University of São Paulo
“Luiz de Queiroz” College of Agriculture

Statistical models for genomic selection in *Panicum maximum*
considering allelic dosage

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Thesis presented to obtain the degree of Doctor in Science.
Area: Genetics and Plant Breeding

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versão revisada de acordo com a resolução CoPGr 6018 de 2011

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1. Melhoramento de plantas 2. Forrageira 3. Autotetraploides 4. Modelos lineares mistos 5. Predição I. Título.
DEDICATORY

To my parents José Maurício and Liberaci,
My angels João Gabriel and Benjamin,
And my fiancé Thiago.
I love you.
I thank God and “Nossa Senhora da Aparecida” for being presence in my life.

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I hope everything I learned is not an obstacle to humility, but an incentive to achieve great dreams.
CANÇÃO ETERNA

Lá estavam os pássaros, sorrindo sob o som da lua
Era primavera, verão, outono e inverno
Tinham vários olhos nos observando do céu
Ícones indignados com o tempo
Cuja pressa era assustadora!
Imersos íamos, porém, sendo levados pela aurora
Agarrados, como se a alegria fosse deveras única!
Andavam e corriam os lábios
Pureza humana! Natural, sedenta...
Amor é a consequência exata
Reflexo de duas almas livres
Esgueiradas de uma sociedade
Cujas diretrizes
Indubitavelmente insanas!

Dias, semanas, meses e anos
A música que a nós toca
Dança sem um registro de fim
E na solidão inexistente
Cercamos os olhos alheios diiscentes
Apáticos, raivosos, deprimentes...
Sarcásticos... Seria a felicidade cobiçada?
Treinados por regras irreais
Respiram os ares que não são deles!
O lado obscuro que nos cerca, então,
Lá fica, longe e decadente
Amor como este não se entende
Respira-se, sente-se e de repente
As vidas que nos cercam sentem

Author: Thiago de Paula Oliveira
SUMMARY

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Diversas espécies de interesse econômico são autotetraploides, como a forrageira *Panicum maximum*, a qual proporciona alta produtividade e qualidade para pastagens tropicais. Os principais acessos na natureza são plantas apomíticas tetraploides, no entanto pode-se encontrar também plantas sexuais diploides. Embora a apomixia seja vantajosa pela facilidade em fixar o vigor híbrido, a reprodução sexual é fundamental por permitir recombinação genética a partir de cruzamentos entre genótipos superiores. Desta forma, o melhoramento nesta espécie consiste em cruzar plantas apomíticas com plantas sexuais tetraploïdizadas. A utilização de parentais sexuais superiores nestes cruzamentos permite aumentar a frequência de alelos favoráveis na progênie. Portanto, programas de seleção recorrente intrapopulacional em populações sexuais tetraploides são fundamentais para programas de melhoramento em *P. maximum*. Além disto, a utilização de estratégias como seleção genômica são promissoras para aumentar os ganhos de seleção, permitindo avançar ciclos de seleção recorrente e lançar cultivares no mercado em menor prazo, quando comparados a programas convencionais. Como *P. maximum* é uma cultura perene, os genótipos são avaliados em sucessivos cortes. Assim, este estudo tem como finalidade avaliar caracteres de produtividade, estruturais e nutricionais em uma população sexual tetraploide de *P. maximum*, investigando diferentes classes de modelos lineares mistos aplicados a dados longitudinais, além de desenvolver modelos de seleção genômica que considere a natureza tetraploide da população. Este trabalho foi dividido em dois capítulos. No primeiro capítulo, três classes de modelos foram analisados: i) Classe A consiste em modelar a interação genótipos por cortes com correlações homogêneas, genótipos não correlacionados entre si e os efeitos residuais são ajustados com homocedasticidade e ausência de correlação; ii) Classe B consiste em grupos de modelos com diferentes estruturas de variância e covariância (VCOV) para efeitos genéticos e residuais e genótipos não correlacionados; iii) Classe C é similar à Classe B, no entanto os genótipos são correlacionados por uma matriz de parentesco aditivo calculado por pedigree. Para todos os caracteres, os modelos da Classe C tiveram melhor ajuste. Portanto, recomenda-se testar matrizes de VCOV que permitam modelar cortes com diferentes níveis de correlações ao longo do tempo bem como incluir informação de parentesco aditivo e, se disponível, matriz de parentesco genômico. No segundo capítulo, marcadores SNPs, obtidos via genotipagem por sequenciamento, foram aplicados em modelos Bayesians e GBLUP os quais foram desenvolvidos para incorporar informação de dosagem alélica tetraploide. Uma vez que as acurácias dos modelos Bayesians não diferiram das acurárias do modelo GBLUP com dosagem alélica, recomenda-se o uso do segundo por requerer menos tempo computacional. A acurácia dos modelos preditivos reforça a vantagem em implementar seleção genômica em programas de melhoramento de *P. maximum*.

Palavras-chave: Melhoramento de plantas; Forrageira; Autotetraploïdes; Modelos lineares mistos; Predição
ABSTRACT

Statistical models for genomic selection in *Panicum maximum* considering allelic dosage

Several species of economic interest are autotetraploid, such as the forage *Panicum maximum*, which is responsible for high productivity and quality of tropical pastures. The main accessions in nature are autotetraploid apomictic plants, on the other hand, diploid sexual plants may also be found. Although apomixis is advantageous because it fixes hybrid vigor, sexual reproduction is fundamental to allow genetic recombination by crossing among superior genotypes. Thus, genetic breeding consists of crossing apomictic plants with tetraploidized sexual plants. In these crosses, the use of superior sexual parents allows to increase the frequency of favorable alleles in the progeny. Therefore, recurrent selection programs in tetraploid sexual populations are fundamental to *P. maximum* breeding programs and strategies such as genomic selection can increase the accuracy of selection, allowing shorter breeding cycles and release cultivars in the market in the short term when compared to conventional programs. As *P. maximum* is a perennial crop, genotypes are evaluated in successive harvests. Thus, the study goals are to evaluate nutritional, structural, and yield traits in a sexual tetraploid population of *P. maximum*, investigating different classes of linear mixed models applied to longitudinal data, as well as to develop genomic selection models which consider tetraploid allelic dosage. This work was split into two chapters. In the first chapter, three classes of models were analyzed: i) Class A consists in modeling the interaction of genotypes and harvests with homogeneous correlations, genotypes were assumed not correlated, and residual effects were assumed homocedastic and not correlated; ii) Class B consists of groups of models in which genetic and residual effects were fitted with different variance and covariance (VCOV) structures and genotypes were not correlated; and iii) Class C is similar to Class B, however genotypes were correlated by an additive relationship matrix based on pedigree values. For all traits, Class C models performed better based on goodness of fit of the models. Therefore, we recommend to incorporate additive relationship matrix besides to model harvests with different levels of correlations over time. In the second chapter, SNP markers, obtained by genotyping-by-sequencing (GBS) technique, were used to develop Bayesian and GBLUP models that consider tetraploid allelic dosage. Bayesian models accuracies did not differ from the accuracy of GBLUP model and, we recommend the latter because it requires less computational time. The accuracy of genomic selection models reinforces the advantage of implementing this strategy in *P. maximum* breeding programs.

**Keywords:** Plant breeding; Forage; Autotetraploids; Linear mixed models; Prediction
1 INTRODUCTION

Brazil is a leader in beef production, being the largest or second-largest producer in the world, competing only with the United States (JANK et al., 2014). This position is due to vast pastures and cattle herds present in the country. Its native and cultivated pasture area is equivalent to its agricultural plus planted and natural forest areas (JANK et al., 2011). The greatest cultivated area in the country is represented by Brachiaria (Syn. Urochloa) spp., where Brachiaria brizantha cv. Marandu grass is the predominant forage (JANK et al., 2014). As this cultivar is produced on a large scale, Brazilian pastures are classified as extensive monocultures. Therefore, the use of several species and cultivars is recommended to mitigate problems caused by monoculture, such as break of resistance to known diseases (JANK et al., 2011). Panicum maximum (Syn. Megathyrsus maximum) Jacq. is an excellent option for diversification and intensification of Brazilian pastures because it is very productive and has excellent nutritive quality, providing high animal production per hectare (JANK et al., 2014).

The main reproduction strategies is apomixis (autotetraploid plants), but sexuality occurs sporadically in diploid plants. Apomixis is a clonal propagation by seeds, in which offsprings genetically identical to the female parent are produced. According to SAVIDAN et al. (1989), the apomixis has several advantages in breeding programs, such as hybrid vigor fixation, simplification in obtaining hybrids, and low cost of seed production. However, the main disadvantage is that it does not allow recombination of superior individuals, avoiding the exploration of genetic variability. Thus, the improvement in this species was made possible from the chromosomal duplication of sexual diploid plants and, later, crossing with apomictic plants (SAVIDAN et al., 1989).

Main tropical forage breeding programs in Brazil are at Embrapa Centers, which hold the main germplasm banks in the country. The P. maximum forage breeding is mainly coordinated by Embrapa Beef Cattle, with the goals of increasing leaf and seed yield, disease resistance, and nutritive quality (JANK et al., 2011). The breeding program uses recurrent selection methods where each cycle requires three to five years of evaluation. According to RESENDE et al. (2014), the process of development, testing, and recommendation of a new cultivars span over approximately fifteen years.

Genomic selection is an effective method to explore genetic variation in breeding programs, from the prediction of breeding values based on markers distributed throughout the genome. It can increase the accuracy of selection, reduce evaluation costs per genotype and get shorter breeding cycles than phenotypic selection (LIPKA et al., 2014; RESENDE et al., 2014). Its potential to increase the efficiency of breeding programs has been shown in several crops (CROSSA et al., 2013; GOUY et al., 2013; LIPKA et al., 2014). Therefore, the application of genomic selection in forage breeding is promising, since many of the main traits have high assessment costs, as well as evaluation after flowering time in the breeding cycle.

This work consists of a partnership between the Embrapa Beef Cattle (Campo Grande, MS, Brazil), the Graduate Program in Genetics and Plant Breeding at ESALQ / USP (“Luiz de Queiroz” College of Agriculture / University of São Paulo - Piracicaba, SP, Brazil), and the Bioinformatics Research Center at NC State (North Carolina State University - Raleigh, NC, USA). The work was organized in two chapters. The first one has the goal of to evaluate
nutritional, structural, and yield traits in a tetraploid sexual *P. maximum* population using linear mixed models and to estimate genetic parameters and canonical correlation between sets of traits. The second one aims to develop statistical models in genomic selection considering tetraploid allelic dosage for the same population of *P. maximum*.

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2 CONCLUSION

Recurrent selection (RS) is an approach that aims to improve breeding populations by increasing the frequency of favorable alleles, while maintaining genetic variability (Hallauer and Carena, 2012). In perennial crops, such as Panicum maximum, the evaluation process is performed by repeated measures in the same individual, in order to infer the genotype’s ability to repeat its performance over successive evaluations (Braz et al., 2015). Analysis of mixed models allows modeling different structures of variance and covariance (VCOV) to investigate different assumptions of heterogeneity and dependence between observations (Crossa et al., 2006). In addition, these analyzes allow the inclusion of information from relatives, exploring the genetic correlation between individuals by an additive relationship matrix estimated by pedigree values or, when available, estimated by molecular markers.

In the first chapter, three classes of models were consider to evaluate nutritional, structural, and yield traits. For all traits, models of the class that consider different VCOV matrices and include pedigree information had a better fit. Therefore, incorporating the additive relationship matrix and modeling the genotype by harvest interaction with VCOV structures allows better selection of individuals of the breeding population.

Genomic selection (Meuwissen et al., 2001) is an approach that can accelerate future genetic progress in plant breeding (Heffner et al., 2010), as well as accelerate recurrent selection cycles (Lipka et al., 2014). Although some crops have benefited from this methodology, studies in autotetraploids (and polyploids in general) are still modest. Thus, the main objective of the second chapter was to develop models that consider tetraploid allelic dosage to estimate genetic values for genomic selection in P. maximum, an autotetraploid forage. These models can be applied to other autotetraploid species, as well as extended to species with other ploidy level.

Therefore, we compared six different GS models and presented strategies for modeling phenotypic residual effects and performing SNP calling in autotetraploids. The GBS and allelic dosage strategies shown to be promising for genomic analyzes in P. maximum and the accuracy of predictive models justifies the implementation of this approach in breeding programs.

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