# **Increasing Accuracy and Throughput in Large-Scale Microsatellite Fingerprinting of Cacao Field Germplasm Collections**

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Abstract In this study we report on increasing the rate and accuracy of microsatellite fingerprinting of accessions in Theobroma cacao L. field germplasm collections with a medium-throughput capillary sequencing system. We examined the use of a reduced number of microsatellite loci to decrease the time and materials required for fingerprinting and determined the best available microsatellite loci for accurately separating accessions. A subset of nine informative loci that could separate sixty accessions into the same genetic groupings as a complete set of 37 loci was found. Stringent probability of identity values (i.e. chance of unique accession) was highly influenced (r=-0.996; P<0.001) by the number of alleles utilised in the fingerprinting set but the composition of the primer set was more important when choosing discriminatory loci. DNA pooling to reduce the number of samples was also investigated. There was a relatively high level of mixture within plots (59% of 54 plots examined) of the field genebank, which opposed the

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M. Boccara CIRAD - UMR DAP TA A96/03, 34398 Montpellier cedex 5, France use of a pooling strategy to fingerprint the multiple trees of an accession plot in the collection.

**Keywords** Discriminatory loci · Germplasm · Identity resolution · SSR fingerprinting · *Theobroma cacao* L.

#### Abbreviations

ADO	allele drop out
AFLP	amplified fragment length polymorphism
ANOVA	analysis of variance
CATIE	Centro Agronomico Tropical de
	Investigacion y Enseñanza, Turrialba
CE	capillary electrophoresis
DNA	deoxyribonucleic acid
EMBL	European Molecular Biology Laboratory
H <sub>o</sub>	observed heterozygosity
H <sub>e</sub>	expected heterozygosity
ICG,T	International Cocoa Genebank, Trinidad
MPP	microsatellite primer pair
PCR	polymerase chain reaction
PIC	polymorphism information content
P <sub>(ID)sib</sub>	probability of identity among full siblings
P <sub>(ID)sib</sub> COM	overall probability of identity among full
	siblings
RAPD	randomly amplified polymorphic DNA
SDW	sterile deionised water
SSR	simple sequence repeat

# Introduction

Cacao (*Theobroma cacao* L.), the source of chocolate and cocoa butterfat, possesses recalcitrant seeds and is an out-crossing species [75] leading to its conservation as

clonally propagated trees within field genebanks. Since the 'discovery' of this crop, cacao germplasm has been sought from South and Central America, wherein resides the centre of diversity [16, 75, 13, 7]. Over fifty-four cacao germplasm collections are present worldwide and only two of these are Universal Collections (representing nearly all of the known genetic diversity) - Centro Agronomico Tropical de Investigacion y Enseñanza, Turrialba (CATIE) in Costa Rica and the International Cocoa Genebank, Trinidad (ICG,T) in Trinidad and Tobago [51, 73]. The ICG,T is one of the largest cacao germplasm collections containing over 2,000 accessions, with each accession being represented by a maximum of 16 trees and an average of six trees. Formally planned in 1982, the gene bank was assembled from germplasm material collected in multiple expeditions (1930 onwards) from Amazonian South America, Central America and the West Indies [31]. The ICG,T contains wild, semi-wild and cultivated material and includes germplasm from the recognised Criollo, Forastero, Refractario and Trinitario germplasm groupings.

Mislabelled plants have been identified as a serious problem in germplasm collections [26]. The impact of duplication in germplasm collections and its estimation was discussed by Van Hintum [72]. Errors within germplasm collections have been reported for French olive [32], Tunisian fig [57] and persimmon [3] using Randomly Amplified Polymorphic DNAs (RAPDs); Cicer using Amplified Fragment Length Polymorphisms (AFLPs; [62]); Eucalyptus with AFLPs and RAPDs [70]; Moroccan fig using inter-simple sequence repeats and Simple Sequence Repeats (SSRs; [33]) and poplar germplasm with AFLPs and SSRs [21]. Cacao germplasm collections were likewise shown to contain mislabelled individuals [20, 12, 56, 51, 50, 65]. In Trinidad, mislabelling can be attributed to the multiplicity of introductions and transfers of plants from point of collection to establishment in early holding sites, and to the subsequent recollection of budwood and repropagation of material for establishment of the ICG,T. The potential for human error during plot demarcations and planting may have also led to identity confusion among and within accession plots.

Mislabelling issues can be resolved by multilocus fingerprinting. A variety of molecular markers are available, and microsatellite markers are well suited since they are co-dominant (thus allowing the detection of heterozygotes), found throughout the genome and have high allelic variability [55]. In addition, SSRs are relatively fast and easy to analyse compared to other DNA markers [48]. Cacao microsatellites developed by Lanaud et al. [37] have been extensively utilised for cacao clone identification [20, 11, 56, 50, 15] with 15 microsatellite primer pairs (MPPs; [58]) being recommended

for resolution of identity issues. Cacao germplasm have been assessed with eleven loci ([60]; 94 accessions), 14 loci ([29]; 69 Trinitario accessions) and 15 loci ([77]; 141 accessions).

Unambiguous identification of individuals within a germplasm collection is, however, a large-scale project requiring allocation of substantial resources and time. There are approximately 12,000 trees (2,000 accessions  $\times$  six propagated trees) to be evaluated over 15 loci (resulting in 360,000 data points). Hence, any effort to resolve identity issues in a more timely and cost-effective manner would be welcomed by users of the ICG,T.

One way of achieving this is by reducing the number of microsatellite loci required for detecting mislabelling in the germplasm collection. Microsatellites with many alleles per locus would favour unique fingerprinting of a large number of accessions by relatively few loci [43]. Kottapalli et al. [36] in a study of 112 peanut accessions with 67 SSR markers found that a subset of 12 markers was sufficient for identification and gave similar clustering pattern. Sixteen SSR loci were sufficient to completely identify 96 sorghum accessions [1]. Ninety cowpea [*Vigna unguiculata* (L.) Walp] breeding lines could be 97.8% resolved with five SSR primers [38]. In addition, many other cacao microsatellites have been identified since the recommendation of Saunders et al. [58], and therefore other MPPs may be more useful than the set that was initially proposed.

Another method involves bulking samples from trees of a given accession thereby reducing the total number of samples to be identified. DNA pooling is a useful, costeffective tool for large scale studies including association studies [61, 5, 46], single nucleotide polymorphism identification [4, 52, 76], mapping of quantitative trait loci [28]; identification of markers for disease resistance [45] and determination of frequencies of microsatellite alleles [53, 17, 63]. DNA pooling was also applied in genetic diversity studies with RAPDs [23], restriction fragment length polymorphisms [18] AFLPs [35] and in cacao microsatellite characterization [14].

Large datasets of natural populations are prone to error accumulation [41] even with high-throughput genotyping [19]. Microsatellite genotyping errors are recognised to occur [9, 22, 25] and several approaches to reduce this error have been documented [25] including repeat genotyping [68, 69] and the mismatch tolerant approach [44, 30] with a combination of these two being recommended for cacao germplasm management [77]. Barratt et al. [6] recommended pooling 50 individuals to balance accuracy and cost for SNP genotyping. Zou and Zhao [78] demonstrated that measurement errors had a greater effect on DNA pooling than genotyping errors had on individual genotyping and that there was a high degree of false positives with DNA pooling. This study was therefore undertaken to determine (i) the composition and minimum number of microsatellite loci required for an accelerated yet reliable fingerprinting protocol of individuals within a cacao germplasm collection and (ii) the feasibility of using DNA pooling for assessing the homogeneity of accession plots that contain more than one tree.

# Results

## Microsatellite Assessment

Under the experimental conditions used in this study, all PCR products of the 37 MPPs were devoid of false alleles. Six SSRs (mTcCIR 11, 17, 33, 45, 56 and 210) had an ADO of at least 0.5, but 14 SSRs with an ADO of not more than 0.15 were relatively error-free. Additional characteristics of the individual SSR loci based on the sixty accessions utilised in this study are provided in Table 1. Allele number, observed heterozygosity ( $H_o$ ), expected heterozygosity ( $H_e$ ), PIC, P<sub>(ID)sib</sub> and ln S<sup>2</sup> were not significantly affected by interruption of the repeat (imperfect vs. perfect) or complexity of the repeat (compound vs. simple) in dimeric loci.

When microsatellite sets were compared, the P(ID)sibCOM and the separation ability were found to be significantly (P < 0.001) dependent on the number of alleles involved (Figs. 1 and 2). Correlation coefficients of -0.996 and 0.997 were obtained respectively for these comparisons. A total of 326 alleles were obtained from 37 loci, which resolved the 60 cacao accessions into 54 (90%) groups. Six pairs of accessions were unresolved: AC 20 vs. IB 9, BC 3 vs. HF 8, CRIOLLO 22 vs. IB 2, NA 184 vs. NA 331, NA 432 vs. NA 680 and NA 831 vs. NA 833. The set of primers currently in use for cacao fingerprinting [58] separated the 60 accessions into 47 (78.3%) groups. The additional accessions that were unresolved with the latter primer set were comprised of only NA accessions, which are known to be comprised of several sib families. One additional pair was added (NA 406 vs. NA 528) and six other NA accessions (NA 266, NA 435, NA 504, NA 734, NA 773 and NA 860) were lumped into the same group as NA 184 and NA 331.

The separation ability of a primer set was also influenced by its composition, and several sets yielded the same separation for a wide range of allele numbers (Table 2). Primer combinations comprising the most informative loci, as ranked by GIMLET v.1.3.3 [71], performed as well as those recommended by Saunders et al. [58] even though the set included less loci and had a lower total number of alleles (Fig. 3). An equivalent separation of the sixty accessions with nine loci, as compared to that with 37 loci, was achieved. These loci were: (a) Group 1 — mTcCIR15, mTcCIR26, mTcCIR37; (b) Group 2 — mTcCIR33, mTcCIR57, mTcCIR42 and (c) Group 3 — mTcCIR12, mTcCIR243, mTcCIR244. Each group represent a trio suitable for post-PCR multiplexing based on allele ranges obtained in this study. In this set of nine loci, a total of 101 alleles were found, with a combined  $P_{(ID)}$  of  $8.886 \times 10^{-12}$ and a  $P_{(ID)sib}COM$  of  $1.437 \times 10^{-4}$ . The latter was a hundredfold increase compared to that obtained ( $2.233 \times 10^{-6}$ ) from the set of 15 loci recommended by Saunders et al. [58]. The relationship among the sixty accessions is represented in Fig. 4.

Effect of DNA pooling on Allele Sizing

All DNA pools were amplified with minimal background noise. Selecting peaks for binning analysis was primerdependent, being easiest with mTcCIR37 and most difficult with mTcCIR15. A high level of consistency was achieved between scorers (Table 3). Type I error was low and was encountered by one scorer for mTcCIR37 when four DNA samples were pooled (three of eight assessments) and for mTcCIR15 for pools of two (three of 15 assessments) and three (three of 16 assessments) individuals. Type II error was more common being reported, by both scorers, in seven out of nine MPP-pool combinations. Type II errors were attributed to the type of MPP (P=0.001), number of DNA samples pooled (P < 0.01) and the interaction of these two terms (P < 0.05). The SSRs mTcCIR15 and mTcCIR26 were primarily responsible for these effects when amplifying pools of three or four individuals. The total number of alleles called was significantly affected by the MPP used (P < 0.05) but all alleles scored were correctly identified (Table 4).

# ICG,T Plot Homogeneity

The 54 plots examined contained 22 homogenous samples (40.7%) and 32 (59.3%) mixed plots with sixteen plots (29.6%) having two genotypes (Table 5). Analysed plots that contained at least three trees had a mixed composition of genetic identities in 65.9% (29 of 44 plots) of the plots (Fig. 5).

# Discussion

This study examined the possibility of accelerating verification of identities in gene banks, with reduced genotyping error, by (1) decreasing the number of loci by identification of loci which were most efficient at differentiating cacao accessions and (2) decreasing the number of samples by utilising pooled samples

Table 1 Attributes of microsatellite loci assessed in sixty cacao accessions

<sup>1</sup> Locus	<sup>2</sup> ADO	<sup>3</sup> Rk	<sup>4</sup> Sepn	<sup>5</sup> N <sub>a</sub>	Allele range	<sup>6</sup> ln S <sup>2</sup>	<sup>7</sup> PID <sub>sib</sub>	<sup>8</sup> PIC	<sup>9</sup> H <sub>o</sub>	$^{10}\mathrm{H}_{\mathrm{e}}$
CIR1	0.42	31	8 (13.3)	7	127-151	3.81	0.51	0.52	0.28	0.60
CIR3	0.19	1	21 (35.0)	15	211-279	5.47	0.33	0.85	0.33	0.86
CIR6	n.d.	23	14 (23.3)	8	229-251	3.29	0.43	0.66	0.30	0.70
CIR7	n.d.	29	11 (18.3)	6	148-162	2.51	0.50	0.55	0.25	0.60
CIR8	n.d.	25	15 (25.0)	7	289-307	3.07	0.46	0.63	0.32	0.66
CIR9	0.00	12	15 (25.0)	9	258-296	4.30	0.39	0.73	0.45	0.77
CIR10	0.45	17	12 (20.0)	6	206-216	1.95	0.41	0.70	0.42	0.74
CIR11	0.62	13	20 (33.3)	13	282-320	4.23	0.39	0.73	0.37	0.76
CIR12	n.d.	11	18 (30.0)	14	164-216	4.78	0.38	0.75	0.47	0.77
CIR15	0.33	4	27 (45.0)	14	232-260	3.76	0.35	0.80	0.48	0.82
CIR17	0.67	35	7 (11.7)	5	271-289	3.35	0.63	0.39	0.17	0.42
CIR18	n.d.	14	17 (28.3)	9	331-355	3.50	0.39	0.73	0.37	0.76
CIR22	n.d.	28	12 (20.0)	8	273-291	3.06	0.50	0.57	0.37	0.60
CIR24	n.d.	33	11 (18.3)	7	186-204	2.99	0.55	0.49	0.28	0.53
CIR26	0.10	15	12 (20.0)	8	272-308	4.33	0.40	0.71	0.38	0.75
CIR29	0.04	21	15 (25.0)	9	159-187	3.57	0.42	0.68	0.42	0.71
CIR30	0.08	18	10 (16.7)	5	172-186	2.80	0.41	0.69	0.33	0.74
CIR33	0.50	3	25 (41.7)	15	273-347	5.54	0.35	0.81	0.43	0.82
CIR37	0.06	6	25 (41.7)	14	134-178	4.54	0.36	0.78	0.45	0.80
CIR40	0.32	16	21 (35.0)	12	258-296	4.04	0.41	0.71	0.43	0.74
CIR42	0.06	5	20 (33.3)	11	202-238	4.39	0.35	0.80	0.57	0.82
CIR43	0.00	8	17 (28.3)	8	202-216	2.48	0.38	0.75	0.42	0.78
CIR45	0.52	36	8 (13.3)	4	288-294	1.20	0.64	0.37	0.08	0.41
CIR55	0.33	34	5 (8.3)	3	240-252	2.89	0.60	0.40	0.30	0.47
CIR56	0.50	22	14 (23.3)	10	314-364	5.00	0.43	0.67	0.15	0.71
CIR57	0.00	24	10 (16.7)	5	247-257	2.00	0.46	0.62	0.38	0.67
CIR58	0.13	7	22 (36.7)	15	208-324	6.33	0.38	0.76	0.38	0.78
CIR60	0.11	10	18 (30.0)	10	189-215	3.84	0.38	0.75	0.42	0.78
CIR184	0.21	20	16 (26.7)	8	117-147	4.21	0.42	0.68	0.35	0.72
CIR210	0.46	26	10 (16.7)	7	138-152	2.58	0.47	0.60	0.35	0.65
CIR229	0.08	27	16 (25.0)	8	309-325	2.66	0.47	0.60	0.37	0.64
CIR243	0.00	9	16 (26.7)	7	125-141	2.81	0.38	0.75	0.33	0.78
CIR244	0.19	2	21 (35.0)	13	240-270	3.85	0.34	0.82	0.47	0.84
CIR274	0.22	19	21 (33.3)	11	186-224	4.28	0.42	0.70	0.47	0.72
CIR278	0.08	37	5 (8.3)	4	98-118	3.83	0.65	0.34	0.25	0.42
S012	0.07	32	9 (15.0)	6	264-285	3.44	0.53	0.52	0.28	0.55
S016	0.20	30	8 (13.3)	5	201-221	3.44	0.51	0.52	0.30	0.60
Average $\pm$ Sem			. /	$8.8{\pm}0.6$		$3.63 {\pm} 0.17$	$0.44{\pm}0.01$	$0.65 {\pm} 0.02$	$0.35 {\pm} 0.02$	$0.69{\pm}0.02$

<sup>1</sup> Microsatellite code, additional information in Table 8; <sup>2</sup> Allele Drop Out, n.d. = not determined; <sup>3</sup> Rank; <sup>4</sup> Separation ability; <sup>5</sup> Number of alleles;  ${}^{6}S^{2} = \sum_{i \neq j} (x_{i} - x_{j})^{2}/2n(n-1)$  where n = number of alleles

<sup>7</sup> Probability of identity of siblings [74]

<sup>8</sup> Polymorphism information content [8]; <sup>9</sup> Observed Heterozygosity; <sup>10</sup> Expected Heterozygosity;

N<sub>a</sub>, range, PIC and H<sub>e</sub> obtained from PowerMarker v3.25 [39]

ADO, P(ID)sib and separation ability obtained from GIMLET v1.3.3 [71]

Successful and efficient verification of identities in a germplasm collection relies on the judicious use of microsatellite loci. The loci chosen must be able to differentiate among existing and future accession holdings. Furthermore, loci should be used that would maximise differences among accessions. Microsatellite structure is known to affect polymorphism. Microsatellites with high tandem repeat numbers had more alleles [49] and higher mutation rates [59]. In quinoa (*Chenopodium quinoa* Willd.), polymorphism in microsatellites was significantly affected by tandem repeat length, motif type and the interaction of these two parameters [42]. Although a small dataset of microsatellites was utilised in the present study, our finding, that complexity and presence or absence of interruption of the repeat did not significantly influence microsatellite variability, agreed with Mason et al. [42].

Microsatellites with many alleles per locus would favour unique fingerprinting of a large number of accessions by Fig. 1 Probability of identity among relatives ( $P_{(ID)sib}$ ) as a function of allele number. Thirty-seven loci in 244 differing combinations were assessed on sixty *Theobroma cacao* L. accessions



relatively few loci [43]. Abu Assar et al. [1] found that 16 loci with 117 polymorphic bands separated each of 96 sorghum *(Sorghum bicolour (L.) Moench)* accessions. Khadari et al. [33] found that 18 genotype pairs of 75 fig *(Ficus carica L.)* accessions were unresolved when 38 SSR alleles from six loci were utilised. Aranzana et al. [2] obtained at least 87% individual identification with 113 alleles from 16 loci on 212 peach *(Prunus persica (L.) Batsch)* cultivars. These results and the present study confirm that the number of alleles is a good discriminatory factor for individual resolution. In addition, our findings indicated that while a high probability value was strongly correlated with increased allele number this was not the

exclusive factor in separating accessions from each other. The combination of loci chosen was also important which agreed with that found for closely related bread wheat cultivars [40].

Results of the primer survey revealed that the 15 primers recommended by Saunders et al. [58] were good discriminatory loci and that eight of these (mTcCIR 11, 12, 15, 18, 26, 33, 37 and 60) were the most useful. However, this set of 15 primers was not the best set to detect mislabelled individuals in the ICG,T where each germplasm group is represented by several families, as exemplified by the reduced ability to resolve NA genotypes. Risterucci et al. [56] suggested, from a small study of 20 genotypes and 19





Table 2 Allelic range and resolution ability of primer sets

Number of primer sets	Range of number of primers used	<sup>1</sup> Resolution (%)	Allele range
16	3–12	75.9	45–90
19	3-7	77.8	36-55
16	3–7	79.6	37–48
14	4-12	83.3	42-117
13	5-12	85.2	47-120
9	5-15	87.0	42-152
39	9–34	94.4	101-314
46	9–37	100	101-326

<sup>1</sup>Groups resolved among sixty *Theobroma cacao* L. accessions as a relative percentage of that obtained with the full complement of 37 loci used in this study.

loci, that 15 loci would be sufficient for cacao fingerprinting. Swanson et al. [66] revised this downwards to 11 loci. Recently, Zhang et al. [77] suggested that seven of the fifteen loci proposed by Saunders et al. [58] were sufficient for cacao fingerprinting. We demonstrated that a substantially reduced subset of cacao microsatellite loci (and number of alleles) of nine recommended loci (mTcCIR12, 15, 26, 33, 37, 42, 57, 243 and 244) would give speedier and more definitive resolution of identities in cacao germplasm collections.

However, an increased (less favourable)  $P_{(ID)sib}COM$  of  $1.437 \times 10^{-4}$  was obtained in comparison to the set of fifteen

recommended by Saunders et al. [58]. The random match probability ( $p_x$ ) for unrelated individuals can be found from  $p_x \le 1$ - $(1-\alpha)^{1/N}$  where  $\alpha$  is the confidence level and N the population size [10]. Wild cacao trees in Belize were found to have a canopy diameter of up to 12 m [47] and the cacao Amazonian area was estimated as 7,700,000 km<sup>2</sup> [7] giving a conservative *T. cacao* population size of  $6.81 \times 10^{10}$ . Setting  $\alpha = 0.01$ ,  $N = 6.81 \times 10^{10}$  gives  $p_x$  of  $1.48 \times 10^{-13}$  which agreed well with the combined P<sub>(ID)</sub> of  $8.886 \times 10^{-12}$  from the recommended set of nine loci. Accessions declared as similar under these circumstances may be considered to be genetically similar and unless morphological or agronomic evidence indicates otherwise may be lumped together as the same accession.

Increasing throughput by bulking was inconvenient due to the non-detection of alleles, an effect that was dependent on both the primer and number of pooled samples. A similar effect was theoretically modelled [78]. The use of pooled samples for determination of mislabelled plants in large-scale verification work was further limited by the degree of mislabelling. A lower efficiency results when there is a high degree of mislabelling [23] due to additional genotyping since the pools must be decomposed into individual samples and re-assayed. Furthermore, these workers demonstrated that an error rate of 20% results in a pooling strategy being less efficient than genotyping individual samples. This study found a high level (59%) of plots in the ICG,T that



Fig. 3 Comparison of resolution ability with allele number in ten primer combinations. Resolution ability was relative to that obtained with all (37) loci on sixty *Theobroma cacao* L. accessions. Top and bottom loci are as ranked with GIMLET v.1.3.3 [71]. Saunders' 15 is

the set recommended by Saunders et al. [58]. The set "Most Alleles" consists of three loci with 15 alleles each. Rec9 is the recommended set of nine primers from this study

Fig. 4 Genetic relationships among sixty *Theobroma cacao* L. accessions with 37 microsatellite loci. Criollo (C), Forastero (F), Refractario (R) and Trinitario (T) groups are indicated



allegedly contained replicated clonal material but instead contained more than one genotype. A misidentification level of around 30% was reported for cacao germplasm collections [51, 20, 56] including the ICG,T [64]. Hence,

Table 3 Correlation of scorer assessment for three microsatellites

Microsatellite	Spearman's r <sub>s</sub>
mTcCIR15	0.641 <sup>a</sup>
mTcCIR26	$0.849^{a}$
mTcCIR37	0.947 <sup>a</sup>

<sup>a</sup> significant at 0.1%

the use of pooled samples for verification work in the ICG, T is not recommended.

The unrooted NJ dendrogram based on 37 microsatellite loci clustered the 60 individuals within distinct groups roughly according to their genetic origin and hidden genealogical relationship. All of the Criollo clones formed a tight cluster at the bottom of the dendrogram and were close to the Trinitario ICS clones. The Trinitario accessions were joined by the "Refractario" accessions (e.g. JA, B, MOQ) from Ecuador. The French Guiana accessions (YAL, GU and ELP clones) formed a small tight group associated with several Upper Amazon Forastero clones. It indicated

Table 4	ANOVA	analyses	for pooled	DNA	samples
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Analysis	Degrees of freedom	F-ratio
% Total Alleles Detected		
Microsatellite	2	11.13 <sup>a</sup>
DNA pool	2	4.28 <sup>n.s.</sup>
Scorer	1	$0.00^{n.s.}$
Microsatellite × DNA pool	4	$0.37^{n.s.}$
Microsatellite × Scorer	2	0.41 <sup>n.s.</sup>
DNA pool × Scorer	2	0.03 <sup>n.s.</sup>
Microsatellite × DNA pool × Scorer	4	$1.00^{n.s.}$
% Correct Alleles of Detected Peaks		
Microsatellite	2	5.44 <sup>n.s.</sup>
DNA pool	2	$1.02^{n.s.}$
Scorer	1	0.14 <sup>n.s.</sup>
Microsatellite × DNA pool	4	2.87 <sup>n.s.</sup>
Microsatellite × Scorer	2	2.09 <sup>n.s.</sup>
DNA pool × Scorer	2	0.21 <sup>n.s.</sup>
Microsatellite × DNA pool × Scorer	4	$1.00^{n.s.}$

<sup>a</sup> Significant at the 5% level

the likely origin of these French Guiana clones from Upper Amazon. All of the twenty Nanay clones were grouped together (Fig. 4). Those individuals that formed small clusters were likely siblings derived from same trees (Zhang et al., unpublished data).

In conclusion, nine discriminant *T. cacao* microsatellite loci (mTcCIR12, 15, 26, 33, 37, 42, 57, 243 and 244) were identified for fingerprinting cacao germplasm collections. These primers are recommended for the detection of mislabelling within collections by assessing individual trees.

# Methods

Plant Material, DNA Extraction and Quantification

Cacao (*T. cacao* L.) leaf tissue was collected from accessions in the ICG,T. DNA was extracted using the Kobayashi protocol [34]. Leaf samples (~0.1 g as thin strips) were placed in Lysing matrix A tubes (MP Biomedicals, Solon, Ohio, USA) with 500  $\mu$ L of Buffer I and maceration was effected in a FastPrep 120V machine (Qbiogene, Inc., California, USA) at speed = 4, *t*=20 sec, thrice for each sample. Precipitated DNA was re-suspended in sterile deionised water (SDW) and kept as stock solutions. Dilutions (×100 or as required) of the stock DNA solutions were prepared with SDW and assayed with PicoGreen<sup>®</sup> (Molecular Probes, Eugene, Oregon, USA) in a Fluoroskan Ascent (Labsystems, Finland). Two samples (H1 and U1), extracted using a Plant Mini Kit (Qiagen, Valencia, California, USA), were obtained from a Peruvian

 Table 5 Plot homogeneity of accessions in the International Cocoa
 Genebank, Trinidad

Accession	UCRS Plot	# Trees	# Trees	# Genotypes
	location	in piot	studied	
AM 1/19	Field 5B, 1771	8	8	1
AM 1/28	Field 6A, A1	8	7	1
AM 1/53	Field 6A, A2	7	5	1
AM 1/54	Field 5B, 1811	11	5	2
AM 1/60	Field 5A, A26	3	3	1
AM 1/70	Field 4A, F549	2	2	1
AM 1/85	Field 4A, F538	3	2	1
AM 2/12	Field 5B, B95	4	4	1
AM 2/18	Field 5B, H679	2	2	1
AM 2/61	Field 5B, H716	3	2	1
AM 2/62	Field 5B, B105	13	9	3
AM 2/65	Field 5B, I810	8	6	2
AM 2/82	Field 5B, 1806	4	4	1
AM 2/83	Field 5B, B108	15	9	1
AM 2/96	Field 5B, 1819	8	3	2
B 12/1	Field 6B, F461	9	9	4
B 13/7	Field 5B, 1728	12	11	2
B 17/17	Field 5B, 1784	10	10	2
B 18/4	Field 6B, F457	14	11	3
B 4/8	Field 6B, F439	5	3	3
B 7/21	Field 6B. F438	9	8	7
CL 10/5	Field 5B, A4	4	4	2
CL 10/14	Field 5A, A1	11	7	4
CL 13/27	Field 5B, A24	9	8	2
CL 27/50	Field 5B, 1774D	12	9	1
CL 91/5	Field 5B, A64	2	2	2
CL 9/17	Field 5B, A24	12	12	4
CRUZ 7/8	Field 6B, B83	6	3	2
DOM 27	Field 4A, B203	2	2	2
ICA 70	Field 4A, C290	3	3	1
JA 1/9	Field 6A, A51	3	3	1
JA 4/17	Field 5B, E425	4	2	1
JA 5/27	Field 5B, F483	6	5	5
JA 5/39	Field 5B, D234	14	11	2
JA 8/33	Field 5B, E378	3	2	1
JA 10/16	Field 5B, E411	2	2	1
LP 1/21	Field 5B, 1746	4	4	2
LP 1/21	Field 5B, 1779	5	4	3
LP 3/4	Field 5B, A33	16	14	4
LP 4/12	Field 5B, 1803	10	12	1
LP 4/48	Field 5B, B140	10	9	8
LP 5/19	Field 6A, B95	3	3	1
LX 38	Field 5B, C206	8	7	4
LX 43	Field 5B, C201	16	14	3
MOO 6/95	Field 5B, C221	5	3	3
NA 176	Field 4A, D389	3	3	1
NA 669	Field 4A, D418	4	3	3
PA 169	Field 6B, C180	11	8	5
PA 293	Field 4A F516	4	3	2
SLA 16	Field 5B D242	8	6	2
SLC 4	Field 5B A39	6	5	1
SLC 18	Field 5B A13	9	5	2
TRD 15	Field 4A A43	2	2	2
TRD 111	Field 4A A87	3	3	- 1
		2	2	-



**Fig. 5** Plot homogeneity assessment in the International Cocoa Genebank, Trinidad. Forty-four plots with genotype data from six microsatellite loci on at least three trees were evaluated with GIMLET v.1.3.3 [71]. Plot homogeneity assessed by the number (and percentage) of genotypes within a plot that should contain only one genotype

collection held at the Sustainable Perennial Crops Laboratory of the Agricultural Research Service (United States Department of Agriculture; USDA-ARS) in Beltsville, Maryland. The accessions utilised (Table 6) contained representatives of Criollo, Forastero, Refractario and Trinitario material.

#### Polymerase Chain Reaction (PCR) Process

The PCR reaction mix was composed of 4 µL Eppendorf HotMaster Mix (Brinkmann Instruments Inc., New York, USA) giving 2.5 mM magnesium ions, 2 mM total deoxynucleotide triphosphates, 0.2 Units Taq polymerase at final composition in the reaction mix; 0.5 µL of a MPP solution in Tris-EDTA buffer (10 µM each primer; reverse unlabeled primers from Operon Technologies, Inc., Alabama, USA; forward, WellRed-labelled primers from Proligo, Boulder, Colorado, USA); and 5.5  $\mu$ L of 0.05 ng  $\mu$ L<sup>-1</sup> appropriate DNA solution. Each MPP was amplified from separate reaction mixes. Cycling was carried out in a GeneAmp PCR System 9,700 thermal cyclers (Applied Biosystems, Foster City, California, USA) with a touchdown protocol: 94°C, 5 min; eight cycles with denaturation at 95°C for 30 sec, annealing at 55°C for 60 sec with reduction by 0.5°C after every cycle and extension at 72°C for 1 min; 25 cycles with denaturation at 94°C for 30 sec, annealing at 51°C for 1 min and extension at 72°C for 1 min; final extension at 60°C for 15 min to ensure complete A addition followed by retention at 4°C until recovery. Each combination of DNA-MPP was repeated at least once. Several combinations were identified for at least three PCR determinations based principally on the need for confirmation of allele sizes.

# Capillary Electrophoresis (CE)

Post-PCR bulking was conducted by pooling 1.5 µL of each MPP-PCR product. Sample loading buffer containing 29.8 µL of Hi-Di formamide (Applied Biosystems, Warrington, UK) and 0.2 µL of GenomeLab<sup>™</sup> DNA size standard-400 (Beckman Coulter Inc., Fullerton, California, USA) was added to each well. Samples were overlaid with one drop of mineral oil (Beckman Coulter Inc.). Fragments were separated on an 8-capillary CEQ<sup>™</sup> 8,000 or 8,800 (Beckmann Coulter Inc.). Products with poor standard profiles (missing bands; improper sizing) were discarded and the appropriate PCR product pools were recomposed and run again to ensure that fragment profiling was suitable for allele sizing. Preliminary allele binning using a 1.5 nucleotide window was performed with the bundled CEQ<sup>™</sup> fragment analysis software (Beckman Coulter Inc.). Allele sizes were assessed graphically and manually to remove spurious and stutter alleles. Final allele bins were decided by repeat number using the deposited sequence length as the starting point for stepwise increase or decrease in repeat number.

#### MPP Assessment

The fifteen recommended MPPs [58] and twenty-two additional MPPS (Table 7) were assessed on a set of 60 cacao accessions (Table 6). PCR and CE were repeated at least once as described above. Allele binning was as described earlier. Summary statistics including the polymorphism information content (PIC; [8]) were obtained with PowerMarker v3.25 [39]. The probability of identity among full siblings ( $P_{(ID)sib}$ ; [74]) from each SSR was obtained with GIMLET v.1.3.3 [71]. The statistic ln S<sup>2</sup> was also calculated where

$$S^{2} = \sum_{i \neq j} (x_{i} - x_{j})^{2} / 2n(n-1)$$
  $n =$  number of alleles

The error level of each locus (allele drop out, ADO; false allele) was determined with GIMLET v.1.3.3 [71] from samples with at least three separate PCR and corresponding CE determinations. The effect of microsatellite structure (perfect vs. imperfect; compound vs. simple) on ln S<sup>2</sup>, PIC,  $P_{(ID)sib}$ , and number of alleles for SSRs classified as dimers, was analysed by ANOVA using IRRISTAT v.5.0 [27].

 Table 6
 Cacao accessions used for determination of best microsatellites for verification studies in field germplasm collections

Accession	<sup>1</sup> Position	Group	Country of origin (Status)
AC 2 [BLZ]	T1 (FP1026)	Criollo	Belize (wild)
AC 20 [BLZ]	T1 (FP1032)	Criollo	Belize (wild)
B 9/10-25 [POU]	Marper Farm, C1078	Refractario	Ecuador (cultivated)
BC 3 [BLZ]	T1 (FP1019)	Criollo	Belize (wild)
COCA 3348/44 [CHA]	UCRS, Field 6B, E374 T2 (FP1047)	Forastero	Ecuador (wild)
CRIOLLO 22 [CRI]	UCRS, Field 4A, C276 T3	Criollo	Costa Rica (cultivated)
EET 400 [ECU]	UCRS, Field 6B, F455 T1	Forastero	Ecuador (cultivated)
ELP 1/S6	(FP950)	Forastero	French Guiana (wild)
GU 241/P	UWI, Campus Field 1A, x2y33 (FP500)	Forastero	French Guiana (wild)
GU 300/P	UCRS, Field 4A, B197 T4	Forastero	French Guiana (wild)
H 1	Not available	Forastero	Peru (cultivated)
HF 8 [BLZ]	T1 (FP987)	Criollo	Belize (wild)
IB 2 [BLZ]	T1 (FP1020)	Criollo	Belize (wild)
IB 9 [BLZ]	T1 (FP996)	Criollo	Belize (wild)
ICS 75	San Juan Estate Block 2	Trinitario	Trinidad (cultivated)
ICS 97	San Juan Estate Block 1	Trinitario	Trinidad (cultivated)
ICS 100	San Juan Estate Block 2	Trinitario	Trinidad (cultivated)
IMC 3	UWI, Campus Field 3, x1y3	Forastero	Peru (wild)
IMC 12	Marper Farm, C1056	Forastero	Peru (wild)
IMC 16	Marper Farm, D603	Forastero	Peru (wild)
IMC 67	La Reunion Estate	Forastero	Peru (wild)
JA 5/4 [POU]	Marper Farm, C526 (FP2307)	Refractario	Ecuador (cultivated)
JA 5/5 [POU]	Marper Farm, C324 (FP1351)	Refractario	Ecuador (cultivated)
LCT EEN 31	UCRS, Field 6A, A6 T3 (FP450)	Forastero	Ecuador (wild)
LCT EEN 162/S-1010	UCRS, Field 5B, C216 T2 (FP2945)	Forastero	Ecuador
MO 9	Marper Farm, D835 (FP253)	Forastero	Peru (wild)
MO 20	Marper Farm, D809 (FP254)	Forastero	Peru (wild)
MOO 6/95	Marper Farm, C1 (FP582)	Refractario	Ecuador (cultivated)
MXC 67	UWL Campus Field 12, x3v6	Criollo	Mexico (cultivated)
NA 184	UCRS, Field 5B, G612 T1	Forastero	Peru (wild)
NA 241	UCRS, Field 4A, D383 T4 (FP2716)	Forastero	Peru (wild)
NA 244	UCRS, Field 5B, E400 T3 (EP16)	Forastero	Peru (wild)
NA 266	UCRS, Field 5B, G634 T3 (FP25)	Forastero	Peru (wild)
NA 331	Marper Farm, $D477$ (FP383)	Forastero	Peru (wild)
NA 406	UCRS, Field 5B, F447 T1 (FP23)	Forastero	Peru (wild)
NA 432	Marper Farm, $D717$ (FP271)	Forastero	Peru (wild)
NA 435	Marper Farm, D760 (FP260)	Forastero	Peru (wild)
NA 504	Marper Farm, $D465$ (FP167)	Forastero	Peru (wild)
NA 528	Marper Farm, D774 (FP112)	Forastero	Peru (wild)
NA 680	UCRS. Field 5A, $D337$ T3 (FP649)	Forastero	Peru (wild)
NA 702	Marper Farm, $D104$ (FP819)	Forastero	Peru (wild)
NA 705	Marper Farm, C102 (FP1280)	Forastero	Peru (wild)
NA 733	Marper Farm D721 (FP274)	Forastero	Peru (wild)
NA 734	Marper Farm D546 (FP377)	Forastero	Peru (wild)
NA 771	UCRS Field 5B F478 T4 (FP27)	Forastero	Peru (wild)
NA 773	UCRS Field 5B F547 T3 (FP1266)	Forastero	Peru (wild)
NA 831	Marner Farm $D741$ (FP267)	Forastero	Peru (wild)
NA 833	Marper Farm $D640$ (FP297)	Forastero	Peru (wild)
NA 860	Marper Farm $D240$ (FP1167)	Forastero	Peru (wild)
NAPO 2 [CHA]	LIWI Campus Field 7 x8v9 (FP1922)	Forastero	Feuador (wild)
PA 279 [PER]	Marner Farm $D50$ (FP426)	Foractero	Peru (wild)
PA 299 [PER]	Marper Farm $C936$ (FP571)	Forastero	Peru (wild)
POR 1 [TTO]	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Criollo	Venezuela
	UCRS Field 6R E $/07$ T2 (ED521)	Eorastoro	Port (wild)
SCA 12	Marror Form $D205$	Forastero	reiu (wiid) Dom (wild)
SCA 12	Marper Farm D560	Forestore	r ciu (wiid) Domi (wild)
SUA 24	waiper raini, D309	rotastero	reiu (wiiu)

# Table 6 (continued)

Accession	<sup>1</sup> Position	Group	Country of origin (Status)
SPA 5 [COL]	UWI, Campus Field 2, x1y15 (FP1817)	Forastero	Colombia or Peru
U 1	Not Available	Forastero	Peru (cultivated)
UF 613	UCRS, Field 4A, A93 T2 (FP1237)	Trinitario	Costa Rica (cultivated)
YAL 6	Not Available	Forastero	French Guiana

 $^{1}$  FP = fingerprinting number; numeric code given to sampled reference trees

Table 7 Characteristics of the microsatellite primers for Theobroma cacao L. utilised in this study

$MPP^1$	EMBL no. <sup>2</sup>	5'-3' Forward primer	5'-3' Reverse primer	$LG^3$	ASL <sup>4</sup> (bp)	Repeat motif	Clsn <sup>5</sup>
mTcCIR1 <sup>a</sup>	Y16883	GCAGGGCAGGC	TGGGCAACCA	8	143	(CT) <sub>14</sub>	DI, P,S
		TCAGTGAAGCA	GAAAACGAT				
mTcCIR3	Y16977	CATCCCAGTATCT	CTGCTCATTTC	2	249	$(CT)_{20}(TA)_{21}$	DI, P,C
		CATCCATTCAGT	TTTCATATCA				
mTcCIR6 <sup>a</sup>	Y16980	TTCCCTCTAAA CTACCCTAAAT	TAAAGCAAAGC	6	231	$(TG)_7(GA)_{13}$	DI, P,C
			AATCTAACATA				
mTcCIR7 <sup>a</sup>	Y16981	ATGCGAATG ACAACTGGT	GCTTTCAGT CCTTTGCTT	7	160	(GA) <sub>11</sub>	DI, P,S
mTcCIR8 <sup>a</sup>	Y16982	CTAGTTTCC CATTTACCA	TCCTCAGCA TTTTCTTTC	9	301	$(TC)_5TT(TC)_{17}$ T <sub>3</sub> (CT) <sub>4</sub>	DI, I,C
mTcCIR9	Y16983	ACCATGCTT CCTCCTTCA	ACATTTATA CCCCAACCA	6	274	(CT) <sub>8</sub> N <sub>15</sub> (CT) <sub>5</sub> N <sub>9</sub> (TC) <sub>10</sub>	DI, I,C
mTcCIR10	Y16984	ACAGATGGC CTACACACT	CAAGCAAGC CTCATACTC	5	208	(TG) <sub>13</sub>	DI, P,S
mTcCIR11 <sup>a</sup>	Y16985	TTTGGTGAT TATTAGCAG	GATTCGATT TGATGTGAG	2	298	(TC) <sub>13</sub>	DI, P,S
mTcCIR12 <sup>a</sup>	Y16986	TCTGACCCC AAACCTGTA	ATTCCAGTT AAAGCACAT	4	188	(CATA) <sub>4</sub> N <sub>18</sub> (TG) <sub>6</sub>	DI, I,C
mTcCIR15 <sup>a</sup>	Y16988	CAGCCGCC TCTTGTTAG	TATTTGGGA TTCTTGATG	1	254	(TC) <sub>19</sub>	DI, P,S
mTcCIR17	Y16990	AAGGATGAAGG ATGTAAGAGAG	CCCATACGA GCTGTGAGT	4	271	$(GT)_7N_4(GA)_{12}$	DI, I,C
mTcCIR18 <sup>a</sup>	Y16991	GATAGCTAAG GGGATTGAGGA	GGTAATTCAAT CATTTGAGGATA	4	345	(GA) <sub>12</sub>	DI, P,S
mTcCIR22 <sup>a</sup>	Y16995	ATTCTCGCAA AAACTTAG	GATGGAAGGA GTGTAAATAG	1	289	$(TC)_{12}N_{146}$ $(CT)_{10}$	DI, I,C
mTcCIR24 <sup>a</sup>	Y16996	TTTGGGGTGA TTTCTTCTGA	TCTGTCTCGTC TTTTGGTGA	9	198	$(AG)_{13}$	DI,
P.S							
mTcCIR26 <sup>a</sup>	Y16998	GCATTCATC AATACATTC	GCACTCAAA GTTCATACTAC	8	298	(TC) <sub>9</sub> C(CT) <sub>4</sub> TT(CT) <sub>11</sub>	DI, I,C
mTcCIR29	AJ271822	CGACATTTCG ACTTTCATC	TTTTGTTTCTT TCTTTTTCATT	1	173	$(CA)_{10}$	DI, P,S
mTcCIR30	AJ271823	TGAAGATCCT ACTGTTGAG	TGATAATAAC TGCTTAGTGG	9	184	(CA) <sub>11</sub>	DI, P,S
mTcCIR33 <sup>a</sup>	AJ271826	TGGGTTGA AGATTTGGT	CAACAATGA AAATAGGCA	4	285	(TG) <sub>11</sub>	DI, P,S
mTcCIR37 <sup>a</sup>	AJ271942	CTGGGTGCT GATAGATAA	AATACCCTC CACACAAAT	10	150	(GT) <sub>15</sub>	DI, P,S
mTcCIR40 <sup>a</sup>	AJ271943	AATCCGACAG TCTTTAATC	CCTAGGCCAG AGAATTGA	3	286	(AC) <sub>15</sub>	DI, P,S
mTcCIR42	AJ271944	TTGCTGAAGT ATCTTTTGAC	GCTCCACC CCTATTTG	5	232	$(CA)_{21}$	DI, P,S
mTcCIR43	AJ271945	TCATGAGA ATGCATGTG	CTGGACATGA AGAAGTTAT	4	206	$(TG)_5TA(GA)_{15}$	DI, I,C
mTcCIR45	AJ271947	GTCATTG CTGTGTG	CATAGCATTA ACTGTGTCTG	8	284	(GT) <sub>9</sub>	DI, P,S
mTcCIR55	AJ271954	GATATTGTCC CATTATTTG	TTCCGCC TTGTTCTC	7	234	(CACACG) <sub>4</sub>	HE. P.S
mTcCIR56	AJ271955	ATACTTTTA CTTCCTTTTG	TCTTATTTT TCTTTACCAG	7	354	(TC)14N(TG)15	DI. I.C
mTcCIR57	AJ271956	TGTAGATGTGA TTTTATAGTTTG	GGAGGGATA AGAAGCAG	4	253	$(AC)_{13}$	DI, P,S

MPP <sup>1</sup>	EMBL no. <sup>2</sup>	5'-3' Forward primer	5'-3' Reverse primer	LG <sup>3</sup>	ASL <sup>4</sup> (bp)	Repeat motif	Clsn <sup>5</sup>
mTcCIR58	AJ271957	TTTTTGGTGA TGGAACTAT	TGGTTAAGCA	9	266	(GT) <sub>40</sub>	DI, P,S
mTcCIR60 <sup>a</sup>	AJ271958	CGCTACTAAC AAACATCAAA	AGAGCAACCA TCACTAATCA	2	207	(CT) <sub>7</sub> (CA) <sub>20</sub>	DI, P,C
mTcCIR184	AJ566512	GGTTTTCTA GCTCCTCC	AGGAAAGAAT GACTCATACTA	1	139	$(CA)_8(CT)_{13}$	DI, P,C
mTcCIR210	AJ566533	CAAACCCCA AACCTCAA	CAGTTATGGAA ATTATTGCTCTA	1	146	(AG) <sub>11</sub> N <sub>7</sub> (AAG) <sub>4</sub>	DI, I,C
mTcCIR229	AJ566550	ATCTCGGTAA TAGCACATAA	CGCAATCC TACAACACA	10	307	$(TC)_8$	DI, P,S
mTcCIR243	AJ566564	ACAGCAGTA GACGCATTC	AAAAGGCT TGGCACAG	4	141	(TC) <sub>9</sub> N <sub>20</sub> (CA) <sub>11</sub>	DI, I,C
mTcCIR244	AJ566565	TGGCAATAA CAATGAACA	ATTTTGATGA TTGATGAAGA	1	264	$(TA)_4CATA$ $(CA)_{17}(TA)_4$	DI, I,C
mTcCIR274	AJ566593	GAAAGGTAA ATGGCTGAA	CGATCATCA CGACTGCT	5	184	$(CT)_6CACG$ $(CA)_6(CT)_2$	DI, I,C
mTcCIR278	AJ566597	TGGCATCT GTCTGTC	GTATATGAC CGTTTGTAG	3	100	$(TCTG)_3$ $(TC)_{10}(TA)_8$	DI, P,C
S012	AY389500	CCACCACCCTT ACCTTTGAAAC	GGGAAAGGGAAA GGCTGACATC	5	285	$(CAA)_{11}$	TR, P,S
S016	AY389503	GGCCCCTAGCA AAGAAAACC	TGTGCGAAGAC GCAATCTAAC	7	216	(CCTTT) <sub>6</sub>	PE, P,S

<sup>1</sup> Microsatellite primer pair entries followed by <sup>a</sup> are recommended by Saunders et al. [58]; <sup>2</sup> European Molecular Biology Laboratory code for deposited microsatellite; <sup>3</sup> linkage group; <sup>4</sup> published amplified sequence length; <sup>5</sup> classification (C = compound, I = imperfect, P = perfect, S = simple, DI = dimer, HE = hexamer, PE = pentamer, TR = trimer)

Varying combinations of MPPs (244 sets) containing 3-326 alleles were prepared and included single primers; two random set of six primers; three sets of least informative loci containing nine, 12, or 15 primers; seven sets each containing nine primers that were most promising; combinations (three, six, nine, twelve, fifteen primers) from within the set of 15 primers recommended by Saunders et al. [58]; and combinations selected to obtain 40-330 alleles in interval classes of five alleles. Twelve of these classes (61-65, 76-80, 81-85, 126-130, 136-140, 146-150, 166-170, 206-210, 216-220, 226-230, 246-250 and 291-295) were not represented. Allelic datasets were analysed with GIMLET v.1.3.3 [71]. The overall P(ID)sib (P(ID)sibCOM) was log-transformed and assessed by regression analysis, with the total number of alleles as the independent variable. The separation success of each of the 244 MPP sets as a function of the separation ability of the full complement of the 37 MPPs was calculated. Percentage data and allele number were subjected to arcsine and natural log transformation respectively before regression analysis. The 244 datasets were examined for the minimal combination of loci that would give resolution identical to the full complement of 37 loci.

The program PAUP\* (Version 4.0; Swofford[67]) was used for a neighbor-joining analysis of Euclidean distance matrices from all primers and to calculate bootstrap consensus results with 100 replications. The majority-rule

Table 7 (continued)

consensus tree was displayed and printed using TreeView v1.6.6 [54].

#### DNA Pooling Assessment

Three MPPs (mTcCIR15, mTcCIR26 and mTcCIR37) were chosen for amplification and post-PCR bulking for capillary

Table 8 Composition of pooled samples

Pool	DNA <sup>1</sup>	Pool	DNA	Pool	DNA
1	AB	14	DF	27	BEF
2	AC	15	EF	28	CDE
3	AD	16	ABC	29	CDF
4	AE	17	ABD	30	CEF
5	AF	18	ABE	31	DEF
6	BC	19	ABF	32	ABCD
7	BD	20	ACF	33	ABCE
8	BE	21	ADF	34	ABCF
9	BF	22	AEF	35	ACDE
10	CD	23	BCD	36	ACDF
11	CE	24	BCE	37	ACEF
12	CF	25	BCF	38	BCDE
13	DE	26	BDF	39	BCDF

 $^{1}$  A = B 9/10–25 [POU], Marper Farm, C1078; B = H 1, C = PA 279 [PER], Marper Farm, D59; D = SCA 12, Marper Farm, D205; E = SPA 5 [COL] University of the West Indies Campus Field 2, x1y15; F = U 1.

electrophoresis. The characteristics of these primers are provided in Table 7. Pooling was composed of two, three or four samples as outlined in Table 8. Each accession contributed equivalently to the final DNA quantity in each pool. MPP-PCR and CE were conducted as before. Separation patterns were examined for allele pattern and two independent scorers assessed the resultant electrophoretograms. Spearman's r<sub>s</sub> for scorer consistency, was determined using the freeware program PAST [24]. Scorer accuracy was then assessed by comparing the number of called alleles to the expected based on individual contributions to each pool. Band intensity (as measured by peak height) was not scored as this parameter is influenced by numerous factors including chance uptake of PCR products by capillaries. Appropriate quantification of this parameter would require additional multiple CEs from the same PCR product and hence would increase the number of runs to be done for any DNA-MPP combination. The possibility of overlooking heterogeneity is recognised for cases when only two peaks are obtained. The experimenter cannot determine whether this is a result of a heterozygous genotype at this locus or whether it originated from a mixed set of homozygous genotypes. The contribution of Type I (real allele not identified) and Type II (false allele identified) error effects was assessed. Analysis of variance (ANOVA) analyses were performed with the freeware program IRRISTAT v5.0 [27] for Type II error, and on arcsine-transformed data for the percentage of total alleles and the percentage of correctly identified alleles detected by scorer.

# Plot Homogeneity Assessment in the ICG,T

Fifty-four plots (53 accessions) containing at least two trees were assessed with six MPPs (mTcCIR1, mTcCIR6, mTcCIR7, mTcCIR8, mTcCIR33 and mTcCIR60). Trees with missing data were excluded from subsequent analysis. Genotype data were analysed with GIMLET v.1.3.3 [71] for individual plot homogeneity using the regroup option.

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