Genetic Diversity among Brazilian Soybean Cultivars Based on SSR Loci and Pedigree Data

Regina Helena Geribello Priolli1*, José Baldin Pinheiro1, Maria Imaculada Zucchi2, Miklos Maximiliano Bajay2 and Natal Antonio Vello1
1Departamento de Genética; Escola Superior de Agricultura “Luiz de Queiroz”; Universidade de São Paulo; Av. Pádua Dias, 11 C.P: 83; 13400-970; Piracicaba - SP - Brasil. 2Centro de Pesquisa e Desenvolvimento de Recursos Genéticos Vegetais; Instituto Agronômico de Campinas; Av. Theodoreto de Almeida Camargo, 1500; C.P.: 28, 13001-970; Campinas - SP - Brasil

ABSTRACT

In this study, simple sequence repeats (SSR) loci and pedigree data were used to investigate the genetic relationship in a group of 168 Brazilian soybean cultivars. Eighteen SSR loci produced an average of 5.06 alleles and a mean gene diversity of 0.58 for the cultivars studied. Genetic distance (GD) was determined using the modified Roger’s Wright distance, and a final dendrogram was in agreement with the cultivar pedigree. A distance matrix based on the coefficient of parentage scores was also generated for the cultivars, which ranged from 0 to 1, with a mean of 0.18, whereas SSR-based genetic similarity (1-GD) ranged from 0.01 to 0.90, with a mean of 0.25. Mantel’s Z test showed that the similarity matrices generated from both the data sets were low, but significantly correlated (r = 0.31, p<0.001). The results showed that SSR data and pedigree analyses could help to quantify more accurately the degree of relationship among the soybean cultivars.

Key words: Coefficient of parentage, Glycine max (L.) Merr., microsatellite markers

INTRODUCTION

Soybean, Glycine max (L.) Merrill is one of the most important crop species worldwide and it has been the subject of intensive breeding. One of the pre-requirements for successful breeding strategies is the complete understanding of the genetic diversity of the crop plant. Several methods have been used to investigate the genetic variation in soybean. Morphological and agronomic traits have been employed (Perry and McIntosh, 1991; Sneller et al., 1997). Although the evaluation based on these data are essential for applied soybean breeding, genotype x environment interactions greatly limit the soybean lines that can be directly compared based on agronomic and morphological data. The Malécot coefficient of parentage (COP) (Kempthorne 1957) is an important method to estimate the genetic diversity based on pedigree analyses. It measures indirectly the genetic diversity among the cultivars by estimating, from pedigree records, the probability that alleles, in a locus, are identical by descent. Its use has been widespread for several plant cultivars, including soybean (Cox et al., 1985; Vello et al., 1988; Gizlice et al., 1996). However, pedigree records are not always available or detailed enough for this type of analysis, especially when large numbers of breeding lines or cultivars are being assessed.

* Author for correspondence: rpriolli@esalq.usp.br
DNA marker analyses are an alternative method to estimate the diversity of soybean in breeding programs, especially when the details of pedigree information are lacking. These markers include SSR, RFLP (restriction fragment length polymorphism), AFLP (amplified fragment length polymorphism) and RAPD (random amplified polymorphic DNA) markers (Akkaya et al., 1992; Keim et al., 1992; Maugham et al., 1996; Li and Nelson, 2001). Each strategy is based on different principles to detect the variation and they present distinct advantages and limitations in terms of costs, time consumption, accuracy and efficiency. Microsatellites or simple sequence repeat (SSR) markers have been especially valuable to estimate genetic diversity in soybean due to their abundance, the high level of polymorphism, the codominant characteristic, PCR based detection and the fact that they have known positions in the genome. Furthermore, the use of fluorescence labeled primers and DNA sequencers make it possible to accurately identify and easily score alleles at a number of loci at once. In soybean, high levels of polymorphism at SSRs have been reported for the number of alleles per locus and gene diversity (Akkaya et al., 1992; Narvel et al., 2001). Each strategy is based on different principles to detect the variation and they present distinct advantages and limitations in terms of costs, time consumption, accuracy and efficiency. Microsatellites or simple sequence repeat (SSR) markers have been especially valuable to estimate genetic diversity in soybean due to their abundance, the high level of polymorphism, the codominant characteristic, PCR based detection and the fact that they have known positions in the genome. Furthermore, the use of fluorescence labeled primers and DNA sequencers make it possible to accurately identify and easily score alleles at a number of loci at once. In soybean, high levels of polymorphism at SSRs have been reported for the number of alleles per locus and gene diversity (Akkaya et al., 1992; Narvel et al., 2001; Priolli et al., 2002; Hadcovicová and Kraikl, 2003). It was also found that RFLPs, AFLPs and SSRs in soybean were highly correlated, while SSRs generated hypervariable polymorphisms (Rongwen et al., 1995).

The estimation of relationships among the soybean accessions using both the molecular markers and COPs has been tested to predict the progeny variance and to study the heterosis (Helms et al., 1997, Manjarrez-Sandoval et al., 1997). In soybean, high heterosis values were estimated for seed yield and their components (Pandini et al., 2002), but commercial F1 hybrids were not economically practicabile. The association between the COP and molecular markers (RFLP and AFLP markers) has been shown to be statistically significant; however, the correlations were demonstrated to range from low (Bonato et al., 2006) to moderate (Kisha et al., 1998).

In order to evaluate whether SSRs and COP determined similar genetic relationship estimates, the present study considered: (1) assess the genetic diversity among a group of Brazilian soybean cultivars using SSR markers; and to (2) determine the correlation between estimates of genetic diversity measured by COP and SSR markers.

**MATERIALS AND METHODS**

A group of 168 soybean elite cultivars, developed and released by Brazilian public and private institutions, was selected to represent the range of cultivars grown in Brazil. The pedigree and seeds of each cultivar were obtained according to Priolli et al., (2002). Thirty to fifty plants of each soybean cultivar were grown in a greenhouse for DNA isolation. The equivalent of 30 leaves of tissue samples were collected from each cultivar, frozen in liquid nitrogen and lyophilized for 1-2 days. DNA was isolated from the bulked lyophilized leaf tissue of the plants of each cultivar by a mini-prep procedure based on Doyle and Doyle (1990). The DNA concentration was estimated by comparison to known concentrations of lambda phage DNA on 0.8% agarose gel.

Eighteen SSR loci with either di- or tri-nucleotide repeats were selected for this study. They were chosen for their distribution across the soybean genome and amplification quality. All the SSR primer sequences, except for the RGA loci shown below, have been published previously (Morgante and Olivieri, 1994; Cregan et al., 1999). The sequence of RGA primers is: F: 5’TTGGCAAGGGAGAGAA3’ and R: 5’TCTGAAATGGCCTTCTTGACCC3’.

Fluorescent labeling of the alleles with the universal tail primers was used (Missiaglia and Grattapaglia, 2006). The PCR reaction was prepared in 20-µL total volume mix, containing 1X PCR buffer (10 mM Tris-HCl, pH 8.9, 50 mM KCl), 3.0 mM MgCl2, 0.2 µM dNTP, 1.0 U Taq DNA polymerase (LabTrade, Brazil) and 30 ng soybean DNA. For each locus SSR, 0.1 µM of each one of the two primers (i.e., forward tailed primer and reverse primer) with 0.01 µM labeled universal primer were added to the PCR mix. All amplifications were carried out using a PTC-100 thermal cycler (MJ Research, Inc) in a “touchdown” program as follows: 94°C (4 min), then 10 cycles at 94°C (40 sec) /50°C decreased 1°C/cycle (40 sec) /72°C (1 min), and 30 cycles at 94°C (40 sec) /40°C (40 sec) /72°C (1 min) with a final extension at 72°C for 5 min. After the amplifications, the PCR product was combined with loading buffer (1.7 µL) containing a fragment size standard RED-labeled (Genescan 500 ROX - Applied Biosystems) and denatured at 95°C for 5 minutes. Electrophoresis was conducted in 1 X TBE (Sambrook et al., 1989), 5% polyacrilamide.
gels for 2.5 hours using ABI377 DNA sequencer (Applied Biosystems). Data were collected automatically by detecting the distinct fluorescence and analyzed by GeneScan/Genotyper softwares (Applied Biosystems). The amplified fragments produced by SSR were considered as alleles of a single locus. Genetic diversity of each SSR loci was calculated according to Weir (1990).

The analyses of diversity among the cultivars were based on a modified Roger’s distance method using TFPGA software, version 1.3 (Miller 1997). The consensus UPGMA (unweighted pair-group method with arithmetic averages) dendrogram was obtained using NTSYS-PC software, version 2.01 (Rohlf 2000). A bootstrap procedure with 1,000 units of re-sampling was carried out for SSR data using BOOD software (Coelho 2000).

The values of the coefficient of parentage (COP) were obtained for pair-wise combinations of two genotypes and corresponded to the probability of an allele in a given locus to be identical by descent to alleles in the same locus from another cultivar (Kempthorne 1957). The coefficient of parentage was calculated using the procedure ‘proc inbreed’ of the SAS software, version 8. Using the assumptions suggested by Cox et al., (1985), COP was considered to range from 0 (cultivars with completely different pedigrees) to 1 (cultivars with the same genetic constitution). For two genotypes related by three or more generations of backcrossing, the coefficient was assumed to be 1. The matrix of genetic distance obtained from the SSR data was converted into a genetic similarity matrix (1- modified Roger’s distance). The genetic similarity matrices obtained from the SSR and COP data were compared by measuring the degree of correlation (r) between them by computing the product-moment correlation and by the Mantel test statistic (Z) using NTSYS-PC software, version 2.1 (Rohlf 2000). The comparisons were carried out to verify whether genetic similarities generated by the SSR and COP data provided similar genetic measures.

**RESULTS**

The investigated 18 SSR loci were polymorphic, as shown in Table 1. The number of alleles per locus ranged from two to eight, with an average of 5.06 alleles per locus distributed in a total of 91 alleles. The genetic diversity (GD), indicative of the effectiveness of SSR loci information, showed broad amplitude, ranging from 0.070 to 0.826, with 0.58 ± 0.20 mean value. However, except for the aforementioned two loci, the majority of values exhibited moderate (0.431 for locus Satt102) to high (0.826 for locus Satt308) ranges. A comparison between the similarity coefficients calculated using COP and SSR marker data is presented in Fig. 1. The distribution of values for both the methods was slightly different. Genealogical similarities covered a greater range (0 to 1) - average 0.18 - in 168 pair-wise cultivar comparisons. However, distribution was skewed towards lower values (89% of the similarity values from 0.01 to 0.30), and sister lines with a COP value of one were shown by SSR data to be not completely similar. The SSR genetic similarity covered a lower range, from 0.01 to 0.90, although skewed towards higher values (91% of the SSR value from 0.11 to 0.40), average 0.25 in 168 pair-wise cultivar comparison. These data suggested that SSR markers were more effective to differentiate the pairs of accessions with lower levels of genetic variation. It was possible to differentiate the totality of the investigated cultivars with SSR markers, even those with the highest identity values from the genealogy analyses.

The correlation (r) between the genetic similarity and the coefficient of parentage was highly significant (p<0.001); however, with low value (r = 0.31). A dispersion graph (Fig. 2) showed that the points presented a grouping trend approximately as a straight line. However, several points were located outside the trend, representing, among others, the pairs of cultivars that showed COP=1, but with similarity coefficient from SSR of approximately zero.
**Table 1 - Polymorphism characterization for 18 SSR loci in 168 Brazilian soybean cultivars.**

<table>
<thead>
<tr>
<th>Locus</th>
<th>Allele number</th>
<th>Allele size (bp)</th>
<th>Genetic diversity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Satt186</td>
<td>8</td>
<td>225,247,250,253,256,258,262,265</td>
<td>0.766</td>
</tr>
<tr>
<td>Satt173</td>
<td>8</td>
<td>205,217,220,232,244,250,266,281</td>
<td>0.752</td>
</tr>
<tr>
<td>Satt005</td>
<td>8</td>
<td>163,172,175,181,184,187,205,208</td>
<td>0.619</td>
</tr>
<tr>
<td>Satt308</td>
<td>7</td>
<td>154,169,172,175,178,190,193</td>
<td>0.826</td>
</tr>
<tr>
<td>Satt009</td>
<td>7</td>
<td>178,205,226,229,241,247,262</td>
<td>0.781</td>
</tr>
<tr>
<td>Satt154</td>
<td>6</td>
<td>275,281,284,290,293,296</td>
<td>0.697</td>
</tr>
<tr>
<td>Sct_189</td>
<td>5</td>
<td>178,181,202,205,211</td>
<td>0.729</td>
</tr>
<tr>
<td>Satt263</td>
<td>5</td>
<td>221,227,239,242,251</td>
<td>0.707</td>
</tr>
<tr>
<td>SOYHSP176</td>
<td>5</td>
<td>112,224,228,230,234</td>
<td>0.576</td>
</tr>
<tr>
<td>Satt406</td>
<td>4</td>
<td>257,263,350,353</td>
<td>0.638</td>
</tr>
<tr>
<td>Satt002</td>
<td>4</td>
<td>121,133,142,148</td>
<td>0.586</td>
</tr>
<tr>
<td>Satt335</td>
<td>4</td>
<td>150,156,162,168</td>
<td>0.562</td>
</tr>
<tr>
<td>Satt045</td>
<td>4</td>
<td>154,160,165,176</td>
<td>0.560</td>
</tr>
<tr>
<td>Satt307</td>
<td>4</td>
<td>166,175,181,193</td>
<td>0.512</td>
</tr>
<tr>
<td>Satt102</td>
<td>4</td>
<td>152,167,170,176</td>
<td>0.431</td>
</tr>
<tr>
<td>Satt309</td>
<td>4</td>
<td>144,147,153,156</td>
<td>0.550</td>
</tr>
<tr>
<td>SOYGY2</td>
<td>2</td>
<td>170,174</td>
<td>0.131</td>
</tr>
<tr>
<td>RGA</td>
<td>2</td>
<td>184,186</td>
<td>0.070</td>
</tr>
</tbody>
</table>

Total 91
Mean 5.06

**Figure 1 - Distribution of the genetic similarity coefficients by SSR analyses and coefficient of parentage calculated for 14,028 pairs of genotypes.**
All of the cultivars were able to be uniquely identified by SSR (Figs. 3 and 4). The results also showed that the cultivars were grouped in six closely related clusters, presenting 75% of the dendrogram distance between each other represented by the letters A to F. The cultivars EMBRAPA 1 (IAS 5 RC) and RS 9 (Itaúba) had the smallest genetic distance (0.13). These cultivars had a similar genetic origin, where the first resulted from a backcross of ‘IAS 5’ for five generations and, the second, from a cross between ‘FT 2’ and ‘IAS 5’.

Bootstrap results of the molecular analyses showed that SSR data were not robust (22 nodes with significant support; >50%). This observation could be due to the amount of data generated from the SSR data (18 loci, 91 alleles). However, the results were highly coherent in relation to the ancestral descent of the groups, and identified groups with variable degrees of parentage. For instance, the cultivars BRSM A Sambaíba, BR28 Seridó, Dourados, FT 5 Formosa, and almost all cultivars in the group named as “D”, were in the same group as the Santa Rosa cultivar. Furthermore, all of them were either descended from “Santa Rosa” or from a selection of it. For the same reason, ‘BR6 Nova Bragg’, ‘BR36’ and ‘CEP 12 Cambará’ were in the same group B as their ancestor ‘Bragg’. Similarly, groups A, C, E and F had small sets of cultivars, such that all of them were related to the same ancestral genotype.

The dendrogram relating the 168 cultivars based on the parentage (Figs. 5 and 6) expressed the distinction of groups with maximum and minimum similarities. Values 1 (maximum parentage) were observed in 17 pair-wise comparisons, while 0 (absence of parentage) was obtained for 31 comparisons. The analysis of the pedigrees and COP of each group showed that clustering was determined by shared ancestors. ‘UFV 1’, for instance, was a selection of ‘Viçoja’, which in turn resulted from ‘D49-2491’ X ‘Improved Pelican’. ‘Bragg’ and ‘FT Cristalina’ shared a common parent, ‘D492491’. ‘Paraná’ and ‘IAS 5’ also shared the common parents, ‘Hill’ and D52-810. All were ancestors of other groups. Thus, it could be inferred that the ancestral cultivars originating the present group of 168 cultivars were ‘Davis’, ‘D492491’ (sister line of ‘Lee’), ‘Hill’, ‘Hood’, ‘Improved Pelican’ and ‘Santa Rosa’.

**Figure 2** - Plot of SSR based genetic similarity and coefficient of parentage for 14,028 pairs of the soybean cultivars.
Figure 3 - Part I of dendrogram of 168 soybean cultivars revealed by UPGMA cluster analysis of 18 SSR loci based on a modified Roger’s Wright distance. All bootstrap values of 1,000 replicates are shown at the corresponding knots. The brackets on the right indicate the common parent identified in a cluster. X marks refer to cultivars present in a cluster not corresponding to their common parent.
Figure 4 – Part II of dendrogram of 168 soybean cultivars revealed by UPGMA cluster analysis of 18 SSR loci based on a modified Roger’s Wright distance. All bootstrap values of 1,000 replicates are shown at the corresponding knots. The brackets on the right indicate the common parent identified in a cluster. X marks refer to cultivars present in a cluster not corresponding to their common parent.
Figure 5 - Part I of dendrogram of 168 soybean cultivars revealed by UPGMA cluster analysis of the coefficient of parentage. The brackets on the right indicate the common parent identified in a cluster. X marks refer to cultivars present in a cluster not corresponding to their common parent.
Figure 6 - Part II of dendrogram of 168 soybean cultivars revealed by UPGMA cluster analysis of the coefficient of parentage. The brackets on the right indicate the common parent identified in a cluster. X marks refer to cultivars present in a cluster not corresponding to their common parent.
DISCUSSION

Polymorphism and gene diversity for the majority of the SSR loci in this study were consistent with previous studies of soybean elite cultivars (Akkaya et al., 1992; Diwan and Cregan; 1997; Song et al., 1999; Narvel et al., 2000; Priolli et al., 2002; Hudcovicová and Kraikl, 2003); however, the loci SOGY2 and RGA presented only two alleles and low gene diversity (Table1). One possible reason for the reduced variability observed was that the loci were either associated with DNA sequences coding for proteins or were derived from the expressed sequence tag simple sequence repeats (EST-SSR) obtained from the sequencing of partial complementary DNA (cDNA). Markers derived from genomic libraries also contained more repeat units as well as a greater range of allele sizes and genetic diversity than markers isolated from EST libraries. The difference in polymorphism between the soybean SSRs derived from the two sources was consistent with the differences reported in rice (Cho et al., 2000), sugarcane (Cordeiro et al., 2001), tomato (Arshchenkova and Ganal, 2002), wheat (Eujayl et al., 2002), and barley (Thiel et al., 2003).

In soybean mapping populations, 133 EST-SSR loci were characterized but only 18% were polymorphic. Alternatively, studying 1,002 SSR loci derived from genomic libraries, 43% primer sets were polymorphic among parental genotypes (Song et al., 2004).

Based on the results of this study, 18 microsatellite loci did not produce scorable bands (91) to detect the genetic relationships with strong bootstrap support among soybean cultivars (only 22 of 154 nodes at a bootstrap level of 50% confidence). The low level of bootstrap support for SSR data indicated that a higher number of loci were required to generate robust relationships. The number of SSR loci necessary to generate sufficient SSR data to accurately detect the genetic relationships remained unclear. Some studies have indicated that a relatively small number of SSRs was sufficient to obtained high resolution estimates of relatedness in soybean (Powell et al., 1996). However, some others have demonstrated that reliable estimates involving many individuals could require the screening of several hundred loci (Zhiyotovsky and Feldman, 1995).

The similarity coefficients obtained by the SSR analyses in the present study (average of 0.25), although higher than the COP values (average of 0.18), were smaller than those obtained in previous studies using other molecular markers. Therefore, it suggested that the SSR markers were highly effective in distinguishing the genetic material with a narrow genetic base, such as the Brazilian soybean germplasm. The average similarity coefficient was 0.82 for 38 soybean cultivars using RAPD markers (Abdelnoor et al., 1995), and 0.61 in 317 soybean cultivars using AFLP markers (Bonato et al., 2006). The higher effectiveness of SSR markers in distinguishing closely related soybean cultivars has already been shown in a previous study. Twelve SSR loci were able to distinguish morphologically similar groups of 186 soybean cultivars, presenting a mean similarity coefficient of 0.46 (Priolli et al., 2002).

A significant correlation (p<0.001) with low value was observed (r = 0.31) between the measurements of the SSR-based genetic similarity and COP. These results were expected since SSR markers identify alleles that were different in state and they reflected the evolution of neutral alleles in the absence of selective pressure. COP estimates the amount of alleles that are identical by descent from common ancestors selected in breeding programs. In germplasm with a restricted genetic base, such as the Brazilian soybean germplasm, the divergence among parents is obtained by their pedigree analyses and on many occasions is a prior requirement in cultivar development.

Similar results of correlation have been demonstrated by several studies with other markers. The comparison of RFLP-based genetic similarity with COP estimates for a set of barley cultivars showed low correlation (r = 0.21) for the winter type (Graner et al., 1994). The correlation between isozyme-based genetic similarity and COP estimates for wheat cultivars also showed a low value (r = 0.27) (Cox et al., 1985). Lower values (r = 0.12) were found in soybean by comparing estimates of the genetic similarity based on AFLP markers and COP (Bonato et al., 2006). Moderate correlation (r = 0.61) was obtained of COP and simple matching coefficient based on RFLP data in elite North American soybean, previously grouped according to their origin (Kisha et al., 1998). The authors suggested that selection or drift appeared to have influenced the correlation, since each gene pool derived from a few major ancestors with gene frequency differences. Moderate correlations were found based on SSR data and COP in other crops, as
Genetic Diversity among Brazilian Soybean Cultivars Based on SSR Loci and Pedigree Data

observed in maize, with $r = 0.55$ (Plaschke et al., 1995) and in sugarcane, where $r = 0.42$ (Lima et al., 2002).

The dispersion graph, obtained by analyzing the correlation between the SSR-based genetic similarity and COP estimates for all the investigated genotypes, showed a clustering trend according to a regression line (Fig. 2). However, the genotypes that presented the highest values were the ones located furthest from the regression line. Thirty-one soybean cultivars with values COP=1 and genetic similarity of approximately 0, belonged to this category. The group also consisted of cultivars whose genetic similarity values based on SSR markers were high, but pedigree information showed very distant relationship (COP = 0 or close to 0). The results suggested that cultivar pedigree-derived information was not always accurate and that the association with molecular data could provide more precise information to soybean breeding programs.

Dendrograms generated by different methods (SSRs and COPs) were partially coincident. Both the methods successfully distinguished similar groups and the variation at SSR loci was in agreement with the cultivar pedigree information. EMBRAPA 1 (IAS 5 RC) and RS 9 (Itaúba) cultivars, which had the smallest genetic distance, could not be distinguished by 12 SSR loci (Priolli et al., 2002) and presented the maximum relatedness coefficient (COP=1). Moreover, several cultivars clustered very closely, thus demonstrating the narrow genetic basis of the gene pool from Brazilian soybean germplasm. It was estimated in the 1980s that approximately 70% of the cultivars developed for the Southern regions of Brazil were descendents from the cultivars ‘Hill’, ‘Hood’ or both (Bonetti 1983). The recommended cultivars for the aforementioned growing season were descendents from 26 cultivars and from that total, only four were responsible for approximately half of the gene pool (Hiromoto and Vello, 1986). The narrow genetic basis of the Brazilian gene pool could also be observed by the COP values. In 69 soybean cultivars, a mean COP value of 0.16 was estimated (Vello et al., 1988). Similar COP values of 0.21 were obtained in larger samples of 100 and 90 soybean cultivars by Bonato et al., (2006) and Miranda et al., (2007), respectively. Actually, the parental selection of elite cultivars based on COP estimates was the first procedure for the enhancement of the genetic basis of cultivated germplasm. Furthermore, the soybean type used in oriental countries for human consumption, for example, vegetable soybean or edaname (Yokomizo and Vello, 2003) also could be selected as parents in breeding type soybean, since the genetic flux has been restricted between both the soybean types.

It could be concluded that the COP provided a good estimation of genetic diversity when the complete pedigree was known and SSR markers also provided a good estimation of genetic diversity when sufficient numbers were used. Despite the low correlation between both the estimates, that, as mentioned above, presented closely related formation of clusters, it could be suggested that the use of these SSR loci associated to pedigree information would allow the breeders to make reliable crossings or to strategically plan the breeding program.

ACKNOWLEDGEMENTS

The authors thank The State of São Paulo Research Foundation (FAPESP) for financial support for this study. R.H.G.P received post-doctorate fellowship from FAPESP. N.A.V. and J.B.P. are recipients of research fellowships from National Council for Scientific and Technological Development (CNPq).

RESUMO

Locos microsatélites e dados de genealogia foram utilizados para avaliar a diversidade genética de um grupo de 168 cultivares brasileiras de soja. Os dezoito locos utilizados apresentaram em média 5,06 alelos por loco e coeficiente de diversidade genética médio de 0,58. O dendrograma final resultante da matriz de distância genética de Roger modificada por Wright, apresentou boa concordância com a ancestralidade dos grupos formados. Também foi estimado os coeficientes de parentesco entre as cultivares, sendo observada variação de 0 a 1 com média de 0,18, enquanto que as similaridades para os locos microsatélites (1- GD) variou de 0,01 a 0,90 com média de 0,25. A correlação entre as duas matrizes obtidas determinada pelo teste Z de Mantel apresentou valor baixo, 0,31, mas significativo (p<0,001). Os resultados obtidos sugerem que os locos...
microssatélites aliados às informações de genealogia proporcionam melhor análise da diversidade genética de cultivares de soja.

REFERENCES


Cordeiro, G.M., Casu, R., McIntyre, C.L., Manners, J.M., Henry, R.J. (2001), Microsatellite markers from sugarcane (Saccharum spp.) ESTs cross transferable to erianthus and sorghum. Plant Sci., 160, 1115-1123.


Received: January 09, 2008; Revised: May 21, 2008; Accepted: June 17, 2009.