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1 Reproductive response to male joining with ewes with different allelic variants of the
2 *MTNR1A* gene

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13

14 **ABSTRACT**

15 The aims were to evaluate the reproductive response to ram placement with Sarda ewes
16 with different allelic variants at position g.15099485A>G of the *MTNR1A* gene. Ram
17 placements occurred between the early and late spring and there was analysis of whether
18 this polymorphism is associated with other nucleotide substitutions. In each of the eight
19 farms where the study was conducted (named F1-F8), 150 ewes (50 with A/A, A/G and
20 G/G genotypes) were selected. In each group of 150 ewes, eight males were joined with
21 ewes on the following dates: 25 March (T1) for F1-F2, 15 April (T2) for F3-F4, 5 May
22 (T3) for F5-F6, and 1 June (T4) for F7-F8. The lambing dates and number of new born
23 lambs were recorded until 220 days after joining rams with ewes. The ewes with G/G or
24 A/G genotypes had a greater fertility rate compared with those with A/A in T1, T2
25 ($P<0.01$), and in T3 and T4 ($P<0.05$). The duration of time in days from the time of ram
26 joining with ewes to lambing was less in the ewes with G/G and A/G compared with those
27 with A/A ($P<0.01$). The g.15099485A>G variation was always associated with that at
28 position g.15099391G>A. Results indicate there is a positive effect of the g.15099485A>G
29 variant on reproduction when males were joined with ewes in March or April. The
30 association that was ascertained in this study between the g.15099391G>A and
31 g.15099485A>G polymorphisms for the *MTNR1A* gene, could provide new insights to
32 clarify the mechanism of action of melatonin.

33

34 **Keywords:** Seasonality; fertility; Male effect; MTNR1A gene

35

38 1. Introduction

39 Melatonin is a pleiotropic signalling molecule that regulates several physiological
40 functions including synchronisation of biological rhythms, such as seasonal reproduction
41 (Tamura et al., 2014). In ewes at temperate latitudes, the seasonal reproductive functions
42 are characterised by periods when there are ovulations and no ovulation occurring, caused
43 by annual variations in day length (Chemineau et al., 2003). Photoperiodic cues are
44 transduced into neuroendocrine signals by melatonin (Karsch et al., 1984). In sheep, this
45 hormone is secreted during the night by the pineal gland and functions in the
46 premammillary hypothalamus to control the release of the GnRH (Malpoux et al., 1998;
47 Carcangiu et al., 2014). Melatonin exerts its action through receptors distributed in
48 different parts of the body (Migaud et al., 2005); among these, *MTNR1A* appears to be
49 involved in the regulation of reproductive seasonality (Weaver et al., 1996; Dubocovich et
50 al., 2003). This receptor subtype, encoded by the *MTNR1A* gene, is a G-coupled protein
51 receptor, mostly found in the pars tuberalis area of the pituitary and pre-hypothalamic
52 region (Chabot et al., 1998; Dardente et al., 2007; Duprè et al., 2008). Polymorphic sites
53 were detected within exon II of the *MTNR1A* gene in sheep, mouflon, goats and buffalo
54 (Carcangiu et al., 2009a; 2009b; 2010; Luridiana et al., 2012). In different sheep breeds,
55 the allelic variants at positions g.15099491C>T and g.15099485A>G of the *MTNR1A* gene
56 exon II (according to the latest genome version Oar4.0, GenBank accession number
57 NW_014639035.1) have been associated with the seasonal reproductive traits (Pelletier et
58 al., 2000; Chu et al., 2006; Luridiana et al., 2015a; Giantsis et al., 2016). In the Aragonesa
59 breed, however, only the g.15099491C>T polymorphism was associated with a greater
60 percentage of oestrous cyclic ewes between January and August (Martínez-Royo et al.,
61 2012). In the Ile de France ewes, the g.15099485A>G polymorphism was not associated

62 with a difference in the onset, cessation, or length of the breeding season among the
63 animals of the two homozygous genotypes (maintained in a controlled photoperiodic
64 setting without contact with males; Hernandez et al., 2005). These results indicate that the
65 relationship between the *MTNR1A* gene polymorphism and the reproductive season can
66 change with the breed or be affected by management systems, (e.g., presence or absence of
67 males) or other environmental conditions. Thus, the relationship between polymorphisms
68 of this gene and reproductive resumption could also be masked or affected by the
69 observation period in different studies. Indeed, the effect of presence of a male(s) on the
70 reproduction of ewes is affected by breed and season (Chanvallon et al., 2011), and also
71 within the same breed the presence of males effects reproduction in different ways (Fabre-
72 Nys et al., 2015). The utilisation of the management practice of joining males with ewes is
73 an approach that meets modern criteria for ‘clean, green and ethical’ production methods,
74 but its use is limited by the great amount of variability in the reproductive response of ewes
75 (Brown et al., 2014). This could be due to the extent that photoperiodic cues have on the
76 duration of anoestrus of sheep, which in turn could depend on the genotype at the *MTNR1A*
77 locus. The goal of the present research was to clarify the role of this polymorphism on the
78 variability of the response to the joining of males with ewes in the transition from the
79 period of anoestrus and the initiation of oestrous cyclic functions.

80 Thus, the first aim of the present study was to evaluate, from early to late spring, the
81 reproductive response to rams of Sarda ewes with the different allelic variants -
82 g.15099485A>G of the *MTNR1A* gene. A further aim was to analyse whether this
83 polymorphism is related to other variations in the nucleotide sequence of the *MTNR1A*
84 gene exon II.

85

86 **2. Material and methods**

87 2.1. *Animals and experimental design*

88 All the animals in this research had veterinary care by the National Health
89 Veterinary Service in accordance with the Animal Welfare Act. Blood samples collected
90 by veterinarians of the National Health Service for routine health assessments.

91 Sarda sheep (more than 3.0 million ewes in Sardinia) are the main dairy breed in
92 Italy, and the milk production in dairy enterprises occurs in a pattern consistent with the
93 pasture growth pattern. The ewes of this breed have a short anoestrous period, generally
94 from late winter to mid-spring. For the present research, ewes were on eight farms located
95 in North Sardinia (on the 40° N) that were located in a similar climatic zone and at a similar
96 altitude utilising a similar nutritional and management regimens. On each farm, there were
97 approximately 800 Sarda sheep and the animals had been maintained in conditions with a
98 natural photoperiod from birth. During the day, the animals grazed on leguminous and
99 gramineous grasses, and received 300 g per animal daily of concentrated commercial food
100 (crude protein 20.4% and 12.5 MJ ME/kg DM) at the time of milking. At each farm, the
101 genotype at position g.15099485A>G of the *MTNR1A* gene exon II was identified in all the
102 animals aged 3 to 5 years (approximately 400 ewes in each farm). There were 150 lactating
103 ewes selected at each farm ($n = 50$ with A/A, A/G and G/G genotypes) that lambed in
104 2015, between October 20 and December 1, for a total of 1,200 ewes for the study. The
105 ewes utilized in the study had at least two previous parities, thus, were at least 3 years of
106 age. Nulliparous ewes and those with a single parity that were in their second gestational
107 period were excluded, as in the Sarda breed, the first lambing generally occurs between
108 January and April with the greatest number of lambs being born between February and
109 March. Sarda sheep have an ample milk yield and consequently their reproductive
110 functions are less than that of ewes of many other breeds for at least 2 months after
111 lambing leading to a delay in the time of lambing in second parity ewes (Luridiana et al.,

112 2015b). This is the reason only multiparous ewes that were pregnant with lambs to be
113 delivered at least at their third parturition were selected for the present study.

114 The number in the ear tag of each animal was recorded, and the ewes were
115 individually marked with numbered collars to avoid recognition errors. At each farm,
116 (identified as F1-F8) the 150 ewes utilised for the present study were separated from the
117 rest of the flock. In every group of 150 ewes, eight adult (3 to 6 years old) Sarda males of
118 proven fertility (male/female ratio 1/20) were joined with ewes during the 2016 time
119 period (Table 1). The ewes were previously isolated from rams for 90 days (sound, sight
120 and smell, minimum distance > 1,500 m). The rams were separated from females after 70
121 days of joining of ewes and rams. Gestation was diagnosed by trans-abdominal
122 ultrasonography examination using Esaote Piemedical Tringa linear equipment (Esaote
123 Europe B.V., Maastricht, the Netherlands) with a 5.0 to 7.5 MHz multiple frequency linear
124 probe. Pregnancy diagnosis was performed in all ewes, every week, from 45 days after
125 male joining with ewes to 45 days after male removal. The lambing dates and the numbers
126 of new born lambs were recorded until 220 days after joining of rams with the ewes.

127

128 *2.2. Blood sampling and DNA analysis*

129 To identify the individual allelic variants, DNA analysis was performed using the
130 whole blood from each ewe. Blood samples (10 ml) were collected from the jugular vein,
131 using vacuum tubes with ethylenediaminetetraacetic acid (EDTA) as an anticoagulant (BD
132 Vacutainer Systems, Belliver Industrial Estate, Plymouth, UK). The DNA was extracted
133 from whole blood using a genomic DNA extraction kit (NucleoSpin® Blood, Macherey-
134 Nagel, Germany). An amount of 150 ng of genomic DNA was subjected to polymerase
135 chain reaction (PCR) using specific primers (Sigma Genosys Ltd., Pampisford, Cambs,
136 UK) according to Messer et al. (1997). The primers corresponded to positions 285 to 304

137 (sense primer 5' – TGT GTT TGT GGT GAG CCT GG – 3') and 1,108 to 1,089 (antisense
138 primer: 5' – ATG GAG AGG GTT TGC GTT TA – 3') of the sequence reported by
139 Reppert et al. (1994) (GenBank accession number U14109). The PCR reaction was
140 performed for all the samples using the method reported by Carcangiu et al. (2009b). All
141 the PCR products were subjected to restriction enzyme analysis using the MnlI
142 endonuclease (New England Biolabs, Beverly, MA, USA), which recognises an A to a G
143 substitution at position 612 of the U14109 *MTNR1A* exon II nucleotide sequence,
144 corresponding to position g.15099485A>G in genome version Oar4.0 (GenBank accession
145 number NW_014639035.1). The digestion reaction was performed using the methods
146 described by Carcangiu et al. (2009b).

147

148 *2.3. Sequencing*

149 One hundred PCR products for each genotype were sequenced to determine whether
150 the g.15099485A>G variant was associated with other nucleotide substitutions. A total of
151 300 amplified products was sequenced using Applied Biosystems 3730 DNA Analyzer
152 (Perkin-Elmer Applied Biosystems, Foster City, CA, USA). The sequences were aligned
153 and compared with the ovine sequence GenBank U14109 and NW_014639035.1, to
154 confirm the correspondence of the known nucleotide changes and underscore other
155 possible substitutions. The homology searches were performed using BLAST (National
156 Centre for Biotechnology Information: <https://blast.ncbi.nlm.nih.gov/Blast.cgi>). To align
157 the sequences, the CLUSTALW tool was used (<http://www.genome.jp/tools-bin/clustalw>).

158

159 *2.4. Statistical analysis*

160 Allele and genotype frequencies were determined by direct counting of the observed
161 genotypes. The chi-squared test was used to determine Hardy-Weinberg equilibrium of the

162 mutation (Genepop 4.2). The R statistical software (Version 3.2.2) was used to perform the
163 statistical analysis.

164 The farm effect was not included in the model as the farms were all located within 20
165 km each other, therefore, there were similar climatic conditions and animals had the same
166 veterinary, feed and reproductive farm management. The data for the climatic variables
167 (humidity, environmental temperature), and for the animals (number of lambing and the
168 lambing period, BCS and age) were statistically evaluated and there were no differences
169 detected.

170 A general linear model (GLM) procedure was performed to analyse the effect of
171 treatment period and genotype on the litter size and on the distance in days from ram
172 introduction to lambing, based on the following model:

$$173 \quad Y_{ilmn} = \mu + S_i + G_l + P_m + (G_l P_m) + e_{ilmn}$$

174 where Y_{ilmn} is the variable measured (pregnancy rate or duration of time in days from
175 joining rams with ewes to lambing), μ is the overall mean, S_i is the random effect of the
176 sire, G_l is the fixed effect of the genotype, P_m is the fixed effect of period, $(G_l P_m)$ is the
177 interaction between G_l and P_m , and e_{ilmn} is the error effect. To compare percentages of ewes
178 lambing with each genotype and each time period, a chi-squared test was used. A P
179 value < 0.05 was considered statistically significant.

180

181 **3. Results**

182 The allelic and genotypic frequency was not different of ewes among the eight farms.
183 The means of allele and genotype frequencies are shown in Table 2. The most frequent
184 allele was G (0.68) at position g.15099485 of the latest genome version Oar4.0 (GenBank
185 accession number NW_014639035.1) and, consequently, G/G was the most frequent
186 genotype (53%). The population was not in Hardy-Weinberg equilibrium due to the small

187 number of heterozygotes ($P<0.05$). The DNA sequencing confirmed the presence of the
188 polymorphic site in all the samples. The single nucleotide polymorphism (SNP) position
189 reported in this paper refers to the Oar4.0 genome version (GenBank accession number
190 NW_014639035.1).

191 The alignment of the sequences with that in GenBank indicated there was a total
192 number of eight SNPs: six of which were silent (g.15099644T>G, g.15099491C>T,
193 g.15099485A>G, g.15099314G>A, g.15099296G>A, g.15099206T>C), while with the
194 others there was an amino acid change (g.15099391G>A causing p.Val220Ile, and
195 g.15099204C>A causing a p.Ala282Asp substitution in the amino acid sequence; Table 3).
196 The g.15099485A>G variation was always associated with g.15099391G>A. The number
197 of ewes diagnosed as pregnant differed from the number of ewes that lambed by
198 approximately 3%. In T1, the fertility (percentage of ewes lambed) of ewes between the
199 two farms (F1 and F2) was similar. This trend was similar to those of the other farms
200 during the other periods when frequency of pregnancy was assessed. Considering the
201 g.15099485A>G SNP, the pregnancy rate in the ewes with G/G or A/G genotypes was
202 greater compared to the ewes with A/A in T1, T2 ($P<0.01$) and for ewes in in T3 and T4 (P
203 <0.05 ; Table 4). Some ewes with G/G or A/G genotypes lambed between 150 and 160
204 days after rams were joined with ewes in all the four periods (i.e., for the G/G genotype
205 two lambs in T1 and six in T4 meaning approximately 2% and 6% of the ewes lambed,
206 respectively; Fig. 1). None of the ewes with the A/A genotype lambed before 160 days
207 after ewes being joined with rams for all the periods. For the ewes with a G/G and A/A
208 genotype, there was a difference in lambing rate ($P<0.05$) between the T1-T2 and T3-T4
209 periods, whereas for the ewes with the A/G genotype there was a similar lambing rate in
210 the four periods. The average duration of time in days from joining the rams with ewes to
211 lambing was shorter in the animals with the G/G or A/G than in ewes with an A/A genotype

212 ($P<0.01$). The litter size (number of lambs born per lambing) at the eight farms was
213 affected neither by genotypes nor by the time periods of joining with rams (T1-T4). The
214 total average litter size was similar for ewes with all the three genotypes: 1.20, 1.18 and
215 1.15 for the G/G, A/G and A/A genotype, respectively.

216 The lambing trend is depicted in Figure 1 for the ewes with different genotypes and
217 that conceived during the different periods (from T1 to T4). For all four periods, the ewes
218 with the G/G or A/G genotypes compared with those with the A/A genotype had a greater
219 lambing percentage between 160 and 180 days after ewe joining with rams ($P<0.01$).
220 There was the greatest number of ewes lambing with the G/G genotype that were joined
221 with rams during the T1 and T2 periods between 160 and 170 days after joining of rams
222 with the ewes. For the ewes with the G/A genotype, there was the greatest number of ewes
223 lambing between 170 and 180 days after joining of rams with ewes. For ewes with the
224 A/A genotype that were joined with rams during T1 and T2, there was the greatest
225 numbers of ewes lambing between 190 and 200 days after joining of ewes with males. For
226 ewes with the G/G and G/A genotypes joined with rams during T3 and T4, the greatest
227 number of ewes lambing was between 160 and 170 days and for the ewes with the A/A
228 genotype between 180 and 190 days after joining ewes with males.

229

230 **4. Discussion**

231 The genotypic and allelic frequencies of the analysed locus were similar to those
232 previously reported for the same breed (Carcangiu et al., 2009b; Luridiana et al., 2015a).
233 At position g.15099485A>G of the *MTNR1A* gene exon II sequence, Sarda sheep had a
234 relatively greater frequency of the mutant allele G, similar to that of some other European
235 sheep breeds (Messer et al., 1997; Mateescu et al., 2009). By contrast, in *Ovis gmelini*
236 *musimon* (a wild sheep) the A allele was most frequently present (Carcangiu et al., 2010).

237 Comparing the frequency of occurrence of the G allele in domestic breeds, in the present
238 study there was a lesser rate than in the Aragonesa and in other sub-temperate and sub-
239 tropical Indian sheep breeds (Magra, Marwari, Chokla, Malpura, Patanwadi, Sandyno and
240 Niligiri; Martínez-Royo et al., 2012; Saxena et al., 2014; 2015a, b).

241 The results from the present study indicate that the reproductive response to joining
242 with males of adult Sarda ewes is affected by the polymorphism at g.15099485A>G.
243 Indeed, the ewes with the G/G or A/G genotype had a greater lambing rate and a lesser
244 duration in days between the time of ram joining with ewes compared to those with the
245 A/A genotype, in all the periods where assessments occurred. Results of the present study
246 are consistent with the findings of previous studies in different sheep breeds, in which the
247 G/G genotype at the same locus was associated with breeding and conceiving out of-
248 season and with a shorter interval between first and second lambing (Chu et al., 2006;
249 Mateescu et al., 2009; Carcangiu et al., 2012). Furthermore, Sarda ewes with the G/G
250 genotype had a shorter period of anoestrus in spring after melatonin administration (Mura
251 et al., 2017). The present results indicate that in the Sarda breed, the presence of even one
252 G allele affects the reproductive response to male joining with ewes and these results are
253 consistent with those previously reported for Sarda and other sheep breeds (Mateescu et
254 al., 2009; Mura et al., 2014).

255 In other European sheep breeds the same allelic variant assessed in the present study
256 had no effect on reproductive performance (Hernandez et al., 2005; Martínez-Royo et al.,
257 2012). This difference was attributed to the breed or to the effects of environmental factors.
258 In almost all the studies, the fertility of the sheep was recorded after the joining of males
259 with ewes, while, in the study by Hernandez et al. (2005), reproductive activity was
260 evaluated only by recording progesterone concentrations and associating these with

261 whether ewes were or were not pregnant. The lack of the male effect, therefore, could have
262 affected the results in this previous study.

263 In the present study, the greater lambing rate in the ewes with the G/G or A/G
264 genotype confirmed the hypothesis about a lesser sensitivity to photoperiod of the ewes
265 with these two genotypes. Presumably, this lesser sensitivity resulted in shorter period of
266 anoestrus, which led to a greater response to joining of rams with the ewes, compared to
267 ewes with the A/A genotype, which could have longer periods of anoestrus. The different
268 hypothalamic sensitivity to photoperiodic cues could be the basis for the different
269 reproductive responses among the three genotypes. The difference in fertility rates of the
270 ewes with the A/A genotype between ewes joined with rams in the T1-T2 and T3-T4
271 periods could be due to the gradual transition from anoestrus to the breeding season, with a
272 consequent greater hypothalamic sensitivity to the oestradiol (E₂)-positive feedback signal,
273 as suggested by Fabre-Nys et al. (2015) or a decreased negative feedback to oestradiol as
274 has previously been reported to occur (Karsch et al., 1984).

275 Also, from the lambing trend, it can be hypothesized that the animals with the
276 different genotypes have a different duration of anestrus. This is supported by ewes with
277 the G/G and A/G animals lambing before the ewes with the A/A genotype, and this
278 confirms the earlier response to the joining with the males because of shorter periods of
279 anoestrus.

280 This effect, however, is not easy to explain because this allelic variant is not
281 associated with an amino acid change affecting the trans domains, and therefore it should
282 not involve the receptor functionality. The eight mutations detected in the *MTNR1A* gene
283 exon II (g.15099644T>G, g.15099491C>T, g.15099485A>G, g.15099391G>A
284 g.15099314G>A, g.15099296G>A, g.15099206T>C, and g.15099204C>A) were identical
285 to those reported as a result of previous studies, both in Sarda (Carcangiu et al., 2009b) and

286 in other sheep breeds (Pelletier et al., 2000; Reppert et al., 1994). Furthermore, in these
287 previous studies (Reppert et al., 1994; Pelletier et al., 2000) of the same gene tract there
288 were two other mutations detected, at position g.426C>T and g.555G>A of the melatonin
289 receptor sequence with GenBank accession number U14109 that corresponded to position
290 g.15099671C>T and g.15099575G>A of the latest genome version Oar4.0:
291 NW_014639035.1, respectively. Neither of these allelic variants lead to amino acid
292 changes. Furthermore, Saxena et al. (2014) reported another mutation in the Indian Chokla
293 sheep breed, at position g.931G>C of the U14109 sequence, corresponding to the
294 g.15099166 of the NW_014639035.1 sequence, which also determines an amino acid
295 substitution (p.Ala295Pro). These differences are, due to the different evolutionary
296 pathways of the breeds, based on the focus of selection for different products (milk, meat
297 or wool). In the present study, and consistent with the report of Saxena et al. (2015b), the
298 variation g.15099485A>G was always associated with g.15099391G>A, which determines
299 an amino acid change at position p.Val220Ile (according to the RefSeq record GenBank
300 accession number NP_001009725.1). These data are very important because of the
301 relationship between these variations and may explain the effect on reproductive
302 seasonality of the polymorphism g.15099485A>G. Although the nucleotide substitution
303 g.15099391G>A leads to an amino acid change in the protein sequence, it is not part of the
304 transmembrane domain of the *MTRN1A* gene and this is consistent with the findings of
305 Barrett et al. (2003), consequently there should not be a change in the receptor's
306 functionality. Nevertheless, the position in the protein of this amino acid change is close to
307 the histidine at positions 211 and 195 of the amino acid sequence (NP_001009725.1),
308 which are involved in melatonin signal transduction (Conway et al., 1997; Kokkola et al.,
309 1998). This amino acid change could lead to a modification of the steric conformation of
310 the amino acid chain with a consequent signal alteration (Trecherel et al., 2010). The

311 p.Val220Ile amino acid substitution affected the inhibition of adenylate cyclase, thus
312 suggesting a possible modification in the transmission of the melatonin signal in this
313 variant (Trecherel et al., 2010) and, consequently, the differences observed in the
314 reproductive function among ewes with the G/G, A/G and A/A genotypes (Pelletier et al.,
315 2000). Studies on the second messengers involved in the melatonin receptor activation,
316 however, are needed to clarify this hypothesis (Kokkola et al., 1998; Brydon et al., 1999).
317 In a study by Calvo et al. (2018), it was hypothesised that the SNP at position
318 g.15099004C>T could be the causative mutation for the effects on reproductive seasonality
319 traits, as there is an arginine to cysteine substitution in the amino acid sequence at position
320 349. In the present study with the Sarda breed, this variation was not detected, but this
321 finding indicates there are new and interesting possibilities for further investigations.

322 Furthermore, data from the present study indicate that in the animals with the G/G or
323 A/G genotype, the placement of males with ewes induces a greater stimulus on the
324 resumption of reproductive activity if the placement occurred at T1 and T2 compared to T3
325 and T4. This was surprising, as it was expected that the reproductive response in ewes with
326 the G/G and A/G genotype would be increased from March to June. Instead, the effect is
327 greater when the number of ewes that were anoestrus was greatest (March and April),
328 rather than when the time that the initiation of the oestrous cyclic functions was
329 approaching at the end of the anoestrous period when induction of oestrous cyclic
330 functions is usually easier to induce (May and June). It is speculated that the effect of the G
331 allele is greater when day length is shorter, confirming that ewes with one or two G alleles
332 are less sensitive to the photoperiod inhibition of reproductive functions.

333 The lambing data in the present study was variable when there was assessment of the
334 different genotypes and periods. In fact, the ewes with the G/G or A/G genotype had the
335 peak of lambing earlier than the ewes with the A/A genotype for all the periods when

336 assessments occurred. It is clear that the trend in lambing rate is associated with the trend
337 in mating, so that the ewes with the G/G and A/G genotype had a greater response to the
338 joining of males with ewes. In several studies, there has been an investigation of the origin
339 of the variability in the reproductive response to joining of males with ewes.

340 Chanvallon et al. (2011) reported that there was a greater response of joining males
341 with ewes at the end of the anoestrus season. Instead, in the present study, the ewes with
342 the G/G and A/G genotype were not affected by the photoperiodic suppression to as great
343 an extent in all periods when there were assessment and the greater response of ewes to the
344 joining of males could be due to a set of positive factors, such as greater oestradiol
345 secretion along with greater amounts of steroid acute regulatory protein (StAR) in
346 granulosa cells Fabre-Nys et al. (2015). Further studies are necessary to clarify if the G
347 allele may be associated with a greater secretion of E₂ and increased amounts of StAR
348 compared to that of ewes with the A allele.

349

350 **5. Conclusion**

351 In conclusion, the present results indicate that both the *MTNR1A* gene allelic variant
352 and the period of joining rams with ewes affected the reproductive response to males in the
353 Sarda breed of sheep. In the present study, there was a positive effect of the
354 g.15099485A>G variant on the time of cessation of anoestrus when males were joined with
355 ewes in March or April. Furthermore, the association between the SNPs, g.15099485A>G
356 and g.15099391G>A, that was determined to exist in the present study could provide new
357 insights to clarify the mechanism of action of melatonin and the role of its receptors on
358 reproductive seasonality. In addition, studying the effects of the different genotypes on the
359 granulosa and luteal cells may provide useful information on the action of the *MTNR1A*
360 gene at the ovary and on fertility in sheep.

361

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365

366 **Conflict of interest**

367 None of the authors have any conflicts of interest to declare.

368

369 **Author contribution**

370 I certify on behalf of all coauthors that this article has not been presented in any other
371 place for publication. All coauthors have contributed equally to the research (conception,
372 design of study, acquisition and interpretation of data) as well as to article preparation. All
373 coauthors have approved the final draft of this article.

374

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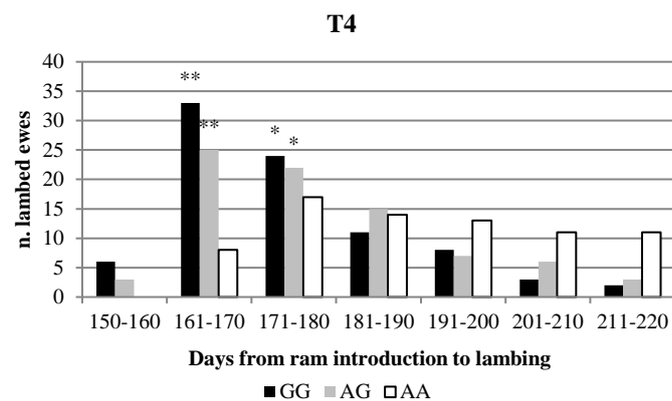
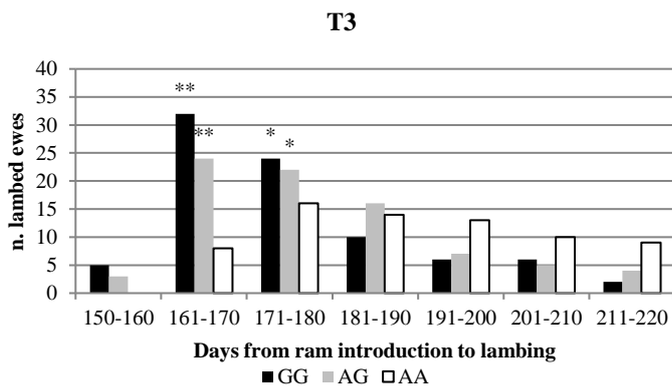
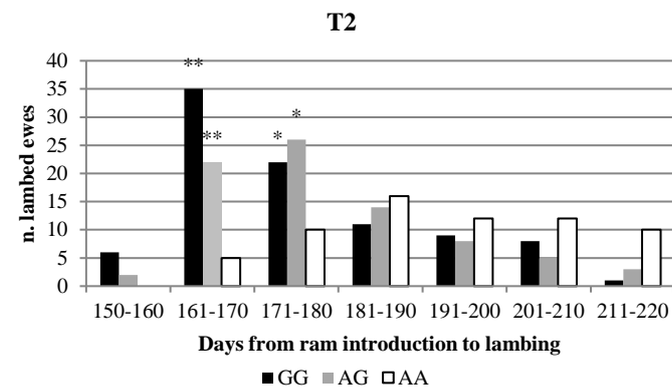
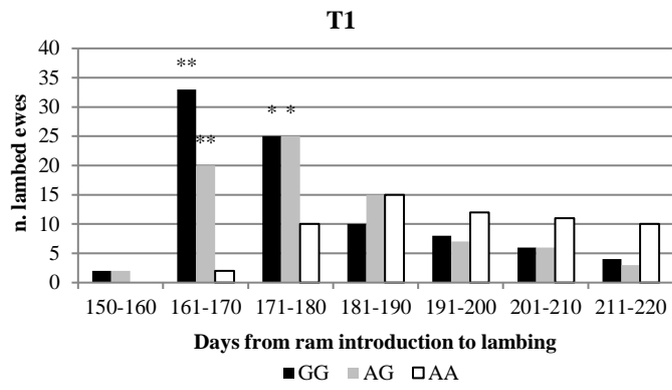


Fig. 1. Depiction of the number of ewes that lambed in in 10-day intervals from 150 to 220 days after the joining of males with ewes in the four time periods, according to genotype at the g.15099485A>G position; T1 = male joining with ewes on March 25; T2 = male joining with ewes on April 15; T3 = male joining with ewes on May 5; T4 = male joining with ewes on June 1; ** $P < 0.01$; * $P < 0.05$

Table 1

Time period of male joining with ewes on eight farms and number of ewes at each farm that was included in the study (1200 ewes)

Time period	Data of male introduction	Farms	<i>n</i>
T1	25 March	F1	150
	25 March	F2	150
T2	15 April	F3	150
	15 April	F4	150
T3	5 May	F5	150
	5 May	F6	150
T4	1 June	F7	150
	1 June	F8	150

Table 2

Genotype and allele frequencies of the *MTNR1A* gene allelic variant in ewes on the eight farms ($n = 1200$ ewes)

Allelic variant	g.15099485A>G		
Genotypes	A/A	A/G	G/G
Genotype frequency	0.14	0.33	0.53
Alleles	A	G	
Allele frequency	0.32	0.68	

Table 3

Nucleotide and amino acid changes within the *MTNR1A* gene exon II in Sarda ewes ($n = 300$ sequenced ewes)

SNP position ^a	Nucleotide change ^b	Codon change ^c	Amino acid change ^d
g.15099644	T>G	ACT/ACG	None: Thr135Thr
g.15099491	C>T	TAC/TAT	None: Tyr186Tyr
g.15099485	A>G	CCA/CCG	None: Pro188Pro
g.15099391	G>A	GTC/ATC	Val220Ile
g.15099314	G>A	CTG/CTA	None: Leu245Leu
g.15099296	G>A	AGG/AGA	None: Arg251Arg
g.15099206	T>C	CCC/CCT	None: Pro281Pro
g.15099204	C>A	GCC/GAC	Ala282Asp

^aAccording to the latest genome version Oar4.0 (GenBank acc. number NW_014639035.1)

^bSequence is in a reverse orientation on the Oar4.0 genome version, so that nucleotide substitution appears in the reverse form compared to the present study

^cNucleotide changes within codons are in bold

^dAccording to NCBI Reference Sequence: NP_001009725.1

Table 4

Lambing rate and duration of time in days from ram joining with ewes to lambing in the four periods (T1 to T4) ewes were with rams based on genotypes at position g.15099485 ($n = 1200$ ewes)

Genotypes	Lambing rate				Duration in days from ram joining with ewes to lambing			
	G/G	A/G	A/A	<i>P</i>	G/G	A/G	A/A	<i>P</i>
T1	88% ^b	78%	60% ^a	<0.01	177.6±14.8	179.5±14.2	193.3±14.3	<0.01
T2	92% ^b	80%	65% ^a	<0.01	176.1±14.5	178.9±14.0	192.1±15.1	<0.01
T3	85% ^a	81%	70% ^b	<0.01	175.7±14.2	178.8±14.3	189.0±15.6	<0.01
T4	85% ^a	81%	74% ^b	<0.05	174.8±13.5	178.6±14.6	189.7±15.9	<0.01

Date of ram joining with ewes - T1: 25 March; T2: 15 April; T3: 5vMay; T4: 1 June

Different lowercase letters in columns differ $P < 0.05$; P value refers to the significance within the row

Conflict of interest statement

None of the authors of this manuscript has a financial or personal relationship with other people or organisations that could inappropriately influence or bias the content of this paper.