Molecular Characterization of Genetic Variability among Sudanese Baggara Cattle within Kappa Casein CSN3 Gene (Exon V)

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Abstract

The present work was conducted to detect possible genetic variants of kappa casein (CSN3) gene in Western Baggara cattle in Sudan. This gene is associated with higher fat, protein, and has a significant influence on milk processing properties in comparison to other casein variants. 70 blood samples were collected from Western Baggara cattle in three areas (Hai Elgabal, Bulbul and Umdamm). Genomic DNA was extracted from the whole blood using guanidine chloroform method. A 495 bp fragment containing exon V of kappa casein gene was amplified by PCR using specific set of primers. The amplified PCR product was digested by Hind I; the bands were visualized under UV-light to observe the polymorphic locus by the size change of DNA fragments. The results revealed that the restriction analysis produced two fragments of 424 and 71 bp indicating the presence of kappa casein B genotype (monomorphic) in all population. The present study is the first report on kappa casein genotype of Baggara cattle in South Darfur State, Sudan, which elucidated uniform and homozygous population for kappa casein B allele.

Keywords: Molecular characterization; CSN3; Exon4; Baggara cattle; Sudan

Introduction

Milk is very important in human nutrition because, it provides energy, high quality protein, vitamins and minerals requirements. However, among the different composition milk protein is considered the most important component therefore, investigation on milk protein particularly caseins is still arousing research interest due to their crucial role in milk quality, composition and processing properties [1].

Out of fourteen variants of κ-CN have been explained i.e. A, A1, B, B2, C, D, E, F1, F2, G1, G2, H, I and J; the A and B are most common in the majority of cattle breeds followed by κ-CN: H [2]. The majority of researchers believe that the kappa casein B variant is associated with higher fat, protein, and casein in the milk and has a significant influence on cheese making properties of milk and superior rennet coagulation properties in comparison to AA or AB variants. The genotypes BB and AB are used in artificial insemination programs to obtain a greater increase of the frequency of these alleles in cattle populations of commercial interest [3].

Western Baggara cattle is belonging to Northern Sudan zebu cattle which represent (33%) of the national herd of the Sudan and contribute effectively in Sudan Gross Domestic Product (GDP) and providing meat source for both export and local consumption and also playing a vital role in improving the livelhoods and enhancing the food security for nomadic tribes in their natural habitat in Kordofan and Darfur [4].

The classifications of Sudanese cattle are based on phenotypic characteristics or geographic origin and are not related to genotype except in as much as the phenotype is in part a reflection of genotype. With the advent of molecular biology technology, a powerful new tool is available for characterization, classification and estimation of distances between breeds and strains. The investigation of genetic variation is very important for future monitoring of gene flow in populations, conservation of species, determination of the level of inbreeding and crossbreeding within and between breeds [5]. The present work was prepared as first attempt to detect possible genetic polymorphism within CSN3 gene (exon four) of Baggara cattle in South Darfur state, Sudan.

Materials and Methods

Collection of blood samples

Blood samples were obtained from the jugular veins of 70 Western Baggara cows in areas of the study (Bulbul, Hai Elgabal, and Umdamm). 10 mL were taken from each cow from the jugular vein in a clean sterile tube contains EDTA and stored at -20°C until time of DNA extraction.

DNA extraction

Genomic DNA was extracted from the whole blood using guanidine chloroform method as described by Ciulla et al. [6]. Quality and concentration of the DNA was checked by electrophoresis on 2% agarose gel.

PCR amplification

A 495 bp fragment containing exon V of kappa casein gene was amplified by PCR using forward 5’GGATTCTCCAGCAAGGAAATA3’ and reverse 5’ GTGGGCTCTCATAACTTCTTG 3’, PCR was carried into 20 μL volume of PCR amplification cocktail contained 1μL of each primer, 1x PCR master mix (DYNAzyme), 2 μL of template DNA and 16 μL of double distilled water. The amplification cycles were carried out in a
PTC-100 thermocycler. Reaction conditions were 94°C for 5 min as initial denaturation followed, by 30 cycles of 94°C for 1 min, 58°C for 1 min and 72°C for 1 min. a final extension step at 72°C for 5 min was followed.

Restriction fragment length polymorphism (RFLP)

After PCR the product was digested by 7.5 units of haemophilus influenza serotype F restriction enzyme (Hinf1) in a final volume of 25 µL containing 1x of the enzyme buffer (2.5 µL), 5 µL of PCR product and 17 double distilled water. The digestion mixtures were incubated at 37°C overnight in the incubator. After digestion, the resulting fragments were analyzed by electrophoresis on 2% agarose gel using 1x TBE buffer containing 0.2 µL ethidium bromide at 50 V until complete separation of the bands. The bands were visualized under UV-light to observe the polymorphic locus by the size change of DNA fragments.

Results

Restriction Fragment Length Polymorphism (RFLP) said to be one of the most accurate, low cost and reliable technique for the identification of structural gene polymorphism that occur as a result of point mutation [2]. In this work; PCR was used for gene amplification using specific primers yielded a 495 bp fragment of kappa casein gene. Observation of a single band visible under UV light removed the need for PCR purification step before restriction analysis as shown in Figure 1.

RFLP analysis was used for allele typing. Genotyping was performed by the restriction digestion of the amplified products using Hinf1 restriction enzyme followed by agarose gel electrophoresis for analysis of the digestion pattern and the restriction analysis produced two fragments of 424 and 71 bp indicating the presence of kappa casein BB genotype (monomorphic) in all population (Figure 2).

Discussion

The present study was aimed to detect possible kappa casein gene polymorphism within exon four among Western Baggara cattle breed. PCR-RFLP technique was used to detect kappa casein genetic variants. A 495 bp fragment of kappa casein gene was digested by Hinf1 restriction enzyme. The results indicated that all studied population of Baggara cattle was found to be monomorphic genotype (BB) with only two bands (71 and 424 bp). The results in the present study were in agreement with numerous researchers: Riaa et al. [7] who detected one allele (BB) in Nilli Rvi breed of Pakistan and Raj et al. [8] and El Rafey and Darwish [9] detected the gene in Egyptian buffalo, Osman [10] reported one allele BB genotype among River Buffalo breed and also, reported monomorphic (BB) for this gene in female buffaloes. Similarly, Zakizaden et al. [11] reported that the frequency of B allele of kappa casein gene in Holstein breed ranges from (0.06-0.57). Some researchers observed highest frequency of allele (B) in Brown Swiss and Jersey with 0.67 and 0.86, respectively [12-16], however, frequency of this allele (B) in Norwegian cattle breeds and Northern region of Europe was found low [17,18]. Also, Faradi et al. [19] reported genotype AA, AB and BB with frequencies of 35%, 37.5% and 27%, respectively among Najdi cattle breed this indicated that BB homozygous frequency was low.

On the other hand, the present findings were not in line with Rahamtalla et al. [20] who studied polymorphism of kappa casein gene
in Butana and Kenana dairy cattle and found frequencies of allele A was higher than allele B but, where, BB genotype was not observed [21]. Also, the present results were not agree with Memon et al. [2] who found the genotype frequency for homozygote AA and the allelic frequency of allele A were higher than the same for homozygote genotype BB and the allelic frequency of allele B, respectively in Pakistani Sindhi cattle. Deb et al. [21] reported that the A allele was more frequent than the B allele and no BB homozygous was observed among the Frieswal (HF × Sahiwal) cross breed of Indian origin. Also, the present results were in disagreement with Mitra et al. [22] who observed low frequency of B allele and absence of homozygous BB animals in Sahiwal cattle. Ng-Kwai-Hang et al. [23] claimed that the frequency distribution of kappa casein genotypes shows rare occurrence in Holstein breeds.

Conclusion

The present study is the first report on kappa casein genotype of Baggara cattle in South Darfur State, Sudan, which indicated uniform and homogenous population for kappa casein B allele.

The majority of researchers believe that the kappa casein B variant is associated with higher fat, protein, and casein in the milk and has a significant influence on cheese making properties of milk and superior rennet coagulation properties in comparison to AA or AB variants, therefore, In future genetic improvement strategies; homogenous population of kappa casein B allele for western Baggara cattle should be considered.

References