A hypomorphic mutation in the mouse Csn1s1 gene generated by CRISPR/Cas9 pronuclear microinjection

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Abstract. Caseins are major milk proteins that have an evolutionarily conserved role in nutrition. Sequence variations in the casein genes affect milk composition in livestock species. Regulatory elements of the casein genes could be used to direct the expression of desired transgenes into the milk of transgenic animals. Dozens of casein alleles have been identified for goats, cows, sheep, camels and horses, and these sequence variants are associated with altered gene expression and milk protein content. Most of the known mutations affecting casein genes' expression are located in the promoter and 3'-untranslated regions. We performed pronuclear microinjections with Cas9 mRNA and sgRNA against the first coding exon of the mouse Csn1s1 gene to introduce random mutations in the α-casein (Csn1s1) signal peptide sequence at the beginning of the mouse gene. Sanger sequencing of the founder mice identified 40 mutations. As expected, mutations clustered around the sgRNA cut site (3 bp from PAM). Most of the mutations represented small deletions (1–10 bp), but we detected several larger deletions as well (100–300 bp). Functionally most mutations led to gene knockout due to a frameshift or a start codon loss. Some of the mutations represented in-frame indels in the first coding exon. Of these, we describe a novel hypomorphic Csn1s1 (Csn1s1C4-SinsTCC) allele. We measured Csn1s1 protein levels and confirmed that the mutation has a negative effect on milk composition, which shows a 50% reduction in gene expression and a 40–80% decrease in Csn1s1 protein amount, compared to the wild-type allele. We assumed that mutation affected transcript stability or splicing by an unknown mechanism. This mutation can potentially serve as a genetic marker for low Csn1s1 expression.

Key words: casein; CRISPR; pronuclear microinjection; hypomorphic mutation.

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Гипоморфная мутация гена Csn1s1 мыши, полученная пронуклеарной микроинъекцией CRISPR/Cas9

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Аннотация. Казеины – это основные молочные белки, которые играют важную эволюционно консервативную роль в питании. Вариации последовательности казеиновых генов влияют на состав молока у животных. Регуляторные элементы казеиновых генов можно использовать для управления экспрессией желаемых трансгенов в молоке трансгенных животных. Десятки аллелей казеина идентифицированы у коз, коров, овец, верблюдов и лошадей, и эти варианты связаны с измененной экспрессией генов и содержанием молочного белка. Большинство известных мутаций, влияющих на экспрессию генов казеина, находится в промоторных и 3'-нетранслируемых областях. Мы выполнили пронуклеарные микроинъекции с мРНК Cas9 и sgRNA против первого кодирующего экзона гена Csn1s1 мыши, чтобы ввести случайные мутации в последовательность сигнального пептида α-казеина (Csn1s1) в начале гена мыши. Секвенирование мышей-основателей выявило 40 мутаций. Как и ожидалось, мутации группировались вокруг сайта разреза sgRNA (3 п. н. от PAM). Большинство мутаций представляют собой небольшие делеции (1–10 п. н.), но мы также обнаружили несколько более крупных делеций (100–300 п. н.). Функционально большинство мутаций приводило к нокаутам генов из-за сдвигов рамки считывания или потери стартовых кодонов. Некоторые из мутаций представлены инделями в рамке считывания в первом кодирующем экзоне. Из них мы описываем новый гипоморфный аллель Csn1s1 (Csn1s1C4-SinsTCC). Мы измерили уровни белка Csn1s1 и подтвердили, что мутация отрицательно влияет на состав молока, который показывает снижение экспрессии гена на 50% и уменьшение количества белка Csn1s1 на 40–80% по сравнению с аллелем дикого типа. Мы предположили, что мутация влияет на стабильность транскрипта или сплайсинг по неизвестному механизму. Эта мутация потенциально может служить генетическим маркером низкой экспрессии Csn1s1.

Ключевые слова: казеин; CRISPR; пронуклеарная микроинъекция; гипоморфные мутации.

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Introduction
Caseins are major milk proteins that have an evolutionary conserved role in nutrition (Rijnkels et al., 2003). Casein locus has been studied for a long time to understand the principle of gene regulation and hormone-induced expression (Rijnkels et al., 2013; Dos Santos et al., 2015; Lee et al., 2017). At the same time, regulatory elements of the casein genes could be used to direct the expression of desired transgenes into milk of transgenic animals (Houdebine, 2009; Kim et al., 2015). This strategy is frequently employed to create “enriched” milk with improved composition (An et al., 2012; Wan et al., 2012; Yuan et al., 2014), or to achieve large scale production of recombinant human proteins in mouse models (Wu et al., 2012; Burkov et al., 2013; Qian et al., 2014) and in livestock species (Kalds et al., 2019). Milk-specific signal peptides are used in biotechnology to enhance recombinant protein secretion during lactation (Yu et al., 2006; Liu et al., 2014; Lu et al., 2019).

During breeding, casein genes acquired many sequence variations that can lead to altered gene expression and are characteristic of some goat and cow breeds (Yue et al., 2011; Fomichev et al., 2012; Guan et al., 2019). Many of these sequence variants could be used as markers for breeding. In dairy industry casein composition is an important milk trait directly influencing quality of dairy products (Sanchez et al., 2018; Cieslak et al., 2019). Hypomorphic casein mutation are also associated with other traits such as litter size (Wang et al., 2018). For example, K. Wang with colleagues showed that 11 bp del in the intron 8 of the goat Csn1s1 negatively affects the expression of the gene (Wang et al., 2018). Other known hypomorphic mutations are located in the promoter and 3’-untranslated regions (UTRs) of casein genes (Huang et al., 2012; Noce et al., 2016).

Novel CRISPR methods greatly facilitate genome editing in farm animals (Kalds et al., 2019), including targeted transgene integration (Park et al., 2017) and mutation modeling (Li et al., 2017; Zhou et al., 2019). The latter approach has potential to explain the molecular mechanism of how hypomorphic mutations affect milk proteins. In this report, we used CRISPR/Cas9 to create a set of mutations within a signal peptide sequence of the α-casein (Csn1s1) gene in mice. One of the mutants was chosen to study effects of a small in-frame insertion on the Csn1s1 expression during lactation.

Materials and methods
Generation and genotyping of the Csn1s1 mutant mice. In vitro transcription and purification of the gRNA were performed with MEGAscript™ T7 Transcription Kit (Thermo Fisher Scientific) and MEGAclear™ Transcription Clean-Up Kit (Thermo Fisher Scientific) according to the manufacturer’s protocol. Cas9 mRNA (GeneArt™ CRISPR Nuclease mRNA) was purchased from Thermo Fisher Scientific. 50 ng/μL Cas9 mRNA and 25 ng/μL gRNA (5’-GGGAGGATGAGAGGATTTCA-3’) were mixed in RNase-free water, backfilled into an injection needle with positive balancing pressure (Transjector 5246, Eppendorf) and injected into the cytoplasm of zygotes (C57BL/6 × CBA background). After injections, the embryos were cultured for 1 hour in drops of M16 medium at 37 °C and an atmosphere of 5% CO₂. The viable microinjected zygotes were transplanted the same day into oviducts of pseudopregnant CD-1 females (0.5 days after coitus). Isoflurane inhalation anesthesia was applied in these experiments.

Mutations were detected using PCR and Sanger sequencing of the target region of the Csn1s1 exon 2 (Supplementary 1). Primers for PCR were as follows: 5’-GCGGACATACTAAGCATTCTTATGCT-3’ (forward primer), 5’-TGAATGGAGTTTAGATTGGACA-3’ (reverse primer). Selected male founders were crossed with C57BL/6 females. For mutation c.4-5insTCC described in this paper, founder male was crossed with two F₁ heterozygous daughters. Offspring was genotyped and two sibling females were selected for each group (wt, heterozygous or homozygous mutation) for further analysis.

All experiments were conducted at the Centre for Genetic Resources of Laboratory Animals at the Institute of Cytology and Genetics, SB RAS (RFMEFI61914X0005 and RFMEFI61914X0010). All experiments were performed in accordance with protocols and guidelines approved by the Animal Care and Use Committee Federal Research Centre of the Institute of Cytology and Genetics, SB RAS operating under standards set by regulations documents Federal Health Ministry (2010/708n/RF), NRC and FELASA recommendations. Experimental protocols were approved by the Bioethics Review Committee of the Institute of Cytology and Genetics, SB RAS.

Droplet digital PCR analysis. Total cellular RNA was extracted from mouse mammary glands at day 8 of lactation using TRI Reagent (Sigma-Aldrich). 1 μg of total RNA was used to generate cDNA in a 20 μl reaction using RevertAid RT Kit (Thermo Fisher Scientific) with random hexamer primers according to the manufacturer’s instructions. Droplet Digital PCR (ddPCR) was performed using a QX100 system (Bio-Rad) with primers and probes specific for the Csn1s1 and Csn2 mouse transcripts (Supplementary 2). The primers and probes sequences were as follows: 5’-TGATGTGGGAGGCAGGCACCTGG-3’ (Csn1s1 forward primer), 5’-TCCTTGGAGACGATAATGGGACTT-3’ (Csn1s1 reverse primer), 5’-HEX-CCAGCTTCTGTCGACCCCTCACA-BHQ2–3’ (Csn1s1 probe), 5’-AGGACCTTGGCAGCACAGTGAAGG-3’ (Csn2 forward primer), 5’-ATGGTCACAGACAGCTCTCCTGACTGA-3’ (Csn2 reverse primer), 5’-FAM-ATCTTGCCGCCTTTTGCGCTTGC-BHQ2–3’ (Csn2 probe). ddPCR reactions were set in 20 μl volumes containing 1× ddPCR™ Supermix for Probes (no dUTP), 900 nM primers and 250 nM probes, and 1 μl of 5000-fold diluted cDNA. ddPCR reactions for each sample were performed in duplicates. PCR was conducted according to the following program: 95 °C for 10 min, then 40 cycles of 95 °C for 30 s and 61 °C for 1 min, with a ramp rate of 2 °C per second, and a final step at 98 °C for 5 min. The results were analyzed using QuantaSoft software (Bio-Rad). Concentrations of cDNA copies of Csn1s1 and Csn2 were derived from ddPCR and relative expression of Csn1s1 to Csn2 was calculated for each animal.

Milk and mammary gland protein analysis. Milk was obtained from narcotized female mice at day 8 of lactation after oxytocin administration (Uusi-Oukari et al., 1997). The milk was collected with a pipette attached to an aspiration

3 Supplementary materials 1 and 2 are available at: http://www.bionet.nsc.ru/vogis/download/pict-2021-25/appx8.pdf
Results

Generation of the Csn1s1 mutant mice

We performed pronuclear microinjections with Cas9 mRNA and sgRNA against the first coding exon of the mouse Csn1s1 gene (Fig. 1, a). Cas9-induced mutations in this region could potentially affect signal peptide coding sequence and lead to altered milk composition. We screened founder mice by Sanger sequencing and identified multiple random mutations at the Cas9 cut site (presented in Supplementary 1). The sgRNA targeted the Csn1s1 site with high efficiency as we detected 41 mutant alleles, 4–5 mosaic alleles (additional background signal) and only 4–5 wild-type alleles in 20 founder mice (~90 % allele mutation efficiency). As expected, mutations clustered around the sgRNA cut site (3 bp from PAM). Most of the mutations represented small deletions (1–10 bp), but we detected several larger deletions as well (100–300 bp). Of note, some of the unique mutation variants had increased incidence rate. For example, 12 bp deletion (GAAACTCCTCAT) arose independently four times and another 10 bp deletion (CCATGAAA) – three times (see Supplementary 1). We suspect this bias towards some variants is caused by microhomology-mediated end-joining (MMEJ), since these two mutations are flanked with 3 and 2 similar nucleotides (CAT and CC, respectively) (see Supplementary 1). Although mutations were mostly deleterious for the gene expression and led to a Csn1s1 knockout (KO) by frameshift, several in-frame mutation variants resulted in subtle changes in signal peptide coding sequence without KO (see Supplementary 1). We selected one of the mutants, tagged Csn1s1<sup>1-4.5insTCC</sup>, which had a 3 bp insertion following the start codon (see Fig. 1, b). To study gene expression and milk composition we chose 6 female siblings (2 wild-type, 2 heterozygotes, 2 homozygotes) from the Csn1s1<sup>1-4.5insTCC</sup> line for milk and mammary glands collection.

Mutation c.4-5insTCC leads to reduced expression of the Csn1s1 gene

We estimated the Csn1s1 gene expression in mammary glands of wild-type and mutant mice at day 8 of lactation using droplet digital PCR (ddPCR). We selected Csn2 (β-casein) as a reference gene as it has a similar expression profile in mammary gland (see Supplementary 2). ddPCR analysis revealed that Csn1s1:Csn2 ratio was roughly 1:3 (0.338) in wild-type siblings (Fig. 2), which is in agreement with published data (Yamaji et al., 2013). Heterozygous and homozygous Csn1s1<sup>1-4.5insTCC</sup> siblings showed lower Csn1s1 expression with ratios around 1:4 (0.248) and 1:6 (0.168), respectively (see Fig. 2), compared to wild-type siblings. We also used females from the parental strains C57BL/6 and CBA as controls for normal caseins level (see Fig. 2). In rare cases, Cas9 activity can provoke rearrangements near the target locus.

![Fig. 1. Generation of the Csn1s1 mutant mice by CRISPR/Cas9 pronuclear microinjection.](image-url)

Fig. 1. Generation of the Csn1s1 mutant mice by CRISPR/Cas9 pronuclear microinjection.

- a – sequence of the first coding exon (63 bp) of the Csn1s1 gene (exon 2, NM_007784.3). Black arrowhead indicates Cas9 cleavage site. Complementary 20 bp sgRNA sequence with PAM is shown above; b – sequence of the mutated Csn1s1 exon with a 3 bp insertion in the second codon (framed). Start codon (ATG) is highlighted.
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We sequenced mutation-flanking regions, including the Csn1s1 promoter and surrounding introns (2.3 kb + 1.1 kb) from homozygous mice (data not shown). We also sequenced top two off-targets for the Csn1s1 sgRNA in the F0 founder (Chr1:24388301; Chr4:140044179). No mutations were found in the examined sequences. Thus, we could confirm that the 3 bp in-frame insertion in the first coding exon led to a 30–50 % decrease in the Csn1s1 gene expression.

**Discussion**

We report a novel in-frame Csn1s1 hypomorphic mutation that leads to a 50 % gene expression decrease in mice. In most cases, mutation effect is tied to disruption of a regulatory element (enhancer, promoter, UTR, miRNA site) (Hogg, Harries, 2014). Frame-shifting indels in the coding sequence could initiate transcript surveillance pathway called non-

**Fig. 2.** Droplet digital PCR analysis of the Csn1s1 and Csn2 expression in mammary gland. Data presented as ratios of Csn1s1/Csn2 transcripts. Error bars – standard deviations. Bold numbers at the bottom represent mean ratios for two females of each group. KO – Csn1s1 knockout female; WT, Het, Hom – siblings with corresponding mutation genotype; C57BL/6, CBA – females from the inbred strains. Statistics: one-way ANOVA.

**Fig. 3.** Detection of the Csn1s1 protein in milk and mammary glands of the mutant mice.

We measured Csn1s1 protein levels and confirmed that mutation has a negative effect on milk composition. We collected milk and mammary glands from lactating females at the day 8 of lactation. Csn1s1 knockout mouse was taken from another experiment (“KO”) as additional control for the Csn1s1 levels. Separation of milk proteins on 12 % SDS-PAGE resulted in a typical band pattern for mouse milk (Fig. 3, a). In wild-type mice, Csn1s1 represents a major protein fraction and corresponds to approx. 30 % of total milk protein (Kolb et al., 2011). In homozygous mutants, loss of Csn1s1 could be observed both at the Coomassie-stained gel (see Fig. 3, a) and after western blotting (see Fig. 3, b). Csn1s1 levels fell down to 30 % in homozygotes, both for milk and for mammary gland lysates (see Fig. 3, c). However, exact ratios varied depending on the control protein band used for calculations. For instance, we performed the following calculations for the milk Csn1s1: Csn1s1 (gel) vs total protein (gel) – 40 % reduction; Csn1s1 (gel) vs albumin (gel) – 70 % reduction; Csn1s1 (western blot membrane) vs total protein (gel) – 80 % reduction. This effect was even more pronounced in mammary gland lysates (intracellular casein levels): Csn1s1 (western blot membrane) vs total gel – 80 % reduction.

**Table 3**

| Condition       | Csn1s1 (gel) vs total (gel) | Csn1s1 (gel) vs albumin (gel) | Csn1s1 (WB) vs total (gel) | Csn1s1 (WB) vs total (gel) |
|-----------------|-----------------------------|--------------------------------|---------------------------|----------------------------|
| KO              | 0.338                       | 0.248                          | 0.168                     | 0.319                      |
| WT              | 0.251                       | 0.246                          | 0.182                     | 0.273                      |
| Het1            | 0.154                       | 0.182                          | 0.289                     | 0.258                      |
| Hom1            | 0.400                       | 0.348                          | 0.308                     | 0.352                      |
| Hom2            | 0.351                       | 0.326                          | 0.298                     | 0.348                      |
| C57BL/6         | 0.214                       | 0.196                          | 0.205                     | 0.223                      |
| CBA             | 0.182                       | 0.168                          | 0.154                     | 0.130                      |

Error bars – standard deviations. Bold numbers at the bottom represent mean ratios for two females of each group. KO – the Csn1s1 knockout female; WT, Het, Hom – siblings with corresponding mutation genotype; C57BL/6, CBA – females from the inbred strains. Statistics: one-way ANOVA.
sense-mediated mRNA decay (NMD) (Popp, Maquat, 2016; Lindeboom et al., 2019). Alternatively, in-frame mutations can lead to exon removal by alternative splicing (Mucaki et al., 2020; Thompson et al., 2020). Essentially, exon skipping could be promoted by internal exon splicing enhancers and suppressors (ESEs and ESSS) which are hard to predict (Sterne-Weiler, Sanford, 2014; Tuladhar et al., 2019), unlike typical splice site mutations (Cartegni et al., 2002). In our mutant mice, promoter had no alterations as the mutation happened in the coding sequence, quite far from a transcription start site. We assumed that it affected transcript stability or splicing by unknown mechanism. It should also be noted that mutated Csn1s1 protein was still secreted in milk, thus the function of N-terminal signal peptide was not critically affected by the mutation.

**Conclusion**

We demonstrated that CRISPR/Cas9 approach could be conveniently exploited to induce a spectrum of mutations in the Csn1s1 gene either by random mutagenesis, or, ideally, by a set of single-stranded oligo DNA nucleotides (ssODNs). Our results warn that careful examination of the gene’s expression is required in addition to protein analysis.

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