

Genetic parameters for stayability to consecutive calvings in Zebu cattle

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Longer-lived cows tend to be more profitable and the stayability trait is a selection criterion correlated to longevity. An alternative to the traditional approach to evaluate stayability is its definition based on consecutive calvings, whose main advantage is the more accurate evaluation of young bulls. However, no study using this alternative approach has been conducted for Zebu breeds. Therefore, the objective of this study was to compare linear random regression models to fit stayability to consecutive calvings of Guzerá, Nelore and Tabapuã cows and to estimate genetic parameters for this trait in the respective breeds. Data up to the eighth calving were used. The models included the fixed effects of age at first calving and year-season of birth of the cow and the random effects of contemporary group, additive genetic, permanent environmental and residual. Random regressions were modeled by orthogonal Legendre polynomials of order 1 to 4 (2 to 5 coefficients) for contemporary group, additive genetic and permanent environmental effects. Using Deviance Information Criterion as the selection criterion, the model with 4 regression coefficients for each effect was the most adequate for the Nelore and Tabapuã breeds and the model with 5 coefficients is recommended for the Guzerá breed. For Guzerá, heritabilities ranged from 0.05 to 0.08, showing a quadratic trend with a peak between the fourth and sixth calving. For the Nelore and Tabapuã breeds, the estimates ranged from 0.03 to 0.07 and from 0.03 to 0.08, respectively, and increased with increasing calving number. The additive genetic correlations exhibited a similar trend among breeds and were higher for stayability between closer calvings. Even between more distant calvings (second v. eighth), stayability showed a moderate to high genetic correlation, which was 0.77, 0.57 and 0.79 for the Guzerá, Nelore and Tabapuã breeds, respectively. For Guzerá, when the models with 4 or 5 regression coefficients were compared, the rank correlations between predicted breeding values for the intercept were always higher than 0.99, indicating the possibility of practical application of the least parameterized model. In conclusion, the model with 4 random regression coefficients is recommended for the genetic evaluation of stayability to consecutive calvings in Zebu cattle.

Keywords: genetic parameters, legendre polynomials, longevity, reproductive traits, Zebu cattle

Implications

The importance of Zebu is related to their adaptation to the tropical climate, to pasture-based rearing systems and to high infestation with parasites. Stayability, a trait related to longevity, is one of the most important trait in beef cattle under the economic aspect. However, the phenotype of a cow for this trait is obtained around 7 years of age. Thus, bulls will have an advanced age when their daughters

manifest the phenotype, resulting in low accuracy of genetic evaluations for younger bulls. The methodology proposed here could overcome this problem, promoting higher genetic improvement at a shorter period of time.

Introduction

In beef cattle farming, the cow is responsible for the effective size of the herd in subsequent years and special attention should therefore be paid to this component of the production system. The time cows stay in the herd is generally determined by a set of productive and reproductive factors (Bertazzo *et al.*, 2004) that influence the decision of

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producers to keep or cull a cow. Longer-lived cows tend to be more profitable because of their greater capacity to return the capital invested in their rearing (Queiroz *et al.*, 2007). For a herd to be considered profitable, the number of cows that stay after the period of payment of their production cost must be greater than that of females that have not yet provided return on the capital invested (Snelling *et al.*, 1995).

Stayability (STAY) is defined as the ability of a cow to survive until a specific age given the opportunity to reach this age (Hudson and Van Vleck, 1981). It is a particularly important trait in beef cattle and has received major attention from researchers in recent years because its inclusion in breeding programs could allow the selection of sires whose daughters have a greater probability of remaining productive in the herd for a long period of time (Van Melis *et al.*, 2007). The main problem that has been pointed out for this trait is that a female should reach a reference age (between 6 and 7 years depending on the definition) to manifest a phenotype. Thus, bulls will have an advanced age when their daughters manifest the phenotype, a fact resulting in low accuracy of genetic evaluations for younger bulls (Hudson and Van Vleck, 1981; Jamrozik *et al.*, 2013).

An interesting alternative to defining and analyzing STAY has been reported by Jamrozik *et al.* (2013). The authors proposed the use of a random regression model for the evaluation of STAY to consecutive calvings and demonstrated its application using data from Canadian Simmental cattle. The advantages of this approach include the fact that the phenotypes can be assigned to each period in the lifetime of the cow, the breeding values for STAY can be estimated for each point of this trajectory, and the time/age-dependent environmental effects are easily implemented in the model (Jamrozik *et al.*, 2013).

The primary importance of Zebu breeds in the tropics is related to their adaptation to the tropical climate, to extensive rearing systems and to high infestation with ecto- and endoparasites (Albuquerque *et al.*, 2007). To our knowledge, no study has so far evaluated STAY to consecutive calvings in Zebu breeds. Therefore, the objective of the present study was to compare linear random regression models to fit STAY to consecutive calvings of Guzerá, Nelore and Tabapuã cows and to estimate genetic parameters for this trait in the respective breeds.

Material and methods

The databases analyzed in the present study were from the National Zootechnical Archive, managed by Embrapa Gado de Corte, Brazil, and contained data of genealogical records of three *Bos indicus* breeds: Guzerá, Nelore and Tabapuã cattle. The herds are located throughout all regions of Brazil. In Brazil, the animals of all Zebu breeds are raised mainly on pastures, regardless the fact they are selected for milk production, meat production or both. The database has records from farms who perform a mating season (the mating occurs only in a small season of the year) and farms where the mating occurs along the entire year. Therefore, all of the

consistency of the records was performed taking into account this particularity.

The trait studied was STAY to consecutive calvings as defined by Jamrozik *et al.* (2013). The following restrictions were applied to the inclusion of cows in the study: age at first calving between 20 and 60 months, age at calving between 20 and 240 months, and an interval between consecutive calvings of 305 to 820 days (take into account cow that failed to conceive in some year and stayed in the herd, a common practice in the herds studied). Cows submitted to embryo transfer or *in vitro* fertilization at some point in their life were excluded from the analyzes. After applying these restrictions and considering the number of records for all calvings (Supplementary Material Table S1), data up to the eighth calving were used as the relative frequency of animals was very low in subsequent calvings.

The phenotypes were assigned by observing whether the cow met the conditions cited above and had the opportunity to stay in the herd up to a given calving. Each cow could have up to seven repeated observations (stayability or not to calving 2, 3, ... and 8). A phenotypic value 1 (success) was established for cows that had the respective calving and 0 (failure) for those that had the opportunity but did not calve. The last calving record (LCR) on the database was the most recent information available. For a cow without a record of it next calving until the date of the LCR, with an interval between the LCR and that date of the LCR available for this cow equal to 820 days or less, the subsequent phenotypes were considered 'censored' and treated as missing data in the model. To illustrate the assignment of these phenotypes, we may consider the example of four cows, called A, B, C and D. Cow A had eight calvings, so its vector of observations would be A = [1, 1, 1, 1, 1, 1]. For cow B, four calvings (first to fourth) were observed and no further information about this cow was found given the possibility of manifesting the phenotype to subsequent calvings. Thus, its vector of observations would be B = [1,1,1,0,0,0,0]. In the case of cow C, the existence of four calvings was also verified, but this cow was found to be still young in relation to the LCR of the database (did not have the opportunity to manifest the remaining calvings). It was therefore not possible to determine whether or not the cow would stay in the herd. Consequently, the data of cow C were censored after the fourth calving (considered missing data) and its vector of observations was C = [1,1,1]. For cow D, only the record of the first calving was found given the possibility of the cow to manifest the phenotype to subsequent calvings. Its vector of observations would therefore be D = [0,0,0,0,0,0,0]. The description of distribution of observations according to calving and breed are shown in Table 1. The final data sets had 61 650 (1130), 153 709 (898) and 75 193 (727) cows (herds) with phenotypic records, daughters of 2811, 7295, 3961 sires, for the Guzerá, Nelore and Tabapuã cattle, respectively. The number of animals in the pedigree file for the Guzerá, Nelore and Tabapuã cattle were 85757, 332 616, 90 100, respectively. The average equivalent complete generation (Maignel et al., 1996) were 4.05, 3.21

Table 1 Distribution of the cows according to the number of phenotypic records (0 or 1)

	Guz	erá	Nelo	re	Tabapuã		
Number of records	NC	RP	NC	RP	NC	RP	
1	4838	7.85	13 122	8.53	5437	7.23	
2	3122	5.06	9827	6.40	3715	4.94	
3	2110	3.42	6868	4.46	2913	3.87	
4	1471	2.39	4813	3.13	2216	2.95	
5	938	1.52	3160	2.05	1495	1.99	
6	631	1.02	2115	1.37	996	1.32	
7	48 540	78.73	113 804	74.04	58 421	77.69	

NC = number of cows; RP = relative percentage.

and 2.69, for Guzerá, Nelore and Tabapuã, respectively, indicating reasonable knowledge of the pedigree of animals participating in the analyses.

Four seasons of birth were defined for analysis: February, March and April; May, June and July; August, September and October; November, December and January. The cows were divided into classes according to their age at first calving as follows: 20 to 24, 25 to 27, 28 to 30, 31 to 33, 34 to 36, 37 to 39, 40 to 42, 43 to 45, 46 to 48, 49 to 51, 52 to 54, 55 to 57 and 58 to 60 months. Random regression models for fitting the trait over the interval from the second to eighth calving were compared. The general model used was:

$$y_{ijklt} = \mathsf{YS}_{it} + \mathsf{AFC}_{jt} + \sum_{m=1}^{o} \beta_{km} \varphi_{mt} + \sum_{m=1}^{o} \alpha_{lm} \varphi_{mt} + \sum_{m=1}^{o} \delta_{lm} \varphi_{mt} + \epsilon_{ijklt},$$

where y_{iiklt} is the phenotype for STAY of cow / in calving t, YS_{it} the systematic effect of year-season of birth subclass *i* in calving t; AFC_{jt} the systematic effect of age at first calving class j in calving *t*, β_{km} the regression coefficient for the random effect of each contemporary group k, defined as herd, year and season of birth; α_{lm} and δ_{lm} the regression coefficients for random additive genetic and permanent environmental effects of each cow *l*; ϕ_{mt} the covariate of the Legendre polynomial for each regression coefficient in calving t, o the number of regression coefficients of the orthogonal Legendre polynomial used and ε_{iiklt} the random error associated with each observation. The random regression models were fitted using orthogonal Legendre polynomials of order 1 to 4 (2 to 5 coefficients, respectively). In each model, the same order was used for contemporary group, additive genetic and permanent environmental effects, so that each effect had the same opportunity to explore the spaces of the curves (Jamrozik et al., 2013). A heterogenous residual variance structure was adopted, with one residual variance component for each calving.

The matrix representation of the model is:

$$\mathbf{y} = \mathbf{X}\mathbf{b} + \mathbf{C}\mathbf{g} + \mathbf{Z}\mathbf{a} + \mathbf{W}\mathbf{p} + \mathbf{e}$$

where **y** is the vector of observations; **b** the vector of systematic effects (YS, AFC); **g**, **a**, **p** and **e** the vectors of

random contemporary group, additive genetic, permanent environment and residual effects, respectively; **X**, **C**, **Z** and **W** the incidence matrices corresponding to the observations for systematic, contemporary group, random additive genetic and permanent environmental effects, respectively. The following assumptions were defined for this model:

$$\mathbf{y} \mid \mathbf{b}, \mathbf{g}, \mathbf{a}, \mathbf{p}, \mathbf{R} \sim \mathsf{NMV}(\mathbf{X}\mathbf{b} + \mathbf{C}\mathbf{g} + \mathbf{Z}\mathbf{a} + \mathbf{W}\mathbf{p}, \mathbf{R}),$$

$$\mathsf{Var} \begin{bmatrix} \mathbf{g} \\ \mathbf{a} \\ \mathbf{p} \\ \mathbf{e} \end{bmatrix} = \begin{bmatrix} \Lambda_{\mathbf{G}} \otimes \mathbf{I} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \Lambda_{\mathbf{A}} \otimes \mathbf{A} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \Lambda_{\mathbf{P}} \otimes \mathbf{I} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{0} & \mathbf{R} \end{bmatrix},$$

where Λ_{G} , Λ_{A} and Λ_{P} are contemporary group, additive genetic and permanent environmental covariance matrices between the regression coefficients, respectively; **R** the 7 × 7 diagonal residual variance matrix, with one specific variance for each calving; **A** the numerator relationship matrix between animals; **I** the identity matrix and \otimes the direct product operator between matrices.

Analysis was performed by the Bayesian method using the GIBBS3F90 program (Misztal *et al.*, 2002). The following prior distributions were assumed for the model:

$$\mathbf{b} \propto \mathsf{constant}$$

 $\begin{array}{l} \mathbf{g} \mid \Lambda_{\mathbf{G}} \sim \mathsf{MVN}(\mathbf{0}, \Lambda_{\mathbf{G}} \otimes \mathbf{I}), \\ \mathbf{a} \mid \Lambda_{\mathbf{A}} \sim \mathsf{MVN}(\mathbf{0}, \Lambda_{\mathbf{A}} \otimes \mathbf{A}), \\ \mathbf{p} \mid \Lambda_{\mathbf{P}} \sim \mathsf{MVN}(\mathbf{0}, \Lambda_{\mathbf{P}} \otimes \mathbf{I}), \\ \Lambda_{\mathbf{G}} \mid \mathbf{v}_{g}, \mathbf{S}_{g} \sim \mathsf{IW}(\mathbf{v}_{g}, \mathbf{v}_{g}\mathbf{S}_{g}), \\ \Lambda_{\mathbf{A}} \mid \mathbf{v}_{a}, \mathbf{S}_{a} \sim \mathsf{IW}(\mathbf{v}_{a}, \mathbf{v}_{a}\mathbf{S}_{a}), \\ \Lambda_{\mathbf{P}} \mid \mathbf{v}_{p}, \mathbf{S}_{p} \sim \mathsf{IW}(\mathbf{v}_{p}, \mathbf{v}_{p}\mathbf{S}_{p}), \\ \mathbf{R} \mid \mathbf{v}_{e}, \mathbf{S}_{e} \sim \mathsf{IW}(\mathbf{v}_{e}, \mathbf{v}_{e}\mathbf{S}_{e}), \end{array}$

where MVN and IW indicate multivariate normal and inverse Wishart distributions, respectively, and v_{g} , S_{g} , v_{a} , S_{a} , v_{p} , S_{p} , and v_{e} , S_{e} correspond to the degree of confidence and prior values for contemporary group, additive genetic, permanent environmental and residual covariances, respectively.

An additional analysis was performed for each breed using a threshold model for a classical definition of STAY published in one of Zebu sire summaries in Brazil (Lôbo *et al.*, 2016), which is the ability of a cow to stay in the herd up to 76 months of age and to have at least three calvings. The cow received the phenotypic value 1 (success) if it met the cited criterion, and 0 (failure) when it had the opportunity but failed to meet the criterion established. For this trait, an animal model was used and included the systematic effects of age at first calving and year-season of birth class and random contemporary group, additive genetic and residual effects. These analyses were performed with the THRGIBBS2F90 software (Misztal *et al.*, 2002) and the heritability estimates and respective 95% highest posterior density intervals (HPD_{95%}) were calculated.

For comparison with the heritabilities obtained with the threshold models, the estimates of the linear random

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regression models were transformed to an underlying normal scale as suggested by Robertson and Lerner (1949):

$$h_u^2 = \frac{h_1^2 p_i (1-p_i)}{z}$$

where h_u^2 is the heritability on the underlying normal scale; h_1^2 is the heritability obtained by random regression on the binary data; p_i is the frequency of success (phenotype 1) for the trait in calving *i*, and *z* is the height of the ordinate at the point of truncation with area p_i below the normal curve.

The following criteria were used for comparison of the random regression models:

- 1. Correlation (R) between the phenotype predicted with the respective model and the observed binary phenotype.
- 2. Mean absolute value of residuals (RES).
- 3. Deviance information criterion (DIC) proposed by Spiegelhalter et al. (2002): $DIC=\overline{D}(\theta) + p_D = 2\overline{D}(\theta) + D(\overline{\theta})$, where $\overline{D}(\theta) = E_{\theta|Y}[D(\theta)]$ (posterior expectation of Bayesian deviance) and $D(\theta) = -2\ln p(y|\theta)$. The effective number of parameters represents a penalty with increasing complexity of the model: $p_D = \overline{D}(\theta) - D(\overline{\theta})$, where θ is the vector of parameters of the model and $D(\overline{\theta})$ is the Bayesian deviance evaluated at the posterior means of the parameters.
- 4. Bayesian deviance: $D(\theta) = -2\ln p(y|\theta)$, where $p(y|\theta)$ is the posterior marginal density.

For DIC and $D(\theta)$, lower values indicate more adequate models.

As complementary analysis for comparison of the models, the rank correlations between animals were calculated, classifying them according to the predicted breeding values for the intercept coefficient obtained with each model.

A chain with 300 000 samples was generated for each model, with a burn-in period of 30 000 samples and a thinning interval of 10. Convergence was monitored by graphic inspection of the parameter estimates across iterations, as well as by the criteria of Geweke (1992) and Raftery and Lewis (1992), using the *boa* package (Smith, 2007) of the R software (R Development Core Team, 2014). Thus, the variance components for the regression coefficients and genetic parameters (heritability and genetic correlation) were estimated from the remaining 27 000 samples. Heritability was calculated as the proportion of additive genetic variance in relation to the sum of contemporary group, additive genetic, permanent environmental and residual variances.

Results and discussion

The means of STAY to consecutive calvings showed the same trend in the Guzerá, Nelore and Tabapuã breeds (Figure 1). The cull rate of primiparous cows between the first and second calving was 43%, 27% and 34% for Guzerá, Nelore and Tabapuã, respectively. The cumulative cull rate until the eighth calving was ~91%, 97% and 96% for Guzerá, Nelore and Tabapuã cows, respectively. Mean STAY was high in early calvings and similar among breeds, but the cull rates between two consecutive calvings were higher than those



Figure 1 Average stayability to consecutive calvings according to breed.

between later calvings of the longitudinal scale. This higher cull rate in early calvings may be explained by the occurrence of problems mainly related to reproduction of primiparous cows, as well as by low maternal ability of the cows and selective culling for the inclusion of genetically superior females in the herd. Differences in STAY between breeds in each calving may indicate differences in productive and reproductive performance and their influence on the maintenance or culling of cows, or even differences in the criteria or intensities of selection practiced by breeders. In a study involving crossbred Angus × Hereford and Bos indicus × Hereford cows, Riley et al. (2001) observed that reproductive failure was the main cause of culling, regardless of calving number. Martinez et al. (2005), who analyzed STAY up to calvings 2 to 6 as distinct traits in Hereford cows, found results that were similar in magnitude for STAY to early calvings and higher for later calvings when compared with the present study. Jamrozik et al. (2013), evaluating STAY to consecutive calvings in Canadian Simmentals, reported means of STAY that were very similar, both in magnitude and trend across calvings, to those found in the present study for Tabapuã cattle. When analysis was performed using the traditional definition of STAY and the threshold model, the success rate was ~ 34%, 38% and 45% for Guzerá, Nelore and Tabapuã, respectively. These results demonstrate one of the advantages of the definition of STAY to consecutive calvings, which permits a more detailed analysis of the populations studied regarding this trait.

Analysis of R and RES values showed improvement in the goodness-of-fit of the models with increasing order of the polynomial used (Table 2). However, higher-order polynomials result in greater parameterization of the model, a fact that generally increases computational demands to obtain accurate estimates of the parameters of interest such as breeding values of the animals. For this reason, the models were also compared using DIC, a criterion that does not only consider the goodness-of-fit but also penalizes greater parameterization caused by higher-order polynomials. Using DIC as a criterion, the model with 4 regression coefficients (cubic) for each random effect was found to be the most adequate for STAY to consecutive calvings in the Nelore and Tabapuã breeds. However, for Guzerá, the best model according to DIC was that including 5 coefficients (quartic). The Bayesian deviance values ($D(\theta)$) also indicated the models chosen based

Table 2 Number of regression coefficients (k) for the contemporary group, additive genetic and permanent environmental random effects, number of parameters (p), correlation between observed and predicted binary phenotype (R), mean absolute value of residuals (RES), Deviance Information Criterion (DIC) and Bayesian deviance (D(Θ)) for the random regression models

Model ¹	¹ k P R		RES	RES DIC ²		
Guzerá						
LEG2	2	16	0.923	0.116	563 258	649 908
LEG3	3	25	0.955	0.081	278 380	283 903
LEG4	4	37	0.974	0.057	54 045	35 905
LEG5	5	52	0.985	0.039	0	0
Nelore						
LEG2	2	16	0.928	0.113	181 319	1 422 174
LEG3	3	25	0.957	0.077	11 910	324 343
LEG4	4	37	0.975	0.053	0	0
LEG5	5	52	0.986	0.036	15 140	163 044
Tabapuã						
LEG2	2	16	0.928	0.033	563 016	758 274
LEG3	3	25	0.958	0.019	397 099	487 863
LEG4	4	37	0.975	0.012	0	0
LEG5	5	52	0.986	0.006	165 586	83 495

¹LEG*k*, where k = number of regression coefficients for the contemporary group, additive genetic and permanent environmental random effects.

²Values scaled as deviation from the respective 'best' values for each breed.

on DIC as the best models. Jamrozik *et al.* (2013), working with data from Canadian Simmentals, also recommended the use of a linear random regression model in which the random regressions for contemporary group, additive and permanent environmental effects were modeled by cubic orthogonal Legendre polynomials.

Applying the model with 4 coefficients for each random effect (additive genetic, permanent environmental and contemporary group) for Nelore and Tabapuã breeds and a model with 5 coefficients for each random effect for Guzerá, the parameters (variances, heritabilities and correlations) were estimated. Figure 2 shows the proportion of phenotypic variance explained by each random effect. The effect of the permanent environment most contributed to phenotypic variation, regardless of breed or calving, with values ranging from 63% to 83%, 59% to 87% and 55% to 93% for Guzerá, Nelore and Tabapuã, respectively. The trend observed was a decrease between the second and third calving and an increase over subsequent calvings. Analyzing a population of Canadian Simmentals, Jamrozik et al. (2013) found an important contribution of the permanent environmental effect, but the proportions were lower than those observed in the present study. The participation of the contemporary group effect decreased across calvings, with magnitudes of ~ 6% to 13%, 9% to 14% and 8% to 21% for Guzerá, Nelore and Tabapuã, respectively. These findings show that this effect, which is related to the environment during the early stages of animal growth, has a greater influence on STAY during early calvings, a fact also observed by Jamrozik et al. (2013) for Canadian Simmentals. Residual variance

contributed ~0.4% to 17%, 0.005% to 15% and 0.03% to 14% to phenotypic variability in the Guzerá, Nelore and Tabapuã breeds, respectively. Peak values were observed in the third calving and the values at the extremes (calvings 2 and 8) were considerably low. Jamrozik *et al.* (2013) observed a relatively greater contribution of residual variance. This lower ratio of residual to phenotypic variance is an indicator that part of the residual effect was estimated as permanent environmental effect. The trends for the variance estimates were similar among breeds, except for the additive genetic variance that showed a slightly different trend in the Tabapuã compared with the other two breeds studied (Supplementary Material Figure S1).

The posterior means of heritability for STAY to each calving ranged from 0.05 to 0.08 in the Guzerá breed, showing a quadratic trend with a peak between the fourth and sixth calving (Figure 3). For Nelore and Tabapuã, the estimates ranged from 0.03 to 0.07 and from 0.03 to 0.08, respectively, and tended to increase with calving number. Martinez *et al.* (2005) analyzed STAY data to different calvings of Hereford cattle in the United States using linear models for binary data and obtained slightly higher estimates (0.05 to 0.19) that increased from the second to the sixth calving. For Canadian Simmentals, Jamrozik *et al.* (2013) found estimates of a higher magnitude ranging from 0.12 to 0.36, which decreased with increasing calving number.

Transformation of the heritabilities of the linear random regression model to the underlying normal scale (Figure 4) resulted in estimates of 0.10 to 0.24, 0.04 to 0.38 and 0.06 to 0.23 for Guzerá, Nelore and Tabapuã, respectively. For all breeds, these estimates increased with calving number. Using the threshold model to fit binary data to the classical definition of STAY, the posterior means of heritability (and their respective HPD_{95%}) were 0.09 (0.07 to 0.12), 0.09 (0.08 to 0.10) and 0.09 (0.07 to 0.11) for Guzerá, Nelore and Tabapuã, respectively. These values are similar to those obtained with the random regression models for STAY to early calvings, but lower than the estimates obtained for STAY to later calvings. Using the classical definition of STAY (success when the cow stays in the herd until 6 years of age given the opportunity to mate and calve regularly until this age), Santana et al. (2012 and 2015) reported a heritability of 0.11 for Nelore cattle, a similar value estimated in the present study for a classical definition of STAY analyzed using the threshold model. Also in Nelore cattle, Silva et al. (2003) studied some definitions for STAY in which the cow should calve every year until 5, 6 or 7 years of age and reported heritabilities of 0.11, 0.12 and 0.17, respectively, magnitudes similar to those observed in the present study.

The additive genetic correlations showed a similar trend among breeds and were higher for STAY between closer calvings (Table 3). Even for more distant calvings (second and eighth), the additive genetic correlations for STAY were moderate to high (0.77, 0.57 and 0.79 for Guzerá, Nelore and Tabapuã, respectively), indicating that STAY to the second calving is a good indicator of STAY to subsequent

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Figure 2 Contribution (%) of additive genetic, permanent environmental, contemporary group and residual variance to the total (phenotypic) variance of stayability to consecutive calvings according to breed.



Figure 3 Heritability estimates for stayability to consecutive calvings according to breed.

calvings. Thus, selection could be performed in the second calving to reduce the generation interval and to increase annual genetic gain. The same conclusion has been drawn by Jamrozik *et al.* (2013). In general, the additive genetic correlations obtained in this study for Zebu breeds were higher than those reported by Jamrozik *et al.* (2013) for Simmentals and by Martinez *et al.* (2005) for a Hereford herd. The permanent environmental correlations for all breeds decreased with increasing distance between calvings. The



Figure 4 Heritability estimates, on the underlying normal scale, for stayability to consecutive calvings according to breed.

contemporary group correlations exhibited a similar trend and magnitude in the breeds and also decreased with increasing distance between calvings. In contrast, slightly higher phenotypic correlations were obtained for the Tabapuã when compared with Guzerá and Nelore breeds.

The heritabilities of the regression coefficients ranged from 0.01 to 0.09, 0.02 to 0.06 and 0.01 to 0.09 for Guzerá, Nelore and Tabapuã, respectively, with higher values being observed for the first regression coefficients. As the first

Table 3 *Means, standard deviations (SD) and highest posterior density intervals (HPD*_{95%}) for the additive genetic, permanent environmental, contemporary group and phenotypic correlations between the stayability to second calving and stayability to later calvings

	Additive genetic			Permanent environmental			Contemporary group			Phenotypic		
Calving	Mean	SD	HPD _{95%}	Mean	SD	HPD _{95%}	Mean	SD	HPD _{95%}	Mean	SD	HPD _{95%}
Guzerá												
3	0.94	0.019	0.90 to 0.97	0.66	0.004	0.65 to 0.67	0.85	0.010	0.83 to 0.87	0.64	0.003	0.63 to 0.64
4	0.91	0.027	0.86 to 0.96	0.51	0.005	0.50 to 0.52	0.77	0.013	0.74 to 0.79	0.53	0.003	0.52 to 0.53
5	0.89	0.032	0.82 to 0.93	0.44	0.005	0.42 to 0.44	0.67	0.018	0.63 to 0.70	0.46	0.003	0.46 to 0.47
6	0.85	0.034	0.78 to 0.90	0.34	0.005	0.32 to 0.34	0.55	0.022	0.51 to 0.59	0.36	0.004	0.35 to 0.36
7	0.81	0.038	0.73 to 0.87	0.21	0.006	0.20 to 0.22	0.46	0.025	0.41 to 0.51	0.24	0.005	0.23 to 0.25
8	0.77	0.042	0.69 to 0.84	0.32	0.005	0.31 to 0.33	0.51	0.022	0.47 to 0.56	0.36	0.004	0.35 to 0.36
Nelore												
3	0.92	0.019	0.88 to 0.96	0.81	0.002	0.80 to 0.81	0.90	0.003	0.90 to 0.91	0.68	0.002	0.67 to 0.68
4	0.86	0.031	0.79 to 0.91	0.60	0.003	0.59 to 0.60	0.79	0.006	0.77 to 0.80	0.57	0.002	056 to 0.57
5	0.80	0.035	0.73 to 0.87	0.48	0.004	0.47 to 0.48	0.70	0.008	0.69 to 0.72	0.48	0.003	0.47 to 0.48
6	0.73	0.041	0.64 to 0.80	0.39	0.004	0.38 to 0.40	0.63	0.010	0.61 to 0.65	0.41	0.003	0.40 to 0.41
7	0.63	0.048	0.54 to 0.72	0.32	0.004	0.32 to 0.33	0.56	0.011	0.54 to 0.59	0.35	0.003	0.34 to 0.35
8	0.57	0.048	0.48 to 0.66	0.29	0.004	0.28 to 0.29	0.55	0.012	0.53 to 0.57	0.32	0.003	0.31 to 0.33
Tabapuã												
3	0.94	0.018	0.90 to 0.98	0.78	0.002	0.77 to 0.78	0.91	0.004	0.90 to 0.92	0.82	0.002	0.81 to 0.82
4	0.92	0.029	0.85 to 0.97	0.57	0.004	0.56 to 0.58	0.81	0.008	0.79 to 0.82	0.64	0.003	0.63 to 0.65
5	0.91	0.032	0.84 to 0.96	0.45	0.005	0.45 to 0.46	0.73	0.011	0.71 to 0.75	0.53	0.004	0.52 to 0.54
6	0.90	0.033	0.82 to 0.94	0.36	0.005	0.35 to 0.37	0.66	0.014	0.63 to 0.68	0.44	0.004	0.43 to 0.45
7	0.86	0.034	0.79 to 0.91	0.29	0.005	0.28 to 0.30	0.57	0.017	0.54 to 0.61	0.36	0.004	0.35 to 0.37
8	0.79	0.038	0.70 to 0.85	0.22	0.005	0.22 to 0.24	0.54	0.019	0.50 to 0.58	0.29	0.004	0.28 to 0.30

coefficient has the same weight for STAY in all calvings and showed the highest heritability in all breeds, selection for STAY using this coefficient may result in a favorable change in the average of the population in all calvings studied. To support this conclusion, the correlations between breeding values for STAY to each calving and for the first regression coefficient were estimated and the values observed were high for all breeds and in all calvings (Supplementary Material Figure S2). Analysis of rank correlations between predicted breeding values for the intercept coefficient obtained with the models studied (Table 4) generally showed high correlations (>0.99) between models with 3, 4 or 5 coefficients. On the other hand, genetic evaluation using the model with 2 coefficients resulted in considerable alterations in the ranking of animals when compared with the other models, demonstrating that this model would be inadequate for the genetic evaluation of this trait. For the Guzerá breed, the high rank correlations

Table 4 Rank correlations using the breeding values for the first regression coefficient predicted with the studied models¹

Model	All animals				Bulls ²		Cows ³		
	LEG3	LEG4	LEG5	LEG3	LEG4	LEG5	LEG3	LEG4	LEG5
Guzerá									
LEG2	0.978	0.976	0.980	0.980	0.963	0.974	0.975	0.974	0.975
LEG3	_	0.992	0.991	_	0.961	0.976	_	0.990	0.990
LEG4	_	_	0.990	_	_	0.987	_	_	0.993
Nelore									
LEG2	0.900	0.861	0.890	0.832	0.790	0.866	0.900	0.861	0.895
LEG3	_	0.986	0.986	_	0.988	0.971	_	0.988	0.988
LEG4	_	_	0.996	_	_	0.968	_	_	0.990
Tabapuã									
LEG2	0.954	0.953	0.961	0.951	0.942	0.941	0.951	0.950	0.958
LEG3	_	0.997	0.996	_	0.993	0.994	_	0.996	0.995
LEG4	-	-	0.996	-	-	0.995	-	-	0.995

 1 LEGk, where k = number of regression coefficients for the contemporary group, additive genetic and permanent environmental random effects.

²Sires of cows with phenotipic records.

³Cows with phenotipic records.

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between models with 4 and 5 coefficients indicate that the simpler model with 4 coefficients could be used in genetic evaluations, replacing the model with the best fit.

Selection for STAY to a specific age, generally 76 months, which usually requires a minimum of three calves born, does not promote a complete differentiation of productivity in genetic evaluations as these cows may have a larger number of calves during the same period, and are therefore more productive for the herd, but eventually receive the same phenotype as less productive cows. The definition of STAY to consecutive calvings using random regression models permits to identify and separate the effects that influence each calving of a female. As emphasized by Jamrozik et al. (2013), the use of this definition in breeding programs also has the advantage that phenotypes can be obtained earlier, as well as a larger number of phenotypic records per cow, increasing the accuracy of dam and sire predicted breeding values. Using this approach, selection could be performed using breeding values for STAY to second calving because of its moderate to high additive genetic correlations with STAY to eighth calving, increasing genetic gain by reducing the generation interval.

The random regression methodology applied to the definition of STAY to consecutive calvings can be used for the genetic evaluation of Zebu beef cattle. A model that uses 4 regression coefficients for additive genetic, contemporary group and permanent environmental effects is recommended.

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Supplementary material

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