Genetic polymorphism among 14 elite *Coffea arabica* L. cultivars using RAPD markers associated with restriction digestion

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Abstract

Knowledge of the genetic variability among genotypes is important for the transfer of useful genes and to maximize the use of available germplasm resources. This study was carried out to assess the genetic variability of 14 elite *Coffea arabica* cultivars using random amplified polymorphic DNA (RAPD) associated with a prior digestion of genomic DNA with restriction endonucleases. The accessions were obtained from the *Coffea* collection maintained at the Instituto Agronômico do Paraná (IAPAR), located in Londrina, Paraná, Brazil. Twenty-four informative RAPD primers, used in association with restriction enzymes, yielded 330 reproducible and scorable DNA bands, of which 224 (68%) were polymorphic. The amplified products were used to estimate the genetic variability using Dice’s similarity coefficient. The data matrix was converted to a dendrogram and a three-dimensional plot using principal coordinate analysis. The accessions studied were separated into clusters in a manner that was consistent with the known pedigree. The associations obtained in the dendrogram and in the principal coordinate analysis plot suggest the probable origin of the Kattimor cultivar. The RAPD technique associated with restriction digestion was proved to be a useful tool for genetic characterization of *C. arabica* genotypes making an important contribution to the application of molecular markers to coffee breeding.

Key words: *Coffea arabica*, RAPD, restriction endonucleases.

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Introduction

The wild coffee plant (*Coffea arabica* L.) is indigenous to Ethiopia. It was discovered in about 850 AD and was spread to other parts of the Islamic world by pilgrims. During the 17th century, coffee beverage consumption spread all over Europe, with great profit for the only producing country, the Yemen. In 1616, a Dutch trader succeeded in stealing seeds and in cultivating them in the colony of Java. Progenies from this first introduction were planted in the Surinam colony in South America (Berthaud and Charrrier, 1988) and brought from there to French Guiana. The introduction of some coffee plants to Northern Brazil in 1727 was an important landmark in the economic activity of Brazil. In few years with the establishment of plantations in many states such as Rio de Janeiro, São Paulo, and Minas Gerais, Brazil became the main coffee producer and exporter in the world. Coffee was very important for the foundation and development of several new cities at the end of the nineteenth and during the last century. Nowadays, coffee is the second most important agricultural product of Brazil.

Until recently, genetic diversity among species or cultivars was determined with morphological or isozyme markers. However, these markers are often unsuitable to measure genetic variation. The analysis of six isozyme patterns in different *C. arabica* accessions revealed absence of polymorphism contrasting with the high level of morphological variation and suggesting that isozymes are not appropriate for the study of genetic diversity and for *C. arabica* accession identification (Berthou and Trouslot, 1977). Located in southern Brazil, Paraná state is an important coffee producer. In this state, the Research Center named Instituto Agronômico do Paraná (IAPAR) maintains a germplasm bank with eight *Coffea* species and several varieties, cultivars, and more than a thousand progenies of *C. arabica* and *C. canephora*. Despite its great importance, this *Coffea* collection lacks information about its genetic variability. Thus, much of the applied research is conducted without the genetic knowledge mainly when it concerns the DNA level.
In the last ten years, techniques based on DNA markers have been used to detect variation at DNA level and have proven to be very effective for distinguishing between closely related genotypes. The development of the random amplified polymorphic DNA (RAPD) markers (Williams et al., 1990; Welsh et al., 1991) allowed genome characterization in several plant groups. The RAPD technique has been useful in studying polymorphism, identifying genes of interest, and characterizing genetic resources. In Coffea, RAPD markers were used to identify polymorphism in different coffee accessions (Orozco-Castillo et al., 1994; Lashermes et al., 1996) and to analyze spontaneous and subs spontaneous C. arabica trees from Ethiopia (Anthony et al., 2001). In this study, the RAPD technique were associated with restriction digestion of genomic DNA and used to identify the genetic polymorphism among 14 C. arabica cultivars acquired from the IAPAR Coffea Germplasm Collection.

Material and Methods

Fourteen C. arabica genotypes representing elite cultivars widely used in breeding programs were used in the current study. The accessions were obtained from the Coffea Germplasm Collection of the Instituto Agronômico do Paraná (IAPAR), Londrina, Brazil (Table 1).

Genomic DNAs were isolated from fresh leaf tissue following the CTAB method (Doyle and Doyle, 1987), except that CTAB was replaced by MATAB (Mixed Alyltrimethylammonium Bromide, Sigma) in the extraction buffer. DNA concentration was estimated using a fluorometer (DNAQuant 200, Höefer-Pharmacia), according to the manufacturer’s instructions. DNA samples, obtained from at least five different plants of each accession, were adjusted to 10 ng/µL. The individual DNA samples were bulked (Michelmore et al., 1991) by accessions and used in the amplification reactions in a 15 µL volume containing 1x PCR buffer (75 mM Tris/-HCl, 50 mM KCl, 2.0 mM MgCl₂, and 20 mM (NH₄)₂SO₄); 0.2 mM each of dATP, dTTP, dCTTP, and dGTP; 0.4 µM primer (Operon Technologies); 0.9U Taq DNA polymerase (Biotools); and 20 ng template DNA. For restriction digestion, genomic DNA was incubated for 1 h with one of the following enzymes, Bam HI, Eco RI, or Hae III, which was added directly in the amplification reaction. DNA amplification was carried out using a PTC 100 (MJ Research) thermal cycler, programmed with 3 min at 94 °C for initial DNA denaturation, followed by 48 cycles of 1 min at 94 °C, 1 min 45 s at 38 °C, and 2 min at 72 °C. The final cycle was followed by a 7 min extension at 72 °C. The samples were stored at 4 °C until electrophoresis. Amplified products were resolved in 1.2 % agarose gel in TAE buffer (40 mM Tris-acetate, 1 mM EDTA, pH 8.0) at 120 V for 3 h and stained with ethidium bro-

Table 1 - Genotypes, accession numbers and origin types of 14 arabica coffee studied.

<table>
<thead>
<tr>
<th>Genotypes Accession numberb</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 - Mundo Novo IAC 376-4</td>
<td>Improved line resulted from Bourbon</td>
</tr>
<tr>
<td>2 - Catuaí Vermelho IAC-81b</td>
<td>Catuca x Mundo</td>
</tr>
<tr>
<td>3 - Caturra Vermelho</td>
<td>Mutant of a Bourbon type coffee</td>
</tr>
<tr>
<td>4 - Caturra Amarello</td>
<td>Mutant of a Bourbon type coffee</td>
</tr>
<tr>
<td>5 - Villasarchi</td>
<td>Mutant of Bourbon</td>
</tr>
<tr>
<td>6 - Colombia Amarello</td>
<td>Catimor germplasm</td>
</tr>
<tr>
<td>7 - IAPAR 77028</td>
<td>Sarchimor germplasm</td>
</tr>
<tr>
<td>8 - IAPAR-59</td>
<td>Sarchimor germplasm</td>
</tr>
<tr>
<td>9 - Tupi LC 1669-33</td>
<td>Sarchimor germplasm</td>
</tr>
<tr>
<td>10 - IAPAR 75163-21-10</td>
<td>Sarchimor germplasm</td>
</tr>
<tr>
<td>11 - IAPAR 75163-12</td>
<td>Sarchimor germplasm</td>
</tr>
<tr>
<td>12 - Kattimor</td>
<td>Unknown origin</td>
</tr>
<tr>
<td>13 - F₁</td>
<td>(IAPAR 59 x Mundo IAC-81)</td>
</tr>
<tr>
<td>14 - F₂</td>
<td>(IAPAR 59 x Mundo IAC-81)</td>
</tr>
</tbody>
</table>

aIAC - Instituto Agronômico de Campinas.  
b Number in the active germplasm collection at Instituto Agronômico do Paraná (IAPAR), Londrina, Paraná, Brazil.

The RAPD profiles were visualized under UV light and photographed with a video camera.

Data analysis

DNA markers were scored for the presence (1) and absence (0) of homologous DNA bands. The genetic similarity among accessions was estimated using the Dice coefficient of the NTSYS package (Numerical Taxonomy and Multivariate Analysis for personal computer), version 2.1 (Rohlf, 2000). A dendrogram was constructed using the UPGMA (unweighted pair-group method using arithmetic averages) method. The matrix of genetic similarity was also used to obtain a principal coordinate analysis (PCOORDA) plot to resolve the patterns of variation among genotypes. The bootstrap method was employed to evaluate the reliability of tree topology. The calculations were performed with the BOOD software, version 3.0 (Coelho, 2001). The cophenetic coefficient between the matrix of genetic similarity and the dendrogram were computed using an appropriate routine of the NTSYS-pc software. The significance of the cophenetic correlation was tested by using the Mantel correspondence test (Mantel, 1967).
Results and Discussion

The RAPD technique associated with prior digestion of genomic DNA with restriction enzymes was used for detection of polymorphism in 14 elite *C. arabica* cultivars (Table 1). A total of 330 highly reproducible markers was analyzed of which 224 (68.0%) were polymorphic. The data matrix with the values of genetic similarities, the resulting dendrogram, and the plot of the principal coordinate analysis with the graphic distribution of the genotypes, are shown in Table 2 and Figures 1 and 2, respectively. The high value of cophenetic correlation (r = 0.91) between the matrix of genetic similarity and the dendrogram indicates the extent to which the clustering of genotypes accurately represented the estimates of genetic similarities among the coffee accessions studied.

The results revealed that the combination of restriction digestion and RAPD amplification of genomic DNA represented an excellent approach for the identification of polymorphism in elite *C. arabica* cultivars. According to Williams *et al.* (1990), restriction digestion of genomic DNA prior to PCR amplification simplifies the pattern of amplified products and changes the relative intensities of the bands. Moreover, digestion of genomic DNA may reveal a restriction site polymorphism located between the primer-binding sites in an otherwise monomorphic band. Koebner (1995) showed that in wheat, restriction digestion of genomic DNA with different endonucleases before PCR amplification improved the level of polymorphism. These results were probably due to a greater efficiency in primer annealing along shorter DNA fragments, where a simplified secondary DNA structure is less likely to interface with the process. In this way, different priming sites may become accessible depending on the restriction enzyme em-

![Figure 1](image1.png)

**Figure 1** - UPGMA dendrogram of 14 *C. arabica* accessions based on Dice’s genetic similarity. Numbers on branches are bootstrap values (%) generated after 1000 permutations.

![Figure 2](image2.png)

**Figure 2** - Principal coordinate analysis of genetic distance among 14 elite *C. arabica* cultivars. The first, the second, and the third principal coordinates explain 22.6%, 11.6%, and 10.6% of the total variation, respectively. The numbers correspond to those listed on Table 2.

Table 2 - Genetic similarity based on the Dice coefficient applied to 14 *C. arabica* cultivars.

<table>
<thead>
<tr>
<th>Accessions</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14</th>
</tr>
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<td>1-Mundo Novo IAC 376-4</td>
<td>1.00</td>
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<tr>
<td>2-Catuai Vermelho IAC-81</td>
<td>0.81</td>
<td>1.00</td>
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<tr>
<td>3-Caturra Vermelho</td>
<td>0.81</td>
<td>0.87</td>
<td>1.00</td>
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<tr>
<td>4-Caturra Amarelo</td>
<td>0.78</td>
<td>0.83</td>
<td>0.87</td>
<td>1.00</td>
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<tr>
<td>5-Villasarchi</td>
<td>0.75</td>
<td>0.79</td>
<td>0.79</td>
<td>0.81</td>
<td>1.00</td>
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<td>6-Colombia Amarelo</td>
<td>0.74</td>
<td>0.79</td>
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<td>1.00</td>
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<tr>
<td>7-IAPAR 77028</td>
<td>0.73</td>
<td>0.80</td>
<td>0.82</td>
<td>0.79</td>
<td>0.86</td>
<td>0.85</td>
<td>1.00</td>
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<td>8-IAPAR-59</td>
<td>0.76</td>
<td>0.81</td>
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<td>0.82</td>
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<tr>
<td>9-Tupi LC 1669-33</td>
<td>0.78</td>
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<td>0.80</td>
<td>0.86</td>
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<td>10-IAPAR 75163-21-10</td>
<td>0.76</td>
<td>0.80</td>
<td>0.82</td>
<td>0.80</td>
<td>0.85</td>
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<td>0.93</td>
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<tr>
<td>11-IAPAR 75163-12</td>
<td>0.71</td>
<td>0.80</td>
<td>0.81</td>
<td>0.79</td>
<td>0.80</td>
<td>0.83</td>
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<td>0.88</td>
<td>0.86</td>
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<tr>
<td>12-Kattimor</td>
<td>0.75</td>
<td>0.80</td>
<td>0.83</td>
<td>0.85</td>
<td>0.84</td>
<td>0.85</td>
<td>0.86</td>
<td>0.89</td>
<td>0.89</td>
<td>0.87</td>
<td>1.00</td>
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<tr>
<td>13-F1</td>
<td>0.74</td>
<td>0.81</td>
<td>0.81</td>
<td>0.80</td>
<td>0.82</td>
<td>0.85</td>
<td>0.86</td>
<td>0.89</td>
<td>0.89</td>
<td>0.89</td>
<td>0.90</td>
<td>0.90</td>
<td>1.00</td>
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<tr>
<td>14-F2</td>
<td>0.70</td>
<td>0.74</td>
<td>0.77</td>
<td>0.79</td>
<td>0.79</td>
<td>0.82</td>
<td>0.81</td>
<td>0.87</td>
<td>0.85</td>
<td>0.87</td>
<td>0.82</td>
<td>0.83</td>
<td>0.82</td>
<td>1.00</td>
</tr>
</tbody>
</table>

* IAPAR-59 x Mundo Novo IAC 376-4.
Orozco-Castillo et al. markers to discriminate *Coffea* between the Sarchimor-derived genotypes and other acces-
sions of the Sarchimor (IAC LC 1669) origin of these cultivars since all of them were derived from 75163-12. These values are in agreement with the common LC 1669-33, to 0.85, between IAPAR 77028 and IAPAR IAPAR 75163-21-10 and the cultivars IAPAR 59 and Tupi ues of genetic similarities ranging from 0.93, between the IAPAR cultivars, which clustered together with the val-
germplasm.

Genetic variability

The molecular profile obtained by combining restric-
tion DNA digestion and RAPD was found efficient enough to reveal usable levels of DNA polymorphism among 14 elite *C. arabica* cultivars. The dendrogram (Figure 1) generated from the matrix of Dice’s genetic similarity revealed two genetic groups comprising the Bourbon-derived geno-
types and the IAPAR cultivars derived from the Sarchimor germplasm.

A relatively close relationship was revealed among the IAPAR cultivars, which clustered together with the values of genetic similarities ranging from 0.93, between IAPAR 75163-21-10 and the cultivars IAPAR 59 and Tupi LC 1669-33, to 0.85, between IAPAR 77028 and IAPAR 75163-12. These values are in agreement with the common origin of these cultivars since all of them were derived from selected progeny of the Sarchimor (IAC LC 1669) germplasm. Lower values of genetic similarity were found between the Sarchimor-derived genotypes and other acces-
sions (Tables 1 and 2, Figures 1 and 2). The genetic similari-
ty of the IAPAR cultivars ranged from 0.71 to 0.78 with the cultivar Mundo Novo, 0.80 to 0.81 with Catuai Vermelho, 0.79 to 0.82 with the Caturra cultivars. As expected, there was a higher genetic similarity among cultivars of the same gene pool, such as the Sarchimor-
derived genotypes, even though they still display enough usable genetic variation.

A relatively high variation was detected among the Bourbon-type coffee (Tables 1 and 2). The values of gen-
etic similarities ranged from 0.79 between Villasarchi and both Catuai Vermelho IAC 81 and Caturra Vermelho to 0.87 between the Caturra genotypes and between Catuai Vermelho IAC 81 and Caturra Vermelho. The genetic simi-
larity (0.87) between Caturra Vermelho and Caturra Amarelo points to the existence of genetic variation that is not reflected in plant morphologies. The Caturra coffee dis-
plays the compact architecture (CtCt) that was originated by spontaneous mutation in plants of Bourbon Vermelho (ctct) (Carvalho et al., 1991). While the Caturra germplasms still demonstrate a heritability of 0.03 for pro-
ductivity (Sera, 1980), Caturra Vermelho and Caturra Amarelo are very similar and can only be separated by the fruit color (red or yellow).

Mundo Novo IAC 376-4 was the most divergent cultivar, showing a genetic similarity ranging from 0.75 to 0.81 with cultivars belonging to the Bourbon gene pool and from to 0.71 to 0.78 with the cultivars of the Sarchimor germplasm. The closer relationship between the cultivars of the Bourbon-type coffee and Mundo Novo IAC 376-4 (Bourbon x Sumatra) is well supported by the origin of this cultivar. Moreover, these associations open the possibility for promising crosses between the Mundo Novo IAC 376-4 with selections of the Sarchimor germplasm and the Bour-
bon-derived genotypes that could result in hybrids with better performance and productivity. Fontes et al. (2000) pointed out that high genetic divergence between *C. arabica* parents increased the possibility of combining dif-
ferent alleles, resulting in higher heterozigosity. Hybrid plants derived F1 and F2 generations of the Mundo Novo IAC 376-4 x IAPAR 59 cross were also included in this study. The F1 showed similarity coefficients of 0.74 and 0.89 with Mundo Novo IAC 376 and IAPAR 59, respec-
tively. Similar genetic similarities values were observed between the F2 and Mundo Novo IAC 376 (0.70) and IAPAR 59 (0.87). The results of molecular data are in agreement with morphological characters. The F1 hybrid displays a compact architecture and is resistant to all known leaf rust physiological races which are characters of the IAPAR 59 cultivar while Mundo Novo IAC-376-4 exhibits normal architecture and presents high susceptibility to leaf rust. The F2 exhibits compact architecture for 75% of the plants and high productivity. The high performance of the F1 and F2 hybrids was also demonstrated with the genetic analysis that showed a heterosis of 25% for the F1 and about a half for the F2 plants (Sera, personal communication).

The considerable polymorphism detected in this study also illustrated that it is possible to find genetic diver-

gence among arabica accessions of the same origin. Impor-
tant characteristics have been incorporated into arabica germplasms. For instance, the leaf rust factors SH6, SH7, SH5, SH2, SH5, and SH (?) were selected in the F3 and F4 generations from crosses between *C. arabica* cv Villasarchi x Híbrido de Timor CIFC 832-2 (Bertrand et al., 1999). The cultivar Villasarchi belongs to the Caturra germplasm and, as a Caturra variety, it is a mutant for small stature (CtCt). However, the RAPD analysis revealed a considerable low genetic similarity between Villasarchi and the Caturra-
derived cultivars. On the other hand, the results are in con-
formity with the higher genetic similarities observed between Villasarchi and most selections of the Sarchimor...
Genetic polymorphism among 14 elite *Coffea arabica* L. cultivars

(Villasarchi x Timor Hybrid) germplasm (Table 1; Figures 1 and 2).

All of the arabica genotypes used in the present study were distinguished from each other. The examination of the dendrogram showed that the molecular data are consistent with pedigree information. For instance, the cultivar Colombia Amarelo is a multilina that resulted from a mixture of several progenies of the Hibrido de Timor CIFIC 1343 x Caturra CCC 135 which were selected for leaf rust resistance and other important agronomic traits, such as productivity, plant architecture, quality, and precocity for fruit maturation (Bertrand et al., 1999). The estimative of genetic similarity revealed a close relationship between Colombia Amarelo and the Sarchimor-derived germplasm (IAPAR cultivars). On the other hand, the values of genetic polymorphism was also revealed, suggesting that even the elite cultivars, which have been exposed to intensive selection, still show potential for genetic studies and for use in breeding programs.

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