSCP) and DNA sequencing methods were employed in screening for genetic variation. Two novel mutations were detected in the 5'-flanking region and in intron 4 of *IGF-I* gene, viz., g.1617G > A and g.5752 G > C (accession D26119.2), respectively. The associations of the g.1617G > A mutation with milk yield and the body size were not significant (p > 0.05). However, in the case of g.5752 G > C, Xinong Saanen dairy goats with the CG genotype presented longer bodies (p < 0.05). Chest circumference (p < 0.05) was larger in Guanzhong goats with the GG genotype. In Xinong Saanen dairy goats with the CC genotype, milk yields were significantly higher during the first and second lactations (p < 0.05). Hence, the g.5752G > C mutation could facilitate association analysis and serve as a genetic marker for Chinese dairy-goat breeding and genetics.

Key words: *IGF-I* gene, dairy goat, milk yield, PCR–SSCP.

Received: August 19, 2009; Accepted: December 9, 2009.

Animal growth is controlled by a complex system, in which the somatotropic axis plays a key role. Genes that operate in this axis are responsible for postnatal growth, mainly by way of growth hormones acting on bones and muscles mediated by insulin-like growth factor I (*IGF-I*) (Sellier, 2000). Thus, *IGF-I* is an important growth factor involved in a variety of physiological processes including cell differentiation, embryogenesis, and regulation of metabolism, reproduction, fetal development, and growth (Adam et al., 2000; Shen et al., 2003). The *IGF-I* gene was mapped on chromosome 5 in bovine (Miller et al., 1991). The *IGF-I* gene is expressed in most tissues, but mainly in liver (Wang et al., 2003). It encodes a polypeptide of molecular weight of 7.5 kDa with 70 amino acids (Daughaday and Rotwein, 1989). The amino acid sequence of *IGF-I* is identical in human, cattle, dog, and pig (Nixon et al., 1999).

In the nineties, several lines of evidence revealed an association of the high level of *IGF-I* in the blood with fast growth in cattle (Barash et al., 1998; Sirotin et al., 2000). In the following decades, cattle *IGF-I* gene mutations (C/T mutation in the 5'-flanking region) were found to be associated with the percentage of fat and protein in milk (Ge et al., 2001; Eulalia et al., 2006). However, these results were quite different from Hines’ discovery whence no association was found between C/T mutation in 5'-flanking region and dairy production traits in Holstein cattle (Hines et al., 1998). The association of *IGF-I* gene polymorphism (short random repeat in the 5'-flanking region) with body weight, both at birth and weaning, has also been described (Li et al., 2004; Moody et al., 1996), although such an association was not found by Curi et al. (2005). Recently, an *IGF-I* intron 2 polymorphism was found to correlate positively with the twinning rate (Kim et al., 2009).

Dairy goats are numerous in northwest China, among which, the Xinong Saanen (SN) and Guanzhong (GZ) breeds are the best-known, due not only to their tolerance of crushed feed and the local harsh weather, but also to their high milk yield. As it is difficult to culture high quality domestic breeds over a short period by the traditional breeding and genetics methods, the major breeders recently focused on using DNA markers for developing breeds through marker-assisted selection (MAS). Thus, it is important to identify significant associations between polymorphisms within crucial candidate genes and production traits. The aim was to find *IGF-I* gene SNPs in two Chinese dairy goat breeds and search for the possible associations of these with milk yield and body size.
708 blood samples were collected from Shaanxi province (440 for GZ breed from Sanyuan Guanzhong dairy goat breeding base in Shaanxi province; 202 for SN breed from Qianyang Saanen dairy goat breeding base in Shaanxi province; 66 for SN breed, also from animal farm in Northwest A&F University in Shaanxi province). All goats were 2 to 3 years old adults. Measurements were taken for all the individuals (BH = body height; BL = body length; ChC = chest circumference). Based on a 305-day lactation period, the first and second lactation data from each of 202 SN dairy goats from the Qianyang Saanen dairy goat breeding base, were recorded for every goat.

Blood samples were drawn from the jugular vein into curettes containing ACD (anticoagulant citrate dextrose). The samples were stored at -80 °C for several weeks, whereupon genomic DNA was extracted with the use of phenol/chloroform (Sambrock et al., 2001).

Based on the goat IGF-1 gene sequence published in the GenBank (D26119.2), five pairs of primers were designed. Primer sequences, annealing temperatures and amplicons are listed in Table 1. The P5 primers for the 5'-flanking end in Table 1 referred to those used in cattle (Ge et al., 2001), whereas the remainder were designed in this study.

The final volume of 25 μL contained 50 ng of genomic DNA, 0.5 μL of each primer (10 pM), 2.5 μL of a 10xbuffer (including 20 mM Mg²⁺), 0.5 μL of dNTPs (10 mM) and 0.25 μL of Taq DNA polymerase (5 U/μL Dongsheng, Guangzhou, China). Amplification was carried out using PCR, under the following conditions: 5 min at 96 °C for initial DNA denaturation, 35 cycles of denaturation at 94 °C for 40 s; annealing at X °C (shown in Table 1) for 40 s for each pair of primer; extension at 72 °C for 40 s. The final cycle was followed by a 10 min extension at 72 °C.

Aliquots of 4 μL of PCR products were mixed with 8 μL of denaturing solution (95% formamide, 25 mM EDTA, 0.025% xylene-cyanole and 0.025% bromophenol blue) and heated for 10 min at 98 °C and chilled on ice. Denatured DNA was subjected to 10% PAGE (80 mm x 73 mm x 0.75 mm) in 1x TBE buffer and constant voltage 250V for 15 min, then 160V for 2 h. The gel was stained with 0.1% silver nitrate (Lan et al., 2007). After the polymorphisms were detected, the PCR products of different electrophoresis patterns were sequenced by the Huada Genetic Sequence Company (Beijing, China). The sequences were analyzed with DNASTAR software (Version 7.10, DNASTAR, Madison, USA).

Based on the individual number of different genotypes in analyzed breeds, Hardy-Weinberg equilibriums, genotypic frequencies and allelic frequencies were directly calculated. The statistical software SPSS (Version 16.0) was used to analyze the relationships between the genotypes and traits in goats. The adjusted Linear Model I with fixed effects was used to evaluate the relationships between genotypes and milk yield of Xinong Saanen dairy goats. The model applied was: $Y_{ijk} = \mu + A_i + G_j + (AG)_{ij} + e_{ijk}$, where $Y_{ijk}$ was the trait measured in each $ij$th animal, $\mu$ the overall population mean, $A_i$ the fixed effect due to the $ith$ age, $G_j$ the fixed effect associated with the $j$th genotype, $(AG)_{ij}$ the interaction between the $ith$ age and the $j$th genotype, and $e_{ijk}$ the random error. Effects associated with farm, sex and season of birth (spring versus fall), and the ages of dams and sires, were not considered in the linear model, since preliminary statistical analysis indicated that these effects did not have significant influence on the variability of traits in SN female dairy goats. The adjusted Linear Model II with fixed effects was used to analyze the relationships between genotypes and body size in these two breeds. The model applied was: $Y_{ijk} = \mu + B_i + G_j + (BG)_{ij} + e_{ijk}$, where $Y_{ijk}$ was the trait measured in each $ij$th animal, $\mu$ the overall population mean, $B_i$ the fixed effect due to the $ith$ breed, $G_j$ the fixed effect associated with $j$th genotype, $(BG)_{ij}$ the interaction between the $ith$ breed and the $j$th genotype, and finally $e_{ijk}$ the random error. Effects associated with farm, sex and season of birth (spring versus fall), as well as the ages of both dams and sires, were not used in the linear model, as the preliminary statistical analyses indi-

| Gene loci | Primer sequences | Annealing temperatures (X °C) | Locations a |
|----------|----------------|-----------------------------|-------------|
| P1       | F: 5'-ATTACAAAGCTGCCTGCCCC-3' | 60.3 | 1407-1655 (249 bp) |
|          | R:5'-ACCTTACCCTGTAGAAGAAATACGT-3' | 54.7 | 1942-2231 (290 bp) |
| P2       | F: 5'-TTCTTCTAAATCCCTTCCTGTGGT-3' | 65.7 | 4728-4956 (229 bp) |
|          | R: 5'-CCATGTGTTAGCAGCCAGTG-3' | 60.3 | 5471-5790 (320 bp) |
| P3       | F: 5'-ACCGAGAGAGAGAGCAGG-3' | 54.7 | 1942-2231 (290 bp) |
|          | R: 5'-ATCCACAGAGCAGTG-3' | 65.7 | 4728-4956 (229 bp) |
| P4       | F: 5'-GCTGGGTGTAGCAGGCAACA-3' | 55.8 | 6030-6580 (550 bp) |
|          | R: 5'-GTGGGGTCAAGGACTCAGTG-3' | 60.3 | 5471-5790 (320 bp) |

aBased on GenBank accession: D26119.2.
cated that these effects did not have significant influences on variability of traits in this two breeds. All the goats were adults, so the age did not influence on the body size. The least square means (LSMs), with standard errors and multiple range tests for IGF-I genotypes and traits, were calculated.

Using PCR-SSCP method with five pairs of primers of the goat IGF-I gene, only two (P1 and P4) of them demonstrated polymorphic patterns in both breeds. In the P1 locus (fragment amplified by the P1 pair of primers), there were only two kinds of SSCP polymorphic patterns detected in all the 708 samples. We named them GG, with three bands, and AG, with five bands (Figure 1a). In the P4 locus (fragment amplified by the P4 pair of primers), three SSCP polymorphic patterns were found in the 708 samples, which were named GG pattern, with four bands, CG pattern, with five bands, and CC pattern also with four bands (Figure 1b).

In order to clarify the nucleotide mutations in these two loci, nine PCR products, each with a different pattern, were sequenced at each locus. The three remaining products (P2, P3, and P5) were also sequenced. On comparing with GenBank sequence D26119.2, 249 bp PCR products, including the 5’-flanking region upstream from the start codon (ATG), were amplified by the P1 pair of primers. There was a G to A mutation, namely g.1617G>A (accession D26119.2). The P4 primers were used to amplify the whole of exon 4 and part of introns 3 and 4. There was a G to C variation in both breeds, viz., g.5752 G > C (accession D26119.2), located in intron 4. No mutation was detected in the remaining loci (P2, P3, and P5).

The allelic and genotypic frequencies of each locus obtained for the different genetic groups, can be seen in Table 2. For the g.1617 G > A mutation (accession D26119.2), frequencies of the IGF-I-A allele in the analyzed populations were 0.08 and 0.11 for GZ and SN dairy goats, respectively, with genotype distribution of both breeds in agreement with Hardy-Weinberg equilibrium (p > 0.05). For the g.5752 G > C (accession D26119.2), significant statistical results were encountered in milk yield and BL (body length) of individuals with the CC and CG genotypes (p < 0.05 and p < 0.05, respectively) in Saanen dairy goats. The BL of SN breed with CG genotype was significantly longer than in the others. Individuals with genotype CC of the IGF-I gene had a superior milk yield at the first and second location when compared to those of individuals with genotype CG and GG (p < 0.05). In the Guanzhong population, chest circumference (ChC) of GZ goats with the GG genotype was the largest of the three genotypes, whereas ChC for individuals with CG genotype was larger than that for CC carriers (p < 0.05).

When focusing on the IGF-I gene, an attempt was made to detect nucleotide mutations in primer amplified

| Gene loci | Breeds | Genotypes | Genotypic frequencies | Allelic frequencies | Hardy-Weinberg equilibrium ($\chi^2$ test) |
|-----------|--------|------------|-----------------------|---------------------|------------------------------------------|
|           |        | GG | AG | AA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N | PGG | PAG | PAA | N |
| P1        | GZ     | 373 | 67 | 0  | 440 | 0.85 | 0.15 | 0   | 0.08 | 0.92 | 2.98 |
|           | SN     | 211 | 57 | 0  | 268 | 0.79 | 0.21 | 0   | 0.11 | 0.89 | 3.79 |
|           |        |     |    |    |    |     |     |     |     |     |     |
|           |        |     |    |    |    |     |     |     |     |     |     |
| P4        | GZ     | 47  | 69 | 324 | 440 | 0.10 | 0.16 | 0.74 | 0.19 | 0.81 | 101.56 (p < 0.05) |
|           | SN     | 22  | 116| 130 | 268 | 0.08 | 0.43 | 0.49 | 0.30 | 0.70 | 0.30 |

$\chi^2_{0.05}$ (df = 1) = 3.84, $\chi^2_{0.01}$ (df = 1) = 6.63.
fragments, and to analyze their associations with body size and milk yield in Chinese dairy goats. The analyzed length of the IGF-I gene was 1588 bp, including part of the 5’-flanking, and almost the entire coding region except for exon 2 which is only 15 bp long. In the g.1617G>A mutation (accession D26119.2), the lack of AA homozygotes indicated that the mutation in the goat IGF-I gene might give rise to diminished production in the population under analysis. These two dairy goat breeds are major milk producers, originating from a varied dairy-goat breeding basis in Shanxi province (P.R. China). As the two populations were over one year old, individuals with low production had possibly already been eliminated. Hence, it is reasonable to surmise that AA homozygotes is almost completely lost in quite large local populations. Obviously, the reason of the lack of genotype AA in these breeding farms requires further investigation.

Significant statistical results were found as regards body size and milk yield among different genotypes in g.5752 G > C (accession D26119.2) (p < 0.05) (Table 3). Being located in one of the intron regions, the SNP studied here may not be a causal mutation. Thus, we suggest that it could be linked to another mutation in the coding or regulatory regions of a gene which is a causal mutation for production traits. Introns have been shown to affect the transcriptional efficiency of numerous genes in a variety of organisms (Greenwood and Kelsoe, 2003; LeHir et al., 2003). Further analysis is required to validate both the association found in this study and the physiological significance of the intron 4 mutation in IGF-I gene.

In conclusion, our study revealed the first known significant associations of IGF-I gene polymorphisms with milk yield and body size in dairy goats. The CC genotype at the g.5752 G > C (accession D26119.2) locus could be used as a molecular marker for superior milk yield. Furthermore, this study will be of practical use for the improvement of Chinese native dairy goats and the local breeding of those with higher milk production.

Acknowledgments

This study was supported by the National 863 Program of China (No. 2008AA10Z138), the Young Topnotch Researcher Support Project of Northwest A&F University (No.QNGG-2009-007), the “13115” Sci-Tech Innovation Program of Shaanxi Province (No. 2008ZDKG-11), the Research Fund for the Doctor Program of Higher Education of China (No.20080712001), and the Natural Science Foundation of Jiangsu Province (No. BK2008120).

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