Novel Single Nucleotide Polymorphisms of Lysozyme (C-Type) Gene in Donkey (Equus Asinus) Populations in Marmara Province of Turkey

Raziye Işık¹,a,*

¹Department of Agricultural Biotechnology, Faculty of Agriculture, Tekirdağ Namık Kemal University, 59030 Tekirdağ, Turkey
*aCorresponding author

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ABSTRACT

The major antimicrobial proteins in donkey milk are lysozyme, lactoferrin, lactoperoxidase and immunoglobulins. Lysozyme has an important role in the host defense by way it inhibits the pathogenic bacteria. The aim of this study is to investigate the Lysozyme (LYZ) gene polymorphism in 82 donkeys reared in Thrace region of Turkey. 716 bp long partial 5' UTR, exon 1, intron 1, exon 2 regions of LYZ gene were amplified and PCR products were analyzed via DNA sequencing. Three novel single nucleotide polymorphisms (SNPs) were identified as g.1782775A>G, g.1782924A>G and g.1782960T>C in the first intron of LYZ gene. The partial DNA sequence of LYZ gene in donkeys was reported in the present study and sequences of LYZ were entered to NCBI Genbank database with the accession number: MK984689-MK984692. This SNP may have an effect on immune system and milk traits in donkeys and additional studies are needed to confirm this assumption for donkey breeding.

Keywords: LYZ Gene, Equus asinus, Donkey, DNA sequencing, Turkey

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Türkiye Marmara Bölgesi Eşek (Equus Asinus) Popülasyonlarında Lizozim (C-Tipi) Genindeki Yeni Tek Nükleotid Polimorfizmeleri

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ÖZ

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Bu çalışma, eşek sütündeki başlıca antimikrobiyal proteinler lizozim, laktoferrin, laktoperoxidaz ve immunoglobulinerdir. Lizozim, konakçı savunmasında patojenik bakterileri inhibe etmede önemli bir rol sahiptir. Çalışmanın amacı, Trakya bölgesinde yetiştirilen 82 eşekte lizozim (LYZ) geni polimorfizmini araştırmaktır. LYZ geninin 716 bp uzunluğunda kısmi 5' UTR, ekzon 1, intron 1, ekzon 2 bölgeleri çoğaltılmış ve PCR ürünlerini DNA dizisi analizi yoluya analiz edilmiştir. LYZ geninin birinci intronunda g.1782775A>G, g.1782924A>G ve g.1782960T>C olmak üzere üç yeni tek nükleotid polimorfizmi (SNP) tanımlanmıştır. Bu çalışmadan eşeklerde LYZ geninin kısmi DNA dizisi belirlenmiş ve LYZ dizileri NCBI Genbank veritabanına MK984689-MK984692 erişim numarasıyla girilmiştir. Bu SNP'lerin eşeklerdeki bağılılık sistemi ve süt özelliklerini üzerine bir etkisi olabilecek ve eşek yetiştiriciliği için bu varsayımı doğrulamak için ek çalışmalara ihtiyaç duyulmaktadır.
Introduction

Donkey milk has a great nutritional value and many functional features such as antimicrobial, antioxidiant, antiviral, anti-inflammatory and hypoallergenic. These features are associated with lysozyme, immunoglobulins, lactoferrin, bioactive peptides, α-3 fatty acids of donkey milk (Tanaka, 2007; Tidona et al., 2011; Brumini et al., 2016; Stefaniuk-Szmukier et al., 2018). Thus, it has some differences from other ruminant milk such as fat ratio, minerals content, lactose and the presence of some antimicrobial proteins (Nayak et al., 2017). Donkey milk contains three main proteins; β-lactoglobulin (β-lg), α-lactalbumin (α-La) and lysozyme (Chiavari et al., 2005).

Lysozyme has an significant role in host defense by the way of hydrolyzing the polysaccharides via 1,4-beta-linkages between N-acetylmuramic acid and N-acetyl-D-glucosamine residues in peptidoglycan of bacterial cell walls. It inhibits the pathogenic bacteria and also it has a role as antimicrobial agent in milk (Palumbo et al., 2010).

The lysozyme gene family contains three different genes and proteins such as lysozyme c (LYZ), lactalbumin which is also present in mammals and calcium-binding lysozyme which is also present in some birds and mammals (Irwin et al., 2011). Three different kinds of lysozymes have been determined such as chicken type (c-type), invertebrate type (i-type) and goose type (g-type). Equine milk lysozyme and ruminant lysozymes are taken part of the c-type family because of the resemblance with chicken egg white lysozyme. Equine lysozyme belongs to the family of c-type lysozymes, but has some structural characteristics typical to α-lactalbumins (Cosenza et al., 2017).

Lysozyme has an important role in reducing the gastrointestinal infections in the infants along with immunoglobulins, lactoferrin and lactoperoxidase. While donkey milk contains around 1 g/L lysozyme c, bovine (0.13 mg/L) and human (0.4 mg/l) contain very low concentration of lysozyme c in milk (Stelwagen, 2003; Gubić et al., 2015). Donkey milk lysozyme molecular weight is approximately 14.9-15.4 kDa (Gubić et al., 2015). LYZ gene is very conservative among the species such as bovine and goat LYZ gene have %91 DNA sequence identity. LYZ gene is situated on chromosome 5 in goat-cattle, chromosome 6 in horse and on chromosome 3 in sheep. However, the location of LYZ gene is not clarified in donkeys. Also, it contains 4 exon counts in donkey, sheep, cattle and goat. The donkey lysozyme gene consists of 4 exons, 148 amino acids and 5193 bp long (NCBI, 2019).

Lysozyme c protein and gene sequences have been investigated from many vertebrate and non-vertebrate species (Irwin et al., 1996; Qasba and Kumar, 1997; Callewaert and Michiels 2010), but few data are available for donkeys (Cosenza et al., 2017; 2018). The aim of this study is to investigate the LYZ gene polymorphisms in donkeys via DNA sequencing method.

Materials and Methods

Experimental Animals And Tissue Collection

In the present study, a total of 82 blood samples were taken from Kırklareli (36), Tekirdağ (28) and İstanbul (Çatalca) (18) Provinces of Turkey. Blood sample were collected to 10 mL of vacuum tubes, including EDTA as anticoagulant and stored at −2°C till the DNA extraction. Genomic DNA was isolated by using a commercial DNA isolation kit (GeneJET Whole Blood Genomic DNA Purification Mini Kit, Thermo Fisher Scientific) according to the manufacturer’s instructions.

Primer Design, Total DNA Extraction and PCR

Primer sequence of LYZ gene was designed based on the donkey sequence retrieved from GenBank (Accession number NW_014638180.1) using Primer-BLAST algorithm. Primer sequences of LYZ gene are F: 5’-AGCTCTCTGCTGCAAGATGACA-3’, and R: 5’-GCTGAAGAGGACACTCCTTC-3’. For amplification reactions, the 25 µL PCR volume contained: 50 ng genomic DNA, 0.5 µM of each primer, 1× PCR Buffer ((NH₄)₂SO₄), 200 µM dNTP, 2.0 mM MgCl₂ and 0.5 U of TaqDNA polymerase (Taq DNA Polymerase, Thermo Scientific, US). The cycling protocol was 5 min at 95°C for initial denaturation, 35 cycles of amplification; 94°C for 45 s, 60°C annealing for 45 s, 72°C for 45 s and 10 min at 72°C for final extension. The PCR products were checked on 1 % agarose gel using horizontal electrophoresis and the gels were stained using SafeView™ Classic (Applied Biological Material Inc. Canada).

716 bp of LYZ gene was sequenced by Applied Biosystems 3500XL Genetic Analyzer System (Applied Biosystems, USA) in order to identify the LYZ gene sequence. The electrophenograms were carefully checked by using ChromasPro Version 2.1.8 (Technelysium Pty. Ltd. Australia). The trimmed sequence file consisting of LYZ fragments were analyzed by the MEGA7 software (Molecular Evolutionary Genetics Analysis, version 6.0) (Kumar et al., 2016).

Results and Discussions

In this study we have identified the gene composition of LYZ gene in donkeys where LYZ was found in relation with immune system in farm animals. The genetic characterization of LYZ gene was analyzed by DNA sequencing and the matching with the DNA sequences taken from GenBank were conducted (Accession number NW_014638180.1). LYZ gene sequence is located between 1781536-1786728 bp at NCBI GenBank database records (Accession number NW_014638180.1). In our study, the studied LYZ gene region has spanned between 5’ UTR 1st – 2nd exon, partial 1st intron and regions in donkeys and it includes 47 amino acids. It is shown the studied 716 bp long partial 3’ UTR, exon 1 (yellow section), intron 1, exon 2 (blue section) regions and SNPs on the LYZ gene in Figure 1.

Three novel SNPs were identified as g.1782775A>G, g.1782924A>G and g. 1782960T>C in the first intron region of LYZ gene (Figure 2). These novel SNPs were identified and the partial DNA sequence of LYZ gene in donkeys was entered for the first time in this study and these sequences were enrolled to NCBI Genbank database with the accession number: MK984689-MK984692.
The studied 716 bp long partial 3' UTR, exon 1 (yellow section), intron 1, exon 2 (blue section) regions and SNPs on the LYZ gene

Cosenza et al. (2018) found two variants as A and B which differ at 32 positions and a SNP g.203C>T that alters a putative binding site of the transcription factor NF-1 in the promoter region of LYZ gene. Cosenza et al. (2017) identified a transition G>A at the 160th nucleotide of exon 2 (NW_014638180:g.1784688C>T) caused an amino acid change from Arg90 to Gln. Šarić et al. (2012) analyzed the antibacterial properties of the lysozymes in raw donkey milk of Serbian breed. They found that the protective antimicrobial factors are very high in donkey milk which has positive effect on gut health and immune system. Nayak et al. (2017) clarified that lysozyme could be remarked as the notable antibacterial agent in donkey milk, because it is present in high concentration similar with Šarić et al. (2014). Aspri et al. (2018) reported that antimicrobial activity of donkey milk samples is high because it is digested with gastrointestinal enzymes similar as reported in Tidona et al. (2011). This antimicrobial effect in donkey milk might be partly because of high antimicrobial proteins as lactoferrin and lysozyme, the reason of this is predict as resistance to degradation by gastrointestinal enzymes.

Nayak et al. (2017) reported that lysozyme is an antimicrobial agent in donkey milk that effect gram positive bacteria, as well as a complex medium involving a large number of different compounds that may contribute to general antimicrobial potential.

Conclusion
Donkey milk has some differences from other ruminant milk with fat ratio, minerals content, lactose and the presence of antimicrobial proteins. The three crucial whey proteins in donkey milk are lysozyme, β-lactoglobulin and α-lactalbumin. One of these antimicrobial proteins is lysozyme in donkey milk that is effective in the elimination of some viruses, anti-microbial activity, antitumor and inhibition.

Very few studies have been carried out on LYZ gene variations in donkeys. In this study, new polymorphisms are found which will implement be beneficial information for improving food intake and immune response in donkeys based on marker-assisted selection. Thus, in further studies it is needed to perform more research and to realize the mechanism and pathways of lysozyme.

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