Indices estimated using REML/BLUP and introduction of a super-trait for the selection of progenies in popcorn


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ABSTRACT. Selection indices commonly utilize economic weights, which become arbitrary genetic gains. In popcorn, this is even more evident due to the negative correlation between the main characteristics of economic importance - grain yield and popping expansion. As an option in the use of classical biometrics as a selection index, the optimal procedure restricted maximum likelihood/best linear unbiased predictor (REML/BLUP) allows the simultaneous estimation of genetic parameters and the prediction of genotypic values. Based on the mixed model methodology, the objective of this study was to investigate the comparative efficiency of eight selection indices estimated by REML/BLUP for the effective selection of superior popcorn families in the eighth intrapopulation recurrent selection cycle. We also investigated the efficiency of the inclusion of the variable “expanded popcorn volume
per hectare” in the most advantageous selection of superior progenies. In total, 200 full-sib families were evaluated in two different areas in the North and Northwest regions of the State of Rio de Janeiro, Brazil. The REML/BLUP procedure resulted in higher estimated gains than those obtained with classical biometric selection index methodologies and should be incorporated into the selection of progenies. The following indices resulted in higher gains in the characteristics of greatest economic importance: the classical selection index/values attributed by trial, via REML/BLUP, and the greatest genotypic values/expanded popcorn volume per hectare, via REML. The expanded popcorn volume per hectare characteristic enabled satisfactory gains in grain yield and popping expansion; this characteristic should be considered super-trait in popcorn breeding programs.

Key words: Zea mays L.; Mixed model; Selection index

INTRODUCTION

The breeding programs have been dynamic and have tested some proposals of changes, always seeking to improve the efficiency of the selection process and facilitate assessments. In this sense, the combined selection is important; also important is the acquisition of superior materials from the selection and recombination of superior families and individuals for a set of features (Hallauer et al., 2010; Cruz et al., 2014). From this perspective, the main objective is to select cultivars with high yields and other desirable traits (Baker, 1986; Magnussen, 1990; Falconer and Mackay, 1996). Thus, selection should be based on a set of traits that enables the simultaneous acquisition of satisfactory genetic gains. The aim of this selection is the identification of genotypes that meet the modern demands of the consumer and producer markets (Hallauer et al., 2010; Cruz et al., 2014).

The use of a selection index, which is established using the optimum combination of several traits, is a biometric technique that allows for the efficient and simultaneous selection of multiple traits (Cruz et al., 2014). The application of selection indices enables the identification of superior genotypes established by the optimal linear combination of various traits; it is preferred that these traits be uncorrelated (Magnussen, 1990; Falconer and Mackay, 1996; Hallauer et al., 2010). The use of selection indices was originally reported by Smith (1936) and Hazel (1943). Other indices were later proposed and are currently used in plant breeding. A special emphasis has been placed on the indices developed by Williams (1962), Pesek and Baker (1969), and Mulamba and Mock (1978). Regardless of the index considered, the use of indices allows the objective determination of the relative merits of a series of genotypes and provides a basis for their differentiation (Carena, 2009).

The REML method, developed by Patterson and Thompson (1971), has been routinely used to estimate variance components in mixed models. Its wide use can be attributed to the fact that it considers all the desirable properties of an estimator. The BLUP procedure (Henderson, 1973) minimizes the variance of the prediction error and maximizing precision. BLUP is a linear function of the observations and does not account for patterns; therefore, it maximizes the accuracy of non-biased predictors and predicts random variables (Viana et al., 2011; Resende et al., 2012).
Resende (2007a) reported that the simultaneous analysis of various traits could be efficiently performed using the multivariate REML/BLUP approach. In this case, the multivariate model is specified to address the existing environmental covariance between traits. The use of the sum of the predicted breeding values by a multivariate model, an additive index with or without economic weights, is an efficient selection strategy. BLUP increases selective accuracy; therefore, it is superior to any other combined selection index (Resende, 2007a).

The selection indices have been widely used in popcorn (Daros et al., 2004; Amaral Júnior et al., 2010) and genetic gains were predicted with satisfactory accuracy, which led to an appropriate performance of breeding programs for this culture. However, the use of selection indices is limited, in some situations, by the difficulty in establishing economic weights for the various traits. Due to these difficulties, Cruz (1990) proposed that economic weights be estimated based on experimental data. A multitrait selection procedure is needed to select simultaneously for improvement in more than one trait. The most important characteristics evaluated in popcorn breeding programs are grain yield (GY) and popping expansion (PE) (Silva et al., 2010; Moterle et al., 2011). PE is the primary quality characteristic of popcorn; this trait comprises the ratio between the volume of expanded popcorn and the initial volume or mass of grains before popping (Moterle et al., 2011; Rodovalho et al., 2014). The negative correlation between the two major characteristics of popcorn (Dofing et al., 1991; Daros et al., 2004) makes breeding difficult, but not impossible. Nevertheless, the selection process becomes biased. This is because it is not possible to uniquely and reliably associate the best genotypes for GY with the best genotypic expressions of PE. In the present study, we present the use of the following alternative trait: expanded popcorn volume per hectare (PV). We evaluated the convenience of its use as a super-trait in breeding programs.

Research into the use of selection indices via REML/BLUP in popcorn breeding programs has been limited. We compared eight selection indices based on mixed modeling to determine the reliability of the selection of the best full-sibling families in the eighth cycle of an intrapopulation recurrent selection program.

MATERIAL AND METHODS

Study population

A total of 200 full-sib families were obtained in the eighth intrapopulation recurrent selection cycle and evaluated. The evaluated families belong to the Popcorn Genetic Breeding Program of the State University of North Fluminense Darcy Ribeiro (UENF).

Experimental design

The experiments were planted in two environments in April 2013. The first experiment was conducted at the Antônio Sarlo State Agricultural College in the municipality of Campos dos Goytacazes, which is in the Northern region of the State of Rio de Janeiro, Brazil. The climate is classified as Aw (tropical savanna climate), with dry in winter according to the Köppen classification. The average annual rainfall is 1023 mm, and the average annual temperature is 23°C. The second experiment was conducted at the Agricultural Research Company of the State of Rio de Janeiro (Empresa de Pesquisa Agropecuária do Estado do Rio de Janeiro - PESAGRO-Rio). The Agricultural Experiment Station at Ilha do Pomba is in...
the municipality of Itaocara, which is in the Northwest region of the State of Rio de Janeiro, Brazil. The climate is classified as Awi (tropical wet-dry isotherm), with rainy summers and dry winters according to the Köppen classification. The average annual temperature is near 22.5°C, and the average annual rainfall is 1041 mm.

The experiments were arranged in a randomized block design with three repetitions. The plots were comprised of one row measuring 5.0 m in length, with 0.90 m spacing between the rows. The spacing between the plants was 0.20 m. Three seeds were planted per hole at a depth of 0.05 m. Thinning was performed at 21 days after emergence; one vigorous plant was left per hole. Fertilization was performed at planting with 800 kg/ha mineral fertilizer in the formulation 04-14-08 (N-P-K). Topdressing was conducted at 15 days after sowing, with 300 kg/ha mineral fertilizer in the formulation 20-00-20 (N-P-K). A second top-dressing was performed at 30 days after emergence; nitrogen in the form of urea was applied at 200 kg/ha. All fertilizers were recommended according to an analysis of the soil in the plantation areas. The management of weeds, pests, and diseases was carried out according to standard recommendations for the region. The GY was corrected to 13% of moisture content.

Traits evaluated

Six important agronomic characteristics were evaluated for popcorn. a) The average plant height (PH) of six competitive plants in a plot was measured from the ground to the insertion point of the flag leaf and was expressed in meters (m). b) Prolificacy (PROL) reflected the average number of ears per plant. c) The mass of 100 grains (M100) was expressed in grams (g) and obtained by weighing 100 grains randomly. Three repetitions were performed per plot. d) The GY was determined by weighing the grains from each plot and was expressed as kg/ha. e) PE was determined by placing 30 g seeds in kraft paper bags; the bags were placed in a microwave oven for 3 min. Three repetitions were performed per experimental unit. The volume of the expanded popcorn after microwaving was quantified in a 2000-mL beaker. PE was expressed as mL/g. f) The PV was obtained by multiplying the average yield of the plot and the PE. This measurement represents the volume of expanded popcorn by hectare planted and was expressed as m³/ha.

Genetic-statistical analyses

The genetic-statistic model utilized was as follows:

\[ Y = X_r + Z_g + W_p + T_i + e \]  

(Equation 1)

where \( Y \) is the vector of the data, \( r \) is the vector of the repetition effects (assumed to be fixed) added to the overall average, \( g \) is the vector of individual genotypic effects (assumed to be random), \( p \) is the vector of environmental effects (assumed to be random), \( i \) is the vector of genotype x environment interaction effects (random), and \( e \) is the vector of errors or residues (random).

The procedures for estimating the breeding values and for executing the additive (Smith, 1936; Hazel, 1943) and multiplicative selection indices (Subandi and Empig, 1973) and the index weighted by the average rank of the families were adapted from Mulamba and Mock (1978); these procedures were performed using the SELEGEN-REML/BLUP software (Resende, 2007b).
After the genotypic values (GV) for each family and variable were estimated, different models of the selection indices were compared and examined for the best population selection gains. The following models were used: i) selection via major GV associated with the variables GY and PE (GV/GY-PE); ii) selection via the greatest GV for the variable PV (GV/PV); iii) the use of the classical selection index (CI), according to the model:

\[
CI = \left( \left( p \cdot \overline{X_i} \right) \cdot \left( GV \cdot \overline{X_i} \right) \right) + \ldots + \left( \left( p \cdot \overline{X_n} \right) \cdot \left( GV \cdot \overline{X_n} \right) \right)
\]  
(Equation 2)

iv) the use of the multiplicative index (MI), according to the model:

\[
MI = \left( GV \cdot \overline{X_i} \right) \cdot \left( GV \cdot \overline{X_n} \right)
\]  
(Equation 3)

and v) the use of the index weighted by the average rank of the families households (IWR). IWR is described as follows:

\[
IWR = \left( r \cdot GV \cdot \overline{X_i} \right) + \left( r \cdot GV \cdot \overline{X_n} \right)
\]  
(Equation 4)

where \( p \) corresponds to the economic weight attributed to the trait, \( GV \) is the predicted genotypic value, \( r \) corresponds to the family rank, and \( \overline{X} \) corresponds to the phenotypic average of the genotype for each variable analyzed (Resende, 2007b).

For the execution of the classical index, economic weights were assigned to the characteristics using estimates of the following parameters: i) genetic coefficient of variation (CVg); ii) genetic standard deviation (SDg); iii) heritability (h^2); and iv) values attributed by trial (BT) of magnitude, as described by Santos et al. (2007a). Genetic parameters were assigned economic weights according to the needs of the employee selection index. The averages for these parameters were obtained by performing analysis of variance using the Genes software (Cruz, 2013); random environmental effects and the genotypes analyzed were considered.

We selected 30 superior families according to each index and fixed a 15% selection pressure. The efficiency of the selection indices was obtained using the coincidence index of the families selected by the eight selection methods employed. The index of coincidence was obtained by the ratio between twice the number of families, in which both selection indexes coincide, and the sum of the total number of families containing the selection index A plus the total number of progenies containing the selection index B (Pedrozo et al., 2009; Freitas et al., 2013). Predicted selection gains were also obtained for each characteristic evaluated using the different selection indices.

RESULTS

Based on the scores related to the additive genetic values of the 200 families, it was possible to select the 30 best families based on the largest reliably estimated genotypic values for the characteristics PE and GY (Figure 1).
For most of the coefficients, an increased degree of coincidence was observed among the families selected using the eight different selection indices. The highest coincidence (96.67%) was observed between the indices CI/CVg and MI (Table 1).

### Table 1. Coincidence coefficients (%) for 30 superior full-sibling families identified using eight selection indices based on genotypic values for six characteristics (the traits were evaluated in the eighth intrapopulation recurrent selection cycle of the popcorn population UNB-2U).

<table>
<thead>
<tr>
<th>Selection indexes</th>
<th>CI/CVg</th>
<th>CI/SDg</th>
<th>CI/h²</th>
<th>CI/BT</th>
<th>MI</th>
<th>IWR</th>
<th>GV/GY-PE</th>
<th>GV/PV</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI/CVg</td>
<td>73.33</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CI/SDg</td>
<td>73.33</td>
<td>86.67</td>
<td>83.33</td>
<td>96.67</td>
<td>80.00</td>
<td>53.33</td>
<td>76.67</td>
<td></td>
</tr>
<tr>
<td>CI/h²</td>
<td>83.33</td>
<td>86.67</td>
<td>90.00</td>
<td>86.67</td>
<td>83.33</td>
<td>76.67</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CI/BT</td>
<td>90.00</td>
<td>80.00</td>
<td>80.00</td>
<td>86.67</td>
<td>66.67</td>
<td>90.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MI</td>
<td>90.00</td>
<td>50.00</td>
<td>66.67</td>
<td>76.67</td>
<td>73.33</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IWR</td>
<td>60.00</td>
<td>73.33</td>
<td>86.67</td>
<td>73.33</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GV/GY-PE</td>
<td>76.67</td>
<td>66.67</td>
<td>66.67</td>
<td>73.33</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GV/PV</td>
<td>90.00</td>
<td>50.00</td>
<td>43.33</td>
<td>70.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CI/CVg: a classical index assigning genetic variation coefficient values as economic weights; CI/SDg: a classical index assigning values of the genetic standard deviation as economic weights; CI/h²: a classical index assigning values of heritability as economic weights; CI/BT: a classical index assigning values by trial as economic weights; MI: a multiplicative index; IWR: an index weighted by the average rank of families; GV/GY-PE: selection via the highest genotypic values assigned to the variables GY and PE; GV/PV: selection via the highest genotypic values for the variable PV.

The GV/ PV index showed values higher than 70% of coincidence with the other selection indices used in the selection of families. The highest coincidence based on GV/PV index (90%) was observed with the CI/BT index. The index based on GV/GY-PE showed the lower values of coincidence when compared to the other indices.
The GV/PV index showed the highest predicted percentage gain for the super-trait PV, with an increase of 10.29% in the next cycle for this population. Furthermore, the highest value for GY was CI/SDg (9.88%), and PE was GV/GY-PE (2.29%) (Table 2).

<table>
<thead>
<tr>
<th>Selection indexes</th>
<th>PH</th>
<th>PROL</th>
<th>M100</th>
<th>GY</th>
<th>PE</th>
<th>PV</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI/CVg</td>
<td>2.14</td>
<td>6.58</td>
<td>4.57</td>
<td>8.46</td>
<td>1.47</td>
<td>9.43</td>
</tr>
<tr>
<td>CI/SDg</td>
<td>2.03</td>
<td>5.13</td>
<td>1.90</td>
<td>9.88</td>
<td>0.53</td>
<td>9.38</td>
</tr>
<tr>
<td>CI/h²</td>
<td>2.74</td>
<td>5.40</td>
<td>5.15</td>
<td>8.03</td>
<td>1.57</td>
<td>9.16</td>
</tr>
<tr>
<td>CI/BT</td>
<td>2.21</td>
<td>4.55</td>
<td>3.64</td>
<td>8.43</td>
<td>2.08</td>
<td>10.19</td>
</tr>
<tr>
<td>MI</td>
<td>2.33</td>
<td>7.08</td>
<td>4.77</td>
<td>8.26</td>
<td>1.23</td>
<td>8.96</td>
</tr>
<tr>
<td>IWR</td>
<td>2.60</td>
<td>6.03</td>
<td>3.53</td>
<td>7.85</td>
<td>1.90</td>
<td>9.43</td>
</tr>
<tr>
<td>GV/GY-PE</td>
<td>1.95</td>
<td>1.46</td>
<td>2.04</td>
<td>5.93</td>
<td>2.29</td>
<td>8.14</td>
</tr>
<tr>
<td>GV/PV</td>
<td>2.13</td>
<td>4.38</td>
<td>2.29</td>
<td>8.40</td>
<td>2.05</td>
<td>10.29</td>
</tr>
</tbody>
</table>

CI/CVg: a classical index assigning genetic variation coefficient values as economic weights; CI/SDg: a classical index assigning values of the genetic standard deviation as economic weights; CI/h²: a classical index assigning values of heritability as economic weights; CI/BT: a classical index assigning values by trial as economic weights; MI: a multiplicative index: IWR: an index weighted by the average rank of families; GV/GY-PE: selection via the highest genotypic values assigned to the variables GY and PE; GV/PV: selection via the highest genotypic values for the variable PV. PH: average plant height; PROL: prolificacy; M100: a mass of 100 grains; GY: grain yield; PE: popping expansion; PV: expanded popcorn volume per hectare.

The predicted percentage gains (Table 2) revealed that the selection performed using genotypic values based on the GV/GY-PE index resulted in the identification of plants with lower average heights when compared to the other indices. However, this index was not suitable for the prediction of high gains in general; rather, it was suitable for PE only where the prediction gains were satisfactory.

DISCUSSION

Several studies have reported the selection for the most important characteristics in popcorn breeding; PE and GY (Santos et al., 2007a; Ribeiro et al., 2012; Freitas et al., 2014). Matta and Viana (2003) evaluated different selection methodologies among and within half-sibling families in the popcorn population; the values observed demonstrated the zero correlation between PE and GY. These authors suggested that selection for both characteristics is feasible. When selecting the best families for PE, the families with high GY are selected simultaneously, despite the aggravating null correlation factor presented by these traits. It should also be said that, among full-sib progenies, which is the case of this study, part of the dominance variance is also capitalized, thus allowing higher gains for selection, compared to the use of half-sib families (Souza Júnior and Pinto, 2000; Santos et al., 2007b; Hallauer et al., 2010). Vilarinho et al. (2002) selected for PE in S₁ and S₂ popcorn progenies and obtained significant gains for both PE and GY. When performing selection directly for GY, simultaneous selection for PE was not possible due to negative PE values. Because these traits are negatively correlated, direct selection via genotypic values of the characteristics becomes compromised (Dofing et al., 1991; Daros et al., 2004; Santos et al., 2007a; Rangel et al., 2011a).

Among the coincidence ratios obtained, the higher values were found when comparing the CI/CVg and MI (Table 1); these findings confirmed those obtained by Pedrozo et al. (2008).
These authors compared the efficiency of selection indices via the REML/BLUP methodology in three sugarcane populations. The greatest coincidence was observed between the high indices, although the coincidence values were relatively low. Freitas et al. (2013) performed a study of popcorn that compared selection indices and prediction methods via BLUP for the evaluation of genetic gain. These authors observed low coincidence values among the selection indices. The high degree of coincidence between the indices is considered a great estimate, as the results of selection between the different indices tend to be more consistent with a greater degree of coincidence between the families and the selected genotypes (Pedrozo et al., 2009).

It can be observed that the lowest coincidence rates were found when comparing selection via the genotypic values of GY and PE. Imbalance occurs in the selection of superior genotypes for other characteristics because this method considers only the two variables as a source of classification. These variables express null or negative correlations (Daros et al., 2004; Santos et al., 2007a; Rangel et al., 2011a); selection based only on these characteristics results in a lower coincidence index compared to other indices.

The predicted gain for GY (Table 2) was verified based on the GV/GY-PE index. It was observed that selection via the genetic values of GY and PE was not appropriate; compared to other indices, high gains were not observed. This can be explained by the null or negative correlations between these characteristics observed in other studies (Daros et al., 2004; Santos et al., 2007a; Rangel et al., 2011a).

Considering the predicted selection gains for GY and PE, the selection index via the genotypic values of popcorn volume (GV/PV) and the classical index with weights attributed by trial (CI/BT) showed the highest estimates for the two characteristics together. Using the CI/BT index, the estimates were 8.43% for GY and 2.08% for PE. Using the GV/PV index, the estimates were 8.40% for GY and 2.05% for PE. The last index resulted in greater selection gains for the most important characteristics. This form of selection can be easily employed for the acquisition of gains in popcorn breeding programs. This efficiency is because important characteristics simultaneously combine both PE and GY. This form of selection is therefore of use in future recurrent selection cycles.

The effectiveness of gains predicted by the GV/PV can be compared with the gains found by the classical index (CI/BT). Santos et al. (2007a) predicted genetic gains in a popcorn population and found that the indices of Smith (1936) and Hazel (1943) resulted in the highest percentage gains for the characteristics GY and PE when arbitrary weights were assigned to the traits. Use of the CI/BT index with weights attributed by the genetic standard deviation (CI/SDg) resulted in the highest percentage gain for GY. This procedure contemplates the greater weight attributed to GY, as the genetic standard deviation used as an economic weight is far superior to other deviations of the characteristics evaluated. In general, this index did not show efficiency in gains for the other characteristics. This resulted in a lower magnitude for PE compared to all indices, with an estimate of 0.53% (Table 2).

In previous recurrent selection cycles of the UNB-2U population, the largest simultaneous gains for GY and PE were obtained using the index described by Mulamba and Mock (1978) with economic weight attributed by the trials. Gains were 8.53, 15.30, and 7.78% for GY and 6.01, 10.97, and 5.11% for PE in the fifth, sixth, and seventh selection cycles, respectively (Rangel et al., 2011b; Ribeiro et al., 2012; Freitas et al., 2014). In the eighth selection cycle, the gains obtained by the same index were considerably lower; the percentage gains were 4.6% for GY and 3.61% for PE (Guimarães AG, Amaral Junior AT, Lima VJ, Leite...
JT, et al., unpublished results). Therefore, CI/BT and GV/PV were satisfactory in optimizing the selection of superior families for GY and PE. Good results were observed; the predicted gains for GY using the CI/BT and GV/PV indices were 8.43 and 8.40%, respectively (Table 2).

The gains predicted using the GV/PV index confirm that PV serves as a super-trait in the selection of popcorn progenies. This trait can be used to discriminate, via unification, families that are simultaneously superior for GY and PE.

CONCLUSIONS

The indices CI/BT, obtained via REML/BLUP, and GV/PV, obtained via REML, provided the most satisfactory results for the selection of superior GY and PE gains.

The insertion of the characteristic PV resulted in simultaneously optimized gains for GY and PE without the need to use arbitrary weights assigned to the trial. PV should be used as a “super-trait” in recurrent popcorn selection programs.

Conflicts of interest

The authors declare no conflict of interest.

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REFERENCES


Genetics and Molecular Research 16 (3): gmr16039769
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