Identification of molecular markers on bovine chromosome 18 associated to calving interval in a Brangus-Ibagé cattle herd

Identificação de marcadores moleculares no cromossomo bovino 18 associados ao intervalo entre partos em um rebanho Brangus-Ibagé

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ABSTRACT

In the detection phase of a bovine marker assisted selection program, this paper investigated the genetic variability of three microsatellites on the chromosome 18 (BTA 18). The possible associations between genotypes or alleles of these markers versus weight at first calving and a lifetime calving interval (as indicators of reproductive performance) were evaluated in a beef cattle herd (5/8 Aberdeen Angus x 3/8 Nelore). Eleven alleles were detected in TGLA227 and ILSTS002 and three in BMS3004, the most frequent being TGLA227*79, ILSTS002*133, ILSTS002*135 and BMS3004*129. Polymorphic information content ranged from 0.41 to 0.84, while heterozygosity ranged from 49% to 86%, with an average value of 77%. The association analyses performed between genotype classes for the genetic markers versus weight at first calving indicated no significant result. Also, no correlation was observed between calving interval (CI) and TGLA227 genotypes. However, positive associations were detected between ILSTS002 and BMS3004 and CI. Animals carrying at least one ILSTS002*135 allele presented a CI about 39 days longer than the individuals with other genotypes; animals heterozygous for BMS3004 presented a CI about 35 days shorter than the homozygous. On these grounds, it can be concluded that these markers can be useful as an aid to fertility selection, in this herd.

Key words: microsatellites, reproduction, marker assisted selection, BTA 18.

RESUMO

Na fase de detecção, em um programa de seleção assistida por marcadores, em bovinos, este trabalho investigou a variabilidade genética de três microssatélites no cromossomo 18 (BTA 18). As possíveis associações entre genótipos ou alelos destes marcadores versus o peso ao primeiro parto e o intervalo entre partos (como indicadores do desempenho reprodutivo) foram avaliados em um rebanho de gado de corte (5/8 Aberdeen Angus x 3/8 Nelore). Onze alelos foram detectados em TGLA227 e ILSTS002 e três em BMS3004, os mais frequentes sendo TGLA227*79, ILSTS002*133, ILSTS002*135 e BMS3004*129. O conteúdo polimórfico de informação variou de 0,41 a 0,84, enquanto a heterozigosidade variou de 49% a 86%, com média de 77%.

As análises de associação efetuadas entre classes genotípicas dos marcadores genéticos versus o peso ao primeiro parto não indicaram resultados significantes. Da mesma forma, nenhuma correlação foi observada entre o intervalo entre partos (IEP) e os genotipos de TGLA227. Entretanto, associações positivas foram detectadas entre ILSTS002 e BMS3004 com o IEP. Animais portadores de pelo menos um alelo ILSTS002*135 apresentaram IEP cerca de 39 dias mais longo que os indivíduos com outros genótipos, e animais heterozigotos para BMS3004 apresentaram IEP cerca de 35 dias mais curto que os homozigotos. Assim, pode-se concluir que esses marcadores podem ser úteis como auxiliares na seleção para fertilidade, o rebanho em questão.

Palavras-chave: microssatélites, reprodução, seleção assistida por marcadores, BTA 18.

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A possible strategy to improve the productive performance in livestock is the use of marker assisted selection (MAS), which means selecting for molecular markers in or around genes involved directly or indirectly in the expression of one desired character, usually a multifactorial trait (DAVIS & DeNISE, 1998; SCHWERIN, 2001). According to DAVIS & DeNISE (1998) there are three phases in the development of MAS programs, the detection phase, the implementation phase; in the first one, direct markers or linked DNA polymorphisms are used to detect QTL segregation in a particular population.

Linkage analysis using short tandem repeats (STR) markers have been able to map a great number of QTL regions including loci responsible for diseases, carcass weight, milk production, conformation traits, and reproduction (for review see SCHWERIN, 2001).

In the detection phase of a bovine MAS program, the present study investigated the genetic variability of three STR mapped at bovine chromosome 18 (BTA 18) and evaluated the possible associations between genotypes of these markers and indicators of reproductive performance in a beef cattle herd.

Blood samples were collected from 166 cows with at least four parturitions records of a Brangus-Ibagé herd, a composite beef cattle (5/8 Aberdeen Angus x 3/8 Nelore) resulting from the crossing between Aberdeen Angus cows (Bos taurus) and Nelore bulls (Bos indicus). The synthetic breed was created by the Brazilian Agricultural Research Corporation (EMBRAPA Pecuária Sul, Bagé, RS, Brasil). Details of the population could be found in DUARTE et al. (2005). Genomic DNA was extracted from total blood by the method of MILLER et al. (1988). Three STR (BMS3004, ILSTS002 and TGLA227) mapped on BTA 18 (http://locus.jouy.inra.fr/cgi-bin/bovmap/intro.pl; access: 2006/04/20), the BMS3004, ILSTS002 and TGLA227 being at 1.7, 54.7, 84.7 cM, respectively (U.S. Meat Animal Research Center, http://www.marc.usda.gov/genome/ cattle/cattle.html; access: 2006/04/20). Among the loci mapped at this chromosome the luteinizing hormone beta polypeptide (LHβ) gene is directly involved on reproductive function. It is mapped at 48.2 cM (National Center for Biotechnology Information, http://www.ncbi.nlm.nih.gov; access: 2006/04/20), and therefore 46.5 and 6 cM distant from BMS3004 and ILSTS002, respectively. Many studies have indicated that STR downstream or upstream gene sequences of the luteinizing hormone beta gene are associated with reproductive function (for review see OLIVEIRA et al., 2002).

Allele and genotype frequencies were determined by gene counting, the parameters of genetic diversity within population [heterozygosity (H) and polymorphic information content (PIC)] were estimated according to NEI (1978) and BOTSTEIN et al. (1980), respectively. One-way analysis of variance was employed to test associations between STR genotypes and average lifetime CI or WFC, according to the model:

\[ Y_{ij} = \mu + A_i + e_{ij} \]

where: \( Y_{ij} \) is the CI or WFC phenotype of the jth individual; \( \mu \) is the effect of the population mean; \( A_i \) is the effect of the ith genotype class; and \( e_{ij} \) is the random error component.

Calving interval did not present normal distribution, and it was previously corrected by using natural logarithm transformation. When a significant association was detected, the Tukey post hoc test was used to identify the significant group. To compare the significant group detected by ANOVA with the other genotypes, CI was dichotomized into classes consisting of animals with CI below or above the anti-mode value (549 days) and the frequencies were compared between these two groups by a \( \chi^2 \) test. The odds ratio (OR) with 95% confidence interval (ci) was obtained by logistic regression analysis. All the statistic analyses were performed using the SPSS® for Windows™ software (SPSS Inc, version 10.0.5 (1999)).

Allele frequencies, genetic diversity parameters (PIC and H) and descriptive statistics for the CI and WFC are presented in table 1. Eleven alleles were detected in TGLA227 and ILSTS002 and three in BMS3004. The PIC and H values were relatively high, the average heterozygosity being 77%. The CI and WFC values did not differ among STR groups.

No association was found between genotype classes and WFC as well as between TGLA227 genotypes and CI. However, positive associations were detected between ILSTS002 and BMS3004 and CI. Animals carrying at least one ILSTS002*135 allele presented a CI about 39 days longer (\( P < 0.05; \) Table 2) than animals with the other genotypes (OR = 2.0, ci = 1.1-3.8, \( P < 0.05 \)). With respect to the BMS3004, heterozygous animals showed a CI about 35 days shorter (\( P < 0.054; \) table 6) than homozygous (OR = 1.8, ci = 1.0-3.5, \( P < 0.10 \)). The reliability of the employment of CI to classify beef cattle mating seasonally remains unclear as observed in other studies in the same population (OLIVEIRA et al., 2002) and WFC may not be useful as a marker for fertility because it can be affected by environmental conditions that heifers were exposed to before the first mating. Despite these limitations CI is the most helpful criterion available to minimize environmental effects on cow fertility.

According to BOVMAP, there are 123 loci mapped on BTA 18 (http://locus.jouy.inra.fr/cgi-bin/bovmap/intro.pl; access: 2006/04/20), the BMS3004, ILSTS002, TGLA227 being at 1.7, 54.7, 84.7 cM, respectively (U.S. Meat Animal Research Center, http://www.marc.usda.gov/genome/ cattle/cattle.html; access: 2006/04/20). Among the loci mapped at this chromosome the luteinizing hormone beta polypeptide (LHβ) gene is directly involved on reproductive function. It is mapped at 48.2 cM (National Center for Biotechnology Information, http://www.ncbi.nlm.nih.gov; access: 2006/04/20), and therefore 46.5 and 6 cM distant from BMS3004 and ILSTS002, respectively. Many studies have indicated that STR downstream or upstream gene sequences...
play a significant role in gene regulation, by altering the primary, secondary or tertiary structure of DNA, by binding to transcription or translation factors, or by affecting RNA edition; they could be also in enhancer regions, and influence gene expression by altering the transcription complex assemblage (review in Li al., 2004). Considering the pivotal role of LH on reproduction, we cannot avoid to speculate whether the associations observed in the present study might be reflecting the effect of these STR on regulation of the LH β-chain gene or the even the possibility that these associations would be due to linkage disequilibrium with some LH β-chain gene mutation. It is also possible that these markers are influencing other gene loci indirectly involved on reproduction and mapped at BTA 18.

The association observed with the BMS3004 suggests the occurrence of molecular heterosis, which occurs when heterozygous for a specific polymorphism show a significantly greater or lesser effect for a quantitative or qualitative trait than homozygous, probably because heterozygotes show a broader range of gene expression (Comings & MacMurray, 2000).

Although this paper developed only the MAS detection phase, and need go through the evaluation and implementation phases before its commercial use, it seems reasonable to suggest that ILSTS002 and BMS3004 markers could be useful to allow early selection of Brangus-Ibagé cows, improving the reproductive performance by reducing calving interval.
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