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Spatially structured genetic diversity of the Amerindian yam (Dioscorea trifida L.) assessed by SSR and ISSR markers in Southern Brazil

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| | species, whose origin and dom maintained by traditional farr São Paulo, Santa Catarina, M Repeats (SSR) and 16 Inter S the accessions was high, 95 % power among accessions com SSR and ISSR markers led to main groups: I—Ubatuba-SP Amazonas State were classifi principal coordinate analyzes three main groups. Higher var and 60.6 % for ISSR and SSR SSR. Significant but low corr p = 0.0007 for SSR and $r = 0$. spatially structured genetic di | nestication are still unresolved issues. In order to estimate the genetic diversity ners in Brazil, 53 accessions of <i>D. trifida</i> from 11 municipalities in the states of ato Grosso and Amazonas were characterized based on eight Simple Sequence imple Sequence Repeats (ISSR) markers. The level of polymorphism among 6 for SSR and 75.8 % for ISSR. The SSR marker showed higher discrimination pared to ISSR, with <i>D</i> parameter values of 0.79 and 0.44, respectively. Although dendrograms with different topologies, both separated the accessions into three ,II—Iguape-SP and Santa Catarina; and III—Mato Grosso. The accessions from ed in group II with SSR and in a separate group with ISSR. Bayesian and conducted with both molecular markers corroborated the classification into riation was found within groups in the AMOVA analysis for both markers (66.5 ,, respectively), and higher Shannon diversity index was found for group II with relations were found between genetic and geographic distances ($r = 0.08$; 16; $p = 0.0002$ for ISSR). Therefore, results from both markers showed a slight versity in <i>D. trifida</i> accessions maintained by small traditional farmers in Brazil. |
| Keywords (separated by '-') | Dioscorea trifida - Genetic di | versity - Genetic structure - Molecular markers - Traditional agriculture - Yams |
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Spatially structured genetic diversity of the endangered Amerindian yam (*Dioscorea trifida* L.) assessed by SSR and ISSR markers in Southern Brazil

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6 F. Rodrigues · Samantha Koehler · Paul Gepts ·
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10 Abstract Dioscorea trifida L. (Dioscoreaceae) is 11 among the economically most important cultivated Amerindian yam species, whose origin and domesti-12 cation are still unresolved issues. In order to estimate 13 14 the genetic diversity maintained by traditional farmers 15 in Brazil, 53 accessions of D. trifida from 11 munic-16 ipalities in the states of São Paulo, Santa Catarina, 17 Mato Grosso and Amazonas were characterized based 18 on eight Simple Sequence Repeats (SSR) and 16 Inter 19 Simple Sequence Repeats (ISSR) markers. The level of 20 polymorphism among the accessions was high, 95 % for SSR and 75.8 % for ISSR. The SSR marker showed 21 22 higher discrimination power among accessions com-23 pared to ISSR, with D parameter values of 0.79 and 24 0.44, respectively. Although SSR and ISSR markers 25 led to dendrograms with different topologies, both 26 separated the accessions into three main groups:

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I-Ubatuba-SP; II-Iguape-SP and Santa Catarina; 27 and III—Mato Grosso. The accessions from Amazonas 28 State were classified in group II with SSR and in a 29 separate group with ISSR. Bayesian and principal 30 coordinate analyzes conducted with both molecular 31 markers corroborated the classification into three main 32 groups. Higher variation was found within groups in 33 the AMOVA analysis for both markers (66.5 and 34 60.6 % for ISSR and SSR, respectively), and higher 35 Shannon diversity index was found for group II with 36 SSR. Significant but low correlations were found 37 between genetic and geographic distances (r = 0.08; 38 p = 0.0007 for SSR and r = 0.16; p = 0.0002 for 39 ISSR). Therefore, results from both markers showed a 40 slight spatially structured genetic diversity in D. trifida 41 accessions maintained by small traditional farmers in 42 Brazil. 43

| Keywords | Dioscorea trifida · Genetic diversity · | 44 |
|---------------|---|----|
| Genetic stru | cture · Molecular markers · Traditional | 45 |
| agriculture · | Yams | 46 |

Introduction

47 49

The genus Dioscorea, family Dioscoreaceae, represents an important food source in the humid and50subhumid tropics (Ayensu and Coursey 1972). This52genus consists of more than 600 species, of which only5310 are used for human consumption (Lebot 2009).54Dioscorea trifida L., originating in South America, is55

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among the economically most important species,
including *Dioscorea cayenensis* Lam. and *Dioscorea rotundata* Poir., originating in Africa, and *Dioscorea alata* L., originating in Asia (Coursey 1976).

60 Dioscorea trifida is an herbaceous, autotetraploid 61 (x = 20 and 2n = 4x = 80), viny, and perennial plant 62 (Bousalem et al. 2006), with quadrangular winged 63 stems without spines but with deeply lobed leaves, 64 usually arranged alternately or rarely opposite (Mont-65 aldo 1991). The plants are dioecious, with small 66 unisexual flowers that, when fertilized, produce ined-67 ible encapsulated fruits (Stephens 2009). Its reproduction occurs by allogamy or vegetative propagation 68 69 (Montaldo 1991). The tuber, the edible plant structure, 70 has a high nutritional quality and astringent, antimi-71 crobial and diuretic properties, which allow its use for 72 combating malnutrition and treatment of diseases such 73 as diabetes and high cholesterol levels (Ramos-74 Escudero et al. 2010). The main limiting factor for 75 growing D. trifida is potyviruses (genus Potyvirus, 76 family Potiviridae), which causes a variety of symp-77 toms on the leaves of infected plants (Odu et al. 2004). 78 Potyvirus infection can cause significant economic 79 damage and process of genetic erosion of the crop (Bousalem et al. 2010). 80

81 The evolutionary history of D. trifida is controver-82 sial. Although it occurs very frequently in various 83 countries of Latin America, and the Amazon has been 84 reported as a possible center of origin and diversifi-85 cation of this species (Degras 1993), the lack of information about its origin and domestication process 86 87 is still evident. It is believed that D. trifida originated 88 on the border between Brazil, Guyana, French Guyana 89 and Suriname, and was domesticated by indigenous 90 peoples in these regions (Pedralli 1998). Recent 91 studies conducted in French Guiana revealed the 92 presence of wild relatives of D. trifida, being the first 93 direct genetic evidence of possible places of origin for 94 this species (Bousalem et al. 2010).

95 In the Amazon, Clement (1999) observed the 96 existence of several areas with large concentrations 97 of genetic resources related to different crop species. 98 D. trifida was present in some of these sites, such as in 99 the Northwestern Amazonian Center, the Central Amazonian Center, the Middle Orinoco Minor Centre 100 and the Guiana Minor Centre, indicating the close 101 102 relationship of these areas with the evolutionary 103 history of D. trifida. In archaeological excavations in 104 Panama, D. trifida tubers were found together with

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cassava (Piperno et al. 2000, Dickau et al. 2007). As 105 cassava was domesticated in southwestern Amazon 106 Basin (Olsen 2004) and quickly spread throughout 107 Tropical America (Piperno et al. 2000, Dickau et al. 108 2007), D. trifida could have been domesticated and 109 propagated by the same tribes involved in the process 110 of cassava domestication (Bousalem et al. 2010), 111 possibly being the first yam species cultivated by 112 indigenous peoples in the Amazon (Degras 1993). The 113 cultivation of D. trifida in Brazil has taken place since 114 then, mainly by small rural farmers (Pedralli 1998). In 115 recent surveys, the occurrence of this species was 116 observed in the Central West, South and Southeast 117 Brazil (Bressan et al. 2005; Veasey et al. 2010). It is 118 also an important crop in the Amazon (Velez 1998). 119

Despite the geo-cultural and socioeconomic impor-120 tance of D. trifida, few studies are conducted to 121 explore its potential and to develop conservation 122 strategies for this crop. In Brazil, there are few 123 institutions currently involved in research related to 124 the yam crop; therefore, new studies are important to 125 add information for breeding programs and conserva-126 tion strategies. As the cultivation and consumption of 127 yam are very intense in family agricultural production 128 systems practiced by traditional communities, these 129 systems provide a favorable environment for the 130 generation and maintenance of genetic diversity of this 131 crop (Veasey et al. 2010). However, the socioeco-132 nomic pressures faced by farmers in recent years have 133 caused the loss of plant genetic resources, specifically 134 D. trifida, and biodiversity losses can be severe and 135 irreversible. In this context, there is a need to estimate 136 the genetic diversity of D. trifida maintained by 137 traditional farmers to assist in developing strategies to 138 preserve the species and lessen losses caused by 139 various socioeconomic pressures on the yam crop. 140

Various molecular biological techniques are avail-141 able to detect genetic variability of natural populations 142 and cultivated plants. Among these techniques, micro-143 satellites or Simple Sequence Repeats (SSR) are very 144 effective because they are codominant, multi-allelic, 145 highly polymorphic and show good reproducibility 146 (Oliveira et al. 2006). In order to study the genetic 147 diversity of D. trifida, Hochu et al. (2006) developed 148 eight SSR primers specific for this species, which were 149 used for the analysis of 24 cultivars, showing high 150 polymorphism. These primers were used by Bousalem 151 et al. (2006) to assess the inheritance pattern of 152 D. trifida, from the analysis of parental genotypes and 153

Another marker used in genetic diversity studies is 156 Inter Simple Sequence Repeats (ISSR), which were 157 developed to explore microsatellite repeats without 158 159 the need to use of DNA sequencing (Zietkiewicz et al. 160 1994; Reddy et al. 2002). ISSR markers are very 161 stable, dominant, multi-allelic, present good reproducibility and generate a large number of polymorphic 162 fragments (Mattioni et al. 2002; Wolfe 2005). Few 163 164 genetic diversity studies have been performed in the 165 Dioscorea genus based on ISSR molecular markers. Among them, Zhou et al. (2008) analyzed the level of 166 167 genetic diversity among different cultivars of Diosco-168 rea opposita Thunb., widely used in traditional Chinese medicine and Wu et al. (2009) evaluated the 169 170 relationship and genetic variability among accessions 171 of D. alata. Both studies found that ISSR provided a 172 good assessment of genetic diversity of yam and 173 valuable information to help in selecting parents for 174 future yam breeding programs.

175 The aim of this study was to characterize 53 176 accessions of D. trifida originating in traditional communities in the States of Santa Catarina, São 177 178 Paulo, Mato Grosso and Amazonas, using ISSR and 179 SSR markers, in order to verify the level of genetic 180 diversity maintained by farmers in these regions in Brazil. The study describes the spatially structured 181 182 genetic variation of D. trifida maintained by these farmers and the genetic diversity that is concentrated 183 184 within the different sampling sites.

185 Materials and methods

186 Plant materials

187 We evaluated 53 accessions of D. trifida collected from 11 municipalities in the States of Sao Paulo (SP), 188 189 Santa Catarina (SC), Mato Grosso (MT) and Amazo-190 nas (AM), located between latitudes 14°43'S and 191 26°15'S and longitudes 44°01'W and 62°05'W (Fig. 1; 192 Table 1). In each visited municipality, a collection 193 was conducted so as to seek greater representation of 194 the genetic variability, taking into account morpho-195 logical variation and information from farmers. Three 196 accessions (the two accessions from Amazonas and 197 one from Ubatuba, SP) were acquired in local markets. Accessions were collected in the form of tubers, which 198

were grown in pots placed in a greenhouse at the Luiz199de Queiroz College of Agriculture, University of Sao200Paulo, in Piracicaba, SP, located at 22°43'S latitude201and 47°25'W longitude, where young leaves were202collected for DNA extraction.203

DNA extraction and quantification

Young, newly expanded leaves were collected and 205 stored at 4 °C for 7 days in a CTAB gel, containing 206 30 mg CTAB, 350 mg NaCl and 70 ml of distilled 207 water (Rogstad 1992). After this period, the gel excess 208 was removed from plant tissues with the aid of a paper 209 towel. The fragments were then macerated in 1 mL 210 STE buffer [0.13 mg saccharose, 45 mL of Tris-HCL 211 (1 M), 150 mL of EDTA (0.5 M), completing with 212 distilled water to a final volume of 1.5 mL] and 213 subjected to DNA extraction by the method of Doyle 214 and Doyle (1990). DNA concentration was estimated 215 in a 1 % agarose gel, using a TBE 10X running buffer, 216 stained in ethidium bromide. A final concentration of 217 5 ng/µL was obtained for the PCR analysis. 218

Amplification of SSR and ISSR

For the SSR amplification, 10 primer pairs developed 220 by Hochu et al. (2006) and Tostain et al. (2006) were 221 tested (Table 2). PCR was conducted in a 16 µL 222 reaction volume containing: 5 ng of genomic DNA in 223 a 5 \times reaction buffer, 1.5 mM MgCl₂, 2.5 mM dNTPs, 224 5 pmol of forward primer, 5 pmol of reverse primer, 225 and 5 U/µL Taq DNA polymerase (Promega, Madi-226 son, USA). The amplification reactions were per-227 formed in a MyCycler Thermal Cycler model BioRad 228 thermocycler using the following steps: 1) denatur-229 ation at 94 °C for 5 min, followed by 30 cycles [30 s 230 at 94 °C, 30 s at annealing temperature (touchdown of 231 50-60 °C) and 30 s for 72 °C], and a final stage of 232 extension of 5 min at 72 °C for the PCR reactions with 233 specific primers for *D. trifida* (Hochu et al. 2006); 2) 234 denaturation at 94 °C for 5 min, followed by 35 cycles 235 [30 s at 94 °C, 1 min at a temperature of annealing 236 (touchdown 50-60 °C) and 1 min at 72 °C], and a 237 final extension of 8 min at 72 °C for PCR reactions 238 with heterologous primers (Tostain et al. 2006). 239

Electrophoresis was performed on denaturing 7 %240polyacrylamide gel, with a constant power of 70 W for241the time necessary for separating the amplified frag-242ments in each primer, using 10 and 100 bp DNA243

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Ladder (InvitrogenTM, São Paulo, Brazil) as markers.
The gels were stained using silver nitrate methodology
(Creste et al. 2001) for the revelation of microsatellite
bands, which were photographed with a digital camera
and evaluated in a transiluminator.

249 For the ISSR analysis 20 primers were tested according to the Wolfe (2000, 2005) protocol 250 (Table 3). PCR reactions were performed in a final 251 252 volume of 30 µL containing: 5 ng of genomic DNA in a 5× reaction buffer, 2.0 mM MgCl₂, 2.33 mM 253 254 dNTPs, 10 pmol primer and 5 U/µL Taq DNA 255 polymerase (Promega, Madison, USA). The amplifi-256 cation of the DNA template were performed in 257 MultiGene Thermal Cycler thermocycler (Labnet 258 International, Inc.) according to the following amplification conditions: 90 s at 94 °C, 35 cycles at 94 °C 259 for 40 s, followed by 46 cycles (52 °C for 45 s, 72 °C 260 for 90 s, 94 °C for 45 s, 44 °C for 45 s), and a final 261 262 stage of extension at 72 °C for 5 min (Wolfe 2000).

The products resulting from the amplification
reactions were subjected to electrophoresis on a 2 %
agarose gel in TBE buffer 10X for 140 min at 90 V

and stained with ethidium bromide. A 100 bp DNA 266 Ladder (InvitrogenTM, Carlsbad, USA) was used as a 267 marker. Additionally, we used control samples previ-268 ously amplified with success. The gel was photo-269 graphed over ultraviolet light source with Syngene 270 photodocumentation system (Synoptics Ltda., Cam-271 bridge, United Kingdom). For statistical analysis we 272 considered only robust and unambiguous bands. We 273 discarded the bands that showed low intensity or 274 coalescing with other bands. 275

Statistical analysis

Due to the tetraploid behavior of D. trifida, as 277 described by Bousalem et al. (2006), the band patterns 278 of the SSR and ISSR markers were both interpreted as 279 binary data, presence (1) and absence (0) of bands, 280 generating data matrices that were subjected to the 281 following statistical programs. Genetic diversity anal-282 yses were based on POPGENE Software, version 1.3 283 (Yeh et al. 1997), where we obtained the number of 284 bands observed per primer, number of polymorphic 285

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| Accession | ID | Geographic localities | Popular name | Lat/Long |
|-----------|-----|--|----------------------|-----------------|
| 01 | 180 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°15′S/44°01′O |
| 02 | 181 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará branco | 23°17′S/44°05′O |
| 03 | 182 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°18′S/44°52′O |
| 04 | 183 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°17′S/44°51′O |
| 05 | 184 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°18′S/44°51′O |
| 06 | 185 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°18′S/44°51′O |
| 07 | 187 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará branco | 23°18′S/44°51′O |
| 08 | 191 | Sertão das Cutias, Ubatuba, São Paulo | Cará roxo | 23°22′S/44°58′O |
| 09 | 193 | Rio Escuro, Ubatuba, São Paulo | Cará branco | 23°28′S/45°08′O |
| 10 | 195 | Sertão do Ingá, Ubatuba, São Paulo | Cará cobrinha | 23°31′S/45°13′O |
| 11 | 196 | Sertão do Ingá, Ubatuba, São Paulo | Cará branco | 23°31′S/45°13′O |
| 12 | 197 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°17′S/44°51′O |
| 13 | 198 | Sertão do Ingá, Ubatuba, São Paulo | Cará roxo | 23°31′S/45°13′O |
| 14 | 201 | Sertão do Ingá, Ubatuba, São Paulo | Cará roxo | 23°31′S/45°14′O |
| 15 | 203 | Sertão do Ingá, Ubatuba, São Paulo | Cará roxo | 23°31′S/45°14′O |
| 16 | 204 | Rio Escuro, Ubatuba, São Paulo | Cará roxo | 23°28′S/45°08′O |
| 17 | 208 | Araribá, Ubatuba, São Paulo | Cará roxo | 23°32′8/45°15′O |
| 18 | 210 | Sertão de Ubatumirim, Ubatuba, São Paulo | Cará roxo | 23°29′S/45°10′O |
| 19 | 216 | Fazenda da Caixa, Ubatuba, São Paulo | Cará roxo | 23°31′S/45°14′O |
| 20 | 217 | Feira de Ubatuba, Ubatuba, São Paulo | Cará roxo | 23°27′S/45°09′O |
| 21 | 236 | Feira de Manaus, Manaus, Amazonas | Cara roxo | 03°08′S/60°01′O |
| 22 | 237 | Feira de Barcelos, Barcelos, Amazonas | Cará | 0°58′S/62°55′O |
| 23 | 281 | Pirabeiraba, Joinville, Santa Catarina | Cará | 26°10′S/48°55′O |
| 24 | 282 | Pirabeiraba, Joinville, Santa Catarina | Cará mimoso | 26°09′S/48°56′O |
| 25 | 283 | Pirabeiraba, Joinville, Santa Catarina | Cará | 26°09′S/48°58′O |
| 26 | 285 | Acaraí, São Francisco do Sul, Santa Catarina | Cará pão | 26°11′S/48°53′O |
| 27 | 286 | Pirabeiraba, Joinville, Santa Catarina | Cará mimoso | 26°15′S/48°37′O |
| 28 | 287 | Pirabeiraba, Joinville, Santa Catarina | Carcanhá de nego | 26°09′S/48°59′O |
| 29 | 290 | Pirabeiraba, Joinville, Santa Catarina | Cará mimoso | 26°09′S/48°59′O |
| 30 | 292 | Pirabeiraba, Joinville, Santa Catarina | Cará | 26°09′S/48°59′O |
| 31 | 297 | Pirabeiraba, Joinville, Santa Catarina | Cará | 26°09′S/48°59′O |
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| 35 | 312 | Icapara, Iguape, São Paulo | Cará São João branco | 24°40′S/47°27′O |
| 36 | 313 | Cavalcanti, Iguape, São Paulo | Cará-pipa | 24°43′S/47°45′O |
| 37 | 323 | Icapara, Iguape, São Paulo | Cará São João roxo | 24°40′S/47°27′O |
| 38 | 328 | Momuna, Iguape, São Paulo | Cará São João roxo | 24°42′S/47°40′O |
| 39 | 329 | Momuna, Iguape, São Paulo | Cará São João branco | 24°42′S/48°40′O |
| 40 | 335 | Carumbé, Acorizal, Mato Grosso | Cará roxo | 15°08′S/56°12′O |
| 41 | 336 | Carumbé, Acorizal, Mato Grosso | Cará roxo | 15°08′S/56°12′O |
| 42 | 340 | Rio dos Couros, Cuiabá, Mato Grosso | Cará pé de anta | 15°36′S/55°48′O |
| 43 | 343 | Carumbé, Acorizal, Mato Grosso | Cará branco | 15°08′S/56°12′O |

 Table 1 Dioscorea trifida accessions used in this study collected in Brazil, including accession and identification number (ID) in the Germplasm Bank, origin (community, municipality, state), popular name and geographic coordinates



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Table 1 continued

| Accession | ID | Geographic localities | Popular name | Lat/Long |
|-----------|-----|---|--------------------|-----------------|
| 44 | 344 | Sela Dourada, Nobres, Mato Grosso | Cará do Joaquim | 15°36′S/56°48′O |
| 45 | 345 | Santo Antônio do Barreiro, Jangada, Mato Grosso | Cará roxo | 15°08′S/56°17′O |
| 46 | 350 | Sela Dourada, Nobres, Mato Grosso | Cará branco | 15°34′S/56°46′O |
| 47 | 351 | Sela Dourada, Nobres, Mato Grosso | Cará mão de anta | 15°30′S/56°42′O |
| 48 | 352 | Timbozal, Rosário Oeste, Mato Grosso | Cará mão de anta | 14°51′S/56°23′O |
| 49 | 355 | Chapada Vacaria, Acorizal, Mato Grosso | Cará roxo | 15°03′S/56°08′O |
| 50 | 361 | Sela Dourada, Nobres, Mato Grosso | Cará roxo | 14°43′S/56°15′O |
| 51 | 364 | Barranco Alto, Rosário Oeste, Mato Grosso | Pombinho branco | 15°14′S/57°59′O |
| 52 | 366 | Sela Dourada, Nobres, Mato Grosso | Cará roxo cumprido | 14°43′S/56°15′O |
| 53 | 368 | Barranco Alto, Rosário Oeste, Mato Grosso | Cará roxo | 15°17′S/57°50′O |

Table 2 List of SSR primers used to evaluate 53 *Dioscorea trifida* accessions, including primer sequence, annealing temperature (T_A) ; size range of SSR bands in base pairs (bp),

number of bands (N_B) , number of polymorphic bands (N_{PB}) , percent polymorphism (P) and discriminating power (D)

| Primer code | Sequence $(5'-3')$ | T_A (°C) | Size range (bp) | N_B | N_{PB} | P (%) | D |
|----------------------|----------------------------|------------|-----------------|-------|----------|-------|------|
| Da1A01 ¹ | F: TAT AAT CGG CCA GAG G | 51–53 | 202–205 | 2 | 2 | 100.0 | 0.97 |
| | R: TGT TGG AAG CAT AGA GAA | | | | | | |
| Dab2C05 ¹ | F: CCC ATG CTT GTA GTT GT | 51-52 | 168–192 | 5 | 5 | 100.0 | 0.91 |
| | R: TGC TCA CCT CTT TAC TTG | | | | | | |
| MTI2 ² | F: TCATCAAGAGCATCAAAAAAC | 50-52 | 121-131 | 6 | 6 | 100.0 | 0.71 |
| | R: GCCTCGTCTTTGAAGTTGGT | | | | | | |
| MTI3 ² | F: TAACAAACAAAAAATGAAAC | 55–59 | 156-205 | 13 | 13 | 100.0 | 0.85 |
| | R: TAACAGTGATTGAGCTAGGA | | | | | | |
| MTI4 ² | F: ACTTGGTGTTGTTGGATTGC | 50–58 | 101-111 | 8 | 8 | 100.0 | 0.61 |
| | R: TATCACTCCCCAGACCAGA | | | | | | |
| MTI10 ² | F: TCGTGTCCATCTTGCTGCGT | 55–58 | 143–198 | 11 | 11 | 100.0 | 0.61 |
| | R: GAAAAGCGGAGATGAAGAGCA | | | | | | |
| MTI11 ² | F: CTCTTTTGCTTCTCATTTCA | 55-56 | 124–137 | 5 | 4 | 80.0 | 0.72 |
| | R: ATGTAGCCAATCCAAAATAG | | | | | | |
| MTI12 ² | F: CTGCCAGCGTTCCGATTC | 55-60 | 100-123 | 6 | 5 | 83.0 | 0.92 |
| | R: CGTAGGACCTCTCGCATCAG | | | | | | |
| Average | | _ | _ | 7.0 | 6.75 | 95.0 | 0.79 |

¹ Tostain et al. (2006); ² Hochu et al. (2006)

286 bands, percent polymorphism and estimated the 287 Shannon index according to the following formula: 288 $H' = -\sum_{i=1}^{s} pi \log pi$, where pi is the frequency of 289 each species, for *i* ranging from 1 to *S* (richness).

In order to compare the efficiency of the markers in
the genotypic identification, the discrimination power
(D) (Tessier et al. 1999) was estimated for each primer.

This parameter was calculated according to the 293 formula: $D_j = 1 - C_j = 1 - \sum_{i=1}^{I} pi \frac{(Npi-1)}{N-1}$, where 294 *D* is the probability that two randomly selected 295 individuals have a different and distinct banding 296 pattern from each other; *C* is the probability that two 297 randomly selected individuals have a similar band 298 pattern, and *N* is the number of individuals analyzed. 299

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Table 3 ISSR primers used to evaluate 53 *Dioscorea trifida* accessions, including primer sequence, annealing temperature (T_A) , size range of ISSR bands in base pairs (bp), number of bands (N_B) , number of polymorphic bands (N_{PB}) , percent polymorphism (P), and discriminating power (D)

| Primer code | Sequence $(5'-3')$ | T_A (°C) | Size range(bp) | N_B | N_{PB} | P (%) | D |
|-------------|--------------------|------------|----------------|-------|----------|-------|-------|
| UBC 7 | (CT)8-RG | 48 | 300-1,300 | 13 | 11 | 84.6 | 0.37 |
| UBC 814 | (CT)8-TG | 50 | 500-1,100 | 8 | 6 | 75.0 | 0.76 |
| UBC 843 | (CT)8-RA | 48 | 600-1,200 | 6 | 6 | 100.0 | 0.82 |
| UBC 844 | (CT)8-RC | 50 | 300-1,300 | 9 | 2 | 22.2 | -0.35 |
| UBC 898 | (CA)6-RY | 48 | 300-1,300 | 12 | 6 | 50.0 | -0.48 |
| UBC 899 | (CA)6-RG | 54 | 300-1,300 | 9 | 7 | 77.8 | 0.63 |
| JOHN | (AG)7-YC | 54 | 100-1,200 | 13 | 9 | 69.2 | 0.81 |
| UBC 901 | (GT)6-YR | 50 | 300-1,300 | 8 | 8 | 100.0 | 0.67 |
| UBC 902 | (GT)6-AY | 50 | 500-900 | 4 | 4 | 100.0 | 0.77 |
| AW3 | (GT)6-RG | 54 | 500-800 | 4 | 3 | 75.0 | 0.75 |
| OMAR | (GAG)4-RC | 50 | 300–900 | 8 | 8 | 100.0 | 0.54 |
| DAT | (GA)7-RG | 54 | 300-800 | 9 | 2 | 22.2 | -0.16 |
| TERRY | (GTG)4-RC | 50 | 300-800 | 9 | 5 | 55.6 | 0.49 |
| MAO | (CTC)4-RC | 50 | 400–1,300 | 8 | 8 | 100.0 | 0.36 |
| MANNY | (CAC)4-RC | 48 | 300-1,000 | 11 | 9 | 81.8 | 0.48 |
| GOOFY | (GT)7-YG | 54 | 300–900 | 6 | 6 | 100.0 | 0.56 |
| Average | - | - | - 7 | 8.56 | 6.25 | 75.8 | 0.44 |

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DARwin software, version 5.0 (Perrier and Jac-300 301 quemoud-Collet 2006), was used to perform a cluster 302 analysis, based on Jaccard similarity coefficient and 303 the UPGMA method. The stability of the groupings 304 was assessed based on estimates of genetic dissimilarity through the procedure of resampling with 1,000 305 306 bootstraps. Values higher than 70 % in the nodes that 307 join the groups indicate homogeneity among acces-308 sions. Software NTSYS-pc (Rohlf 1992) was used to conduct a principal coordinate analysis (PCoA) and 309 310 obtain scatter plots.

311 To confirm the reliability of the groups obtained in the cluster analysis and PCoA, we conducted a 312 313 Bayesian analysis using the software Structure (Pritchard et al. 2000; Pritchard and Donnelly 2001; Falush 314 et al. 2007), which does not rely on prior information 315 316 on possible groups, for example based on the origin of 317 the accessions. The Structure software was run using the admixture model, correlated allele frequencies and 318 319 repeated ten times for each K (number of assumed 320 clusters) with a burn-in of 500,000 interactions 321 followed by 500,000 interactions MCMC (Markov 322 Chain Monte Carlo). The most likely number of 323 clusters was chosen using the ΔK method (Evanno 324 et al. 2005).

In order to identify the proportion of genetic
variation between and within groups obtained using
the software Structure, which coincided with the

groups of PCoA and cluster analysis, a molecular 328 variance analysis (AMOVA) was carried out with 329 Arlequin software (Schneider et al. 2000). Another 330 parameter analyzed was the correlation between 331 matrices of genetic and geographic distances, as well 332 as between genetic distance matrices for SSR and 333 ISSR markers, through the Pearson correlation (r), 334 whose significance was evaluated by Mantel (1967) 335 test, using NTSYS-pc software (Rohlf 1992). 336

Results

Eight SSR and 16 ISSR primers were selected based 338 on the presence of well defined and with good 339 resolution bands (Tables 2, 3). We obtained 56 bands 340 or amplification products with sizes ranging from 341 101 pb to 205 pb for SSR and 137 bands ranging from 342 100 bp to 1,300 pb for ISSR, in a total of 193 bands, 343 with an average of 7.0 bands/primer for SSR and 8.56 344 bands/primer for ISSR. The number of polymorphic 345 bands for SSR and ISSR was 54 and 100, with an 346 average of 6.75 and 6.25 polymorphic bands per 347 primer, respectively. The level of polymorphism was 348 high, 95 % for SSR and 75.8 % for ISSR. Parameter 349 D value for SSR was 0.79, while for ISSR was 0.44, 350 demonstrating that although the ISSR marker has 351 generated a greater number of bands, the SSR marker 352

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showed greater discriminatory power between theaccessions.

The Jaccard coefficient among 53 accessions of *D. trifida* ranged from 0.40 to 0.96, with a variation of 56 % similarity for SSR marker and from 0.66 to 0.97, with a variation of 31 % for ISSR. Although the two types of markers are located mostly in neutral regions and related to different sequences of the genome, the correlation between genetic matrices obtained from SSR and ISSR markers was high (r = 0.57; p =0.0002), demonstrating similar relationships between data from both marker classes.

Although ISSR and SSR markers generated dendrograms with different topologies (Figs. 2, 3), in general, both dendrograms showed the formation of the same groups, with a few exceptions. Despite the low bootstrap values, below 60 % and thus not shown in the dendrograms, it was possible to identify three welldefined groups: group I (accessions from Ubatuba-371 SP), group II (accessions from Iguape-SP and Santa 372 Catarina-SC) and group III (accessions from Mato 373 Grosso-MT). The yam varieties collected in Iguape-374 SP and Santa Catarina showed higher genetic similar-375 ity, while the varieties from Ubatuba-SP and Mato 376 Grosso were more divergent and classified into distinct 377 groups. All accessions were grouped according to their 378 collection locations for both markers, except the 379 accessions from Amazonas, which changed their 380 position in the dendrogram according to the molecular 381 marker analyzed. These two accessions were classified 382 into a separate group (group IV) in the ISSR analysis 383 while in the SSR analysis they were classified in group 384 II. Also, within group II, accessions from Santa 385 Catarina were apparently better separated from those 386 from Iguape-SP in the SSR than in the ISSR cluster 387 analysis. Although variations obtained in the PCoA, 388



dendrogram based on eight SSR primers showing the genetic relationships among 53 accessions of *D. trifida*: group I [accessions from Ubatuba-SP (*green*)]; group II [accessions from Iguape-SP (*blue*), Santa Catarina (*pink*) and Amazonas (*yellow*)]; and group III [accessions from Mato Grosso (*red*)]



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Fig. 3 UPGMA

dendrogram based on 16 ISSR primers showing the genetic relationships among 53 accessions of *D. trifida*: group I [accessions from Ubatuba-SP (green)]; group II [accessions from Iguape-SP (blue) and Santa Catarina (pink)]; group III [accessions from Mato Grosso (red)] and group IV [accessions from Amazonas (yellow)]



389 whose first two principal coordinates represented 390 35.7 % of total variation, were not significant, a scatter plot from data obtained with SSR separated the 391 genotypes in the same groups obtained in the scatter 392 393 plot of the data obtained from ISSR, whose first two 394 principal coordinates represented 31.6 % (not shown 395 data). So, both markers seem to be useful in discrim-396 inating the genetic diversity of D. trifida accessions.

The Bayesian analysis performed with Structure software for the SSR and ISSR data confirmed the groups obtained in the SSR cluster analysis and the PCoA, since the value of k was equal to three, showing that the accessions are genetically structured in three groups (Fig. 4). Based on SSR data, the two Amazonian accessions showed more than 90 % of their genetic constitution similar to accessions from Iguape-404 SP and Santa Catarina (Fig. 4a), while based on ISSR 405 data, the same accessions showed more than 60 % 406 similarity to those from Mato Grosso and more than 407 30 % to those from Ubatuba-SP (Fig. 4b). Comparing 408 both analysis, the SSR marker (Fig. 4a) showed some 409 exceptions to the groups formed in the dendrogram for 410 this marker (Fig. 2), such as accessions no. 17 and 18, 411 from Ubatuba-SP, showing more than 50 % similarity 412 to the group from Iguape-SP and Santa Catarina, while 413 accession no. 40, from Mato Grosso, showed more 414 than 80 % similarity to the group from Iguape-SP and 415 Santa Catarina. The ISSR marker (Fig. 4b), on the 416 other hand, showed a group pattern with great 417 similarity to the ISSR dendrogram in Fig. 3. 418

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Fig. 4 Structure of the genetic diversity of 53 *D. trifida* accessions based on Bayesian approach for eight SSR markers (**a**) and 16 ISSR markers (**b**). Each accession is represented by a

vertical bar. Accessions 1–20 originated from Ubatuba-SP; accessions 21–39 originated from Iguape-SP, Santa Catarina and Amazonas; and accessions 40–53 originated from Mato Grosso

419 From the analysis of variance considering the three 420 groups formed in the Bayesian analysis and two 421 comparison levels, within and between groups, dif-422 ferences were found between the genetic material studied for both the SSR data ($\Phi st = 0.39$; p =423 0.0000) and for ISSR data ($\Phi st = 0.33$; p = 0.0000) 424 425 (Table 4). Genetic variation was greater within than 426 between groups for both SSR (60.6 %) and ISSR (66.5 %) markers (Table 4). A low positive correla-427 428 tion was identified between genetic and geographical 429 distances for both SSR data (r = 0.08; p = 0.0007) 430 and ISSR data (r = 0.16; p = 0.0002), demonstrating 431 a slight spatial structure of genetic material in the geographic area sampled. 432

The Shannon diversity index obtained from SSR 433 434 data indicated that the State of São Paulo showed 435 greater genetic diversity (0.40), followed by Mato 436 Grosso (0.31) and Santa Catarina (0.27) (Table 5). 437 Similar results were obtained for ISSR data, but with lower values (0.28, 0.21 and 0.19, respectively). The 438 439 Amazonas State had the lowest diversity indexes (0.09 440 for SSR and 0.03 for ISSR), due to the low number of 441 accessions sampled. Considering the groups formed in 442 the Bayesian analysis and in the cluster analysis with 443 SSR, group II (accessions from Iguape-SP, Santa 444 Catarina and Amazonas) had the highest diversity index (0.40), followed by group III (0.31) with acces-445 sions from Mato Grosso, and group I (0.30) (accessions 446 447 from Ubatuba-SP) for SSR, while lower and similar 448 values for the three groups were obtained for ISSR data.

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Table 4 Analysis of molecular variance (AMOVA) for 53 accessions of *D. trifida* L. considering three groups, according to the Bayesian analysis on Structure software: group I (accessions from Ubatuba-SP), group II (accessions from Iguape-SP, Santa Catarina and Amazonas), and group III (accessions from Mato Grosso)

| Variation | DF | SSR | | ISSR | | |
|-----------------|----|---------|---------------------------|---------|---------------------------|--|
| source | | SQ | Total variation (%) | SQ | Total variation (%) | |
| Among groups | 3 | 150.295 | 39.37 | 221.218 | 33.53 | |
| Within group | 49 | 288.346 | 60.63 | 528.404 | 66.47 | |
| Total | 52 | 438.642 | | | | |

 $\Phi st = 0.3937$ for SSR and $\Phi st = 0.3353$ for ISSR; *DF* degrees of freedom, *SQ* sum of squares

* Value $p^{1}(1,023 \text{ permutations}) = 0.0000$

Discussion

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Genetic diversity and comparative analysis of SSR450and ISSR markers451

The large amount of molecular markers available to
estimate genetic diversity allows us to make compar-
isons in order to determine which technique is best
suited for a particular crop (Biswas et al. 2010).452
453
454
455Choosing the most appropriate technique depends on
the purpose of the research, reproduction mode of the457

Table 5 Shannon diversity index (H') of accessions of *D. trifida* classified according to State, to both municipalities in São Paulo (Ubatuba and Iguape), and to Bayesian analysis on Structure software: group I (accessions from Ubatuba-SP), group II (accessions from Iguape-SP, Santa Catarina and Amazonas), and group III (accessions from Mato Grosso)

| | H' | |
|----------------|------|------|
| | SSR | ISSR |
| States | | |
| São Paulo | 0.40 | 0.28 |
| Mato Grosso | 0.31 | 0.21 |
| Santa Catarina | 0.27 | 0.19 |
| Amazonas | 0.09 | 0.03 |
| Average | 0.27 | 0.18 |
| Groups | | |
| Group I | 0.30 | 0.21 |
| Group II | 0.40 | 0.20 |
| Group III | 0.31 | 0.22 |
| Average | 0.34 | 0.21 |

458 species and its genetic structure (Badfar-Chaleshtori 459 et al. 2012), as well as their ability to estimate heterozygosity (Vogel et al. 1996). However, difficul-460 ties in relating fragment patterns for specific loci and 461 462 genotypes in the genomes of polyploid species such as 463 D. trifida and other tuberous plants limit the use of heterozygosity estimates to assess different molecular 464 465 markers in these species (McGregor et al. 2000).

This study showed that it is possible to use both 466 ISSR and SSR techniques for characterizing and 467 468 discriminating morphologically distinct or similar 469 vam accessions. Also, both ISSR and SSR results 470 highlight the importance of traditional farmers in 471 maintaining high genetic diversity among their local 472 varieties. The eight SSR primer pairs were highly 473 polymorphic and informative among the 53 D. trifida 474 accessions analyzed in this study. The heterologous 475 primers Da1A01 and Dab2C05 developed by Tostain 476 et al. (2006) for D. alata L., Dioscorea abyssinica 477 Hochst. ex Kunth and Dioscorea praehensilis Benth, 478 showed 100 % polymorphism, and high discrimina-479 tion power among accessions, equal to 0.97 and 0.91, 480 respectively. These primers were also suitable for analyzing D. trifida accessions in the transferability 481 tests conducted by Tostain et al. (2006). The primers 482 483 specific for D. trifida, developed by Hochu et al. 484 (2006), besides providing good resolution of bands in 485 the gel electrophoresis, showed high polymorphism, with an average of 93.8 %, and a high number of bands486per primer (7 bands, on average). We highlight here487primers MTI3 and MTI10 that revealed 13 and 11488bands, respectively (Table 2). In contrast, only five489bands were revealed by primer MTI11, similar to490results obtained by Hochu et al. (2006), which found491only three bands for this primer.492

Except for primers UBC 844, UBC 898 and DAT, 493 which showed low polymorphism and low discrimi-494 nation power among the accessions analyzed, all the 495 other ISSR analyzed primers were highly polymorphic 496 with a high number of bands per primer, especially for 497 the UBC 7, UBC 898, JOHN and MANNY primers. 498 These primers revealed more than 10 bands (Table 3), 499 as well as high discrimination power among acces-500 sions, with D ranging from 0.36 to 0.82. The high 501 percentage of polymorphism (75.8 %, on average) 502 observed for ISSR was also reported by Zhou et al. 503 (2008), analyzing 28 cultivars of Dioscorea opposita 504 based on seven ISSR primers, which had a total of 65 505 fragments with 83 % polymorphism. High levels of 506 polymorphism are common in ISSR, such as the 507 results obtained for potatoes, with 90 % polymor-508 phism (Prevost and Wilkinson 1999) and sweet potato, 509 with 62.2 % (Huang and Sun 2000). 510

Mantel test results revealed that data obtained with511the SSR and ISSR markers are correlated (r = 0.5;512p = 0.0002), although they represent different geno-513mic sequences. Several studies have also noted the514existence of high correlation between different tech-515niques of molecular markers in different species (Belaj516et al. 2003; Biswas et al. 2010).517

Genetic structure and conservation strategies

In Brazil, a center of diversity and domestication of 519 various species, studies of genetic diversity are most 520 often associated with economically important crops 521 (Clement et al. 2010). Roots and tuber crops such as 522 yams have been neglected by breeding and conserva-523 tion research (Siqueira 2011). In this context, yam is 524 considered an underutilized crop, subject to selection 525 of interesting characters by the traditional communi-526 ties, where farmers maintain varieties of their prefer-527 ence. Therefore, there is a low level of marketing and 528 exchange of these materials when compared, for 529 example, to other root crops, such as cassava, potato 530 and sweet potato (Siqueira 2011). With this in mind, D. 531 trifida can be affected by the process of genetic drift, 532

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533 whose effect is intensified because of the presence of 534 dioecy, which requires a male and a female plant in the 535 same site for genetic recombination to occur between 536 individuals (Mignouna et al. 2003). Although this species is allogamous in favorable climatic conditions, 537 538 it is strongly maintained by vegetative propagation in 539 cultivated fields, and in most cases, the clones of the 540 same individuals are collected and planted for several 541 years. However, the accessions analyzed in this study 542 showed high genetic variability, with a variation in the 543 similarity coefficient of Jaccard equal to 31 and 56 % 544 for SSR and ISSR data, respectively. Veasey et al. 545 (2012), by analyzing accessions of the same species 546 collected in the Vale do Ribeira, São Paulo, based on 547 isozymes, observed a variation in the similarity coefficient of Jaccard equal to 83 %. 548

549 In the cluster analysis, based on the Jaccard 550 coefficient and UPGMA method, as well as in the PCoA and the Bayesian analyses, three genetically 551 552 distinct and consistent groups were identified, with 553 similar or identical membership. One of the three 554 groups mentioned above consisted of accessions from 555 Iguape municipality in the southern coast of the São Paulo State, which were grouped with accessions 556 collected in the north coast of Santa Catarina, in the 557 558 municipalities of Joinville and São Francisco do Sul. A 559 second group was formed by accessions from the north coast of São Paulo, in Ubatuba municipality, and a 560 561 third group classified the accessions collected in Mato Grosso, a region well apart from the others. The 562 exception was the Amazonian accessions, purchased 563 564 in local markets. Their group membership depended 565 on the marker type or on the genomic region sampled.

566 The interesting fact in this study was the separation 567 of accessions from the south and north coasts of São 568 Paulo. Within Vale do Ribeira. in the south coast of 569 São Paulo, Veasey et al. (2012) had already noticed a 570 spatially structure in genetic variation along a much smaller geographic scale with isozyme markers. 571 572 D. trifida local varieties from Vale do Ribeira were 573 grouped according to their municipalities. Two clus-574 ters (with 100 % bootstrap) were obtained in the 575 cluster analysis, one with varieties from Iguape 576 municipality and the other with varieties from Cana-577 neia municipality. The same genetic structure was 578 observed in the present study, but at a higher 579 geographic scale, separating accessions among the 580 north and south coastal areas in São Paulo State. The explanation for this finding is perhaps the introduction 581

of accessions of this species by waves of migrants582from different regions or even different ethnic groups.583On the other hand, there was a greater similarity of584accessions on the south coast of São Paulo, Iguape,585with those from Santa Catarina, suggesting that586accessions from these two geographically adjacent587regions have the same origin.588

This dynamics can also be related to indigenous 589 influence on the domestication of D. trifida, which has 590 been cultivated by indigenous people from the coastal 591 areas to the Central West region of Brazil (Pedralli 592 1998). Among the indigenous groups involved in this 593 process, we highlight the Guaraní, who are very 594 itinerant and widely scattered throughout Brazil, 595 including the coast of São Paulo (Ladeira 1992). This 596 ethnic group traveled over long distances carrying 597 with themselves various species of edible plants, 598 among which D. trifida (Schmitz and Gazzaneo 1991). 599 This species is, therefore, strongly associated with 600 these indigenous people who, on the other hand, have a 601 strong influence upon the traditional populations in the 602 Vale do Ribeira, São Paulo (Veasey et al. 2012), where 603 the Iguape municipality is located. 604

The maintenance of genetic variation is a major 605 objective for conservation (Hamrick and Godt 1996) 606 and knowledge of variation within and among popu-607 lations provides essential information in the formula-608 tion of appropriate conservation strategies (Francisco-609 Ortega et al. 2000). In this study, most of the genetic 610 variability was observed within groups (60.6 and 611 66.5 % for SSR and ISSR, respectively). In agreement 612 with this result, in a study with wild and cultivated 613 Guinea yams from south and south west Ethiopia, 614 Mengesha et al. (2013) observed that most of the 615 microsatellite diversity was found within rather than 616 among populations. However, our study showed that 617 even among groups the genetic variability was high 618 (39.4 and 33.5 % for SSR and ISSR, respectively). 619 Veasey et al. (2012), comparing two groups of 620 D. trifida accessions analyzed with eight SSR primer 621 pairs, one group including seven accessions from 622 Iguape and Cananéia municipalities in São Paulo and 623 one accession from Mato Grosso, and another group 624 with four accessions from the Amazon, observed that 625 most of the variation was between groups (62.9 %)626 compared with the variation within groups (37.1 %), 627 in contrast to our findings. This result showed that the 628 Amazonian accessions were genetically different from 629 the other accessions. 630

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631 In the present study, we observed a large genetic 632 difference among accessions collected from the different locations analyzed, i.e., between groups, 633 634 since the values of Φst was 0.3937 for SSR and 0.3353 for ISSR, which corresponds to a low gene flow among 635 636 the regions studied. This pattern demonstrates that, 637 unlike other species grown in traditional agriculture, 638 such as cassava (Siqueira et al. 2009) or sweet potato 639 (Veasey et al. 2008), D. trifida is a more regionalized 640 crop or less scattered when compared to other yam 641 species, such as D. alata (Bressan et al. 2011; Siqueira 642 et al. 2012). This observation can be related to 643 historical and socioeconomic factors, such as the 644 different ways of using these tubers and variation in 645 the preference of varieties over time and space (Veasey et al. 2010). The positive correlation between 646 647 genetic and geographical distances obtained in the 648 Mantel test, confirms the structure of these materials in 649 the geographic area sampled. Although the value of 650 this correlation was low, the fact that it is significant at 651 1 % confirms the spatially structure of the genetic 652 variation, which is consistent with the observed Φst 653 value. However, the low correlation values could be 654 due to possible recent exchange of materials among 655 these regions, carried out by non-governmental orga-656 nizations working with indigenous groups, to encour-657 age the exchange of genotypes and inserting new material to prevent genetic erosion of these varieties. 658 659 Thus, exchange fairs are held, allowing the acquisition and supply of new genotypes. There is little informa-660 tion about these indigenous groups, such as the 661 662 Guaraní group mentioned above, who have survived 663 various pressures, such as land struggles, beyond sociocultural pressures of modern societies (Arruda 664 665 1999). These difficulties are common to both indigenous groups and for farmers that grow various species 666

in a traditional way in the tropics. 667 668 Considering the genetic differences among the 669 genotypes grown in the studied locations, the acces-670 sions of São Paulo showed higher diversity, followed 671 by Mato Grosso, Santa Catarina and Amazonas. The 672 average level of genetic diversity present in accessions 673 of D. trifida was 0.27 and 0.18 among states, for SSR 674 and ISSR data, respectively. Considering the groups formed in the Bayesian analysis, using Structure 675 676 software, higher diversity was found among acces-677 sions in group II for SSRs, which includes accessions 678 from three states (São Paulo, Santa Catarina and Amazonas). However, for the ISSR marker, the 679

highest diversity was found for group III (among 680 Mato Grosso accessions), although the values were 681 very similar among the three groups. In general, the 682 average level of genetic diversity was 0.34 and 0.21, 683 for SSR and ISSR data, respectively. These indices are 684 greater than the values found for perennial herbs 685 (H' = 0.17) and to those species with wide geo-686 graphic distribution (H' = 0.20) (Hamrick and Godt 687 1989). However, studies by Zhou et al. (2008) with 688 varieties of D. opposita, widely used in Chinese 689 medicine, showed H' = 0.32, a relatively high value 690 and in most cases superior to that obtained for D. 691 trifida in this study. It is noteworthy that several 692 factors influence the level of genetic diversity of a 693 species, among them, the geographic distribution, life 694 cycle, reproductive system, dispersal patterns, popu-695 lation size, among others (Hamrick et al. 1991; 696 Gaudeul et al. 2000; Zhou et al. 2008). 697

The level of genetic diversity observed among 698 accessions is directly related to the fact that D. trifida is 699 a polyploid species and reproduces both by outcrossing 700 and vegetative propagation. Thus, individuals are 701 usually highly heterozygous, preserving the allelic 702 diversity at the individual level (Veasey et al. 2008; 703 Siqueira et al. 2009). However, the emergence of 704 variant plants arising from seeds, result of genetic 705 recombination, is unlikely, since the occurrence of 706 flowering and fruiting in this species was not reported 707 and detected by farmers during plant collection. 708

The genetic diversity level displayed by the acces-709 sions collected in the States of São Paulo, Mato Grosso 710 and Santa Catarina was expected because of the wide 711 distribution of the crop in these regions. However, we 712 also found genetically similar varieties, which can be 713 related to the fact that the collection has been restricted 714 to a few farmers and a few communities in Ubatuba. 715 SP, where practically all the collection was made in 716 the community of the Sertão de Ubatumirim, and in 717 the community Pirabeiraba in Santa Catarina. The 718 similarity between accessions corroborates the names 719 given by farmers in the studied regions. In São Paulo, 720 except for accession 195, called cara cobrinha (little 721 722 snake yam), varieties were obtained with only two popular names, cara roxo (purple yam) and cara 723 branco (white yam), the latter being represented in 724 smaller numbers. In Mato Grosso, most accessions 725 also received the name cara roxo. In Santa Catarina, 726 the predominant designations for D. trifida were cara 727 and cara mimoso. 728

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729 In conclusion, our results suggest that both markers 730 were useful for evaluation of genetic diversity and 731 assessing differentiation between D. trifida popula-732 tions in Brazil. However, the SSR marker detected 733 higher diversity indices while the ISSR marker 734 seemed more efficient in the clustering of the different 735 genotypes, being able to separate the two Amazonian 736 accessions in the cluster analysis. But both markers 737 detected high levels of genetic diversity for accessions 738 of D. trifida maintained by traditional farmers in the 739 states of São Paulo, Mato Grosso, Santa Catarina and 740 Amazonas. The high within-group variation found is 741 quite interesting for the maintenance of the crop over 742 time. Knowledge of the genetic relationships among 743 accessions is an important information for the efficient 744 use and conservation of this species, both ex situ in 745 genebanks, or in situ, within the aim of conservation in 746 the rural property, known as on-farm conservation. 747 Considering that high genetic diversity was found both 748 within and between groups of accessions from differ-749 ent regions visited, the collection and conservation 750 strategies should consider a large number of individ-751 uals from all regions sampled in order to cover all the 752 genetic diversity present in these materials. On-farm 753 conservation, in the case of D. trifida, is quite 754 interesting for considering the socio-cultural factors 755 involved in the evolution of the species, considering 756 that agricultural practices, through the cultivation and 757 artificial selection, allows the accumulation over time 758 of morphological traits of agronomic interest, provid-759 ing an enrichment of genetic variability not only of 760 this species, but of other crops as well.

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