SHORT COMMUNICATION

A new polymorphism in the *Growth and Differentiation Factor 9* (*GDF9*) gene is associated with increased ovulation rate and prolificacy in homozygous sheep

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Summary

Brazilian Santa Inês (SI) sheep are very well-adapted to the tropical conditions of Brazil and are an important source of animal protein. A high rate of twin births was reported in some SI flocks. *Growth and Differentiation Factor* 9 (*GDF*9) and *Bone Morphogenetic Protein* 15 (*BMP15*) are the first two genes expressed by the oocyte to be associated with an increased ovulation rate in sheep. All *GDF*9 and *BMP15* variants characterized, until now, present the same phenotype: the heterozygote ewes have an increased ovulation rate and the mutated homozygotes are sterile. In this study, we have found a new allele of *GDF*9, named *FecG^E* (Embrapa), which leads to a substitution of a phenylalanine with a cysteine in a conservative position of the mature peptide. Homozygote ewes presenting the *FecG^E* allele have shown an increase in their ovulation rate (82%) and prolificacy (58%). This new phenotype can be very useful in better understanding the genetic control of follicular development; the mechanisms involved in the control of ovulation rate in mammals; and for the improvement of sheep production.

Keywords growth factor, *Ovis aries*, prolificacy.

Some breeds of sheep are naturally prolific and they are very informative for the study of reproductive genetics and physiology. It is postulated that GDF9 and BMP15 may form non-covalent homo and heterodimers in vivo and, in a species-specific way, modulate the ovulation rate in mammals (Moore *et al.* 2004). The TGF β -family member *BMP15* was the first gene to be associated with increased ovulation rate in Inverdale ($FecX^{I}$ polymorphism) and Hanna ($FecX^{H}$) sheep (Galloway et al. 2000). Soon after, the Booroola variant $(FecB^B)$ was found in the *BMPR1B* gene of Merino Booroola (Mulsant et al. 2001; Souza et al. 2001; Wilson et al. 2001). The BMPR1B, together with its partner BMPRII, is responsible for the BMP15 signalling in the ovarian follicles (Moore et al. 2003). The last major gene found to be associated with prolific phenotype was GDF9, in which a polymorphism $(FecG^{H})$ found in Cambridge and F700-Belclare sheep is responsible for an increased ovulation rate in heterozygotes and sterility in homozygotes, in a way very similar to all BMP15 variants (Hanrahan et al.

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2004; Bodin *et al.* 2007; Martinez-Royo *et al.* 2008; Monteagudo *et al.* 2008).

Besides the GDF9 and BMP15 activities during cumulus expansion, oocyte maturation and ovulation (Elvin et al. 1999, 2000; Gui & Joyce 2005; Yoshino et al. 2006), these two paracrine factors play important roles during many steps of follicular development. They influence follicle growth (Dong et al. 1996; Nilsson & Skinner 2002), cumulus and granulosa cell proliferation (Hayashi et al. 1999; Gilchrist et al. 2006; Spicer et al. 2006), cell-survival signalling (Hussein et al. 2005; Orisaka et al. 2006) and act as modulators of many other growth-factors and endocrine hormones (Juengel et al. 2004). As a result of their role in the folliculogenesis, the availability of GDF9 and BMP15 polymorphisms can be very useful in the study of animal reproductive genetics and physiology. In this study we have shown, for the first time, a polymorphism in the GDF9 gene that increases the ovulation rate in homozygotes, without sterility. in sheep.

In this study, 23 ewes (*Ovis aries*) from a Santa Inês (SI) population with a history of multiple births (twin and triplet births) were investigated for SNPs in the *BMP15* and *GDF9* genes. The ewes were genotyped for the Booroola SNP (*FecB^B*) as previously described (Wilson *et al.* 2001). Subsequently, exon 2 of the *GDF9* and *BMP15* genes were

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screened for SNPs by DNA sequencing of PCR amplicons using the following primers: GDF9 (forward 5'-GGAGAAA AGGGACAGAAGC; reverse 5'-ACGACAGGTACACTTAGT); and BMP15 (forward 5'-GGCTGCTTGTCAGTTTGTAC; reverse 5'-GAGCACTTTCAGATTTAA) (see Appendix S1 for details). Seven (GI to GVII) single nucleotide polymorphisms (SNPs) were found in GDF9 (Table A1, additional data). Only the GVII polymorphism is a non-conservative change in position 345 (phenylalanine to cysteine), which was detected in 43% of the sequenced animals, and it is in the mature peptide of GDF9. This polymorphism provokes a change in a residue which is 100% conserved in the sequence of four representative mammalian species (FigureA1 in Appendix S1), and was named $FecG^{E}$ (GenBank FJ429111) according to the previous nomenclature for the high fertility GDF9 allele ($FecG^{H}$) (Hanrahan *et al.* 2004).

To find the frequency of the $FecG^{E}$ polymorphism, a total of 334 animals from a separate flock (Appendix S1) that had not been selected for prolificacy have had their genotypes identified by a PCR-RFLP strategy (Appendix S1). All data about the parturition of these ewes during the period of 2002 to 2008 were collected, and the association between the number of lamb births and genotypes was tested. The genotype distribution and allele frequency were analysed by the Chi-square test. A difference (P < 0.001) in the frequency of $FecG^{E}$ and $FecG^{+}$ alleles, as well as in the genotype distribution, was observed between the randomly selected and the prolificacy-selected flocks (Table 1).

To investigate the association between the $FecG^{E}$ genotypes (E/E, +/E, and +/+) and the ovulation rate, 39 ewes (15 $FecG^{+/+}$, 15 $FecG^{+/E}$ and 9 $FecG^{E/E}$) were selected from the genotyped flocks and submitted to oestrus synchronization. The animals were oestrus synchronized twice, with eCG and PGF2alpha-based protocols in a cross-over design (Appendix S1). Eleven days after the last oestrus detection, laparoscopy was performed as previously described (Killen & Caffery 1982) to infer ovulation rate by counting corpora lutea (CL). At the end of the breeding season, pregnancy status was evaluated by ultrasound. All animals submitted to laparoscopy had their GDF9 and BMP15 exon 2 sequenced to confirm the $FecG^{E}$ genotyping and to verify that there was no other characterized polymorphism

 Table 1 Genotypic and allelic frequencies of FecG^E in Santa Inês flocks.

SI Flock	Genotype	Frequency (<i>N</i>)	Allele	Frequency (<i>N</i>)
Prolific-selected ^a	+/+	0.174 (4)	FecG ⁺	0.478 (22)
	+/E	0.609 (14)	FecG ^E	0.522 (24)
	E/E	0.217 (5)	–	-
Randomly selected ^b	+/+	0.656 (219)	FecG ⁺	0.808 (540)
	+/E	0.305 (102)	FecG ^E	0.192 (128)
	E/E	0.0389 (13)	–	-

Distinct letters are different (P < 0.001) for genotype distribution.

associated with ovulation rate. The animals were handled in accordance with pertinent Brazilian legislation and following Embrapa's procedures for animal care.

The CL number and the number of lamb births were fitted to the GLM (Generalized Linear Model), where the Poisson distribution was attributed to the ovulation rate, pregnancy and lambing data. In this analysis, the lamb count was considered as response variable, measured for each animal in seven different breeding seasons from 2002 to 2008 (time variable). To measure the influence of genotype over the offspring number through the time, a generalized linear mixed model (GLMM. SAS software) was applied. The offspring number y_{ij} of the *i*th animal at the *j*th time was modelled as the per following model:

$$y_{ij}|\gamma_j \sim \text{Poisson}(\mu_{ij})$$

$$\gamma_j \sim N(0, \sigma^2)$$

$$\mu_{ij} = \exp(\beta_0 + \beta_1 X \mathbf{1}_i + \beta_2 X \mathbf{2}_i + \beta_3 Z_{ij} + \gamma_i)$$

$$\text{Var}(y_{ij}|\gamma_j) = \sigma^2 \mu_{ij},$$

where y_{ij} follows the Poisson distribution conditioned to the random effect for animal γ_i , which was assumed to be normally distributed with variance σ^2 . The expected mean μ_{ij} is a non-linear function of the effects of genotype (β_1 and β_2), time when the counting was made (β_3) and the random effect because of each animal. The variance for y_{ij} irrespective of the random effect γ_i is $Var(y_{ij}) = \sigma^2 \mu_{ij}$, where the extra (or sub) variation is taken into account. The estimate for σ^2 is 0.1696 (standard error = 0.0087), indicating a strong under-dispersion, but this is correctly modelled by GLMM (see Appendix S1, SAS output in additional data).

The parturition data of the 334 genotyped ewes showed a difference (P < 0.0001) in the prolificacy amongst the groups (Table 2). Regarding the ovulation rate, it was greater (P < 0.001) in the homozygote (E/E) group, which showed an 82% increase in CL average (2.22 ± 0.12, Fig. 1a), as well as the highest frequency (96.3%) of multiple-ovulating ewes (Fig. 1b), when compared with +/E and +/+ groups. The heterozygote group (+/E) presented no difference (P = 0.612) in CL average (1.34 ± 0.08) or in the frequency (31.8%) of ewes with multiple ovulations (Fig. 1a and b), when compared with the wild-type ewes (1.22 ± 0.11 and 14.6%)

Table 2 The effect of of *FecG^E* in Santa Inês prolificacy.

SI Flock (F1)	Genotype	Prolificacy of F1 (mean; [95% CI])
Randomly selected	+/+ +/E E/E	1.13; [1.11, 1.16] ^a 1.44; [1.41, 1.48] ^b 1.78; [1.69, 1.87] ^c

Distinct letters are different (P < 0.001).

Non-selected SI; N = 334 ewes (219 +/+; 102 +/E and 13 E/E), called F1.

Prolificacy = mean of 764 offspring (F2) from the 334 genotyped ewes (F1); separated according to their genotype category.



Figure 1 The effect of genotypes on ovulation. (a) The average number of corpora lutea (CL) per ewe in each genotype: E/E (*FecG^E* in homozygosis) N = 9, +/E (*FecG^E* in hereozygosis) N = 15 and +/+ (without *FecG^E* allele) N = 15. The CL data are presented as mean \pm SE. (b) The frequency of multiple-ovulating ewes (≥ 2 CL) in each genotype as described in (a). (c) The frequency of twinning scored by ultrasonography at the 45th day of gestation in each genotype: E/E (N = 9), +/E (N = 14) and +/+ (N = 14). Groups with different letters differ (P < 0.001; Figures 1a and 1b) or (P = 0.0136; Figure 1c).

respectively). We observed a genotype effect on the number of twins per ewe (P = 0.0136); E/E ewes showed 44% of twin-pregnancy, while no twin-pregnancy was observed in +/+ ewes (Fig. 1c). Moreover, the E/E ewes presented no observable effect of $FecG^E$ other than the increased ovulation rate and twinning.

It has been suggested that increasing multiple births may be an efficient way to improve meat production per ewe, and an increase of 50% in total weight weaned per ewe lambing twins has been reported (Rajab et al. 1992). The increase of one extra CL and 58% more lambs born observed in E/E ewes compared with +/+ was a strong evidence of the $FecG^E$ effect on ovulation rate control and prolificacy, and represents a new phenotype for GDF9 in sheep. Our parturition data point to an additive effect for the $FecG^{E}$ allele. despite no difference being observed in the ovulation rate between +/+ and E/+ ewes. However, the allele interactions of $FecG^{E}$ are certainly distinct from the over-dominant behaviour observed in $FecG^{H}$ and all FecX alleles described until now. The E/E pregnancy and parturition data confirm that their oocytes were viable and fertile; which correlate with the increased prolificacy (number of lambs/ewe) observed amongst these animals. In this study, for the first time, a new SNP that increased the ovulation rate and prolificacy of homozygote sheep was documented for the GDF9 gene. This new genetic variant, together with the other documented variants in GDF9 and BMP15, can be very useful to obtain a better understanding of the genetic control of ovulation rate in mammals. Moreover, this major gene variant can be applied in breeding programmes by gene-assisted selection (GAS), aiming towards the improvement of sheep reproductive potential and production. However, further investigation is necessary to shed light on the allelic interactions of the $FecG^E$ variant.

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Supporting Information

Additional supporting information may be found in the online version of this article.

Appendix S1 Additional data, materials and methods.

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