

Unique haplotypes of cacao trees as revealed by *trnH-psbA* chloroplast DNA

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About 4000 years ago cacao trees were domesticated in Mesoamerica and are still grown. In this study we analyzed the sequence variation of chloroplast DNA *trnH-psbA* intergenic spacer in 28 cacao trees from different farms in the Soconusco region in southern Mexico. Genetic relationships were established by two analysis approaches, based on geographical origin (five populations) and genetic origin (based on a previous study). In our results we identified six polymorphic sites, where five insertion / deletion (indels) type and one substitution were detected. We also found that the overall nucleotide diversity was low for both approaches (geographic = 0.003197; genetic = 0.003841), conversely was obtained moderate to high haplotype diversity (0.6610 and 0.7949), with ten and 12 haplotypes, respectively. The common haplotype (H2) for both networks involved cacao trees of all geographic locations (geographic approach) and four genetic groups (genetic approach). This common haplotype (ancient) derived a set of intermediate haplotypes and singletons interconnected by one or two mutational steps, which suggested directional selection and event-purifying from expansion of narrow populations. No genetic differentiation (AMOVA, $F_{ST} = 0$) was found, and the F_{ST} value (0.04339) of SAMOVA was not big enough to show moderate differentiation between populations. Only one population showed high frequency of haplotypes, thus it could be considered as an important reservoir of genetic material. The indels located in the intergenic spacer *trnH-psbA* of cacao trees could be useful for markers of DNA Barcoding development.

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24

25 **Abstract**

26 About 4000 years ago cacao trees were domesticated in Mesoamerica and are still grown. In this
 27 study we analyzed the sequence variation of chloroplast DNA *trnH-psbA* intergenic spacer in 28
 28 cacao trees from different farms in the Soconusco region in southern Mexico. Genetic
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 32 detected. We also found that the overall nucleotide diversity was low for both approaches
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 34 haplotype diversity (0.6610 and 0.7949), with ten and 12 haplotypes, respectively. The common
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 39 No genetic differentiation (AMOVA, $F_{ST} = 0$) was found, and the F_{ST} value (0.04339) of
 40 SAMOVA was not big enough to show moderate differentiation between populations. Only one
 41 population showed high frequency of haplotypes, thus it could be considered as an important
 42 reservoir of genetic material. The indels located in the intergenic spacer *trnH-psbA* of cacao trees
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44

45 **Keywords**

46 Chloroplast DNA, Haplotype, Nucleotide diversity, indels, *trnH-psbA*

47

48 Introduction

49 The Neotropical cacao tree (*Theobroma cacao* L.), is widely cultivated in America, Africa and
 50 Asia, it is considered an economical important crop because its seeds are used in the chocolate
 51 industry (Wood & Lass, 1985). Based on morphological traits and the geographical origin, trees
 52 are classified as, Criollo, Forastero and Trinitario (Cheesman, 1944). In Mesoamerica, the
 53 Criollo cacao is being widely used as food for nearly 4000 years ago (De la Cruz *et al.*, 1995;
 54 Whitkus *et al.*, 1998; Powis *et al.*, 2011).

55 Based on simple sequence repeat (SSR) analysis, Motamayor *et al.* (2008) propose ten genetic
 56 groups. But nowadays Criollo retains the identity as a separate group, while the other proposed
 57 genetic groups comprising all trees from South America. In this region, it has reported the
 58 highest genetic diversity of cacao trees.

59 On the other hand, genetic diversity of cacao in the South of Mexico was registered as moderate
 60 to low in natural populations (Whitkus *et al.*, 1998; by using RAPD markers), and cultivated
 61 forms (Vazquez-Ovando *et al.*, 2014, by using microsatellite markers), but a wide diversity in
 62 cacao pod and seed morphology was observed. In Soconusco farms (Chiapas, Mexico) Vazquez-
 63 Ovando *et al.* (2014) found moderate to high allelic richness, however high levels of
 64 homozygosity, they also reported the presence of trees sharing genetic identity with those
 65 considered "Ancient Criollo" but also the presence of private alleles. These alleles may be
 66 associated with commercial interest phenotypic traits, while preserving relation with other
 67 polymorphic regions DNA.

68 The chloroplast DNA (cpDNA) and markers based on it, they are increasingly used for studies of
 69 genetic population structure, evolution, gene flow, haplotype frequency and phylogenetic

relationships. Given its high conservation due to maternal uniparental inheritance, cpDNA is the main source of data for construction of phylogenetic relationships in plants (Shaw & Small, 2005). However, there are DNA regions that have variability, which makes them useful for studies of population genetics and conservation issues (Shaw & Small, 2005; Shaw *et al.*, 2007). These regions have been widely used to establish phylogeography patterns in alpine species (Wang *et al.*, 2008), to gain further insight of the centre of origin of cultivated grape populations in Europe (Arroyo-Garcia *et al.*, 2006) and to explain the diversity and structure population of cultivated Chinese cherry (Chen *et al.*, 2013).

The cpDNA intergenic spacer mostly used is *trnH-psbA*, which has showed high variability and besides useful to elucidate genetic relationships at the intraspecific level (Azuma *et al.*, 2001; Hamilton, Braverman & Soria-Hernanz, 2003). The *trnH-psbA* sequenced region of ten cacao accessions deposited in the NCBI database, expose only one haplotype (Kane *et al.*, 2012), while Jansen *et al.* (2011) showed the presence of polymorphic sites, which set a different haplotype. The main polymorphisms reported in the noncoding cpDNA region are inversions, transitions and transversions (Whitlock, Hale & Groff, 2010; Zeng *et al.*, 2012). Few studies report the presence of insertions or deletions (indels), although indels are probably a common feature in the *trnH-psbA* spacer (Aldrich *et al.* 1988).

Nonetheless, the use of indels for diversity and phylogenetic analysis has been questioned by some authors (Bieniek, Mizianty & Szklarczyk, 2015; Whitlock, Hale & Groff, 2010), because the mechanism by indels are generated remains unclear. However, indels are informative character states, since genetic variability analyzed using polymorphism due to indels or substitutions can be studied without distinction (Nei, 1987) therefore they are used as markers. Moreover, when included in diversity or phylogenetic analysis the discriminant power between species is

enhanced (Raymúndez *et al.*, 2002; Hamilton, Braverman & Soria-Hernanz, 2003; Kress & Erickson, 2007; Sun *et al.*, 2012) and even between conspecific individuals (Pérez-Jiménez *et al.*, 2013). Therefore, the aim of this study was to evaluate the genetic diversity and to describe the relationship between individuals of *Theobroma cacao* L. Criollo type of the Soconusco region (Chiapas, Mexico) by using the variation of chloroplast DNA revealed by *trnH-psbA* spacer sequence.

Material & methods

Plan material and sample collection

A total of 45 samples cacao were including in this study. 38 trees were sequencing and analyzed, and seven sequences accessions were taken from GenBank as references. From 38 trees sequenced, 28 were selected from plantations of Soconusco (Chiapas, Mexico) based on a previous characterization (Vazquez-Ovando *et al.*, 2014) using ten SSR's molecular markers. The individuals were selected based on traits of fruits (pod) and seeds, which resembled those of the Criollo variety. Pods were elongated, deeply grooved, pointed at the pod end, had a lumpy surface with a warty appearance outside, white or slightly pigmented seeds, and sweet mucilage. In agree with that reported by Vazquez-Ovando *et al.* (2014) the individuals were classified as: 12 trees with high Criollo ancestry, ten non Criollo group (hereinafter called Forastero) and six admixtures (hereinafter called hybrid) (Table 1). Additionally, ten accessions were sequenced and included as references: two Forastero variety (Catongo and EET 399), one Trinitario variety (RIM 24) and seven wild Criollo (one collected in the Lacandon rainforest [SL01] and six obtained from germplasm of the Instituto Nacional de Investigaciones Forestales, Agrícolas y Pecuarias, México [Yaxcabá, Xocen, Lacandón 06 Lacandón 28, Lagarto and Carmelo]) (Table

1). *Theobroma bicolor* was used as outgroup. The average age of the trees was 30 year. Leaves were sampled and placed in plastic bags, taken to the laboratory (4°C) and they stored at -20 °C until processing.

DNA extraction, amplification and sequencing

Total DNA extraction was performed by modifying the method described by Doyle & Doyle (1990). Leaves were washed with sterile water and 70% ethyl alcohol. Approximately 200 mg cacao leaves were ground with liquid nitrogen with 60 mg polyvinyl pyrrolidone and 1 mL CTAB buffer [2% CTAB (w/v), 20 mM ethylenediaminetetraacetic acid (EDTA), 1.4 M NaCl, 100 mM Trizma® base, pH adjusted to 8 with HCl, and 1% 2-mercaptoethanol (v/v)]. DNA extractions were performed with chloroform-isoamyl alcohol and precipitation with isopropanol. The extracted DNA was purified with a mixture of phenol:chloroform:isoamyl alcohol (25:24:1). The DNA integrity (dissolved in 60 µL Milli-Q water) was checked by 0.8% agarose electrophoresis and quantified by spectrophotometry at 260 nm (Jenway, Genova), and the purity was inferred by the 260/280 and 260/230 absorbance ratios.

The cpDNA amplification of trnH-psbA intergenic spacer was conducted by using forward primer 5'-CGCGCATGGTGGATTCAACAATCC-3' and reverse primer 5'-GTTATGCATGAACGTAATGCTC-3' (Shaw & Small, 2005). PCR conditions was performed following to described by Shaw & Small (2005) with changes in the concentration of MgCl₂ (2mM) and using the average value of the temperature melting reported. PCR was performed in a 25 µL reaction mixture, containing 100 ng genomic DNA, 4 µL of 10x PCR buffer ViBuffer A (Vivantis™), 1 µL of MgCl₂ (50 mM), 0.5 µL of dNTP Mix (10mM, Promega), 0.05 mM of each primer and 2.5 U of Taq DNA polymerase (Vivantis™). Following one cycle of 5 min at

94 °C, 35 PCR cycles of 30 s at 94 °C, 30 s at 53 °C, 1 min at 72 °C and 10 min at 72 °C final extension were performed in a thermal cycler TC3000 (Techne, Cambridge, UK). PCR products were separated on 6% polyacrylamide gels using 0.5X TBE buffer at 110 V for 210 min, stained with ethidium bromide (0.6 ng/μL) for 15 min, visualized under UV light and photographed with a Gel Doc TM EZ Imager gel documentation system (Bio-Rad, USA). Fragment sizes were estimated using the Image Lab (v. 4.0.1, Bio-Rad Laboratories) and integrating GeneRulerTM 100 bp DNA Ladder Plus (Fermentas®) as a molecular weight marker.

PCR products were directly sequenced by using Dye Terminator Cycle Sequencing with Quick Start Kit (GenomeLabTM) on a CEQTM 8000 automatic DNA sequencer (Beckman CoulterTM). To validate the results, DNA was extracted twice, amplified independently, and sequences were verified by comparison of their forward and reverse sequences when applicable.

Sequence alignment and data analysis

The sequence quality was checked and electropherograms edited by using BioEdit © (Hall, 1999). Sequences were limited at the ends to avoid the presence of variable sites due to artifacts sequencing by polymerase (approx. 40 bp) and aligned with ClustalW 1.81 (Thompson, Higgins & Gibson, 1994). Visual inspection and manual edition of sequences for confirming variable sites was performed. We used two different analytical approaches, based on the geographic origin and the genetic origin of the samples (Table 1). In both approaches molecular diversity indices; the number of segregating sites (S), number of haplotypes, haplotype diversity (Hd) and nucleotide diversity (π d) were estimated following Nei (1987) in DnaSP© 5.1 (Rozas *et al.*, 2010).

In order to infer evolutionary relationships at the intraspecific level, we evaluated network building. The method used was median-joining (MD) based on parsimony criteria (Bandelt, Forster & Röhl, 1999; Polzin & Daneshmand, 2012), which performed with the software Network© 4.6.1.3.

Analysis of molecular variance (AMOVA), pairwise F_{ST} values as well as the statistics of molecular variances F_{CT} (test by permuting individuals within populations), F_{ST} (test by permuting genotypes among populations but within groups) and F_{SC} (test by permuting genotypes among groups) were estimated using the Arlequin© version 3.0 (Excoffier, Laval & Schneider, 2005). Significance was evaluated by 99 999 random permutations of sequences. In order to determine whether sample sites clustered on a population level, a spatial analysis of variance (SAMOVA) was conducted (Dupanloup, Schneider & Excoffier, 2002), using haplotype data and geographic co-ordinates of each of the 5 sample sites. The SAMOVA was run for $K = 2-5$ putative populations to determine the maximum F_{ST} value, the highest proportion of differences between populations due to genetic variation.

The neutral evolution of chloroplast DNA was evaluated to examine whether any population had experienced historical demographic changes using test Tajima's D (Tajima, 1989) by using Arlequin© version 3.0 (Excoffier, Laval & Schneider, 2005). We evaluated for geographical approach, overall as well as populations.

When the analysis was conducted for genetic origin approach, seven accessions from NCBI database as reference were included: Matina-06 (HQ336404.2), Criollo-22 (JQ228379.1) Amelonado (JQ228380.1) Scavina (JQ228382.1), ICS-01 (JQ228381.1), ICS-06 (JQ228383.1), ICS-39 (JQ228387.1).

181

182 Results

183 Sequence characterization and genetic diversity

184 The sequences of *trnH-psbA* intergenic spacer in 45 samples *Theobroma cacao* (Table 1) were
 185 aligned with a consensus length of 526 bp, of which six polymorphic sites (Table 2), including a
 186 substitution of T (A) at position 134, five indels (Figure 1) and six segregating sites were
 187 detected. This resulted in 12 haplotypes, of which four were singletons represented by a unique
 188 sequence in the sample (Table 2). The nucleotide composition of the fragment revealed AT-rich
 189 (A+T, 75.52%).

190 The geographic approach analysis revealed overall the average values of haplotype diversity
 191 (Hd) and nucleotide diversity (μd) were 0.6610 and 0.0031, respectively (Table 3). 10 haplotypes
 192 identified, the most frequent haplotype (H2) was shared by 19 trees of the seven geographic
 193 populations formed *a priori* (Table 2). Four trees that belong to Population 1 (one tree),
 194 Population 3 (one tree) and Population 5 (two trees) were the second most common haplotype
 195 (H1). Overall 50% haplotypes were singletons (Table 2). The analysis showed that most genetic
 196 diversity was found in Population 4 (Mazatán), with the highest values for all indices; also it
 197 included 50% of the haplotypes identified. The other populations maintained moderate Hd and
 198 low πd with similar values each population (Table 3). Yucatán and Selva Lacandona populations
 199 (wild) they exhibited Hd 1 and 0 respectively, although these data are influenced by the low
 200 number reference individuals.

201 Meanwhile when the data analysis was based on the genetic origin, the highest Hd (0.93) was
 202 found in the hybrid group (Table 3). In contrast, the Trinitario-reference group had the lowest

203 value H_d (0.5). The π_d was low (0.0025 to 0.0027) for all groups, similar to another approach.
 204 Forastero-reference and Trinitario-reference groups did not present singletons (Table 3).
 205 Sequences from the NCBI database were grouped into one haplotype (H12), except HQ336404.2
 206 it grouped in H11 whit EET399, that corresponding to Forastero-reference group.

207 Intraspecific relationship haplotype

208 Figure 2 shows the haplotype network built with data from geographic approach (a) and genetic
 209 approach (b). Both networks show a star arrangement. The general base has a common haplotype
 210 for the two networks (H2) that included cacao trees from all geographic populations (a) and four
 211 of six genetic groups (b). From this common haplotype (H2) derive a unique set of intermediate
 212 haplotypes and interconnected by one or two mutational steps, in both networks. H4-H6
 213 haplotypes were farthest from the central clade, i.e. haplotypes newly created (Figure 2).
 214 Haplotypes H3 and H8-H10 were singletons.

215 Population genetic structure

216 The analysis of molecular variance (AMOVA) was not significant and with a value $F_{st}=0$; while
 217 in the spatial analysis of molecular variance (SAMOVA) it was found that the value of $K=2$
 218 extends the F_{st} to 0.04339 (Table 4), generating two clusters; the first contained only the
 219 Population 4 (Mazatán) and the second cluster grouped the other geographic populations (Table
 220 4).

221 Neutrality tests showed non-significant value in the Tajima's D , except for the Population 4, in
 222 which the Tajima's D value was negative ($D = -0.93302$). All other populations showed values
 223 of $D = 0$; however the overall value for this test was $D = -0.13329$ ($P > 0.1$).

224

225 Discussion

226 In this study was found high haplotype variation in chloroplast DNA cacao trees grown in the
 227 Soconusco region. No found inversions, transitions or transversions which reported as commons
 228 in other plants (Whitlock, Hale & Groff, 2010; Zeng *et al.*, 2012). However, we found
 229 polymorphism, type insertions or deletions (indels) in three poly-A regions (Figure 1). This
 230 agrees with that reported by Jansen *et al.* (2011), in HQ336404.2 accession and support the
 231 affirmation Aldrich *et al.* (1988) that indels are a presumably common feature in the region *trnH*-
 232 *psbA*. For the data analysis, we included the indels as informative character states, being as the
 233 high interspecific divergence of the region spacer allow even be used as a marker of DNA
 234 Barcoding (Kress & Erickson, 2007). Molecular diversity indices found in the present study have
 235 similarity to the results of Zeng *et al.* (2012) using the same intergenic spacer, which revealed 11
 236 haplotypes for 35 samples of *Thinopyrum intermedium*, low nucleotide diversity ($\pi_d = 0.00473$)
 237 and moderately high haplotype diversity ($H_d = 0.7331$) (our results for geographic populations
 238 were $\pi_d = 0.003197$, $H_d = 0.6610$). The results of these authors further support the use of one
 239 intergenic spacer to reveal nucleotide polymorphism.

240 Our results of the haplotype diversity are contrary, those reported by Vazquez-Ovando *et al.*
 241 (2014) who reported low genetic diversity when the study conducted with individuals of the
 242 same region (in particular the Population 4 Mazatán) using microsatellite markers. However, the
 243 low nucleotide diversity found in this study is supported by the low genetic variability found
 244 with microsatellites. Individuals included in both studies showed great phenotypic pod variability
 245 resemble to Criollo-type (eg. different degrees of roughness, color, deeply grooved). This could

be revealing greater association between morphological variability cacao pod with reported allelic richness (Vazquez-Ovando *et al.*, 2014) and variability of haplotypes found in our study. The number of haplotypes was higher than polymorphic sites (Table 2) this feature is associated with ancestral species that have diverged enough, accumulating mutations among different haplotypes (Roger, 1995). Population 7 (Selva Lacandona) exhibited no haplotype diversity ($H_d = 0$), however the haplotype (H2) located in this population is considered the common ancestor due to it share all populations (Figure 2A). On the contrary, the two individuals belong the Population 6 (Yucatán), which exhibited each other different haplotypes (H2 and H9), interrelated by only a mutational step (Figure 2A). This shows that in the trees belonging to Yucatán population, an individual eventually descended from other of this region where the Maya people grown cacao.

Low levels nucleotide polymorphism could be explained by rapid population expansion events in its distribution range, whereas high haplotype diversity may be due to continuous introduction of individuals from different locations. In populations recently introduced or expanded from a small number of founders, would have a common haplotype shared by most individuals and many rare haplotypes connected to the main by few independent mutations (Slatkin & Hudson, 1991; Avise, 2000) as in the present study (Figure 2). A similar argument is explained when using microsatellite markers (Vazquez-Ovando *et al.*, 2014).

The relative low variability in cacao cultivated populations is also supported by the lack of neutrality revealed by the global test Tajima. Specifically in the Population 4 (Mazatán) the negative value of Tajima's D (-0.93302), could be related to an event "bottleneck" which would indicate population expansion, not natural because of it is cultivated populations. It is reported that in the past occurred unclearly events (disease, volcanic eruptions or other natural events)

they may have caused the almost complete disappearance of populations established by the people in the Mesoamerican region (De Sahagún, 2009, *Codex Florentino*). A process rapid expansion due to recolonization in populations, probable introduction of other varieties of trees cacao not native in the region, would subject the populations to event bottleneck in very recent periods. However, these are presumptive weak to infer the population history by use a single locus. The bottleneck event could also be related to the loss of alleles (haplotypes), mainly rare alleles, which is much greater than the loss of genetic variance *per se*. Although these rare alleles contribute little to the total genetic variability, can provide unique responses against challenges evolutionary as found in this study a high number of unique haplotypes (3 haplotypes in population 4). The presence of both common and rare haplotypes can be understood by a directional-purifying process selection or expansion events from small populations (Hedrick, 2005). H3 and H6-H10 haplotypes (cultivated populations) are singletons, agree with Crandall & Templeton (1993) the singletons located in this study were connected with haplotypes from the same population. Population 4 (Mazatán), shows the highest haplotype diversity, which makes this population an important reservoir of genetic material at the level of chloroplast, and possibly phenotypic, as also it is the abundance of morphology in pods seen in this population.

Overall, cacao trees with high ancestry was located in the center of haplotype network, this supported by coalescence theory that predicts the ancient haplotype should be the most common and most distributed among populations. In concordance derived haplotypes would be less frequent, and in many cases private; these would be located in regions for cacao cultivated populations latest. H4 and H5 are haplotypes perhaps recently created because of they are located at the ends of the network, which may be due to germplasm exchange with traits of interest of cacao farmers. These anthropogenic activities perhaps had a strong impact on the

levels of variation observed in cpDNA sequences, which explains the observed no differentiation. In addition, migration over long distances by the exchange by farmers contributed to the colonization of new regions founded by few individuals, establishing different alleles by mutation and genetic drift.

Related to the genetic origin MAJH02 and Carmelo individuals are located in the haplotypes 4 and 6 respectively (Figure 2B). They possibly belong to "hybrid" group rather than the Criollo. But also they are contenders for Modern Criollo group i.e. individuals classified as Criollo but which might have been introgressed with Forastero genes (Motamayor *et al.*, 2002) and preserve phenotypic traits of Criollo ancient.

Furthermore the value found for $F_{ST} = 0.00$ by AMOVA reveals that all molecular variance is within populations. Indeed the F_{ST} value of SAMOVA (Table 4) is not enough to show at least moderate differentiation between populations ($F_{ST} \geq 0.05$). This provides some explanations regarding the demographic history of *T. cacao* trees, indicating that populations formed *a priori* have experienced gene flow, which results in population homogenization. Spatial analysis reveals highest differentiation between groups when $K=2$ is tested; meanwhile a $K=3$ ($F_{ST} = 0.00088$) grouped trees of Yucatán, Selva and Cacahoatán in the same genetic population. This is unusual; being that geographic distance is longer among the three localities and may be associated with distribution of trees in the past, i.e. the ancestral haplotype (H2) grouped individuals of Selva and that for one mutational step it originated the individuals of Yucatán, which in turn originated at individuals of Cacahoatán by the same event (network haplotype by genetic origin, Figure 2B). Following this criterion, the H3 has a greater correspondence with the Criollo genotype, rather than it reported previously as hybrid (Vazquez-Ovando *et al.*, 2014).

314

315 Conclusions

316 Indels located in the chloroplast DNA *trnH-psbA* spacer region of cacao trees could allow the
 317 development of genetic markers barcode. The molecular analysis of nucleotide diversity showed
 318 low diversity, but high haplotype diversity, which may be due to events bottlenecks populations,
 319 confirmed with negative Tajima's D and haplotype network in a star arrangement. It also allowed
 320 identifying ten different haplotypes (trees grown) of which H3 and H6-H10 resulted singletons
 321 because they are not associated with other cacao grown or with those reported in the molecular
 322 databases. The presence of these haplotypes, accompanied by the low number of mutational
 323 steps that groups might suggest a very short evolutionary history or events of disappearance-
 324 expanding populations of southern Mexico. A geographical population (Pop 4 Mazatán) was
 325 located high frequency haplotypes, which makes this zone an important reservoir of genetic
 326 material at the level of chloroplast, and possibly phenotypic, since they were also observed in
 327 this population abundance of morphology in pods. The genetic differentiation between
 328 populations was zero, by suggesting that gene flow homogenized populations.

329

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332

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Table 1 (on next page)

Theobroma cacao trees, geographical coordinates and Genbank accessions. Genetic origin (Criollo, Non Criollo and admixtures) based on Vázquez-Ovando et al. (2014).

*Geographic population *a priori*, **reference tree 

1



Pop*	Coordinates latitude (N)/ longitude (W)	Criollo (n=20)	Non Criollo (n=15)	Admixtures (n=10)
1	14°59'28''N, 92°26'44''W (Huehuetán) 14°52'55''N, 92°21'42''W (Tapachula)	HUJF01 TASG12	HUJF03	TASG16 TASG18
2	14°56'41''N, 92°09'59''W (Tuxtla Chico) 14°59'53''N, 92°10'44''W (Cacahotán)	TCHR04	CAAM12	CAAM04
3	14°47'31''N, 92°11'11''W (Frontera Hidalgo) 14°38'27''N, 92°13'47''W (Suchiate)		FHSA06 SUED02 SUED03 SUED06	FHSA02
4	14°48'56''N, 92°29'06''W (Mazatán)	MAMG12	MAMG03 MAMG04 MAMG07 MAMG08	MAMG10
5	15°28'07''N, 92°48'42''W (Mapastepec) 15°10'31''N, 92°38'06''W (Villa Comaltitlán) 15°11'17''N, 92°36'55''W (Villa Comaltitlán)	MAJH02 VCHL01 VCHL02 VCHL03 VCHL04 VCLB02 VCLB03 VCLB04		MAJH03
6**	20°32'29.25''N, 88°50'35.82''W (Yucatán)	Yaxcabá Xocen		
7**	16°06'42.92''N, 90°56'31.28''W (Selva Lacandona)	Lacandón 06 Lacandón 28 SL01		
8**	INIFAP (Several)	Lagarto Carmelo	Catongo EET399	Rim24
9**	GenBank	Criollo-22	Scavina Amelonado Matina-06	ICS-01 ICS-06 ICS-39

2

Table 2(on next page)

Nucleotide polymorphic sites and cpDNA haplotypes in populations of cacao based on variation spacer intergenic *trnH-psbA*

1

Haploty pe	Polymorphic site						Populations								
	22	13 4	20 6	30 9	31 0	48 7	Pop 1	Pop 2	Pop 3	Pop 4	Pop 5	Pop 6	Pop 7	Pop 8	Pop 9
H1	C	T	-	A	A	A	1		1		2				
H2	-	T	-	A	A	A	3	2	3	2	5	1	3		
H3	-	A	-	A	A	-		1							
H4	C	T	A	-	-	-			1		1				
H5	-	T	A	-	-	-				1	1				
H6	-	T	A	-	-	A				1					
H7	-	T	-	A	-	-				1					
H8	C	T	-	A	-	A	1								
H9	-	T	-	A	A	-						1			
H10	-	A	-	A	A	A				1					
H11	-	T	-	A	-	A								1	1
H12	-	T	A	A	-	-									6

2

Table 3(on next page)

Genetic diversity cacao from Soconusco (Chiapas, Mexico) grouped by geographic approach (Pop) and genetic origin approach

1



Pop	Locality	N	S	Sn	H	Hd ± de	πd ± ED
1	Huehuetán, Tapachula	5	2	1	3	0.7000±0.218	0.001905±0.00177
2	Cacahoatán, Tuxtla Chico	3	2	1	2	0.6667±0.314 3	0.002545±0.00261 4
3	Frontera Hidalgo, Suchiate	5	5	0	3	0.7000±0.218 4	0.004183±0.00322 3
4	Mazatán	6	5	3	5	0.9333±0.121	0.004825±0.00347
5	Mapastepec, Villa Comaltitlán	9	5	0	4	0.6944±0.147 0	0.003908±0.00273 5
6	Yucatán	2	1	1	2	1.0000±0.500	0.001908±0.00269
7	Selva Lacandona	3	0	0	1	0	0
Total		3	--	6	--	0.6610±0.089	0.003197±0.00213
 Genetic origin approach							
“Criollo”		1	6	1	4	0.6364±0.127	0.002506±0.00188
“Forastero”		1	5	1	5	0.6667±0.163	0.003295±0.00236
“Hybrid”		6	6	1	5	0.9333±0.121	0.005450±0.00383
Criollo-reference ^a		8	4	1	5	0.7857±0.150	0.003333±0.00245
Forastero-reference ^a		5	3	0	3	0.8000±0.164	0.003053±0.00251
Trinitario-reference ^a		4	4	0	2	0.5000±0.265	0.003802±0.00319
Total		4	--	4	--	0.7949±0.052	0.003841±0.00244
N=Samples sizes, S=Number of segregating, Sn=Singletons, H=Number of haplotypes, Hd=Haplotype diversity, πd=Nucleotide diversity. ^a Including sequences GenBank (Criollo-reference n=1, Forastero-reference n=3, Trinitario-reference n=3).							

2

Table 4(on next page)

Spatial analysis of molecular variance ($K = 2$) for populations cacao and the statistics of molecular variances fixation indices corresponding to groups

1

Source of variation	df	SS	VC	Variation (%)	Fixation indices	<i>P</i> value
Among groups	1	1.613	0.12816	13.98	$F_{SC} = -0.11146$	0.73412
Among populations within groups	5	2.507	-0.08789	-9.59	$F_{ST} = 0.04393$	0.00684
Within populations	26	22.789	0.8765	95.61	$F_{CT} = 0.1398$	0.14956
Total	32	26.909	0.91677			
df= degrees of freedom, SS=Sum of squares, VC=Variance components.						

2

3

1

Location of indels (blue arrows) in a fragment of chloroplast DNA *trnH-psbA* intergenic spacer of *Theobroma cacao* trees. See Table 1 for details of the samples



2

Haplotypes frequency in each geographical population (A) and contribution of genetic groups to the haplotype network (B). Built with Network© 4.6.13 by Median Joining method.

