The Characteristic of Tuvan Short-Fat Tailed Sheep 
(*Ovis aries*) Populations by *GDF9* Gene Polymorphism

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Abstract: Tuvan short-fat tailed sheep is a local breed spread in the Russian Federation. This breed inhabits the Tuva Republic and consists of steppe and mountain breed types. The aim of investigation is the identification of the genetic polymorphism of the growth differentiation factor-9 (*GDF9*) gene in Tuvan local sheep populations. It is a potential genetic marker of prolificacy and the feature of sheep populations by this gene polymorphism can be useful for sheep breeding and breed conservation. Genomic DNA was isolated from samples of blood of 144 sheep of steppe type and 106 sheep of mountain type. Two primers were used to obtain a 462 bps DNA fragment. The polymorphism of *GDF9* gene (CC, CD and DD genotypes) were detected after amplicons were digested with AspLEI restriction enzyme. In mountain type population CC and CD genotypes of *GDF9* gene were identified with 0.886 and 0.114 frequencies and DD genotype was not observed. C and D allele frequencies were 0.94 and 0.06, respectively. The steppe type sheep population included CC, CD and DD genotypes with 0.833, 0.160 and 0.007 frequencies. C and D allele frequencies were 0.91 and 0.09, respectively.

Key words: sheep, genetic polymorphism, PCR-RFLP, *GDF9*, allele frequencies

I. INTRODUCTION
Sheep breeding is one of the main branches of animal husbandry in the Russian Federation and a traditional branch of activity among many peoples of the country. Sheep are raised everywhere. The variety of domestic breed composition is the foundation for creating new breeds, breed types and other pedigree groups of animals with high productivity potential and good fitness to local technological, economic and environmental conditionals. So saving and rational use of all sheep breeds of the country are an important task.

The Tuva Republic is a one of the most important region of the Russian Federation characterized by sheep breeding. That region is situated in southern Siberia, in the geographical center of Asia. Mountains occupy over 80% of the regional territory, however mountain pastures and intermountain basins, characterized by steppe landscape, are the excellent basis for sheep breeding.

The regional climate is sharply continental. The major local sheep breed in that region is Tuvan short-fat tailed breed (Tuvan sheep breed), well adapted to specific regional climate environment. Animals of that breed have a high immunity and stamina (Yuldashbaev et al., 2016).

Tuvan sheep breed contains two breed types. These are steppe and mountain types characterized by a different level of economic
traits. The minimal live weight of adult rams of the steppe type is 78 kg and that of ewes is 56 kg. The natural wool is strong, 12-14 cm long. Adult rams of the mountain type are characterized by 55 kg of live weight and ewes weigh 42 kg. The wool length is 10-12 cm.

The scouring yield is 50-60%. The birth rate of Tuvan sheep of all types is 100-110 lambs per 100 ewes (Yuldashbaev et al., 2016).

One of the ways to intensify sheep breeding is to use genetic markers of productive traits to organize marker selection. Marker selection (MAS) is the use of DNA markers to improve the selection response in an animal population. Markers should be closely related to one or more target loci, which for the most part can be quantitative trait loci.

The discovery of polymorphic variants of various genes is a key moment in the start of breeding programs or work to conserve sheep breeds. Polymorphic variants of genes can be associated with different levels of productive traits.

One of the possible (potential) genetic markers of fecundity is the growth differentiation factor gene (GDF9) located on the fifth chromosome of sheep. This gene contains two exons and one intron (Bodensteiner et al., 1999; Sadighi et al., 2002; Hanrahan et al., 2004).

The aim of this study is to detect polymorphic variants of the GDF9 gene to assess the possibility of its use in breeding programs and work on the conservation of the Tuvan short-fat tailed breed of sheep.

II. MATERIALS AND METHODS

Polymorphism of the studied gene was detected using the PCR-RFLP method. Blood samples for DNA analysis were taken from 250 sheep of Tuvan short-fat tailed breed of sheep belonging to two different intra-breed types: 106 samples of the mountain intra-breed type originated from State Unitary Enterprise “Malchyn” and 144 samples of the steppe intra-breed type originated from Municipal Unitary Enterprise "Despen". About 9 ml of blood per sample was collected in sterile tubes. Good preservation of blood samples was achieved with the help of K3-EDTA-sprayed tubes and sample freezing at -20° C.

Genomic DNA was isolated using commercial kits in accordance with the manufacturer's instructions.

To obtain amplicons of the GDF9 gene, the following pair of primers was used:

GDF9-F: 5’-GAAGACTGGTATGGGGAAATG-3’;
GDF9-R: 5’-CCAATCTGCTCCTACACACC-T-3’.

The amplification reaction was carried out under the following conditions. 35 cycles: 94° C for 2 min., then 94° C for 30 s., 63° C for 40 s., 72° C for 30 s. and final elongation at 72° C for 4 min (Bahrami et al., 2014; Kolosov et al., 2015; Gorlov et al., 2018). Fragments of GDF9 gene obtained in PCR were digested with AspLEI restriction endonuclease at 37° C for 12 hours. All initial amplicons and digested PCR products of GDF9 gene fragments were separated on a 2.0-3.0% agarose gel and visualized after staining with ethidium bromide in a gel documentation system.

The data generated by electrophoresis of AspLEI digested samples were used for estimating the frequency of different restriction fragment patterns.

The genotypes and allelic frequency were estimated by standard procedure. Genotypes frequency was calculated according to following formula:

$$P_i = \frac{n_i}{N}$$

where: $P_i$ is the $i^{th}$ genotype frequency; $n_i$ is a number of samples of the $i^{th}$ genotype; $N$ is a total number of samples of all genotypes.

Allelic frequency was calculated in the following way:

$$p_i = \frac{2n_{(homozygote)} + n_{(heterozygote)}}{2N}$$

where: $p_i$ is the $i^{th}$ allele frequency; $n$ is a number of homozygotes of particular gene and heterozygotes, respectively; $N$ is a total number of individuals.

III. RESULTS

Amplified products of 462 bps fragments of GDF9 gene were obtained after amplification in the analyzed samples (Figure1).
C and D alleles of DGF9 gene were observed after digesting of PCR products by AspLEI restriction enzyme. Three genotypes presented by DNA-fragments of different size were established (Figure 2). The AspLEI digestion of amplified loci of GDF9 gene produced fragments of 254, 156 and 52 bps for allele C. Allele D was identified as a 410 and 52 bps patterns. Homozygous genotype CC was characterized by 254, 156 and 52 bps bands. DD genotype contained 410 bps and 52 bps fragments. Heterozygous genotype CD consisted of 4 bands: 410, 254, 156 and 52 bps (Bahrami et.al., 2014; Kolosov et.al., 2015).

Fragments sized 52 bps were low-observable (Figure 2).
The largest part of investigated populations of sheep was homozygous of C alleles of GDF9 gene (Table 1). According to results of the investigation the largest part of animals in mountain type population was characterized by CC genotype and its frequency was 0.886. The frequency of CC genotype in steppe type population was 0.833. Heterozygous genotype CD was not so widespread in populations and had a frequency of occurrence equal to 0.114 and 0.160 in animals of mountain and steppe intra-breeds, respectively. DD genotype was observed in steppe type sheep population only. Its frequency was the smallest in the result of the investigation (0.007).

Table 1. The genotypes frequency for GDF9 genes in Tuvan short-fat tailed sheep breed in SUE "Malchyn" (mountain type, n=106) and MUE "Despen" (steppe type, n=144)

<table>
<thead>
<tr>
<th>Types of Tuvan sheep breed</th>
<th>Genotypes</th>
<th>The number of animals</th>
<th>Genotypes frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountain type</td>
<td>CC</td>
<td>94</td>
<td>0.886</td>
</tr>
<tr>
<td></td>
<td>CD</td>
<td>12</td>
<td>0.114</td>
</tr>
<tr>
<td></td>
<td>DD</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Steppe type</td>
<td>CC</td>
<td>120</td>
<td>0.833</td>
</tr>
<tr>
<td></td>
<td>CD</td>
<td>23</td>
<td>0.160</td>
</tr>
<tr>
<td></td>
<td>DD</td>
<td>1</td>
<td>0.007</td>
</tr>
</tbody>
</table>

Table 2. The alleles frequency for GDF9 genes in Tuvan short-fat tailed sheep breed in SUE "Malchyn" (mountain type, n=106) and MUE "Despen" (steppe type, n=144)

<table>
<thead>
<tr>
<th>Types of Tuvan sheep breed</th>
<th>Alleles</th>
<th>Allele frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mountain type</td>
<td>C</td>
<td>0.94</td>
</tr>
<tr>
<td></td>
<td>D</td>
<td>0.06</td>
</tr>
<tr>
<td>Steppe type</td>
<td>C</td>
<td>0.91</td>
</tr>
<tr>
<td></td>
<td>D</td>
<td>0.09</td>
</tr>
</tbody>
</table>

C allele of GDF9 gene had the highest frequency in sheep of all bred types of Tuvan sheep (Table 2). The frequency of C allele in sheep population of steppe and mountain types was 0.91 and 0.94, respectively. This mark for D allele of GDF9 gene was 0.06 in mountain type sheep and 0.09 in steppe inter-breed type.

IV. DISCUSSION
The distribution of identified genotypes in Tuvan short-fat tailed sheep breed populations are in agreement with the polymorphism detected in ovine GDF9 gene in other local sheep breeds. One of the previous experiments wild genotype (CC genotype in our investigation) in Hisari sheep was presented with a frequency of 93.64%. The frequency of heterozygous genotype (CD in our investigation) was 6.36% (Bahrami et al., 2014).

The analysis of polymorphism for GDF9 gene in Baluchi sheep indicated all three possible genotypes: FecG+/FecG+ (CC genotype in our investigation) was 0.72, FecG+/FecG1 (CD in our investigation) was 0.20, FecG1/FecG1 (DD genotype in our investigation) was 0.08 (Moradband et al., 2011).

The genotypes frequencies for GDF9 gene in Salsk and Volgograd sheep breeds had the similar character. The frequency of CC and CD genotype in Salsk breed was 88% and 12% respectively and those in Volgograd breed were 84% and 16%. Homozygous DD genotypes were not observed in the studied populations (Gorlov et al., 2018).

In the other investigation Salsk sheep breed had a high frequency of CC genotype equal to 90%. The frequency of CD genotype was 10% (Kolosov et al., 2015).

The frequencies of alleles and genotypes of GDF9 gene showed a higher level of polymorphism in the Romanov sheep population. CC genotype was detected (60.9%), DD genotype was not observed and the frequency of CD genotype was 39.1% (Kolosov et al., 2015).
In this way, the observed character of GDF9 genotypes distribution was typical in many sheep breeds population.

V. CONCLUSION
The characteristic of Tuvan short-fat tailed sheep populations by GDF9 gene is one of the steps in the implementation of the candidate genes approach in sheep breeding of the Tuva Republic and can be the basis of complex projects for conservation of local sheep breeds.

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