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(Article begins on next page)

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Alpha S₁-casein polymorphisms in camel (*Camelus dromedarius*) and descriptions of biological active peptides and allergenic epitopes

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Abstract

Milk samples of 193 camels (*Camelus dromedarius*) from different regions of Sudan were screened for casein variability by isoelectric focusing. Kappa-casein and beta-casein were monomorphic, whereas three protein patterns named α_{s1} -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 Gharbawi) in the different ecotypes. *CSN1S1**C shows a single G > T nucleotide substitution in the exon 5, leading to a non-synonymous amino acid exchange (p.Glu30 > Asp30) in comparison to *CSN1S1**A and D. At cDNA level, no further single nucleotide polymorphisms could be identified in *CSN1S1**A, C, and D, whereas the variants *CSN1S1**A and *CSN1S1**C are characterized by missing of exon 18 compared to the already described *CSN1S1**B, as consequence of DNA insertion of 11 bp at intron 17 which alter the pre-mRNA spliceosome machinery. A polymerase chain-restriction fragment length polymorphism method (PCR-RFLP) was established to type for G > T nucleotide substitution at genomic DNA level. The occurrence and differences of IgE-binding epitopes and bioactive peptides between α_{s1} -casein A, C, and D after digestion were analyzed *in silico*. The amino acid substitutions and deletion affected the arising peptide pattern and thus modifications between IgE-binding epitopes and bioactive peptides of the variants were found. The allergenic potential of these different peptides will be investigated by microarray immunoassay using sera from milk-sensitized individuals, as it was already demonstrated for bovine α_{s1} -casein variants.

Keywords

Milk proteins

Genetic polymorphisms

CSN1S1

IgE-binding epitopes

Bioactive peptides

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Introduction

Camel milk plays an important role as protein source for many humans especially for the people living in the arid lands of the world (Konuspayeva et al. 2009). In addition, there is a growing interest in usage of camel milk as a healthy food (Nikkah 2011) and alternative protein source for humans with milk protein allergy (Hinz et al. 2012). However,

the knowledge in milk protein of this species and the genetic variation is very limited. Kappeler et al. (1998) described two genetic variants of α_{s1} -CN (*CSN1S1**A; *CSN1S1**B) by protein- and mRNA-sequencing within Somali camel (*Camelus dromedarius*). Shuiep et al. (2013) reported a partial genomic DNA sequence for α_{s1} -casein and could demonstrate genetic variation at protein and DNA level in two camel ecotypes of Sudan, whereas in the same camel population, Pauciullo et al. (2013, 2014) have found genetic variants in the regulatory regions of the kappa- and beta-casein (κ -CN; *CSN3* and β -CN; *CSN2*) genes. In sheep, isoelectric focusing of milk samples led to the identification of further alleles at alphas1-casein (α_{s1} -CN; *CSN1S1*) and α_{s2} -CN (*CSN1S2*). In addition, in East Friesian Dairy sheep and Lacaune sheep, *CSN1S1* variants were significantly associated with higher protein content (Giambra et al. 2014). Lisson et al. (2014) could demonstrate differences in the immunoreactivity of α_{s1} -CN epitopes not only between species but also between genetic variants of cows. In bovine and non-bovine milk, ACE-inhibitory properties could be demonstrated (Weimann et al. 2009, Salami et al. 2011, Moslehishad et al. 2013, Jrad et al. 2014) and proposed to be used as functional food with biological activity.

Therefore, the aim of this study was to investigate further the occurrence of polymorphisms in camel α_{s1} -CN especially in Sudan. Furthermore, options to describe the allergenic potential as well as bioactive peptides of those protein variants should be demonstrated.

Material and methods

Milk and blood samples

Milk and blood samples were collected from Lahaoi ($n = 81$), Shanbali ($n = 40$), Arabi Kahli ($n = 51$), and Arabi Gharbawi ($n = 21$) camel (*C. dromedarius*) ecotypes in Sudan. Areas of sampling were El Shuak, El Obied, west Omdurman, and Nyala, respectively (Fig. 1). Fresh milk samples ($n = 5$) of *C. dromedarius* were obtained from Kamelhof Rotfelden (Rotfelden-Ebhausen, Germany) and immediately kept at 4 °C.

Fig. 1

Areas of sampling. The locations (1–4) correspond to Table 1



Isoelectric focusing of milk proteins

Simultaneous phenotyping of camel milk protein variability on protein level was done by isoelectric focusing (IEF) according to Erhardt (1989) using a 0.3-mm polyacrylamide gel ($T = 5\%$, $C = 3.75\%$, and 7.1 M urea). The following mixture of carrier ampholytes was used: 0.4% (w/v) Servalyte pH 3.0–5.0 (Serva Electrophoresis, Heidelberg, Germany), 1.1% (w/v) Pharmalyte pH 4.2–4.9 (GE Healthcare Europe GmbH, Freiburg, Germany) and 1.2% (w/v) Servalyte pH 5.0–7.0 (Serva Electrophoresis, Heidelberg, Germany). After prefocusing for 14 min (20 mA , 3.000 V), milk samples as well as standard samples were separated for 67 min (40 mA , 3.000 V).

Molecular analyses: reverse transcription, PCR amplification, and sequencing

DNA was isolated according to Sambrook et al. (1989) from the blood samples on the filter paper (FTA[®] Classic Card—Whatman[®] BioScience, Maidstone, UK). For mRNA extraction, somatic cells of the fresh milk samples with α_{s1} -CN AA and AC were gained for mRNA extraction by centrifugation (10 min, 2200×g), and washed twice with phosphate buffered saline/0.5 M ethylenediaminetetraacetic acid according to Boutinaud et al. (2002). Invisorb[®] Spin RNA Mini Kit (Invitek GmbH, Berlin, Germany) was used for extraction of total RNA which was reverse transcribed to cDNA using Verso[™] cDNA kit (Thermo Fisher Scientific, Waltman, MA, USA).

For sequencing of *CSN1S1* cDNA, a set of primers for amplification were designed by means of DNAsis-Max ver. 3.0 software (Hitachi Software, San Bruno, CA, USA), using first the complete sequences of camel cDNAs reported by Kappeler et al. (1998) and then the new sequences determined. PCR was used to amplify complete cDNA of CSN1S1.

Primers were designed using GenBank sequence NW_006210750.1 in order to analyze the exons of CSN1S1 from animals carrying α_{s1} -CN*A and α_{s1} -CN*D. Primer pairs used for identifying the causal mutation on DNA level and their relevant annealing temperatures (x°C) are listed in Table 1.

Table 1

Forward and reverse primers, size of PCR products, gene regions, and annealing temp *CSN1S1*. Primers were designed using Genbank Acc. No. NW_006210750.1

PCR no.	Forward primer 5'-3' (name, sequence)	Reverse primer 5' (name, sequence)
1	CD-CSN1S1-5'UTR-F GGGAATCTTATTGATGTAACAGT	CD-CSN1S1-Intron1-R TACAAATGCATATAAAAT
2	CD-CSN1S1-Intron1-F CACTTCATGGATCAAATGCG	CD-CSN1S1-Intron2-R TGACCAAAATGTGAAGC
3	CD-CSN1S1-Intron2-Fb ACTTGGTGTTCAGACAGATTTAGCT	CD-CSN1S1-Intron3-R TGGCTGTTGTAGAAGGC
4	CD-CSN1S1-Intron3-F CCTGTGCCTCCCCTCTCTAT	CD-CSN1S1-Intron4-R TCCTTCTTTACCAGACC

5	CD-CSN1S1-Ex4-F TGAACCAGACAGCATAGAG	CD-CSN1S1-Ex6-R CTAAACTGAATGGGTGA
6	CD-CSN1S1-Intron5-F TGTACCTTTGCAAACACTCATG	CD-CSN1S1-Intron6-R TGCATTAAGTGTCTCCT
7	CD-CSN1S1-Intron6-Fb GCAAGGAGACACTTAATGCAAG	CD-CSN1S1-Intron7-RZW TCAGTAGAATTCATCAG
8	CD-CSN1S1-Intron7-F TTTGGATCTGGATTCTCTACCA	CD-CSN1S1-Intron8-R TGAATTAGAGATATCTG
9	CD-CSN1S1-Intron8-F GGTTATTTACATTGATTTATCACAACA	CD-CSN1S1-Intron9-R GGCTATAATAGTCCTGG
10	CD-CSN1S1-Intron9-F ATAAAAAGGGCCCAAGTGT	CD-CSN1S1-Intron10-R TGGGCAGCACAGATAG
11	CD-CSN1S1-Intron10-F CCCAGGAATTTGTGGCTAAA	CD-CSN1S1-Intron12-R ACCACGTGCTGCTAAAA
12	CD-CSN1S1-Intron12-F ATCCTCGGTCAGAAAGCACA	CD-CSN1S1-Intron13-R ACAGCACATTAGTCTCA
13	CD-CSN1S1-Intron13-F TGCTGGTGTTTAAAGTCCAAG	CD-CSN1S1-Intron16-R TCAGTTGGTTTCTTCACA
14	CD-CSN1S1-Intron16-FZW TTTTAAATAATGCAATTGGCAA	CD-CSN1S1-Intron17-RZW AATCATTCTTTCAAAGG
15	CD-CSN1S1-Intron17-FZW TAGCAAGACTTTTCCATGTAAT	CD-CSN1S1-Intron18-RZW ATTCAAGTAATTCTCTA
16	CD-CSN1S1-Intron18-FZW AGAGTCAATAACTCTTGTGATG	CD-CSN1S1-Intron19-RZW GGTACCAGAAATTATTA
17	CD-CSN1S1-Intron19-F CCTTTGTGAGCTGCATCGTAAG	CD-CSN1S1-3'UTR-R CTCTGTTCCCACACCTT
18	CD-CSN1S1-mRNA-F CTTCTTCCCAGTCTTGGGTTC	CD-CSN1S1-mRNA-R ATAGGCGTGGAGGAGA
19	CD-CSN1S1-Ex15-F GAGCCTGTGAAAGTAGTGA	CD-CSN1S1-Ex17-R CCTTCAGGGGTGTCGTA
20	CD-CSN1S1-Ex4-F TGAACCAGACAGCATAGAG	CD-CSN1S1-Ex6-R CTAAACTGAATGGGTGA



A typical PCR reaction mix (50 μ l) comprised 50 ng of gDNA/total cDNA, 1 \times PCR Buffer (Promega, Madison, WI, USA), 2.5-mM MgCl₂, 5 pmol of each primer, dNTPs each at 200 μ M, and 1 U of Taq DNA Polymerase (Promega). PCR was performed under the following thermal conditions: 95 $^{\circ}$ C for 4 min, 35 cycles at 95 $^{\circ}$ C for 45 s, annealing temperature (x° C)

according to the amplicon (Table 1) for 45 s, 72° C for 60 s, and the final extension at 72 °C for 5 min. The amplified products were analyzed by electrophoresis on 1.5 % agarose gel in 0.5× TBE buffer and stained with Midori Green Advance (Nippon Genetics). The amplified fragments were afterward sequenced using a Big Dye Terminator sequencing kit v.1.1 (Applied Biosystems, Forster City, CA, USA).

Development of a DNA-based test

In order to screen for an identified non-synonymous nucleotide substitution in exon 5 (c.150G > T GenBank ID JF429138) of camel *CSN1S1* on DNA level, a polymerase chain-restriction fragment length polymorphism (PCR-RFLP) method has been established using *SmlI* (New England Biolabs, München, Germany). PCR product no. 20 (Table 1) containing exon 4 to exon 6 was digested in a 10- μ l reaction mixture with *SmlI* (55 °C–180 min). Digested fragments were separated by electrophoresis in a 3 % agarose gel and visualized after staining in Midori Green Advance with ultraviolet light.

Identification of IgE-binding epitopes and bioactive peptides

To determine peptides containing IgE-binding epitopes and differences between the genetic variants after digestion, potential cleavage sites were predicted using the *in silico* ExPASy tool, Peptide Cutter (http://web.expasy.org/peptide_cutter/). Enzymes chymotrypsin (high-specificity) and trypsin were selected. Resulting peptides were compared with epitopes described in bovine α_{s1} -casein (Chatchatee et al. 2001, Lisson et al. 2014). For identification of bioactive peptides, the amino acid sequences of α_{s1} -casein A, C, and D were digested *in silico* (<http://www.uwm.edu.pl/biochemia/>) using the following enzymes: pepsin, trypsin, chymotrypsin A, chymotrypsin C, pancreatic elastase I, pancreatic elastase II, oligopeptidase B, and oligopeptidase II.

Statistical analyses

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 (χ^2) test was performed.

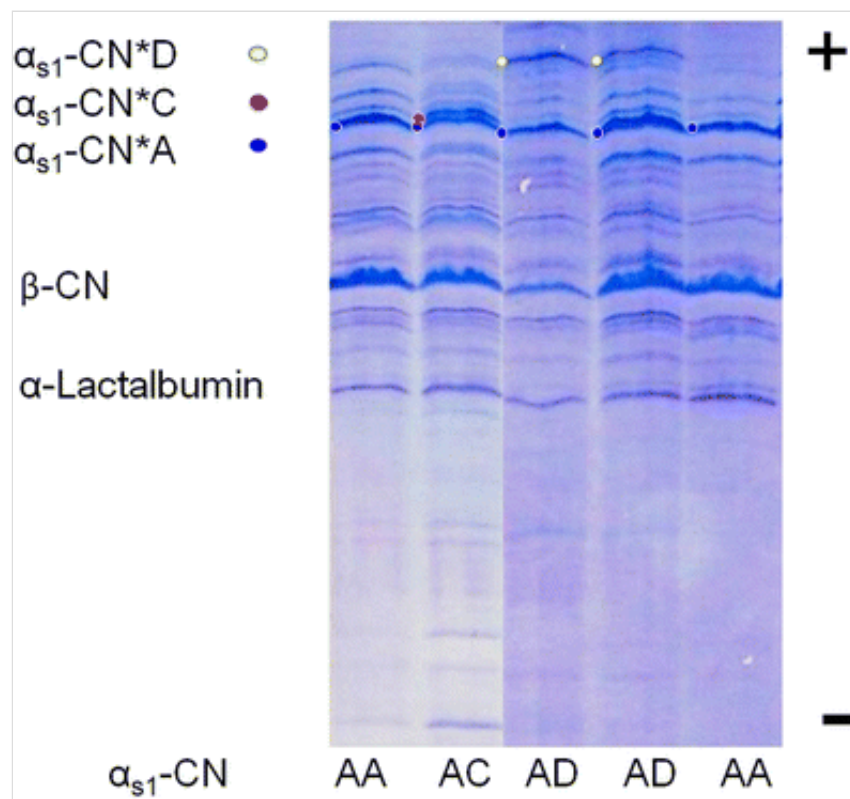
Results and discussion

IEF and allele frequencies

Simultaneous typing of camel milk protein using isoelectric focusing (IEF) revealed polymorphic protein pattern named α_{s1} -CN in the most acidic region of the gel (Fig. 2). Three variants, A, C, and D, were observed in this fraction. Each variant consists of one major and two more alkaline-located minor bands. α_{s1} -casein A revealed the least acidic apparent isoelectric point (pI), and C was just focused in its acidic side, whereas D revealed the most acidic pI. Nomenclature was done considering Kappeler et al. (1998). Due to this, α_{s1} -CN C and α_{s1} -CN D seem to be new variants which are clearly separated by IEF. This shows once more that screening of milk samples still offers the possibility to identify further protein variants (Giambra et al. 2010) as starting point for additional milk protein analyses. Screening of all 193 camel milk samples by IEF revealed dominance of α_{s1} -CN A in comparison to α_{s1} -CN C and α_{s1} -CN D. The major allele A revealed frequencies ranging from 0.750 to 0.902 among Shanbali and Arabi Kahali ecotypes, respectively. While α_{s1} -CN C was found as 0.111, 0.100, 0.078, and 0.071 in Lahaoi, Shanbali, Arabi Kahali, and Arabi Gharbawi, respectively. Allele D revealed the lowest frequencies in all populations except in Shanbali (Table 2).

Fig. 2

Isoelectric focusing of milk samples from *Camelus dromedarius* and the demonstration of different α_{s1} -casein (CN) genotypes. Each variant is characterized by a main (marked and used for phenotyping) and up to three minor bands

**Table 2**

Genotypes and allele frequencies of camel α_{s1} -CN in different ecotypes from different locations of Sudan

No.	Location	Ecotype	N	Genotype frequencies (%)			Allele frequencies		
				AA	AC	AD	A	C	D
1	El Shuak	Lahaoi	81	58.0	22.2	19.8	0.790	0.111	0.099
2	El Obied	Shanbali	40	50.0	20.0	30.0	0.750	0.100	0.150
3	Omdurman	Arabi Khali	51	80.4	15.7	3.9	0.902	0.078	0.020
4	Nyala	Arabi Gharbawi	21	76.2	14.3	9.5	0.881	0.071	0.048

Sequencing of cDNA and genomic DNA

In this study, we found in total 20 exons confirming the results of Pauciullo and Erhardt (2015). Sequencing of the complete coding sequence of camel *CSN1S1* of IEF-pretyped α_{s1} -CN A and D DNA samples showed full sequence similarity to α_{s1} -CN A of Kappeler et al. (1998). Samples with α_{s1} -CN C are characterized by a non-synonymous G > T-

SNP (c.150G > T; GenBank ID JF429138) resulting in the amino acid substitution p.Glu30 > Asp30 in the mature protein. *CSN1S1**C and α_{s1} -CN B (Kappeler et al. 1998) are both characterized by p.Glu30 > Asp30 in the deduced mature protein sequence and are differing by missing or non-missing of exon 18, respectively. Therefore, we postulate a new genetic variant *CSN1S1**C, coding for 207 AAs like *CSN1S1**A, and not for 215 AAs like α_{s1} -CN B.

Removal of intron sequences from pre-mRNA is carried out by the spliceosome machinery, which recognizes specific sites (donor splice site, branch point, polypyrimidine tract, and acceptor splice site) in a complex molecular mechanism. Any deviation from consensus can result in an overall decreased affinity for the spliceosome (Clark and Thanaraj 2002). In order to verify the presence of these sites, the PCR-fragments containing genomic DNA sequence of exon 17 to exon 19 was sequenced. Sequence analyses showed the presence of exon 18 on genomic level in individuals carrying AA, AD, and AC genotypes. No mutation at the splicing donor and acceptor sites in the sequenced animals was identified; however, the following comparison with the genome sequence available in Genbank (acc. no. NW_011591251) showed an insertion of 11 bp (ATTGAATAAAA) at the intron 17 between the branch sequence and the polypyrimidine tract just upstream the exon 18 (Fig. 3).

Fig. 3

Comparison of the DNA sequence spanning from the exon 17 to exon 19 of the dromedary camel *CSN1S1* gene from the genome sequence (*up*) and the same DNA fragment belonging to the variants *CSN1S1* A, C, and D (*down*). Numbering is relative to the gene sequence submitted with acc. no. JF937195. *Dots* indicate identical nucleotides, *dashes* show missing base pairs. Branch sequence is *underlined*, whereas polypyrimidine tract is *double underlined*. Junction of splice are reported in *bold underlined*



The genome sequence might be likely considered as the variant *CSN1S1*B*. In fact, it is characterized by the presence of a thymine at the exon 5 like the variant *CSN1S1*C*, but it does not show the 11-bp insertion at the intron 17. In order to verify whether the 11-bp insertion could affect the secondary structure of the pre-mRNA, both sequences (from the genome and the allele object of the present study) underwent computation analysis by RNAfold server (<http://rna.tbi.univie.ac.at/cgi-bin/RNAfold.cgi>).

The result evidenced that the insertion of 11 bp negatively affect the secondary structure of the pre-mRNA for the investigated alleles (Fig. 4). In fact, the polypyrimidine tract and the following exon 18 appear both extremely hairpin looped and coiled (Fig. 4), whereas the same regions in the pre-mRNA of the putative allele B are well extended (Fig. 5).

Fig. 4

a Secondary structure prediction of the pre-mRNA generated by the transcription of the DNA fragment spanning from the exon 17 to exon 19 of the dromedary camel *CSN1S1* gene alleles A, C, and D. **b** Detail of the structure with relative indication of branch point and 11-bp insertion. The polypyrimidine tract and exon 18 appear hairpin looped and coiled

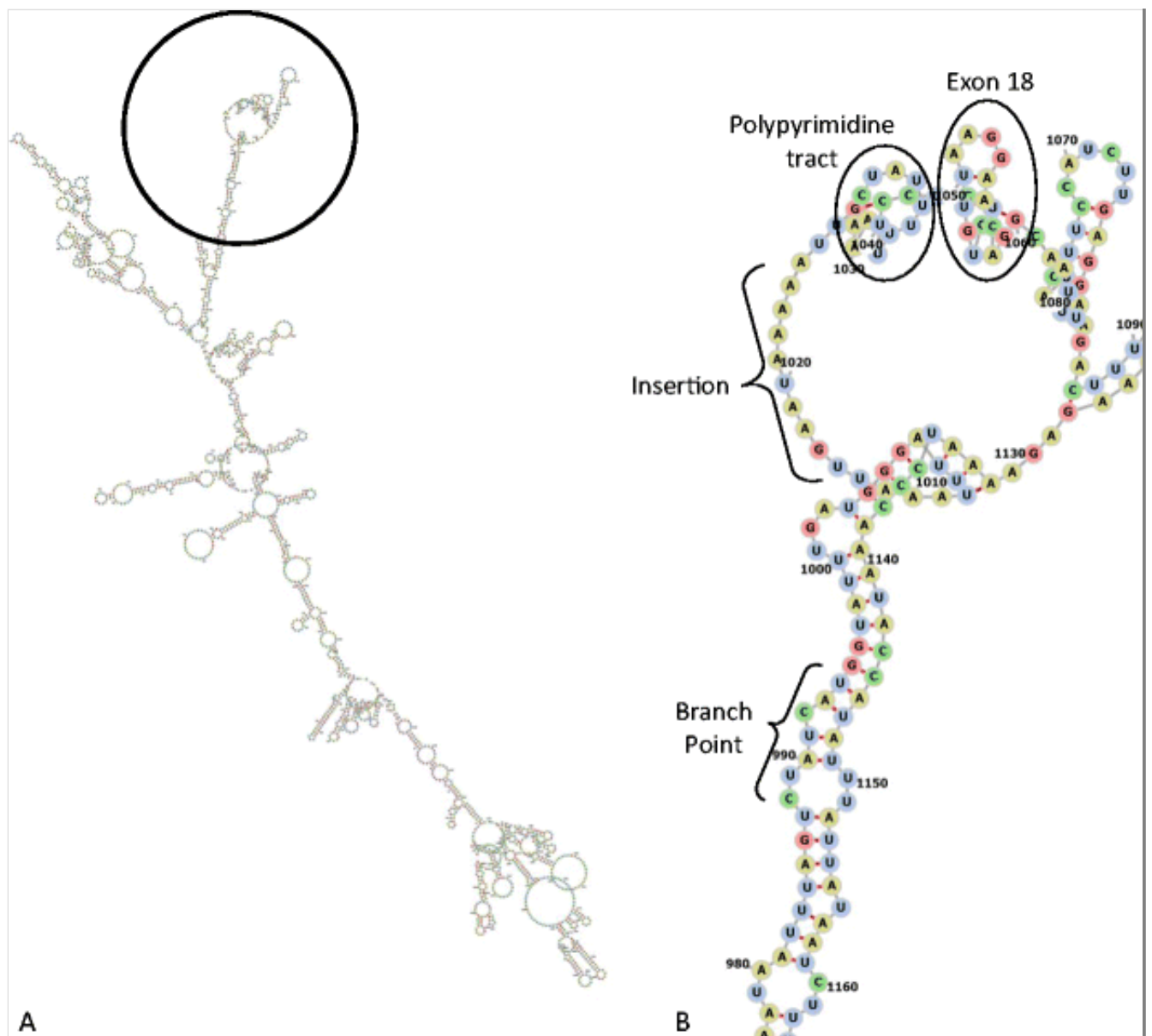
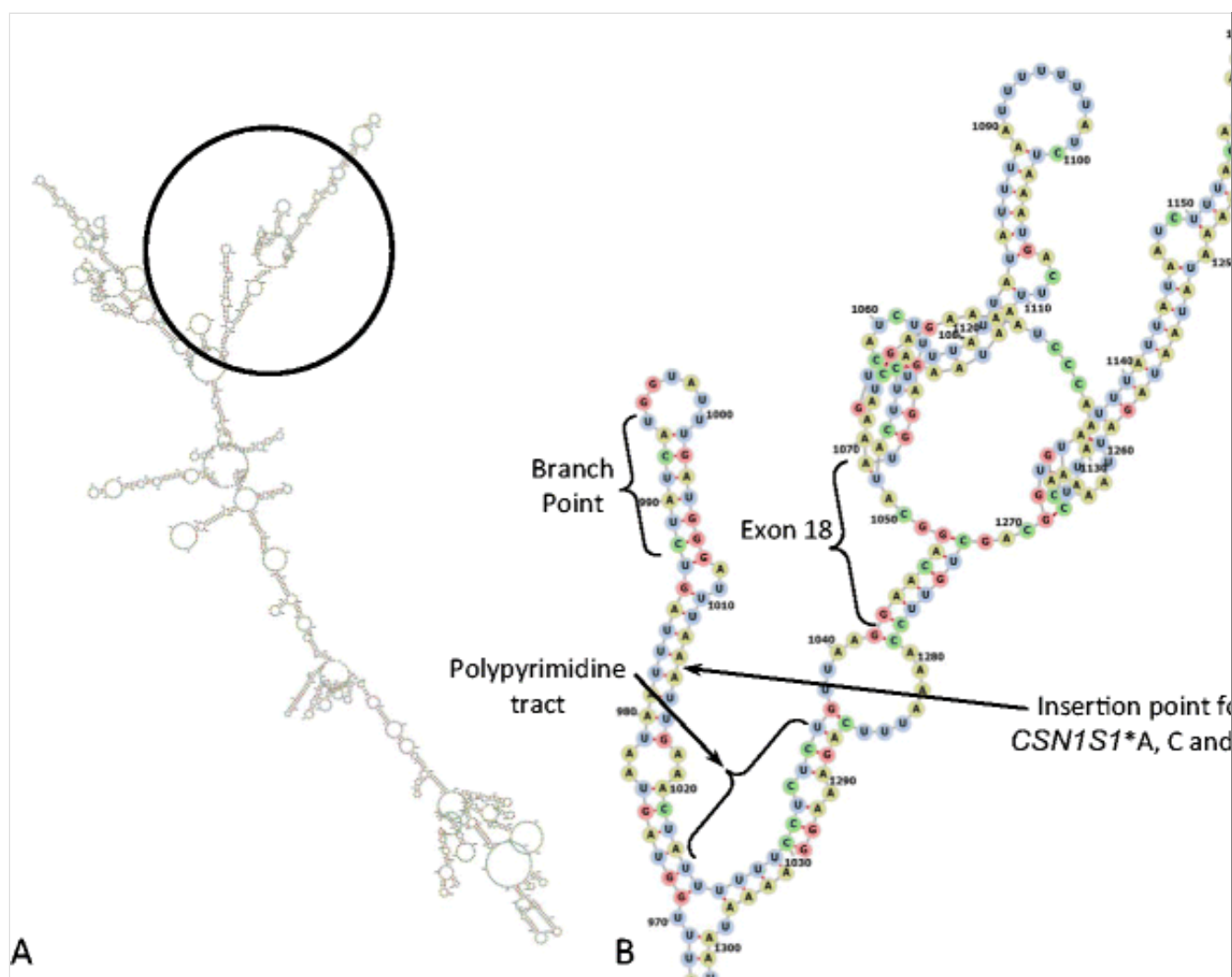


Fig. 5

a Secondary structure prediction of the pre-mRNA generated by the transcription of the DNA fragment spanning from the exon 17 to exon 19 of the allele B for the dromedary camel *CSN1S1* gene. **b** Detail of the structure with relative indication of branch sequence and insertion point. The polypyrimidine tract and exon 18 appear extended



Alterations of the polypyrimidine tract have been shown to inhibit the 5' cleavage reaction, binding of the splicing factor U2AF and the U2snRNP, and splicing complex assembly (Frendeway and Keller 1985).

Specifically, exon skipping as a consequence of the formation of mRNA secondary structure has been already suggested for the goat α_{s1} -casein (Leroux et al. 1992) and human β -casein (Martin and Leroux 1992). Of particular interest is the proposal of Leroux et al. (1992) which is similar to that described here. They suggest that the skipping of exon 9 of the goat α_{s1} -casein gene is caused by an 11-bp insertion within intron 9 base-pairing with the donor site. Therefore, it seems likely that also the variants *CSN1S1**A, *CSN1S1**C, and *CSN1S1**D (all characterized by this 11-bp insertion) were affected by the same alteration of the spliceosome machinery and, therefore, preferentially splice-out of the exon 18.

The lack of the exon 18 is the usual form within camel *CSN1S1* mRNA and α_{s1} -CN protein, as frequencies of the alleles without this exon seems to be higher in *C. dromedarius*. A similar event also characterizes the α_{s1} -CN of South American camelids (Pauciullo and Erhardt 2015). In particular, the

llama α_{s1} -cDNA showed two variants, with and without the exon 18 coding for the same octapeptide (EQAYFHLE) which differentiates the variants A and B found by Kappeler et al. (1998). However, in llama, the shorter variant represents only 26.4 % of the α_{s1} -cDNA transcripts, whereas in the present study, there was no simultaneous occurrence of cDNA containing and missing the exon 18. Therefore, further studies need to clarify such a difference between old world and new world camelids, at least in regard of the efficiency of transcription for short and full-length mRNAs.

AQ3

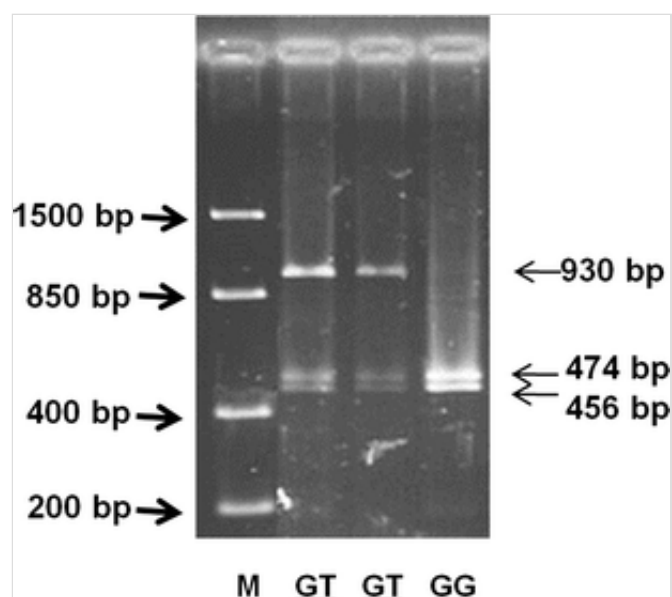
For *CSN1S1**D, we could finally clarify the same genetic event characterizing the alleles *CSN1S1**A and C; however, no further investigation was possible as the necessary mRNA sample was not available.

DNA-based test

Using genomic DNA, a PCR-RFLP test was established for typing the g.942G > T SNP (GenBank ID JF429140) characteristic for the new *CSN1S1**C variant (Fig. 6) The test developed confirmed the IEF and can therefore be used for typing camel *CSN1S1* variability independent of age, gender, and lactation stage. This can be useful for broader analysis of camel milk protein variability and is a requirement for further analyses concerning associations between milk protein variability and milk performance traits in camel, as already done for cattle, goat, and sheep (Martin et al. 2002; Caroli et al. 2009; Giambra et al. 2011).

Fig. 6

PCR-RFLP for genotyping camel *CSN1S1**A and C. M = marker, CSN1S1 AC (GT 930, 474, and 456 bp) CSN1S1 AA (GG, 474 and 456 bp)



IgE-binding epitopes and bioactive peptides

A comparison of the identified peptides after *in silico* digestion of α_{s1} -casein A, B, and C with IgE-binding epitopes described in bovine α_{s1} -casein showed that IgE-binding epitopes f24-43 (NKRKILELAVVSPIQFRQEN) and f153-168 (TQEQAYFHLEPFQQFF) did not resist digestion in intact form and that the amino acid substitutions and deletion (underlined) influence the resulting peptide pattern. Thus, peptide f28-40 (ILELAVVSPIQFR) occurred in all three variants. Peptide f151-159 (VVTQPFQQF) was only identified in variants A and C, whereas peptides f151-158 (VVTQEQAY) and f160-167 (HLEPFQQF) were found in variant B. These peptides comprise major parts of the IgE-binding epitope f153-168. To determine the allergenic potential of these peptides, microarray immunoassay using synthesized peptides and sera from humans with cow milk allergy are currently in progress. A total of 83 bioactive peptides with ACE inhibitory (53), vasoactive/glucose uptake stimulating (8), antioxidative (12), inhibitory (9), and hypotensive (1) activities were found in α_{s1} -casein A, B, and C (Table 3). Due to the deletion of eight amino acids, two further ACE-inhibitory and antioxidative peptides (fAY and fHL) were exclusively identified in α_{s1} -casein B. This confirms that genetic variants of camel casein are a source of different bioactive peptides as already described in cattle (Weimann et al. 2009) and reveals the additional potential of camel milk protein variants for human health.

Table 3

Predictive bioactive peptides from α_{s1} -casein variants A, B, C, and D of camel (in parenthesis is the number of identified peptides) according to the BIOPEP database

(Minkiewicz et al. 2008)

Effect	Peptide sequence	α s1-casein variant
ACE inhibitor (53)	VMP, RL, IR, RY, LY, VF, FP, PR, VSP, YL, YP, PL, AW....	A, C, D,
Vasoactive/glucose uptake stimulating (8)	SSS, VL, IL, LI, LL, EE, SE, YL	A, C, D
Antioxidative (12)	LH, LY, AH, EL, WY, WYY, LHR, VKL, VKV, KD, IR, LK	A, C, D
DPP IV inhibitor (9)	PP, MP, VA, LA, FP, LL, VV, IR, LA	A, C, D
Hypotensive (1)	IR	A, C, D
ACE inhibitor (2), antioxidative (2)	AY, HL	B

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Compliance with ethical standards

Statement of animal rights The manuscript does not contain clinical studies or patient data.

Conflict of interest The authors declare that they have no competing interests.

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