

Research Article

Genetic characterization of Aberdeen Angus cattle using molecular markers

Luciana Pimentel de Mello Klocker Vasconcellos¹, Daniella Tambasco-Talhari¹, Andréa Pozzi Pereira¹, Luiz Lehmann Coutinho² and Luciana Correia de Almeida Regitano³

 ¹Programa de Pós-Graduação em Genética e Evolução, Universidade Federal de São Carlos, São Carlos, SP, Brazil.
 ²Departamento de Produção Animal, Escola Superior de Agricultura Luiz de Queiroz, USP, Piracicaba, SP, Brazil.
 ³Laboratório de Biotecnologia Animal, Embrapa Pecuária Sudeste, São Carlos, SP, Brazil.

Abstract

Aberdeen Angus beef cattle from the Brazilian herd were studied genetically using restriction fragment length polymorphism (RFLP) of the κ -casein - *Hinfl* (*CSN*3 - *Hinfl*), β -lactoglobulin - *Hae*III (*LGB* - *Hae*III) and growth hormone *Alul* (*GH*- *Alul*) genes, as well as four microsatellites (TEXAN15, CSFM50, BM1224 and BM7160). The RFLP genotypes were determined using the polymerase chain reaction (PCR) followed by digestion with restriction endonucleases and electrophoresis in agarose gels. With the exception of the microsatellite BM7160, which was analyzed in an automatic sequencer, the PCR products were genotyped by silver staining. The allele and genotype frequencies, heterozygosities and gene diversity were estimated. The values for these parameters of variability were comparable to other cattle breeds. The genetic relationship of the Aberdeen Angus to other breeds (Caracu, Canchim, Charolais, Guzerath, Gyr, Nelore, Santa Gertrudis and Simmental) was investigated using Nei's genetic distance. Cluster analysis placed the Aberdeen Angus in an isolated group in the *Bos taurus* breeds branch. This fact is in agreement with the geographic origin of this breed.

Key words: Aberdeen Angus, bovine, DNA, characterization, RFLP, microsatellite.

Received: April 22, 2002; accepted: December 4, 2002.

Introduction

Various aspects of a population are useful in its characterization, including phenotypic traits (monogenic and polygenic), reproduction, geographic distribution, origin and habitat. The genetic characterization of populations, breeds and species allows the assessment of genetic variability, a crucial element in determining breeding strategies and genetic conservation programs.

The assessment of genetic variability is especially important in highly specialized livestock breeds since the use of assisted reproduction techniques, such as artificial insemination and embryo transfer, to maintain these breeds can rapidly reduce the genetic variability of the population. Molecular markers have been widely used to access this variability since they provide information on every region of the genome, regardless of the level of gene expression. RFLP and microsatellites (highly polymorphic simple sequence repeats) are currently the most widely used molecular markers, mainly because of the possibility of combining their analysis with the polymerase chain reaction (PCR). These markers have been used to explain bovine domestication and migration patterns (Loftus *et al.*, 1994; Bradley *et al.*, 1994) and to characterize cattle populations (MacHugh *et al.*, 1994; Kemenes *et al.*, 1999; Tambasco *et al.*, 2000). Another application of molecular markers is in uncovering parentage mistakes. This is especially important for guaranteeing the accuracy of breeding programs in which the relationship between individuals is used to estimate breeding value. For this application, the frequencies of each marker in the population must be known.

Aberdeen Angus cattle have been bred in Scotland for more than 400 years, although the presence of polled cattle in the counties of Aberdeen and Angus dates from prehistoric times, as suggested by prehistoric carvings found in these regions (http://www.ansi.okstate.edu/breeds/cattle/). This polled black breed of cattle is considered to have a long life span, high fertility, precocity and easy calving (Hermsdorf, 1941). The latter attribute makes Aberdeen

Send correspondence to Luciana Correia de Almeida Regitano. Rodovia Washington Luiz km 234, Caixa Postal 339, 13560-970, São Carlos, SP, Brasil. Fone: (0xx16) 261-5611. Fax: (0xx16) 261-5754. E-mail: luciana@cppse.embrapa.br.

Angus sires a good choice for first parity females. The polled phenotype is another attractive attribute since it reduces losses from injury to cohorts and allows for denser occupation. Aberdeen Angus cattle were introduced to Brazil in 1906 by Leonardo Colares Sobrinho in Bagé, in the state of Rio Grande do Sul. In southern Brazil, the breed is used in purebred schemes, whereas in southeast and midwest Brazil it is used in crosses with Nelore cattle.

The aim of the present work was to evaluate the genetic variability of the Brazilian herd of Aberdeen Angus cattle and it's genetic relationship to other beef cattle breeds used in Brazil, in order to provide information for beef cattle breeding programs.

Material and Methods

Animals

Fifty-two unrelated Aberdeen Angus animals were sampled from the Bela Vista farm at Pardinho, São Paulo state (N = 25), from artificial insemination centers (N = 4), and from two farms in Paraná state (N = 23). These animals were chosen to represent all of the sire lines used in the Brazilian Aberdeen Angus herd.

Markers

Four microsatellites (TEXAN15, CSFM50, BM1224 and BM7160) and three RFLPs (*CSN3 - Hinf*I, *LGB -Hae*III and *GH - Alu*I) were analyzed by PCR. The corresponding chromosomal location and primer sequences of these markers are given in Table 1.

Table 1 - Description of molecular markers analyzed.

DNA extraction

DNA was extracted from blood or semen samples using the salting out procedures described by Regitano (2001).

Genotyping

PCR reactions were done in a GeneAmp 2400 thermocycler with 200 ng of DNA in PCR solution (20 mM Tris-HCl, pH 8.4, 50 mM KCl) plus 1.5 mM MgCl₂, 0.2 µM of each dNTP, 0.4 µM of each primer and 0.5 U of Taq DNA polymerase in a final volume of 25 µL. For the microsatellite BM7160, the forward primer was labeled with fluorescein at the 5' end. The RFLP cycling conditions were those described in the references listed in Table 1. After digestion with the appropriate restriction enzyme, fragments were analyzed in 3% agarose gels in 1X TBE buffer, stained with ethidium bromide and photographed using an EDAS (Kodak TM) digital system. For microsatellite loci the cycles were the same as described by Bishop et al. (1994) and the PCR products were analyzed in 8% polyacrylamide gels in a manual sequencing apparatus, followed by silver staining (Comincini et al., 1995). Genotypes for the locus BM7160 were determined with an automatic laser fluorescence DNA sequencer (Pharmacia Biotech).

Statistical analysis

Gene and genotype frequencies were estimated according to Weir (1996). Hardy-Weinberg equilibrium was tested using an exact probability test (Guo and Thompson, 1992). Heterozygosity (H) and gene diversity (D) were estimated as genetic variability parameters (Weir, 1996). Polymorphic information content (PIC) was estimated according to Botstein *et al.*, (1980). Nei's genetic distance

Marker Nature CSN3 ² RFLP <i>Hinf</i> I		Chromosome	Sequence 5'- 3' ¹	Reference	
		6	ATCATTTATggCCATTCCACCAAAC gCCCATTTCgCCTTCTCTgTAACAGA	1	
LGB ³	RFLP HaeIII	11	gTACTTgTggTggACACCgACTACA CAgCAgACCggCTCCTggTATATgA	2	
GH^4	RFLP AluI	19	gCTgCTCCTgAgggCCCCTTCg gCggCggCACTTCATgACCCT	3	
BM1224	MS	4	AggAACCACATTgggTAgTCC TCCCTCTCTCCCTgAggC	4	
TEXAN15	MS	5	TCgCAAACAgTCAgAgACCACTC TggATgAgAAAgAAgAgCAgAgTTg	5	
CSFM50	MS	2	AgTTCTCCTCTTgATTTCAgTAAC CCTACTTCCTgCCTTTgTAgCATA	6	
BM7160 MS		7	TggATTTTAAACACAgAATgTgg TCAgCTTCTCTTTAAATTTCTCTgg	7	

¹Sequence of forward primer is given in the upper line of each pair. The bottom line corresponds to the reverse primer. RFLP = restriction fragment length polymorphism, MS = microsatellite. ${}^{2}\kappa$ -casein, ${}^{3}\beta$ -lactoglobulin, 4 Growth hormone.

1. Medrano and Aguilar-Cordova (1990), 2. Ron et al. (1994), 3. Schlee et al. (1994^a), 4. Bishop et al. (1994), 5. Burns et al. (1995), 6. Moore et al. (1994), 7 Stone et al. (1995).

(Nei, 1978) was used to investigate the relationship between the Aberdeen Angus breed and the cattle breeds Caracu, Canchim, Charolais, Guzerath, Gyr, Nelore, Santa Gertrudis and Simmental. This data was used for cluster analysis by the unweighted pair-group method using an arithmetic mean (UPGMA).

Results and Discussion

The allele frequencies observed for the seven markers are listed in Table 2, as are the frequencies reported for other breeds in the literature. The only significant departure from Hardy-Weinberg equilibrium (p < 0.05) was for the GH - Alu locus, which resulted from an excess of homozygotes. This disequilibrium may reflect a series of events such as inbreeding, selection, genetic drift or population subdivision. The latter event could also result from sampling few individuals from each location, although this would be expected to affect more than one locus. Several reports on the effects of GH - AluI polymorphism suggest that this is not a neutral mutation for production traits (Schlee *et al.*, 1994b; Pereira *et al.*, 2002). However, the data obtained here was insufficient to assess this hypothesis in our Aberdeen Angus sample. The microsatellite loci were polymorphic and the number of alleles ranged from 3

Table 2 - Allele frequencies for the seven markers in ten bovine populations.

Marker	Allele	An 52	Si 52	Ne 63	Cn ¹ 30	Gi ¹ 83	Gu ¹ 25	Ch1 ¹ 30	Ch2 36	$\frac{SG^1}{20}$	Ca^1 30
CSN3	А	0.82	0.65	0.91	0.63	0.93	0.92	0.48	0.51	0.85	0.68
GH	L	0.77	0.82	1.00	0.90	1.00	1.00	0.72	0.74	0.97	0.80
LGB	A	0.17	0.46	0.40	0.40	0.37	0.34	0.55	0.54	0.17	0.57
TEXAN15	203	_	0.01	0.57	0.13	0.53	0.28	_	_	0.20	0.02
	205	0.02	0.05	0.03	-	0.19	0.05	-	-	0.32	0.07
	207	0.05	0.16	0.20	0.33	0.07	0.60	0.07	0.08	0.10	0.18
	209	0.19	0.36	0.03	0.17	0.06	0.05	0.08	0.13	0.20	0.25
	217	0.23	0.01	0.09	0.33	-	0.02	0.54	0.53	0.18	0.37
	219	0.14	-	-	-	0.07	-	0.07	0.02	-	-
	221	0.13	-	-	-	0.02	-	-	0.08	-	-
	others	0.24	0.41	0.08	0.04	0.06	0.00	0.24	0.16	0.00	0.11
CSFM50	168	-	0.01	0.32	0.18	0.38	0.47	0.05	0.04	0.15	0.15
	170	-	0.02	0.16	0.07	0.22	0.28	-	-	-	-
	172	0.63	0.11	0.22	0.23	0.07	0.07	0.17	0.19	0.15	0.15
	176	0.29	0.66	0.28	0.40	0.20	0.15	0.53	0.51	0.58	0.57
	others	0.08	0.20	0.02	0.12	0.13	0.03	0.25	0.26	0.12	0.13
BM1224	167	-	-	-	-	0.08	0.25	-	-	0.37	-
	169	0.02	0.11	0.07	0.43	0.20	0.15	0.02	0.04	-	-
	171	0.06	0.11	-	0.03	0.01	0.08	0.03	0.03	0.05	0.43
	173	0.12	0.23	0.13	-	-	0.03	0.15	0.18	0.15	0.03
	175	0.15	0.07	-	-	0.07	0.15	0.03	0.03	0.13	-
	177	0.42	0.11	0.30	0.35	0.04	0.07	0.43	0.42	-	0.35
	179	0.12	0.07	-	-	0.16	0.08	-	-	0.05	-
	181	0.08	0.02	0.45	0.03	0.30	0.10	0.13	0.14	0.15	0.03
	183	-	0.14	0.04	0.15	0.01	0.07	0.15	0.11	0.10	0.15
	others	0.03	0.14	0.01	0.01	0.13	0.02	0.06	0.05	0.00	0.01
BM7160	173	0.06	0.25	-	-	-	-	-	-	-	-
	175	0.12	0.03	-	-	-	-	-	-	-	-
	177	0.31	0.54	-	-	-	-	-	-	-	-
	179	0.33	0.16	-	-	-	-	-	-	-	-
	others	0.18	0.02	-	-	-	-	-	-	-	-

An - Aberdeen Angus, Si - Simmental (Regitano *et al.*, 2000), Ch - Charolais (Ch1 - Garcia, 2001 and Ch2 - Regitano *et al.*, 2000), Ca - Caracu, Cn - Canchim, Ne - Nelore, Gi - Gyr, Gu - Guzerath, SG - Santa Gertrudis. ¹Marker data for RFLP from Kemenes *et al.* (1999) and for microsatellites from Garcia (2001).

(CSFM50) to 12 (TEXAN15). Allele sizes and frequencies are shown in Table 2.

The parameters H and D have been used to assess the genetic variability of a population. The values obtained for Aberdeen Angus cattle based on the seven markers are shown in Table 3. The Aberdeen Angus breed is a highly specialized beef breed in which intense use is made of artificial insemination and embryo transfer, and this could result in a decrease in genetic variability. However, the average heterozygosity (H = 0.491) and the gene diversity (D = 0.554) determined here do not indicate such a reduction. The PIC values for each marker were estimated in order to assess the relevance of each locus for linkage analysis (Table 3). With the exception of CSFM50, the microsatellite loci would provide an excellent contribution to a genome scan with more than 70% of the meiosis expected to be informative in this population.

The genetic distances between the Aberdeen Angus population and eight other breeds used in Brazil were determined using published frequency data - Simmental (Regitano *et al.*, 2000), Charolais (Regitano *et al.*, 1999), Nelore, Gyr, Guzerath, Charolais, Santa Gertrudis, Canchim and Caracu RFLP data from Kemenes *et al.* (1999)

 Table 3 - Genetic diversity (D), heterozygosity (H) and polymorphic information content (PIC) estimates for Aberdeen Angus cattle.

Marker	Number of alleles	Н	D	PIC
CSN3	2	0.288	0.302	0.254
LGB	2	0.346	0.358	0.292
GH	2	0.192	0.289	0.245
TEXAN15	12	0.840	0.866	0.842
CSFM50	3	0.596	0.524	0.447
BM1224	8	0.608	0.769	0.736
BM7160	8	0.569	0.770	0.728

Table 4 - Nei's (1978) genetic of	distances.	
-----------------------------------	------------	--

	An (52)	Si (52)	Ne (63)	Cn (30)	Gi (83)	Gu (25)	Ch1 (30)	Ch2 (36)	SG (29)	Ca (30)
An	0.00									
Si	0.20	0.00								
Ne	0.21	0.24	0.00							
Cn	0.17	0.14	0.16	0.00						
Gi	0.27	0.25	0.05	0.19	0.00					
Gu	0.29	0.26	0.11	0.17	0.09	0.00				
Ch1	0.22	0.17	0.29	0.13	0.39	0.42	0.00			
Ch2	0.20	0.16	0.27	0.12	0.36	0.39	0.01	0.00		
SG	0.18	0.15	0.16	0.19	0.13	0.16	0.28	0.26	0.00	
Ca	0.20	0.10	0.23	0.12	0.29	0.28	0.10	0.09	0.21	0.00

An - Aberdeen Angus, Si - Simmental, Ch - Charolais (Ch1 - Garcia, 2001 and Ch2 - Regitano *et al.*, 2000), Ca - Caracu, Cn - Canchim, Ne - Nelore, Gi - Gyr, Gu - Guzerath, SG - Santa Gertrudis. Sample size in parentheses.

and microsatellite data from Garcia (2001) - (Table 4). The dendrogram obtained for these distances using the UPGMA clustering method (Figure 1) clearly distinguished the two main groups of cattle (*Bos indicus* and *Bos taurus*) and placed the crossbred Canchim (5/8 Charolais+3/8 *Bos indicus*) in the *Bos taurus* group. In this analysis, Aberdeen Angus was placed in an isolated group in the *Bos taurus* branch. Since Aberdeen Angus is a Scottish breed, it was expected to be less related to continental European breeds, such as Charolais and Simmental, because of its geographic separation.

In conclusion, genetic marker analysis showed that the genetic variability of the Brazilian population of Aberdeen Angus was comparable to that of other cattle

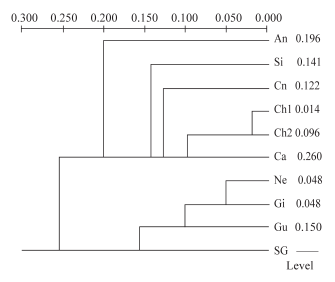


Figure 1 - Dendrogram obtained based on the UPGMA clustering of Nei's (1978) genetic distances. An -Aberdeen Angus, Si - Simmental, Ch1 - Charolais pop 1 (Garcia, 2001), Ch2 - Charolais pop 2 (Regitano *et al.* 2000b), Ca - Caracu, Cn - Canchim, Ne - Nelore, Gi - Gyr, Gu - Guzerath, SG - Santa Gertrudis.

breeds in Brazil. Genetic distance analysis separated the Aberdeen Angus breed from other taurine breeds, a result consistent with their geographic origins.

Acknowledgements

The authors thank Jovelino Carvalho Mineiro (Bela Vista Farm) for providing blood samples. This work was supported by Embrapa and Fapesp.

References

- Bishop MD, Kappes SD, Keele JW, Stone RT, Sunden SLF, Hawkins GA, Solinas Toldo S, Fries R, Grosz MD, Yoo J and Beattie CW (1994) A genetic linkage map for cattle. Genetics136:619-639.
- Botstein D, White RL, Skolnick M and Davis RW (1980). Construction of genetic linkage map in man using restriction fragment length polymorphisms. Am J Hum Genet 32:314-331.
- Bradley DG, MacHugh DE, Loftus RT, Sow RS, Hoste CH and Cunningham EP (1994) Zebu-taurine variation in Y chromosomal DNA: a sensitive assay for genetic introgression in West African trypanotolerant cattle populations. Anim Genet 25:7-12.
- Burns BM, Taylor JF, Herring AD, Holder MT, Collins JF, Guerra TM and Sanders JO (1995). Bovine mirosatellite dinucleotide repeat polymorphisms at the TEXAN11, TEXAN12, TEXAN13, TEXAN14 and TEXAN15 loci. Anim Genet 26:201-202.
- Comincini S, Leone P, Redaelli L, De Giuli L, Zhang Y and Ferretti, L (1995) Characterization of bovine microsatellites by silver staining. J Anim Breed Genet 112:415-420.
- Garcia MCC (2001) Estudo da diversidade genética entre sete raças de bovinos pelo uso de marcadores microssatélites. PhD Thesis, Universidade Estadual Paulista, Botucatu.
- Guo SW and Thompson EA (1992) Performing the exact test of Hardy-Weinberg proportion for multiple alleles. Biometrics 48:361-372.
- Hermsdorff GE (1941) Zootecnia Especial Bovinos. Etnologioa das Raças Européias que mais Interessam ao Brasil. Rio de Janeiro, Imprensa Nacional, v. 2, 387 p.
- Kemenes PA, Regitano LCA, Rosa AJM, Packer IU, Razook AG, Figueiredo LA, Silva NA, Etchegaray MAL and Coutinho LL (1999) κ-casein, β-lactoglobulin and growth hormone allele genetic distances in Nelore, Gyr, Guzerá, Caracu, Charolais, Canchim and Santa Gertrudis cattle. Genet Mol Biol 22:539-541.
- Loftus RT, MacHugh DE, Bradley DG, Sharp PM and Cunningham P (1994) Evidence for two independent domestications of cattle. Proc Natl Acad Sci USA 91:2757-2761.
- MacHugh DE, Loftus RT, Bradley DG, Sharp PM and Cunningham P (1994) Microsatellite DNA variation within

and among European cattle breeds. Proc R Soc Lond B 256:25-31.

- Medrano JF and Aguilar-Cordova E (1990) Genotyping of bovine kappa-casein loci following DNA sequence amplification. Bio/Technology 8:144-146.
- Moore SS, Byrne K, Berger KT, Barendse W, McCarthy F, Womack JE and Hetzel DJS (1994) Characterization of 65 bovine microsatellites. Mamm Genome 5:84-90.
- Nei M (1978) Estimation of average heterozygosity and genetic distance from a small number of individuals. Genetics 89:583-590.
- Pereira AP, Alencar MM, Oliveira HN and Regitano LCA (2002) Association of GH and IGF-1 polymorphisms to growth traits in a synthetic beef cattle breed. In: 28th Conference of the International Society of Animal Genetics, Goettingen, Germany.
- Regitano LCA (2001) Extração de DNA para aplicação em reação em cadeia da polimerase (PCR). In: Regitano LCA and Coutinho LL (eds) Biologia Molecular Aplicada à Produção Animal. Embrapa, Brasilia, DF, pp 179-186.
- Regitano LCA, Azevedo JL, Vencovsky R, Packer IU, Barbosa PF, Rosa AJM, Silva NA, Etchegaray MAL and Coutinho LL (1999) Selection for breed-specific growth hormone and IGF-I alleles in synthetic beef cattle cross, Canchim. Genet Mol Biol 22:531-537.
- Regitano LCA, Vasconcellos LPMK, Jacinto E, Tambasco-Studart M, Tambasco DD, Euclides Filho K, Barbosa PF, Packer IU and Coutinho LL (2000) Genetic distances among Aberdeen Angus, Canchim, Caracu, Nellore and Simmental beef cattle breeds. In: Global Conference on Conservation of Domestic Animal Breeds, 5, 2000, Brasília, DF. Proceedings... Brasília, DF: Embrapa Genetic Resources and Biotechnology, 2000a 1 CD-ROM.
- Ron M, Yoffe O, Ezra E, Medrano JF and Weller JI (1994) Determination of effects of milk protein genotyping on production traits of Israeli Holstein. J Dairy Sci 77: 11061113.
- Schlee P, Graml R, Rottmann O and Pirchner F (1994a) Influence of growth-hormone genotypes on breeding values of Simmental bulls. J Anim Breed Genet 111:253-256.
- Schlee P, Graml R, Schallenberger E, Schams D, Rottmann O, Olbrich-Bludau A and Pirchner F (1994b). Growth hormone and insulin-like growth factor I concentrations in bulls of various growth hormone genotypes. Theor Appl Genet 88:497-500.
- Stone RT, Pulido JC, Duyk GM, Kappes SM, Keele JW and Beattie CW (1995) A small-insert bovine genomic library highly enriched for microsatellite repeat sequences. Mamm Genome 6:714-724.
- Tambasco DD, Alencar MM, Coutinho LL, Tambasco AJ, Tambasco MD and Regitano LCA (2000) Caracterização molecular de animais da raça Nelore utilizando microssatélites e genes candidatos. Rev Bras Zootec 29:1044-1049.
- Weir BS (1996). Genetic Data Analysis: Methods for Discrete Population Genetic Data. 2nd edition. Sinauer Associates, Massachusetts, 445 p.