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#### Short communication

## The analysis of mitochondrial data indicates the existence of population substructure in Karayaka sheep

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#### ABSTRACT

In the current study, we investigated the existence of population substructure in the Karayaka sheep breed from Turkey. A fragment of the mitochondrial D-loop region has been successfully sequenced in 69 Karayaka sheep sampled in the districts of Çarsamba, Bafra and Ladik. Though the B haplogroup was the most abundant in these three subpopulations, haplogroup frequencies and nucleotide diversities were remarkably different. The analysis of molecular variance (AMOVA) showed the existence of a significant (P-value = 0.019) between-subpopulations component representing 5.14% of the total variation. Moreover, phylogenetic analyses showed that the fixation index ( $F_{\rm ST}$ ) and the Gammast genetic distance between Karayaka subpopulations are similar to those observed between certain Turkish sheep breeds. We conclude that there is a significant population substructure in Karayaka sheep. Since breeds do not behave as single panmictic populations, this result could probably be extrapolated to other ovine breeds. Noteworthy, population substructure can have adverse effects on the maintenance of breed diversity and it is an important confounder effect in genome-wide-association studies.

#### 1. Introduction

Karayaka is a sheep breed from Northern Anatolia that is raised for the production of meat, milk and wool (Yilmaz et al., 2013). In general, Karayaka sheep display a white coat, although the legs and head can be black. Moreover, black and brown coated individuals can also be found (Yilmaz et al., 2013). Karayaka can be classified as a long-thin tailed breed, and males usually have spiral horns, while females are polled. The milk production of Karayaka sheep is quite low (121 kg milk in a lactation of 204 days), and this is one of the main reasons why it is not used in large-scale dairy farms, but this flaw is compensated to some extent by heavy fleece production (Yilmaz et al., 2013).

In the 1980–2009 period, the population of ovine local breeds of Turkey steadily decreased from 48 million to 21.6 million heads, due to competition with foreign breeds, lack of government support, the progressive abandonment of rural activities and market changes (Yilmaz et al., 2013). The Karayaka breed has also suffered from population decline. According to the DAD-IS database (http://dad.fao.org), the census of the Karayaka breed has decreased from 1.7 million animals in

1983 to approximately half a million in 2012. Although the accuracy of these numbers cannot be fully confirmed, they clearly indicate a diminishing demographic trend.

Five main mitochondrial haplogroups (A-E) have been defined in sheep (Hiendleder et al., 1998; Tapio et al., 2006; Meadows et al., 2007). All of them segregate at variable frequencies in Turkish sheep, though D and E are relatively rare (Demirci et al., 2013). The genetic diversity of the Karayaka breed has been reported in several studies. Uzun et al. (2006) analyzed the autosomal diversity of several Turkish breeds with a panel of 30 microsatellites and found that fat-tailed sheep showed a significant genetic differentiation when compared to other ovine populations. Demirci et al. (2013) characterized the mitochondrial variation of 13 Turkish native breeds and found that in Karayaka sheep, the most frequent haplogroup was B (74%), followed by A (18%), while C (6%) and E (2%) had low frequencies. These studies provided a valuable picture of the diversity of Karayaka sheep and their relationship with other ovine breeds from Turkey, but the existence of population substructure has not been investigated yet. This is an important issue because the existence of population subdivision can

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augment homozygosity (Garnier-Géré and Chikhi, 2013), thus intensifying the effects of genetic drift and inbreeding. The goal of the current work was to analyze the mitochondrial variation of Karayaka sheep from the Ladik, Çarsamba and Bafra districts with the aim of determining wether these three subpopulations are genetically differentiated. We also wanted to determine whether the sustained demographic decline of the Karayaka and other Turkish breeds has left a detectable genetic signature in their mitochondrial gene pools.

#### 2. Materials and methods

#### 2.1. Total DNA isolation

Blood samples were obtained from the jugular vein under veterinary supervision and collected in 10 mL tubes containing lithium heparin. Blood extraction procedures were carried out in accordance with the ARRIVE guidelines published by Kilkenny et al. (2010). At the time of sampling, identification numbers and morphological characteristics were recorded in each sheep as well as the owner's name. To make sure that sampled animals were unrelated, genealogical records gathered by the Sheep and Goat Breeder's Association of Samsun (Turkey) were checked . In total, blood samples from 100 sheep raised in 19 farms located in the districts of Çarşamba (8 farms), Bafra (8 farms) and Ladik (3 farms) were collected. Blood samples were transported in containers filled with dry ice to the Animal Biotechnology Laboratory at Ondukuz Mayıs University (Samsun, Turkey) and stored at −20 °C. Total DNA was isolated with the IDPURE Spin Column Genomic DNA MiniPrep Kit (Empire Genomics, Buffalo, NY) by following the instructions of the manufacturer.

#### 2.2. Amplification and sequencing of the D-loop region

Two primers were designed to amplify a fragment of 575 bp (positions 221-795 of GenBank sequence AM279285.1) of the mitochondrial D-loop region: FW 5'-GCC CAC ATA ACA ACC CAT AC-'3 and REV 5'-CGG AGC GAG AAG AGG GAT-3'. Polymerase chain reactions were performed in a final volume of  $15\,\mu\text{L}$  containing  $1.5\,\mu\text{L}$  of  $10\times$  PCR buffer,  $1.5\,\mu L$  of MgCl<sub>2</sub>(25 mM),  $0.45\,\mu L$  of each primer (final concentration =  $0.3 \,\mu\text{M}$ ),  $0.15\,\mu L$  of each concentration = 0.25 mM), 0.75 U of AmpliTaq DNA polymerase (Applied Biosystem, Foster City, CA) and 60-100 ng of genomic DNA. Ultrapure water was added to the reaction until reaching a final volume of 15 µL. This reaction mixture was heated to 95 °C for 10 min, followed by 35 cycles of 95 °C for 1 min, 64 °C for 1 min and 72 °C for 1 min. Subsequently, a final extension step at 72 °C for 7 min was carried out. Amplification products were purified with the ExoSAP-IT PCR Cleanup kit (Affymetrix, Santa Clara, CA). Sequencing reactions were prepared with the Big Dye Terminator Cycle Sequencing Kit v1.1 (Applied Biosystems, Foster City, CA) and electrophoresed in an ABI 3730 DNA Analyzer (Applied Biosystems, Foster City, CA). High-quality sequences from 69 Karayaka individuals were obtained (GenBank accession numbers: MG209399-MG209467).

#### 2.3. Data analyses

Mitochondrial D-loop sequences were aligned with the Multalin software (Corpet, 1988) and trimmed to a final size of 511 bp. The number of haplotypes and nucleotide and haplotype diversities were calculated with DnaSP v5.1 software (Librado and Rozas, 2009). The  $F_{ST}$  coefficients among breeds and the tests of neutrality; *i.e.*, Fu and Li D\* and F\* (Fu and Li, 1993) and Tajima's D (Tajima, 1989) were also carried out with DnaSP v5.1 (Librado and Rozas, 2009). A phylogenetic neighbor-joining tree depicting the genetic relationships among Karayaka sheep sampled at the Çarsamba, Ladik and Bafra districts was built with the MEGA 5 software (Tamura et al., 2011) under a Tamura-Nei model (Tamura and Nei, 1993) and 1000 bootstrap replicates.

Gamma $_{\rm ST}$  and  $F_{\rm ST}$  distances among three Karayaka subpopulations (Bafra, Çarsamba and Ladik) and 12 Turkish breeds (Akkaraman, Çineçaparı, Morkaraman, Gökçeada, Sakız, İvesi, Dağlıç, Herik, Hemşin, Karagül, Norduz and Kıvırcık) were estimated using DnaSP v5.1 software (Librado and Rozas, 2009) and 142 mitochondrial sequences retrieved from GenBank (Supplementary Table 1) plus 36 new sequences selected at random (Bafra = 13, Çarsamba = 11, Ladik = 12) from the data set generated in the current work. Subsequently, two neighborjoining trees (Gamma $_{\rm ST}$  and  $F_{\rm ST}$  trees) were built with the MEGA 5 software (Tamura et al., 2011) by considering a 462 bp fragment shared by all sequences. An analysis of molecular variance (AMOVA) based on mitochondrial sequences from three Karayaka subpopulations was carried out with the Arlequin 3.5 software by considering the default parameters (Excoffier and Lischer, 2010).

#### 3. Results and discussion

The genetic diversity of the Karayaka sheep and their genetic relationship with other ovine breeds have been reported and discussed in previous publications (Uzun et al., 2006; Yilmaz et al., 2014; Das et al., 2015). Herewith, we analyzed the mitochondrial variation of Karayaka sheep from the districts of Çarsamba, Ladik and Bafra with the aim of investigating the existence of population substructure. The data presented in Fig. 1 and Table 1 demonstrate that the B haplogroup is the most abundant, followed by A, but haplogroup frequencies were quite different among the subpopulations. Moreover, the neighbor-joining tree based on Gamma<sub>ST</sub> and F<sub>ST</sub> distances showed that genetic distances among these three subpopulations have magnitudes similar to those observed between distinct Turkish breeds (Fig. 2). We have also performed several tests of neutrality in order to assess whether the population reduction suffered by Karayaka sheep in the last 30 years has left a detectable genetic signature at the mitochondrial level. The Fu and Li's D\* and F\* test statistics (D\* = -1.29, F\* = -1.28) and Tajima's D (D = -0.74) were all non-significant at the whole population scale, and we also observed a lack of significance when we calculated these statistics for each subpopulation (data not shown). These results indicated that Karayaka sheep have not suffered a strong genetic bottleneck but a progressive and sustained population decline.

To assess more robustly the existence of population substructure in the Karayaka sheep, an analysis of molecular variance (AMOVA) was carried out (Table 2. Our data showed that the between-populations component of variation was small (5.14%) but significant (Pvalue = 0.019). Based on these results, we can conclude that there is a certain level of population substructure in the Karayaka sheep. In a genome-wide analysis of Sicilian sheep breeds, Mastrangelo et al. (2014) also detected the existence of population substructure in the Comisana breed, probably due to a differential introgression with foreign breeds and/or to geographic isolation. Population substructure was also detected when analyzing the diversity of the Portuguese cattle breed "Brava de Lide" (Mateus and Russo-Almeida, 2014). From a reproductive point of view, breeds do not behave as single panmictic populations. In contrast, there is a certain level of reproductive isolation among farms and regions due to the lack of national breeding programs facilitating the widespread diffusion of the genetic progress by artificial insemination and embryo transfer. The differential admixture of Karayaka subpopulations with highly improved exotic breeds, adaptation to specific environmental or breeding conditions, sampling constraints and the existence of geographic barriers might also explain the weak but significant genetic differentiation that we observed among the Çarsamba, Ladik and Bafra subpopulations.

The existence of population subdivision can have important consequences on the maintenance of genetic diversity by preventing random mating and decreasing heterozygosity; *i.e.*, the so-called Wahlund effect (Garnier-Géré and Chikhi, 2013). From a practical point of view, population stratification can be an important confounder in genetic association analyses (Salmela et al., 2011). While haplotype

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