Molecular characterization and genetic variability of Alpha Casein gene, CSN1S1 in Bikaneri camel (Camelus dromedarius) milk

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ABSTRACT
Camel milk is an important protein source for the nomadic communities living in the arid lands of the world. In recent years there has been an increase in consumption of non-bovine milk as an alternative protein source for humans. Camel milk seems to be containing larger amount of total proteins, such as lactoferrin and immunoglobulins as compared to the cow milk, which may be responsible for the better antimicrobial properties. The casein fraction of milk proteins consists of four caseins, namely αs1-casein, αs2-casein, β-casein, and κ-casein. Casein genetic polymorphisms are important due to their effects on quantitative traits and technological properties of milk. This work was designed to study occurrence of polymorphism of α-casein in native Bikaneri camel (Camelus dromedarius) raw milk sample and to characterize these variants on molecular level.

Key words: Alpha-casein, Casein gene polymorphism, Camel milk production traits, Camelus dromedarius.

INTRODUCTION
Camels belong to the family of Camelidae, were domesticated around 3000–6000 years ago (Sushma et al., 2014). It has two species, double-humped camel (Camelus bactrianus, 70 chromosomes) and single-humped camel (Camelus dromedaries, 74 chromosomes). The single-humped camel inhabits dry, desert habitat of Africa, Arab and west Central Asia while the double-humped inhabits eastern Central Asia and China (Cui et al., 2007). In recent years there has been an increasing interest in consumption of non-bovine milk as an alternative protein source for humans, and is being promoted as healthy food due to its therapeutic value such as antioxidant activity. Camel milk is an important protein source (Kappeler et al., 1998), especially for the people living in the arid lands of the world, where other sources of proteins are scarce (Konuspayeva et al., 2009). In composition camel milk is more similar to goat milk and contains less short-chain fatty acids than cow, sheep and buffalo milks, and it contains about 3 times greater vitamin-C than cow milk.

It is worth noting that, camel milk seems to be containing larger amount of total proteins, such as lactoferrin and immunoglobulins as compared to the cow milk, which may be responsible for the better antimicrobial properties (Farah, 1993). Moreover unlike cow milk, camel milk is reported to have antidiabetic (Pauciullo et al., 2012) and anti-hypertensive effects. Camel milk fat lacks β-lactoglobulin and is rich in immunoglobulins which are compatible with human milk (Shamsia, 2009). Studies in India have revealed that populations who regularly consume camel milk have zero incidences of type 1 diabetics as compared to 5.5 percent in other communities that do not consume this milk (Rachagani et al., 2008). One of the biggest hurdles in camel milk processing is its incompatibility with the Ultra High Temperature (UHT) exposure, which is present day dairy industries use to preserve milk.

The milk composition of camel has been studied previously by many authors. Total protein content ranges from 2.4 to 5.3 percent (Konuspayeva et al., 2009; Nikkah, 2011a, 2011b) and is divided into caseins (CN ~80%) and whey proteins. (Giambra et al., 2013; Ereifej et al., 2011; Ikonen et al., 2008; Medrano et al., 1990). There are four main types of milk caseins, i.e. αs1-casein, αs2-casein, β-casein, and κ-casein, encoded by four genes, CSN1S1, CSN1S2, CSN2 and CSN3, respectively. In camel milk α-casein (22percent) is the second main fraction after β-casein (65percent) (Kappeler et al., 1998; El Agamy et al., 2006). Polymorphism of the blood groups, enzymes and milk proteins could be used as biological tools to improve the genetic merit in animal breeding studies. Casein genetic polymorphisms are important and well-known due to their effects on quantitative traits and technological properties of milk (Frajman et al., 2004). The objective of this study was to investigate genetic and phylogenetic analysis of Bikaneri camel’s milk casein gene.

MATERIALS AND METHODS
Sample collection and DNA isolation: Milk and blood samples from camel (C. dromedarius) were collected from Bikaner region, Rajasthan. Milk samples 10mL was collected

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with Sodium azide and stored at -4°C. Blood samples (10 mL) were obtained and stored at 20°C temperature until DNA extraction. The genomic DNA was isolated from the samples using the standard protocol (Shashikanth, 1999). For mRNA extraction, milk somatic cells, from fresh camel milk samples were gained by centrifugation (10 min, 2200 X g), and washed twice with phosphate buffered saline/0.5 M EDTA according to Boutinaud et al., (2002). Invisorb Spin RNA Mini Kit (Invitek GmbH, Germany) was used for extraction of total RNA.

**PCR Amplification and purification:** To amplify targeted DNA sequences by polymerase chain reaction (PCR), Primer3 v.0.4.0 (Rozen and Skaletsky, 2000) was used to design five set of primer pairs using camel α-casein mRNA-sequence (GenBank ID: AJ012628-AJ012630) as a preliminary template. The PCR amplification was carried out in a volume of 25 μl, containing 100 ng genomic DNA, 0.5 μl m of each primer, 0.3 μl of Tag DNA polymerase (Fermentas), 0.5 μl MgCl2, and 0.5 μl of dNTPs. Amplification was carried out in a GeneAmp9600 thermal cycler (PerkinElmer) employing the following conditions: 94°C for 2 min; 30 cycles at 94°C for 30 s, 50°C for 5 s, and 68°C for 5 min. The amplified PCR product was assessed by agarose gel electrophoresis using 1% EE agarose using 1X TAE buffer and visualized under UV light.

**Sequencing reaction and sequence analyses:** PCR products were sequenced directly using 50 ng (2.0 ul) of PCR product and 4 pM (1.0 ul) of primer, 4 ul of BigDye Terminator ready reaction kit (Perkin Elmer), and 3.0 ul of double distilled water to adjust the volume to 10.0 ul. Cycle sequencing was carried out in a GeneAmp9600 thermal cycler (Perkin Elmer) employing 30 cycles at 96°C for 10 s, 50°C for 5 s, and 68°C for 4 min. Products were purified by alcohol precipitation followed by 70% alcohol washing. Purified samples were dissolved in 10 ul of 50% Hi-Di formamide and analyzed in an ABI3700 automated DNA Analyzer (Perkin Elmer). Resulting chromatograms were compared by alignment using Chromas software ver 2.6.4 (Technelysium Pty. Ltd., Tewantin, Queensland, Australia).

**Statistical analyses:** Sequences used for analysis were downloaded from NCBI, aligned in the ClustalX package (Thompson et al., 1994), Nucleotide BLAST program at NCBI was used for sequence homology searches in public databases. The Sequence data from amplified gene product were edited manually using Chromas Ver.2.6.4. Multiple sequence alignments were performed using MegAlign tool of LASERGENE software (DNA STAR, Inc., Madison, WI, USA). The GENEDOC package was used for formatting the sequences to make them compatible with the desired software. Allele and genotype frequencies were calculated using PopGene program v. 1.31 (Yeh et al., 1999). To test the distribution of genotypes on the base of Hardy-Weinberg, chi-square test was performed. For determination of phylogenetic relationship of casein among different mammalian species, nucleotide sequences from the GenBank and Ensemble databases were used.

**RESULTS AND DISCUSSION**

Sequences were edited and initially aligned using SEQUECHER and then optimally aligned visually. Multiple sequences downloaded from NCBI were aligned by ClustalX package. In the present study direct sequencing of the complete coding region of camel samples revealed full sequence similarity to α-casein A of Kappeler et al. (1998) who reported the occurrence of two variants (A & B) in camel αS1-casein, it did not revealed any variability as reported earlier. The sequencing revealed that the size of alpha (s1)-casein cDNA was of 1087 bp with GC content of 42.5 percent. These results were comparable with other submitted sequences in NCBI for α-casein gene (1094 bp). The gene coded 222 amino acids precursor with a signal peptide of 15 amino acid residues, Coding sequences were translated to amino acids using the EBI online translation tool. On the whole, the camel sequence shares a similar organization with the bovine counterpart, with some differences (AY380228). Genetic polymorphism at α-casein locus the estimated minimum values of x² (0.2082) evidenced for a good relatedness between empirically obtained and theoretically expected genotypes population. The observed H indicates the representativeness of the samples, despite of their limited size as well as it emphasizes that current selection has not eliminated (suspended) any of the alternative forms (alleles) of the α-casein gene.

*Milks protein composition traits are associated with protein genetic variants. Camel α-casein may exist as several variants. Erhardt (2016) and workers have worked on casein gene polymorphism in milk samples of camels from different regions of Sudan (Africa) by isoelectric focusing, where three protein patterns named α-casein A, C, and D were identified. The major allele A revealed frequencies of 0.79 (Lahaoi), 0.75 (Shanbali), 0.90 (Arabi Khali), and 0.88 (Arabi Ghabrui) in the different ecotypes. α-casein C variant shows a single G > T nucleotide substitution in the exon 5, leading to a non-synonymous amino acid exchange (p.Glu30 > Asp30, Genbank ID: JF429138) in comparison to α-casein A and D. At cDNA level, no further single nucleotide polymorphisms could be identified in α-casein A, C, and D. The variants A and C are characterized by missing of exon 18 as compared to CSN1S1* B, due to DNA insertion of 11 bp at intron 17 which alter the pre-mRNA spliceosome machinery, multispecies alignment of Martin et al. (2003) already confirms. Alpha α-casein C (Shuiep et al., 2013) and α-casein B (Kappeler et al., 1998) are both characterized by p.Glu30 > Asp30 in the deduced mature protein sequence, and are only differing by missing or presence of the eight amino acids encoded by exon 16. It is often skipped during the processing of mRNA, as it is known in sheep (Ferranti et al., 1995; 1998).
Beside exon skipping as a reason for casein allele, simultaneous occurrence of skipped and non-skipped forms of the same CSN1S1 allele are usual in sheep, goat, cattle, pig, and human. However, the characteristics of camel \( \alpha \)-casein have not been well-documented. Further research and detailed DNA sequencing studies are needed to clarify complete sequence differences between \( \alpha \)-casein *A, B, and C* in detail, probably the occurrence of this variant is characteristic of other *C. dromedarius* ecotypes.

It is well known that the variation in mRNA and protein is primarily due to alternative splicing, duplication, and insertion/deletion events in addition to nucleotide mutations. Alternative splicing as a reason for the presence of minor and major fractions of camel milk protein is already described for camel lactophorin, a major whey protein in camel milk by Kappeler *et al.*, 1999. About 75 percent of the protein is expressed as a long variant and the minor fraction (25 percent) of camel lactophorin is a shortened variant, characterized by the lack of exon 2. Long and short variants of \( \alpha \)-S1-casein also occur in ovine milk as a result of differential splicing of the heterogeneous nuclear RNA (Ferranti P. *et al.*, 1995), as well as it was also showed in goat (Ramunno L. *et al.*, 2005; Leroux C. *et al.*, 1992) and cattle where, for instance, the skipping of the exon 4 results in the A variant (Mohr U. *et al.*, 1994).

The database search of sequences for a possible match to the DNA sequence of casein gene was conducted using the BLAST (NCBI). In the phylogenetic trees, constructed from the sequences of the alpha (S1)-casein mRNA as well as protein sequences, it has been observed that camel, buffalo, cattle, goat and sheep formed a cluster with a closer relationship between cattle and buffalo followed by goat and sheep based on genetic distance (F, values) using MEGA5 (Tamura *et al.*, 2011). Comparing gene sequence with other reported sequence in this region showed estimated homology as 94, 94, 94, 82, 94, 97 and 96 percent, respectively with that of cattle, goat, sheep, pig, buffalo, equine and human. A similar trend was observed when compared with amino acid sequences of these species.

However, this could be a starting point for further characterization of genetic diversity in camel milk proteins. Milk protein variability would help in studying association concerning milk performance traits in camel. Moreover, at phenotypic level, milk protein could be used to increase economic value of milk, for screening of breeds and population for phenotyping animals for breed characterization.

**CONCLUSION**

Camel milk has developed a high reputation as a healthy nutrition with most of its therapeutic value ascribed to its biological properties such as antioxidant activity. Casein genes have been deeply investigated in ruminants, whereas little information is available in camel, the present study can be a starting point for further characterization of genetic diversity in camel milk proteins. The genetic polymorphism of milk proteins can be studied and used further for better diversity, genetic structure preservation assessment of populations and the relationship with their dairy production traits.

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