

PAPER

Molecular characterisation of κ -casein gene in Girgentana dairy goat breed and identification of two new alleles

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Abstract

The κ -casein fraction plays an important role in the formation, stabilisation and aggregation on casein micelles and thus affects technological and nutritional properties of milk. In this study, exon 4 of κ -casein (CSN3) gene was sequenced and analysed in Girgentana goat breed. Analyses of the obtained sequences showed the presence of A, B, D, and G known alleles and two new genetic variants, named D' and N. The new D' allele differs from D in one transition, $G_{284} \rightarrow A_{284}$, which did not cause amino acid change. The new N allele differs from A in five single nucleotide polymorphisms (SNPs): T₂₄₅/C₂₄₅, G_{284}/A_{284} , G_{309}/A_{309} , G_{471}/A_{471} and T_{591}/C_{591} , while it differs from C in one transition, *i.e.* $T_{583} \rightarrow C_{583}$. Comparing the amino acid sequences of N and A alleles, the first two SNPs caused no amino acid change, whereas the other SNPs produced changes (Val₆₅/Ile₆₅, Val₁₁₉/Ile₁₁₉, and Ser₁₅₉/Pro₁₅₉, respectively). Comparison of N allele with C revealed the amino acid change Val₁₅₆->Ala₁₅₆. The most frequent allele was A (0.480) followed by B(0.363), D (0.112), and N (0.034). The D' and G alleles were identified only in two animals and in heterozygous conditions with a very low frequency (0.005). The most common genotype was AB (39.5%) followed by AA (19.5%), AD (12.7%), and BB (11.7%). Homozygous D'D', GG, and NN individuals were not found. Further analysis will be performed in order to establish associations among genotypes and quantitative and qualitative milk traits.

Introduction

In the milk of ruminants, more than 95% of proteins are synthesised by six structural genes, four caseins [αs_{I^-} , β -, αs_{2^-} and κ -casein (CSN3)] and two whey proteins (α -lactalbumin and β -lactoglobulin). Polymorphisms of the four casein genes have been the focus of considerable research effort because of their potential effects on milk quality. The κ -casein fraction plays an important role in the formation, stabilisation and aggregation of the casein micelles and thus affects the technological (Mariani et al., 1976; Aleandri et al., 1990; Lodes et al., 1996; Falaki et al., 1997) as well as nutritional properties of milk (Mercier et al., 1973, 1976; Malkoski et al., 2001).

The goat CSN3 gene comprises five exons (Coll et al., 1993, 1995) with the mRNA coding region for mature protein (171 amino acids) spanning from exon 3 (9 amino acids) to exon 4 (162 amino acids) (Yahyaoui et al., 2003). The CSN3 gene is considered to be monomorphic in sheep (Moioli et al., 1998) whereas several studies on goat CSN3 showed that this gene is highly polymorphic (Caroli et al., 2001; Yahyaoui et al., 2001; Angiolillo et al., 2002; Chessa et al., 2003; Yahyaoui et al., 2003; Jann et al., 2004; Reale et al., 2005; Prinzenberg et al., 2005; Gupta et al., 2009; Kiplagat et al., 2010). According to the last nomenclature proposed by Prinzenberg et al. (2005), a total of 16 DNA variants have been identified in the domestic goat, of which 13 are protein variants (named in alphabetical order from A to M) and 3 are silent mutations (B', B'') and (B', B'') involving a total of 15 polymorphic sites. Recently, Gupta et al. (2009) in Jakhrana goat breed and Kiplagat et al. (2010) in indigenous Eastern African goat population reported the presence of new genetic polymorphisms at CSN3 gene. These studies reported conflicting results for allele nomenclature because according to Prinzenberg et al. (2005) only missense mutations associated with amino acid changes should be indicated with new allele names (i.e. new letter), while silent mutations should be named with the same letter as the related protein-associated allele followed by prime symbol. The Girgentana goat is an ancient Sicilian goat breed reared in Southern Italy for its good dairy production. Average milk production was 224±66 L in the first lactation, and 320±109 L for later lactations (AIA, 2013). Due to sanitary policies, population size of Girgentana goat breed decreased of almost 90% in 20 years. In 1983, the population consisted of 30,000 individuals but, nowadays, only 374 heads are enrolled in the Herd Book (ASSONAPA, 2013). Corresponding author: Prof. Baldassare Portolano, Dipartimento Scienze Agrarie e Forestali, Università di Palermo, viale delle Scienze 4, 90128 Palermo, Italy. Tel. +39.091.23896069 - Fax: +39.091.23860814. E-mail: baldassare.portolano@unipa.it

Key words: CSN3 gene, Polymorphisms, New alleles. Girgentana goat breed.

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Over the last years this breed has become almost extinct, in part as a consequence of the marked decrease in fresh goat milk consumption. The aim of this work was to investigate the genetic polymorphisms of *CSN3* gene in the *Girgentana* dairy goat breed in order to assess genotypes distribution and to use this information in future conservation programmes for this breed considering that genotype could influence milk properties.

Materials and methods

Sampling and DNA extraction

A total of 205 individuals, all females, of Girgentana goat breed were randomly chosen. They belonged to 15 different herds located in different areas of Sicily. Samples were collected from 10 to 20 unrelated individuals per herd. About 10 mL of blood was collected from jugular vein using vacutainer tubes containing ethylenediamine tetraacetic acid (EDTA) as anticoagulant. Genomic DNA was extracted from buffy coats of nucleated cells using a salting out method (Miller et al., 1988). After checking the quantity and quality of the DNA using a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA), samples were diluted to a final concentration of 50 ng/µL in ultrapure water and stored at 4°C until use.





DNA amplification and purification

A 552 bp fragment of Girgentana goat κ casein exon 4 (GenBank Acc. No. X60763 mRNA goat CSN3) was amplified by polymerase chain reaction (PCR) using the following primers: forward -AGAAATAATACCATTCTG-CAT, and reverse - TCTTTGATGTCTCCTTAGAG. The PCR reaction was performed in a 25 µL of final volume containing 1 µM of each primer, 1 mM of dNTP Mix, 1 U of Taq DNA polymerase (Fermentas, Hanover, MD, USA), 1X PCR buffer with KCl, 1.25 mM MgCl2, and approximately 100-150 ng of genomic DNA. Thermal cycling conditions were 94°C for 90 sec for initial denaturation, 30 cycles of 45 sec each at 94°C, 50°C and 72°C, with a final extension at 72°C for 5 min. The PCR products were checked by electrophoresis on 2% agarose gel stained with SYBR Safe (Invitrogen, Carlsbad, CA, USA).

DNA sequencing reaction

All the collected samples were amplified and the PCR products purified in order to sequence and determine the complete nucleotide sequences. Polymerase chain reaction products were purified using 10 U of Exonuclease I and 1 U of Shrimp Alkaline Phosphatase (Fermentas). DNA sequencing reaction was carried out using BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Carlsbad, CA, USA) with 5 µM of the same primers used in the PCR reaction. Cycle sequencing reaction was performed according to manufacturer's instruction following Ethanol/EDTA/Sodium Acetate precipitation. Sequencing analyses were performed in an ABI PRISM 3130xl Genetic Analyzer (Applied Biosystems).

Sequence data analysis

Nucleotide sequences obtained were checked using Sequencing Analysis Software v5.3.1 (Applied Biosystems) and subsequently analysed with SeqScape v2.5 Software (Applied Biosystems). Polymorphic sites were confirmed by visual examination of the electropherograms. Multiple alignments of the sequences were performed using ClustalW software (Thompson et al., 1994). The translation of DNA sequences to amino acid sequences was performed using ExPASy-Traslate tool. The same software was used to calculate the isoelectric point (IP) of the new genetic variants found in Girgentana goat breed. The exact P value associated with the null hypothesis of Hardy-Weinberg equilibrium (HWE) was estimated using GENEPOP version 4.0.11 (Rousset, 2008). The programme performed a probability test using a Markov Chain

Table 1 K-casein gene variants according to Prinzenberg et al. (2005).

A AFF AND	CSN3 variant	IEF pattern	GenBank Acc. no.							Nuclec	Nucleotide position	sition							Reference
AIFF X69763 C T A G G G G A T				170	245	247	274	284					385	471		550	583	591	
A ^{IE} AF485340 AF478388 AFE AF485341 AFE AF485341 AFE AF7 AFE BEF AF7 AF7 AFF AF7 AF7 AF7 AF7 AF7 AF7 AF	А	AIEF	X60763	C	Т	A	A	9	S	A		g	А	g	А	⊢	၁	⊢	Coll <i>et al.</i> (1993)
AFF	В	A^{IEF}	AF485340											А					Yahyaoui <i>et al.</i> (2001)
Aler AY166706 T T A A A A A A A A A A A A A A A A A			AF434988											А					Jann <i>et al.</i> (2004)
AFF	B,	$A^{\rm IEF}$	AY166706	⊢										A					Jann <i>et al.</i> (2004)
AFF AFASSA425 C A A A A A T(Val) C AFF AFASSA41 C G G(Leu) A A A T C BFF AY090465 C G G(Leu) A A G T C BFF AY090465 C G A(Leu) A A A C C AFF AFS0202 C G A(Leu) A A A C	B'.	A^{IEF}	AY166707						Г					Α					Jann <i>et al.</i> (2004)
A ^{IEF} AF483341 C A A A A G T C B ^{IEF} AY027868 C G G(Leu) A A A T C AY090465 C G G(Leu) A A A C C B ^{IEF} AY486523 C G A(Leu) A A C C C G A C C C C A A A C	C	A^{IEF}	AY350425		ల			A			A			A		- '	r(Val)	၁	Prinzenberg et al. (2005)
BIEFO AY027868 C G G(Leu) A A A C C BIEFO JX889422 C G A(Leu) A A A C	j	A^{IEF}	AF485341		၁			A			A			A	g		Ē	၁	Yahyaoui <i>et al.</i> (2001)
BIEPO AY090465 C G G(Leu) A A A A A BIEFO JX889422 C G A(Leu) A A A A BIEFO AF486523 C G A(Leu) A A A AIFFO AY090467 C C A A AIFFO AY090467 C C A A AIFFO AY166710 C A A AIFFO AY166710 C A A AIFFO AY166709 C A A AIFFO AY18577 C A A AIFFO AY18577 C A A AIFFO AY428577 C A AIFFO AY428577 C A AIFFO AY428577 C AIFFO AY428577 C A AIFFO AY428577 C A	D	B ^{IEF}	AY027868		၁	9					A			А				၁	Caroli <i>et al.</i> (2001)
BIEP			AY090465		೦	9	J	(Leu)			A			A				၁	Yahyaoui <i>et al.</i> (2001)
BIGF AF486523 C A A A A C <th< td=""><td>D,</td><td>B^{IEF}°</td><td>JX889422</td><td></td><td>ပ</td><td>9</td><td>¥</td><td>(Leu)</td><td></td><td></td><td>A</td><td></td><td></td><td>A</td><td></td><td></td><td></td><td>၁</td><td>This paper</td></th<>	D,	B ^{IEF} °	JX889422		ပ	9	¥	(Leu)			A			A				၁	This paper
AIGH AY090466 C C C C A A A A A A A A A A A A A A A	ш	B ^{IEF}	AF486523										9	A					Angiolillo et al. (2002)
AIGH AY090467 C G A A A A A A A A A A A A A A A A A A	ᄕᅭ	A^{IEF}	AY090466		၁									A				၁	Yahyaoui <i>et al.</i> (2003)
A ^{IEF} AF521022 G A A A ^{IEF} AY166710 G A A A ^{IEF} AY166711 G A A B ^{IEF} AY166709 G A A I B ^{IEF} AY428577 C C C A ^{IEF} JX889424 C A A C(Ala) C	g	A^{IEF}	AY090467		ပ						A			A				၁	Yahyaoui <i>et al.</i> (2003)
A ^{IEF} AY166710 G A A A A A A A A A A A A A A A A A A	Н	A^{IEF}	AF521022				9							A					Jann <i>et al.</i> (2004)
A ^{IEF} AY166711 G A A B ^{IEF} AY166709 G A A ^{IEF} AY166708 C C A A A A C A ^{IEF} AY428577 C A A A A A C A ^{IEF} AY428577 C C A ^{IEF} A A C C(Ala) C	Ι	A^{IEF}	AY166710								A			A					Jann <i>et al.</i> (2004)
B ^{IEF}	ſ	A^{IEF}	AY166711							9				A					Jann <i>et al.</i> (2004)
A FF	K	B ^{IEF}	AY166709			9								A					Jann <i>et al.</i> (2004)
AY428577 C A A A C C C C C 3.3889424 C A A A C (Ala) C		$A^{\rm IEF}$	AY166708								A			A				၁	Jann <i>et al.</i> (2004)
JX889424 C A A A C(Ala) C	M	$\mathrm{B}^{\mathrm{IEF}}$	AY428577		ల							A		A		၁		၁	Prinzenberg et al. (2005)
	Z	${ m A^{IEF\circ}}$	JX889424		၁			А			A			А			C(Ala)	С	This paper

g. "The IEF pattern of D" and Nalleles was not experimentally tested but it was estimated using Expasy pl tool. GenBank accession numbers, nucleotide position compared with X60673, and published reference are it skets. Reproduced with permission of the American Dairy Science Association via Elsevier: Journal of Dairy Science, Vol. 88, E.-M. Prinzenberg, K. Gutscher, S. Chessa, A. Caroli, G. Erhardt, Caprine x-casein (CSN3) p knowledge, page 1492, Copyright (2005). CSN3, x-casein gene; IEF; isoelectrofocusing. "The cated. Amino acids are indicated within brackets. I morphism: new developments in molecular knowl





method (1000 dememorisation steps, 100 batches, and 1000 iterations per batch). Moreover, GENEPOP was used to calculate genotype and allele frequencies and fixation index F_{is} (Weir and Cockerham, 1984). Expected (He) and observed (Ho) heterozygosity were calculated using the POPGENE software version 1.31 (Yeh $et\ al.$, 1999).

Results and discussion

Identified alleles in *Girgentana* goat breed

Sequencing analysis and alignment of the obtained sequences of CSN3 exon 4 showed the presence in *Girgentana* goat breed of A, B, D, and G known alleles and two new genetic variants (GenBank Acc. No. JX889419-JX889424). All single nucleotide polymorphisms (SNPs) described by Prinzenberg et al. (2005), including the two new polymorphic sites detected in our samples, are showed in Table 1. Considering the conflicting results reported by Gupta et al. (2009) and Kiplagat et al. (2010), we named D' and N the two new alleles identified in Girgentana goat breed according to Prinzenberg et al. (2005). The new CSN3 D' (GenBank Acc. No. JX889422) allele differing from CSN3 D (GenBank Acc. No. AY027868) in one transition $G_{284} \rightarrow A_{284}$, which did not cause amino acid change (Leu₅₆/Leu₅₆). The new CSN3 N (GenBank Acc. No. JX889424) allele differing from CSN3 A (GenBank Acc. No. X60763) allele in five SNPs $(T_{245}/C_{245}, G_{284}/A_{284}, G_{309}/A_{309}, G_{471}/A_{471})$ and T_{591}/C_{591}), while differing from C (GenBank Acc. No. AY350425) allele in one transition $(T_{583} \rightarrow C_{583})$. Comparing the amino acid sequences of CSN3 N and A alleles, the first two SNPs (T₂₄₅/C₂₄₅ and G₂₈₄/A₂₈₄) caused no amino acidic change, whereas the other SNPs produced changes: Val₆₅/Ile₆₅, Val₁₁₉/Ile₁₁₉, and Ser₁₅₉/Pro₁₅₉, respectively. Comparison of CSN3 N allele with CSN3 C allele revealed the amino acid change Val₁₅₆ → Ala₁₅₆.

Allele frequencies and genetic variability

Table 2 shows genotype and allele frequencies at CSN3 locus in Girgentana goat breed. The most frequent allele was A (0.480) followed by B (0.363), D (0.112), and N (0.034). The D' and G alleles were identified only in two animals and in heterozygous conditions with a very low frequency (0.005). These results are not in agreement with those reported for Girgentana goat breed by Gigli et al. (2008), and by other authors for Italian (Sacchi et al.,

method (1000 dememorisation steps, 100 Table 2. Genotype and allele frequencies at K-casein locus in Girgentana goat breed.

Genotype (n)	Frequency	Allele	Frequency
AA (40)	0.195	A	0.480
AB (81)	0.395	В	0.363
BB (24)	0.117	D	0.112
AD (26)	0.127	D'	0.005
BD (16)	0.078	G	0.005
DD (2)	0.010	N	0.034
D'G (2)	0.010		
AN (10)	0.049		
BN (4)	0.020		

n, number of individuals.

2005), European and African (Prinzenberg et al., 2005) goat breeds, where the most frequent allele was B. Prinzenberg et al. (2005), proposed to differentiate the nomenclature at protein level from the one used at DNA level introducing two codes (AIEF and BIEF) corresponding to two IPs (IP=5.53 and 5.78, respectively) identified using isoelectrofocusing (IEF) method (Table 1). According to this nomenclature, among the CSN3 alleles found in our study, only the D and D' alleles are included in B^{IEF} group, whereas A, B, G and Nalleles belong to A^{IEF} group (Table 1), which represents the less favourable variants group in terms of milk composition and technological properties (Chiatti et al., 2007).

Nine genotypic classes were found in our Girgentana goat samples. The most common genotype was AB (39.5%) followed by AA (19.5%), AD (12.7%), and BB (11.7%). The other genotypes showed a frequency of less than 10% (Table 2). In this study, we found no homozygous D'D', GG, and NN subjects. Caravaca et al. (2009), in a study on the effect of CSN3 genotypes on goat milk composition, showed that AB and BB genotypes were significantly associated with higher levels of total casein and protein content compared with the AA genotype, thus underlining the importance of taking into account the CSN3 genotype when performing selection for milk composition in dairy goats.

Observed and expected heterozygosity, fixation index F_{is} and P value associated with the null hypothesis of HWE were estimated. Significant departure from HWE was observed for *Girgentana* goat breed at *CSN3* locus (P<0.05), probably due to heterozygote excess (Ho=0.6766 vs He=0.6243). This hypothesis could be confirmed considering Ho heterozygosity and F_{is} (-0.0855) values.

Conclusions

Two new genetic variants have been identified and characterised in *Girgentana* goat breed. Currently, phenotypic data are not available for this goat breed; hence, further studies could establish the possible association and the effects of polymorphisms on quantitative and qualitative characteristics of milk. Moreover, it could be useful to take into account *CSN3* gene to use lines of goats producing different types of milk for specific cheese-making technologies or nutritional human needs.

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