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Mediterranean River Buffalo *CSN1S1* gene: search for polymorphisms and association studies

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Abstract. The aim of the present work was to study the variability at *CSN1S1* locus of the Italian Mediterranean river buffalo and to investigate possible allele effects on milk yield and its composition. Effects of parity, calving season and month of production were also evaluated. Three SNPs were detected. The first mutation, located at position 89 of 17th exon (c.628C>T), is responsible for the amino acid change p.Ser178 (B allele)/Leu178 (A allele). The other two polymorphisms, detected at the positions 144 (c.882G>A) and 239 (c.977A>G) of 19th exon respectively, are silent (3’ UTR).

Associations between the *CSN1S1* genotypes and milk production traits were investigated using 4,122 test day records of 503 lactations from 175 buffalo cows. Milk yield, fat and protein percentages

were analyzed using a mixed linear model. A significant association between the c.628C>T SNP and the protein percentage was found. In particular, the CC genotype showed an average value of about 0.04% higher than the CT and TT genotypes. The allele substitution effect of the cytosine into the thymine was -0.014, with a quite low (0.3%) protein percentage (PP) contribution on total phenotypic variance. A large dominance effect was detected.

Furthermore, a characterization of the *CSN1S1* transcripts and a method based on *MboI*-ACRS-PCR for a rapid genotyping of c.628C>T were provided.

Additional keywords: Mediterranean river buffalo, *Bubalus bubalis*, *CSN1S1*, milk protein percentage, marker assisted selection

Introduction

Historically domestic water buffalo were divided into swamp and river subspecies that differ in morphology, behaviour, and chromosome number (2n=48 and 2n=50, respectively) (Ajmone-Masan P. *et al*, 2013). In particular, the Mediterranean river buffalo, is the only indigenous Italian breed of water buffalo. It is of the River subtype of water buffalo, and is genetically and ecological similar to the buffalo breeds of Hungary, Romania and the Balkan countries. Previously considered to belong to the Mediterranean buffalo group, it was officially recognised as a breed in 2000, based on its long isolation period and lack of interbreeding with other buffalo breeds.

The improvement of animal performances represents a priority for the Italian dairy Buffalo industry in order to fulfill the increasing market demand for mozzarella cheese. Italian buffalo stock consists of approximately 344,000 Mediterranean river buffaloes (<http://faostat.fao.org/>). In 2011, the average milk yield per lactation per buffalo cow (35,963 registered cows in the national herd book) was kg 2,223 with 8.49 and 4.65 % of fat and protein content, respectively (<http://www.aia.it/aia->

website/it/home). The milk is almost completely processed into cheese. A breeding program aimed at improving buffalo milk yield and composition is currently operating in Italy, however the low efficiency of the artificial insemination, the difficulties to detect the oestrus and the variability of its length are among the main causes of a very limited impact on the population (Barile, 2005).

Recent advances of molecular genetics offer the possibility to investigate genomic regions that affect traits of economic importance and to identify genetic polymorphisms useful for marker-assisted selection (MAS) programs. In the last decades, several association studies between milk production traits and markers located in milk protein genes have been carried out in cattle, sheep and goat (Ibeagha-Awemu *et al*, 2008; Martin *et al*, 2002; Mroczkowski *et al*, 2004). In particular, the goat α s1 encoding gene (*CSN1S1*) was found to be highly polymorphic with at least 17 alleles associated with qualitative and quantitative differences for the content of α s1casein (Ramunno *et al*, 2004; Ramunno *et al*, 2005), fat (Grosclaude *et al*, 1994; Chilliard *et al*, 2006), urea level (Schmidely *et al*, 2002; Bonanno *et al*, 2007; Avondo *et al*, 2009), fatty acid profile Chilliard *et al*, 2006) and milk yield (Yue *et al*, 2011). Associations between alleles at *CSN1S1* locus and protein content were also observed for other species as for instance in bovine (Rando *et al*, 1998; Prinzenberg *et al*, 2003; Çardak, 2005) and ovine (Pirisi *et al*, 1999; Wessels *et al*, 2004).

In ruminant species, the *CSN1S1* gene is characterized by an extremely split architecture with 19 exons, many of which (exons 5, 6, 7, 8, 10, 13 and 16) of small size (24 bp) (Ramunno *et al*, 2004; Koczan *et al*, 1991; Calvo *et al*, 2011). In buffalo, the α s1-casein gene codes for a precursor of 214 amino acids with a signal peptide of 15 amino acid residues (Ferranti *et al*, 1998; Sukla *et al*, 2007). Currently, several partial or complete bubaline *CSN1S1* cDNA sequences are available in EMBL (FJ392261; AJ005430; AY948385; EF025981; EF025982; EF025983; DQ111783). The similarity between buffalo and cattle, goat and sheep α s1-casein mRNA sequence is 97.2, 93 and 92.3%, respectively. A similar trend was observed comparing amino acid sequences of these species (Sukla *et al*, 2007).

96 Few polymorphisms have been reported for the *CSN1S1* locus in buffalo. The occurrence of a
97 α s1-casein B genetic variant characterized by a single amino acid substitution (p.Leu178Ser) as
98 consequence of single nucleotide substitutions was found for the first time in Romanian Buffalo breed
99 (Balteanu *et al*, 2008) and confirmed at amino acid level in Mediterranean water buffalo (Chianese *et al*,
100 2009). Furthermore in Indian water buffalo a novel *CNS1S1* allele has been characterized by c.620G>A
101 substitution. It led to a p.Gly192Glu replacement in the peptide chain (Sukla *et al*, 2007).

102 Recently, additional SNPs were detected: a transition c.136G>A at exon 5, leading to a
103 p.Val31Met substitution, a transition c.175A>G at exon 7, leading to a p.Ile44Val substitution in the
104 peptide chain, one SNP (g.218T>C) in intron 5 and three SNPs (g.472G>C; g.547C>T; g.856T>C) in
105 intron 6. In particular, it was reported that the g.472G>C substitution inactivates the intron 6 splice donor
106 site promoting the skipping of exon 6 of the buffalo *CSN1S1* mRNA and, as consequence, triggering the
107 synthesis of a defective protein lacking eight amino acids (Balteanu *et al*, 2013).

108 In recent years, many studies have been carried out for the identification of the genetic
109 polymorphisms at the *loci* coding for the buffalo milk proteins (Cosenza *et al*, 2009a, b; Masina *et al*,
110 2007), and candidate genes responsible for the variation of the quali-quantitative characteristics of the
111 Mediterranean water buffalo milk have been found (Cosenza *et al*, 2007; Pauciullo *et al*, 2010). Efforts
112 in this direction also allowed to find significant associations with traits of economic interest, as milk
113 yield (Pauciullo *et al*, 2012a, b) and milk coagulation properties (Bonfatti *et al*, 2012b).

114 The aim of our work was to study the variability at the Italian Mediterranean river buffalo *CSN1S1*
115 cDNA and to investigate possible associations with milk yield and composition.

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117

118 **Materials and methods**

119 *Sampling*

120 Individual blood samples were collected from 175 Italian river buffaloes randomly chosen and belonging
121 to an experimental herd, located in Salerno province (Southern Italy).

122 In order to characterize the *CSN1S1* transcripts and to detected polymorphisms at this *locus*,
123 individual milk samples were collected from 10 animals at comparable age, in third calving, at 120 days
124 in milking and free of clinical mastitis and randomly chosen in different farms located in the province of
125 Salerno and Caserta (Southern Italy). After collection, milk samples were immediately frozen and kept
126 at -80 °C until analysis. Sampling was carried out in collaboration with the Italian National Association
127 of Buffalo Breeds (ANASB).

128

129 *RNA extraction*

130 Total RNA was isolated from somatic cells (SCC range from 10,000 to 12,000/mL) present in individual
131 milk samples by using NucleoSpin® Extract Kits (Macherey-Nagel). A digestion with 2U of DNase I
132 (Ambion) in 1X DNase buffer was carried out according to the manufacture guidelines at 37°C for 30
133 min followed by the enzyme inactivation at 75°C for 5 min. The quantity, quality, purity, and integrity
134 of RNA after DNase treatment were estimated by means of Nanodrop 2000c spectrophotometer (Thermo
135 Scientific, Barrington, IL) and by electrophoresis on a denaturing agarose gel.

136

137 *Reverse transcription, PCR and cloning*

138 Total RNA was converted into cDNA by reverse transcription using Improm- II Reverse Transcriptase
139 (Promega) with a final volume of 20µl. The reaction was performed using cDNA19R (5'-
140 CAAAATCTGTTACTGCACA - 3'), a reverse primer complementary to nt 327–345 of 19th exon of
141 buffalo *CSN1S1* cDNA sequence (accession number AY948385).

142 The PCR was performed using: cDNA19R and cDNA1F (5' - AACCCAGCTTGCTGCTT - 3'),
143 a forward primer corresponding to nt 1–17 of partial 1st exon of buffalo *CSN1S1* cDNA sequence
144 (accession number AY948385). The PCR reaction mix comprised 20 µl of RT reaction product, 50 mM
145 KCl, 10 mM Tris–HCl, 0.1% Triton X-100, 2 mM MgCl₂, 10 pmol of each primer, dNTPs each at 0.2
146 mM, 5 U of Taq DNA Polymerase (Promega, Madison, WI), with a final volume of 100 µl.

147 The amplification protocol consisted of 39 cycles: the first cycle involved a denaturation step at
148 97 °C for 2 min, an annealing step at 57 °C for 30 s and an extension step at 72 °C for 1 min and 30 s.
149 The next 37 cycles were performed under the following conditions: 94 °C for 30 s, 57 °C for 30 s. and
150 72 °C for 1 min and 30 s. In the 39th cycle, the final extension step was carried out at 72 °C for 10 min.

151 The amplified products were first analyzed by electrophoresis on 3% agarose gel in TBE 1X
152 buffer (Bio-Rad) and then cloned in pCR2.1-TOPO plasmid by using the TOPO TA cloning kit
153 (Invitrogen, Pro, Milan, Italy). White recombinant clones were randomly chosen and screened by PCR
154 using standard vector primers M13. Recombinant clones underwent plasmid purification by PureYield™
155 Plasmid Midiprep System (Promega, USA) and then sequenced on both strands at CEINGE -
156 Biotecnologie Avanzate (Naples, Italy).

157

158 *DNA extraction*

159 DNA was extracted from leukocyte, using the procedure described by Gossens and Kan (1981). Briefly,
160 fresh buffy coat samples were washed twice with distilled water and NaCl 1.8% to remove the excess of
161 red cells, protein digestion was carried out with 500 µl of proteinase K solution (2 mg/ml of proteinase
162 K, 1% w/v SDS and 0.02 M EDTA). Proteins were extracted using phenol-chloroform method followed
163 by DNA precipitation with cold isopropanol. The isolated DNA was then resuspended in 100 µl TE buffer
164 pH 7.6 (10 mM Tris, 1mM EDTA). DNA concentration and OD_{260/280} ratio of the samples were then
165 measured by the Nanodrop ND-2000C Spectrophotometer (Thermo Scientific).

166

167 *CSN1S1* locus genotyping

168 In order to genotype 175 individual samples of water buffalo for the c628C>T mutation, a method based
169 on ACRS-PCR (Amplification Created Restriction Site PCR) was developed according to Lien *et al*
170 (1992). The ACRS-PCR was performed using: ACRS17F (5'- CAATACCCTGATGCCCCGAT - 3') as
171 forward and ACRS17R (5'- CACCACAGTGGCATAGTAG - 3') as reverse, corresponding to nt 70–88
172 and complementary to nt 137–155 of the 17th exon of buffalo *CSN1S1* cDNA sequence (EMBL
173 HE573919), respectively. According to the method, the forward primer was modified by changing C→G
174 in position 17 in order to provide a restriction site for the *Mbo*I (!GATC) endonuclease.

175 The amplification protocol consisted of 39 cycles: the first cycle involved a denaturation step at
176 95 °C for 5 min. The next 38 cycles were performed under the following conditions: 95 °C for 45 s, 63.2
177 °C for 45 s. and 72 °C for 20 s. In the 39th cycle, the extension step was carried out at 72 °C for 10 min.
178 PCR reaction mix comprised: 100 ng of genomic DNA, 50mM KCl, 10mM Tris–HCl, 0.1% Triton X-
179 100, 3mM MgCl₂, 5 pmol of each primer, dNTPs each at 400 µM, 2.5U of *Taq* DNA Polymerase
180 (Promega, Madison, WI), and 0.04% BSA, with a final volume of 25 µl.

181

182 *Digestion and electrophoresis condition*

183 Each PCR amplification product was digested with 10U of *Mbo*I after incubation for 5 h at 37 °C;
184 following the supplier's guidelines. The restricted fragments were analyzed directly by electrophoresis
185 on 3% TBE agarose gel in 1 X TBE buffer and stained with ethidium bromide.

186

187 *Association study*

188 Associations between *CSN1S1* genotypes and milk production traits were carried out on 4,122 test day
189 records of 503 lactations from 175 buffalo cows, supplied by the Italian Association of Buffalo Breeders

190 (ANASB). Milk yield (MY), fat (FP) and protein percentages (PP) were tested with the following mixed
 191 linear model:

192

$$193 \quad y_{ijklmno} = \text{Month}_i + \text{Par}_j + \text{Sea}_K + \text{DIM}_l + \alpha_{s1m} + c_n(\alpha_{s1m}) + e_{ijklmno} \quad [*]$$

194

195 where: $y_{ijklmno}$ is the test-day record of MY, FP or PP; Month_i is the fixed effect of the i -th month of
 196 production (12 levels); Par_j is the fixed effect of the j -th parity (6 levels: 1 to 5, >5); Sea_K is the fixed
 197 effect of the k -th calving season (4 levels: autumn, winter, spring, summer); DIM_l (Days In Milk) is the
 198 fixed effect of the l -th stage of lactation (30 levels of 10 days each); α_{s1m} is the fixed effect of the m -th
 199 genotype at the c.628C>T SNP of *CSN1S1* gene (3 levels: CC, CT, TT); c_n is the random effect of
 200 individual cow (175 levels), nested within α_{s1} genotype; and $e_{ijklmno}$ is the random residual. Pairwise
 201 comparisons among different levels of fixed effects included in model were performed using a
 202 Bonferroni adjusted test. (Co)variance matrices of random effects of cow and residual were assumed to
 203 be diagonal, $\mathbf{I}\sigma^2_c$ and $\mathbf{I}\sigma^2_e$, respectively. They allow for the REML estimation of variance components
 204 associated to individual cow (σ^2_c) and residual (σ^2_e). Variance component associated to the α_{s1} locus
 205 ($\sigma^2_{\alpha_{s1}}$) was estimated running a mixed model having the same structure of [*] but with the α_{s1} genotype
 206 treated as random. Contributions of α_{s1} locus ($r^2_{\alpha_{s1}}$) and cow (r^2_c) to the total phenotypic variance of the
 207 trait was calculated as the ratio between $\sigma^2_{\alpha_{s1}}$ and σ^2_c , respectively and the sum of all variance
 208 components (i.e. $\sigma^2_{\alpha_{s1}} + \sigma^2_c + \sigma^2_e$).

209 In order to estimate the average of gene substitution effect (α) and a possible dominance effects
 210 (d), gene effect was treated as a covariable, represented by the number of T alleles at the α_{s1} locus (0, 1,
 211 2). Finally, an interaction between alleles at the SNP locus was considered (Banos *et al*, 2008; Barendse
 212 *et al*, 2008).

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Results and discussion

a) Characterization of CSN1S1 transcripts

Analysis of the cloned RT-PCR fragments

The mRNAs extracted from individual milk samples obtained from 10 Mediterranean river buffaloes, randomly chosen in the province of Salerno and Caserta, was investigated through the clone analysis. 10 positive clones for each individual were screened. The electrophoretic analysis of the PCR products and the subsequent sequencing of clones, showed at least two populations of transcripts for each examined individual. The most represented population (about 90 %) was correctly assembled, followed by that one deleted of the first triplet of the 11th exon. The last event is a constitutive allele independent event which takes place during the maturation of the pre-mRNA. The first codon of exon 11 (CAG), coding for glutamine in position 78, is in fact competitively eliminated from the mRNA because it is recognized as cryptic site of splice instead of the canonic site (AG) located at the end of the 10th intron. This feature was already observed in buffalo (Ferranti *et al*, 1999) and it is common event for the *CSN1S1* gene of other ruminants, like sheep (Ferranti *et al*, 1998), goat (Ramunno *et al*, 2005) and cattle (Ferranti *et al*, 1999). Furthermore, as already observed in the afore mentioned species, also for the river buffalo it is reasonable to hypothesize the existence of other transcripts different from those reported in this study. Their undetectable amounts opens a new opportunity of investigation in the field of the transcript analysis for buffalo milk protein.

Polymorphism detection

In order to detect polymorphisms at the Mediterranean river buffalo *CSN1S1* locus, the correctly assembled transcripts were sequenced. The analysis of the sequences showed a transcript of 1083 bp,

spanning from the 16th nt of the 1st exon to the 345th nt of the 19th exon. The comparison of the obtained sequences (EMBL HE573919 and HE573920) showed 3 transitions. The first was located at the position 89 of the 17th exon (c.628C>T) and the other two at the position 144 and 239 of the 19th exon, respectively (c.882G>A and c.977A>G). The last two mutations are silent because they are located in the 3' UTR, whereas the first is a miss-sense SNP (p.Ser178Leu). This amino acid change at the buffalo *CSN1S1* locus was already observed and it characterizes the genetic variants of the α s1-casein, named B and A, respectively (Balteanu *et al*, 2008; Chianese *et al*, 2009).

Since the presence of the cytosine in position 628 characterizes even other *CSN1S1* sequences of buffalo (FJ392261, AY948385, AJ005430, DQ111783) and of other ruminants, such as goat (AJ504710.2), sheep (NM_001009795), bison (EU862388) and cattle (X59856), its presence might be indicative of an ancestral condition. According to the sequence analysis of the cDNA samples belonging to individuals with informative genotypes, the G in position 882 and the A in position 977 are in *cis* with the T at 89th nucleotide of exon 17. Although the complete genomic sequence of the river buffalo *CSN1S1* is not available, a close distance exists between the exons 17 and 19 in the homologous sequences of the other ruminants. For instance, this DNA region is only 1962 bp long in the bovine *CSN1S1* gene (EMBL acc. No. X59856), therefore, a condition of linkage disequilibrium can be assumed for these SNP in river buffalo.

b) Association of *CSN1S1* polymorphism on milk yield and composition

Genotyping of Mediterranean river buffalo *CSN1S1* alleles

The c.628C>T does not alter or create any restriction site, therefore, we established a screening method based on the *Mbo*I-ACRS-PCR to identify the carriers of this mutation in an easy and rapid way.

The amplified fragment includes the last 86 bp of 17th exon. Therefore, the digestion with such endonuclease produces an undigested fragment of 86 bp in individuals homozygous for the thymine and

two fragments of 70 and 16 bp (not visible electrophoretically) in buffaloes homozygous for the cytosine. The heterozygous individuals produce a pattern characterized by all 3 restriction fragments: 86, 70 and 16 bp (Figure 1). The investigated population was in Hardy–Weinberg equilibrium. Genotype distribution and allele frequencies are reported in Table 1. The frequency of the thymine was 0.33. This result is in agreement with data reported by Chianese *et al* (2009) and Bonfatti *et al* (2012a) and it confirms that the most common genetic variant in Mediterranean breed reared in Campania (Italy) is the $\alpha s1$ -CN B variant.

Association study

All environmental factors included in model [*] affected significantly all the traits considered (Table 2) except the calving season for FP. The genotype at the *CSN1S1* locus was significantly associated with protein percentage ($p < 0.04$). In particular (Table 3), the CC genotype showed an average value of about 0.04% higher than the CT and TT genotypes. The allele substitution effect of a cytosine in a thymine was -0.014 ± 0.014 , with a quite low contribution of the *CSN1S1* locus to the total phenotypic variance of PP ($r^2_{\alpha s1} = 0.003$) PP. A large effect of dominance (-0.028 ± 0.019) was also observed. Often such effect is not detected or considered non relevant because numerically much lower than the additive effect. Although dominance effects are not important in the estimation of breeding values, being not transmitted in the offspring, they might have an impact on allele substitution effect in the population as recently reported (Pauciullo *et al*, 2012a, b).

Recently, Bonfatti *et al* (2012a) estimated the effects of the *CSN1S1* (B and A alleles, c.628C>T transition) -CSN3 (k-casein, X1 and X2 alleles) genotypes on milk production traits and milk coagulation properties in Mediterranean water buffalo. In particular, these authors report that genotypes did not affect milk protein content, but the composite genotype AB-X1X1, compared to genotype AA-X1X1, was associated with an increased fat content and they indicate a role for the casein genes in the variation of

285 the coagulation properties of the buffalo milk. However, the same authors (Bonfatti *et al*, 2012b) reported
286 that the increased proportion of the α s1-CN on the total casein (TCN) content is associated with
287 genotypes carrying the allele *CSN1S1* A. On the contrary, genotypes associated with a marked decrease
288 of the α s1-CN on the TCN (composite genotypes AB-X1X1 and BB-X1X2) are associated with marked
289 increases in the proportion of α s2-CN.

290 Although results of the present work need to be confirmed with large-scale studies, they might be
291 of great economic interest for the buffalo dairy industry. In fact, increases in average protein content
292 would lead consequently to an expected increase of mozzarella production.

293

294 **Conclusions**

295 The present study reports a characterization of the Mediterranean water buffalo *CSN1S1* transcripts. At
296 least two populations of transcripts were detected. The most represented population (about 90 %) was
297 correctly assembled, followed by that one deleted of the first triplet of the 11th exon.

298 Also the study confirms the existence of genetic polymorphisms at this *locus* and it offers a
299 method based on the *Mbo*I-ACRS-PCR for a rapid genotyping at DNA level for the *CSN1S1* A and B
300 alleles. Furthermore, a significant association between the c.628C>T SNP and the protein percentage
301 was found. In particular, the CC genotype showed an average value of about 0.04% higher than the CT
302 and TT genotypes.

303 Therefore, further studies are necessary to better determine the real effects of the transition
304 c.628C>T on milk composition. Besides, an investigation on larger population are needed in order to
305 validate its application.

306 In addition, it is necessary to investigate the remaining polymorphisms detected in the 3' un-
307 translated regions (UTR). In fact, it is well known that the sequences in the 3' UTR can affect the
308 mechanism of mRNA regulation, such as de-adenylation and degradation. Therefore, it is reasonable to

309 hypothesize that these observed mutations might influence, directly or indirectly, the gene expression
310 and, consequently, the milk protein composition.

311

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315

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442

443 **Table 1.** Genotyping data and allele frequency of the c.628C>T SNP at the *CSN1S1* gene in the Italian
444 Mediterranean river buffalo population.

	Genotypes			Total	Allelic frequency	
	CC	CT	TT		C	T
<i>Observed</i>	75	84	16	175	0.67	0.33
<i>Expected</i>	78.22	77.55	19.22			

445 $\chi^2=1.20$ - $p \leq 0.05$ - d.o.f.=1
446
447

448 **Table 2.** Statistical significance of factors included in model

Effect	P-value		
	Milk yield	Fat	Protein
Genotype	0.63	0.93	0.04
Parity	<.001	<.001	<.001
Month of production	<.001	<.001	<.001
DIM	<.001	0.02	<.001
Season	<.001	0.48	0.02

449
450

451 **Table 3.** Least squares means of milk yield (kg/d), fat and protein percentage (%) for the genotypes at
452 the *locus* c.628C>T of river buffalo *CSN1S1* gene estimated with model [*].

Genotype	Animals n.	Milk yield (kg/d)	Fat (%)	Protein (%)
CC	75	7.81±0.17	9.22±0.12	4.72±0.016 ^a
CT	84	7.92±0.17	9.27±0.12	4.68±0.016 ^b
TT	16	7.56±0.36	9.20±0.23	4.69±0.029 ^{ab}

453 ^{a,b} Means within columns with different superscripts differ (P=0.038)
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Table 4. Substitution effect of a cytosine with a thymine at the c.628C>T SNP in the *CSN1S1* gene (mean \pm SE) and contribution of the α s1 polymorphism to the phenotypic variance for protein percentage.

Statistic	Value
α	-0.014 \pm 0.014
d	-0.028 \pm 0.019
$\sigma^2_{\alpha s1}$	0.0004
σ^2_c	0.0034
σ^2_e	0.1596
$r^2_{\alpha s1}$	0.003
r^2_c	0.021

α : Substitution effect;
d: dominance effect;
 σ^2 : variance components associated to the genotype (α s1); to the individual buffalo cow (c), to residuals (e);
 r^2 : contributions of genotype (α s1) and of individual buffalo cow (c) to the total phenotypic variance.

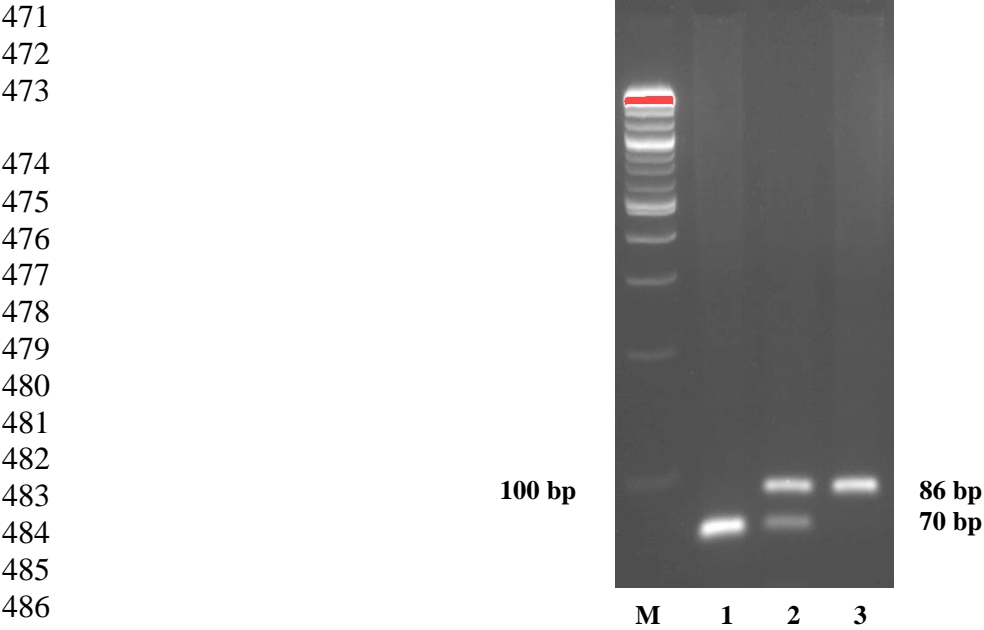


Figure 1. Observed genotypes after *Mbo*I digestion of fragments obtained by ACRS-PCR of a DNA region corresponding to the last 86 bp of the 17th exon into Mediterranean river buffalo *CSN1S1* gene. M=2-Log DNA ladder (0.1-10.0kb) (New England Biolabs); lane 1: *CSN1S1* C/C; lane 2: *CSN1S1* C/T; lane 3 *CSN1S1* T/T