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Discovery and mapping of a new expressed sequence tag-single nucleotide polymorphism and simple sequence repeat panel for large-scale genetic studies and breeding of *Theobroma cacao* L.

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**Abstract**

*Theobroma cacao* is an economically important tree of several tropical countries. Its genetic improvement is essential to provide protection against major diseases and improve chocolate quality. We discovered and mapped new expressed sequence tag-single nucleotide polymorphism (EST-SNP) and simple sequence repeat (SSR) markers and constructed a high-density genetic map. By screening 149 650 ESTs, 5246 SNPs were detected *in silico*, of which 1536 corresponded to genes with a putative function, while 851 had a clear polymorphic pattern across a collection of genetic resources. In addition, 409 new SSR markers were detected on the Criollo genome. Lastly, 681 new EST-SNPs and 163 new SSRs were added to the pre-existing 418 co-dominant markers to construct a large consensus genetic map. This high-density map and the set of new genetic markers identified in this study are a milestone in cocoa genomics and for marker-assisted breeding. The data are available at http://tropgenedb.cirad.fr.

**Key words:** *Theobroma cacao*; genetic map; SNP; molecular marker

1. **Introduction**

*Theobroma cacao* L. is a diploid species (2n = 2x = 20) with a small genome ranging in size from 411 to 494 Mb. According to Cheesman, its centre of origin is at the lower eastern equatorial slopes of the Andes.

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The quality of chocolate is another important trait in cocoa breeding, and consumer demand for high-quality chocolate is increasing. A better understanding of the molecular and genetic bases of these traits is a key goal of cocoa genetic research.

High-density genetic maps are essential tools for trait genetic studies. Several molecular marker types have been developed in *T. cacao* in recent decades: restriction fragment length polymorphism (RFLP), microsatellites or simple sequence repeats (SSRs), random amplified polymorphic DNA, amplified fragment length polymorphism and isozymes. Among them, only RFLP, SSR and single-nucleotide polymorphism (SNP) are co-dominant markers, and therefore more powerful for genetic analyses. Compared with RFLP, the advantage of SSR and SNP markers is that they can be revealed using high-throughput technologies with scant amounts of DNA. Semagn et al. made a detailed comparison of the characteristics of each kind of marker. A high-density cocoa linkage map enriched with SSR genomic markers, including only co-dominant markers, was developed by Pugh et al. More recently, that map was supplemented with 114 EST-SSRs.

In recent years, the use of SNP markers has substantially increased in plant genetics such as in *Arabidopsis*, grapevine, wheat, and also a few woody perennial species. SNP is one of the most abundant types of DNA sequence polymorphism and the SNP markers are suitable for large-scale genome analysis using high-throughput automated genotyping techniques. SNPs have been used to construct high-resolution genetic maps or to trace evolution, particularly in the human genome, using large-scale SNP datasets. Knowledge of nucleotide substitution dynamics is an important basis for molecular evolutionary studies, phylogeny reconstruction and natural selection studies. Transitions are generally observed with higher frequencies than transversions. During natural selection, transitions are better tolerated because they generate more likely synonymous mutations in protein-coding sequences than transversions.

Of existing SNP markers, EST-SNPs (i.e. SNPs located within a gene expressed sequence) are of particular interest for studying functional genetic diversity and identifying candidate genes as the functional base of quantitative trait loci (QTLs). EST-SNPs have been developed for numerous plant models such as melon, *Brassica rapa*, barley, poplar, and sugarcane to detect QTLs for many traits and facilitate the selection of resistant and productive plants. In *T. cacao*, a few SNPs were detected in ESTs from expression libraries representing *T. cacao*/*M. perniciosa* interactions.

In our study, we discovered and mapped several hundred EST-SNP markers detected in an exhaustive collection of cocoa ESTs homologous to genes with a known function. These SNP markers were supplemented by 163 new SSR markers to construct a very high-density genetic map suitable for large-scale genetic studies.

### 2. Materials and Methods

#### 2.1. Plant material

SNP polymorphisms were screened in a collection of diverse germplasm representing the major part of the *T. cacao* diversity and two existing mapping populations denominated UPA402 × UF676 and F2.

The collection of diverse germplasm consisted of 249 genotypes from various genetic groups and geographical origins (Table 1). Most of these accessions are maintained at the International Cocoa Genebank (ICG) at the Cocoa Research Unit (CRU), University of the West Indies, Trinidad and Tobago.

The UPA402 × UF676 mapping population consisted of 264 individuals derived from a cross of two unrelated heterozygous tree accessions; UPA402, an Upper Amazon Forastero from Peru, and UF676, a Trinitario (Forastero × Criollo hybrid) selected in Costa Rica. This progeny was maintained by Centre National de Recherche Agronomique (CNRA) in Bingerville and Divo, Côte d’Ivoire. It was used to

| Accession collection group name | Number of genotypes | Geographical origin |
|---------------------------------|---------------------|---------------------|
| AMAR 2 Ecuador                  | 2                   | Ecuador             |
| APA 1 Colombia                  | 1                   | Colombia            |
| Nacional 3 Ecuador              | 3                   | Ecuador             |
| Criollo 14 Mexico-Belize        | 14                  | Mexico-Belize       |
| EBC 4 Colombia                  | 4                   | Colombia            |
| Trinitario 28 Trinidad          | 28                  | Trinidad            |
| GU 12 French Guiana             | 12                  | Peru                |
| IMC 19 Peru                     | 19                  | Peru                |
| LCTEEN 46 Ecuador               | 46                  | Ecuador             |
| MORONA 3 Peru                   | 3                   | Peru                |
| NANAY 49 Peru                   | 49                  | Peru                |
| PARINARI 40 Peru                | 40                  | Peru                |
| POUND 6 Peru                    | 6                   | Peru                |
| SC 5 Colombia                   | 5                   | Colombia            |
| SCAVINA 8 Peru                  | 8                   | Peru                |
| Amelanoado type 3 Brazil        | 3                   | Brazil              |
| SPEC 6 Colombia                 | 6                   | Colombia            |
| Total 249                       |                     |                     |
establish genetic map as a reference in our laboratory.\textsuperscript{4,6,8,9} We mapped the new SSR and SNP markers in this population.

The F2 second progeny of 132 individuals was obtained by selfing a hybrid between two heterozygous parents: Scavina 6, an Upper Amazon Forastero collected in Peru, and ICS1, a Trinitario selected in Trinidad. This progeny was produced by Comissão Executiva do Plano da Lavoura Caçaueira (CEPLAC) at Itabuna, Brazil.

2.1.1. Genotypes used for EST-SNP detection and selection

Most of the ESTs screened for SNPs had been obtained from the contrasting genotypes Scavina 6, an upper Amazon Forastero genotype from Peru and ICS1, a Trinitario selected in Trinidad, a hybrid between a Criollo from Central America and a Forastero from Lower Amazonia of Brazil.

These two genotypes, which represent the three distinct genetic origins, Upper Amazon Forastero, Lower Amazon Forastero, and Criollo, were also the parents of the F2 population from Brazil used to map SNPs.

Eleven other genotypes were involved in the construction of the cDNA libraries and SNP identification: B97-CC2, a Criollo from Belize, P7, IMC47, UPA 134, Upper Amazon Forastero genotypes from Peru, Jaca, an Upper Amazon Forastero from Brazil, GU255V, collected in French Guiana, B240 and 33–49, two Nacional genotypes from Ecuador, UF676, UF273, two Trinitario, and seedlings from a hybrid selected in Papua New Guinea.

SSRs were screened in three \textit{T. cacao} genotypes: the two parents of the reference map (UPA402 and UF676), and the sequenced Criollo genotype (B97-61/B2).

2.2. DNA extraction and purification

Genomic DNA was extracted according to a protocol using MATAB buffer already described for the isolation of genomic DNA.\textsuperscript{6} DNA was resuspended with 1 ml of TE (10 mM Tris–HCl and 1 mM EDTA, pH 8.0).

DNA was purified with the Nucleobond\textsuperscript{TM} PC 20 kit (Macherey-Nagel, Cat. No. 740.571.100) with the modification that steps 1 and 2 were omitted and the DNA was purified directly after its isolation. A 1 ml mixture composed of 200 \mu l of crude DNA (20 \mu g DNA maximum), 450 \mu l water and 350 \mu l S3 buffer + RNAse (buffers provided with the kit) was passed through the column (step 3). This solution was homogenized on a rocking table for at least 1 h. After precipitation of the eluate with an equal volume of isopropyl alcohol, the pellet was resuspended in 70 \mu l of TE.

The quality and quantity of DNA were first checked on 0.8% agarose gel, compared with a standard range, and then the Quant-it\textsuperscript{TM} PicoGreen\textsuperscript{®} dsDNA Assay from Invitrogen\textsuperscript{TM} was used. A quality test was performed for each sample by amplifying microsatellite markers in a PCR mixture with a high DNA concentration (100 ng DNA in a 10 \mu l reaction volume). The purification step was repeated when the amplification failed.

2.3. In silico SNP discovery and verification

A collection of 149 650 ESTs (EMBL accession number CU469588 to CU633156), corresponding to 48 594 unigenes, was produced after sequencing 56 cDNA libraries constructed from material collected from different organs, genotypes, and under different environmental conditions.\textsuperscript{32}

SNPs were detected in silico and quality checked using the QualitySNP pipeline\textsuperscript{33} as reported in Argout et al.\textsuperscript{32} and in ESTtik (http://esttik.cirad.fr).

QualitySNP uses quality information related to each EST and a haplotype-based strategy to predict reliable SNPs. In order to detect SNPs in known homologous coding sequences, we selected contigs displaying a significant similarity with proteins from a non-redundant protein sequence database (NR), with entries from GenPept, Swissprot, PIR, PDF, PDB, and NCBI RefSeq, and as described in Argout et al.\textsuperscript{32}

2.4. Validation of SNPs via golden gate assay

A total of 30–50 ng of genomic DNA per plant was used for Illumina SNP genotyping with the Illumina BeadArray platform at the French National Genotyping Centre (CNG, CEA-IG, Evry, France), according to the GoldenGate Assay manufacturer's protocol. Three 3-day assays were carried out to genotype the progeny samples for the 1536 SNP set revealed by the GoldenGate assay (Supplementary Table S1). The protocol was similar to that briefly described by Hyten et al.\textsuperscript{34} except for the number of oligonucleotides involved in a single DNA reaction, thus comprising 4608 custom oligos assembled in the oligo pooled assays (OPA) designed by Illumina Inc. Raw hybridization, intensity data processing, clustering, and genotype calling were performed using the genotyping module in the BeadStudio/GenomeStudio package (Illumina, San Diego, CA, USA). Illumina has developed a self-normalization algorithm that relies on information contained in each array, as described by Akhunov et al.\textsuperscript{35}

The clustering and genotype calling of each of the 1536 SNP markers were checked for their conformity and correct genotype distribution using known homozygous and heterozygous genotypes, included in the collection of diverse genotypes, as standards.
2.5. **SSR in silico discovery and genotyping**

The MicroSatellite identification tool (MISA http://pgrc.ipk-gatersleben.de/misa) was used to perform SSR searches, and primers were designed with Primer3 software.36

Primers flanking microsatellite loci were designed at each end of the scaffolds to orient and anchor them to the genetic map.

SSRs identified in the scaffolds were mapped only on the reference map established from the UPA402 × UF676 cross.

A total of 409 primer pairs (Supplementary Table S2) were defined in the 100 larger non-anchored scaffolds using Primer3 software36 and screened for their ability to segregate in the UPA402 × UF676 progeny.

For a given SSR locus, the forward primer was designed with a 5′-end M13 tail (5′-CACGACGTTGTAACAAGAC-3′). PCR amplifications were performed in a Mastercycler ep384 thermocycler (Eppendorf, Germany) with 5 ng of purified DNA in a 10 μl final volume of buffer containing 10 mM Tris–HCl (pH 8), 50 mM KCl, 0.001% glycerol, 2.0 mM MgCl2, 0.08 μM of the M13-tailed forward primer, 0.1 μM of the reverse primer, 200 μM of dNTP, 1 U of Taq DNA polymerase (Life Technologies, USA), 0.1 μM of M13 primer-fluorescent dye 6-FAM™, NED™, VIC®, or PET® (Applied Biosystems, CA, USA). The DNA and buffer were distributed in 384 plates using a Biomek NX automatic pipetting robot (Beckman Coulter, CA, USA). The touchdown PCR programme used was as follows: initial denaturation at 95°C for 5 min, followed by 10 cycles at 95°C for 45 s, Tm of 56–46°C (−1°C/cycle) for 1 min, and 72°C for 1 min 30 s. After these cycles, an additional round of 25 cycles were performed at 95°C for 45 s, Tm of 50°C for 1 min, and 72°C for 1 min, with a final elongation step at 72°C for 30 min.

PCR products were diluted specifically for each dye and pooled for multiplex SSR genotyping (revealing two SSRs having different sizes of amplified product per dye). A mixture of 15 μl of Hi-Di™ formamide (Applied Biosystems) and 0.12 μl of size marker GeneScan™ 600-LIZ-Size® Standard V2.0. (Applied Biosystems) was added to 2 μl of the diluted PCR pool. This pool was then analysed using the ABI 3500xL automatic sequencer (Applied Biosystems).

Images were analysed using Genemapper 4.0 software (Applied Biosystems) and exported as a data table.

2.6. **Genetic mapping**

The UPA402 × UF676 population was the result of a cross between two heterozygous cocoa clones, UPA402 (♂) an Upper-Amazon Forastero and UF676 (♀) a Trinitario. In this case, there were three segregation possibilities: loci that were homozygous for one parent and heterozygous for the other, segregation (1:1), and those that segregated in both parents (1:2:1 or 1:1:1:1).

Segregations were checked for goodness-of-fit to the expected Mendelian ratio using a chi-square test at significance levels of 0.05 and 0.01.

Individual and consensus maps were constructed using Joinmap software, version 4.0.37

Joinmap is able to combine data of several segregation types to construct a consensus genetic map. Here we used population type CP for the UPA402 × UF676 map, and population type F2 for the F2 population from Brazil. A lod score of 6 was used to identify 10 linkage groups (LGs) independently for each map. A consensus genetic map was established from the two distinct genetic maps. The corresponding groups were associated in pairs with JoinMap software. The Kosambi mapping function, with a lod score of 5 and a jump threshold of 3, was used to convert recombination frequencies into map distances.38

This consensus map combined the new EST-SNPs and genomic SSRs defined from the scaffolds, in addition to the previously mapped markers.9 This map contained only markers with a known nucleotide sequence.

3. **Results**

3.1. **Identification of SNPs and development of the golden gate assay**

The assembly made from the 149 650 T. cacao EST sequences (see Materials and Methods) generated 12692 T. cacao contigs. The number of ESTs per contig ranged from 2 to 5102. To detect good quality in silico SNPs, we assumed that contigs with more than 100 members contained paralogous sequences.13,39 We therefore first selected 4818 contigs that contained at least 4 but no more than 100 EST members. A total of 5246 SNPs were identified in silico in 2012 contigs.

We selected 4150 in silico SNPs detected in 1834 contigs that had a significant BlastX annotation similarity with known proteins of the NCBI non-redundant protein sequence database (NR) with entries from GenPept, Swissprot, PIR, PDF, PDB, and NCBI RefSeq, and as described in Argout et al.52

3.2. **SNP performance and quality**

The set of 4150 in silico SNPs was selected in the EST contigs and the SNP-harbouring sequences were then submitted to Illumina for processing using the Illumina® Assay Design Tool (ADT). ADT generates scores for each SNP that can range from 0 to 1;
SNPs with scores >0.6 have a high probability of being converted into a successful genotyping assay. In the set of 4150 submitted SNPs, 83.5% showed a high conversion success rate (>0.6), 9.2% showed a moderate conversion success rate (between 0.4 and 0.6), and 7.3% showed either a low conversion success rate or no score. A total of 1536 SNP sites having ADT scores >0.4 and without any other SNPs within the adjacent 60 bp was selected for the OPA design (Supplementary Table S1).

3.3. Analysis of base changes

One thousand and forty-four in silico SNPs (68%) were transitions and 462 (32%) were transversions (Table 2). This ratio of transition/transversion SNPs tallies with the results observed in other plant species, where transition SNPs are always more frequent than transition SNPs.

3.4. SNP polymorphism

From the 1536 SNPs, 841 (55%) with a non-ambiguous polymorphic pattern across accessions were retained as true and verified SNPs and denominated TcSNP. Of the rest, 113 (7%) failed to be genotyped, 436 (28%) had a monomorphic pattern, and 146 (10%) were polymorphic but did not show any clear fluorescent pattern suitable for reliable genotype classification.

Of the 841 polymorphic SNPs, 461 segregated in the mapping population (UPA402 × UF676) and could be mapped on the reference map. Five hundred and thirty-one were polymorphic and mapped on the F2 population map. Two hundred and thirty-nine SNP markers were segregating in both maps, thus enabling construction of a consensus map between them.

3.5. SSR polymorphism

A high-density genetic map is a key tool to order the scaffold assembly needed to generate a complete cocoa genome sequence. SSR markers were defined in the largest non-anchored scaffolds in order to improve anchoring of the *T. cacao* genome assembly provided by the International Cocoa Genome Sequencing consortium on the genetic map.

From the 409 screened SSRs (Supplementary Table S2), 163 were polymorphic for the UPA402 × UF676 progeny and could be mapped.

The new SSR markers defined from scaffolds were named mTcCIR450 to mTcCIR613 to extend the previously identified SSR marker series; mTcCIR 1 to mTcCIR 291 from genomic DNA and mTcCIR 292 to mTcCIR 447 from ESTs.

3.6. Individual genetic linkage maps

3.6.1. Map of the UPA402 × UF676 population

A new set of 624 markers with their corresponding sequences, including 461 EST-SNPs and 163 new SSR markers located on scaffolds of the genome assembly, were added to the reference map (Fig. 1, Supplementary Table S3).

The new UPA402 × UF676 map contained 1043 markers, including 461 EST-SNPs and 524 SSRs and 58 RFLPs (Table 3). Of the 1043 markers, 571 corresponded to gene markers. The length of this map was 751.7 cM having an average distance of 0.7 cM between adjacent markers.

Skewed segregation was observed for 118 markers (11.3%). The skewed markers were mainly located in LGs 2, 3, 6 and 10, as is shown in Fig. 3.

3.6.2. Map of the F2 population

The F2 map (Fig. 2, Supplementary Table S4) contained 531 EST-SNP markers. This map had a total length of 753.9 cM, with an average distance of 1.4 cM between neighboring markers. The marker density varied, with an average distance between neighboring markers ranging from 0.9 cM in LG 9 to 2.7 cM in LG7 (Table 4).

Skewed segregation was observed in 97 markers (18.3%). The skewed markers were mainly located in LGs 1, 3 and 4, as is shown in Fig. 3.

3.7. Consensus genetic linkage map

Two hundred and thirty-nine SNP markers were mapped in both populations.

The complete consensus map (Table 5) contained 1262 codominant markers including 681 EST-SNPs, 523 SSRs (163 scaffold-tagged-SSRs, 110 EST-SSRs, 250 SSRs from genomic DNA) and 58 RFLPs including 14 resistance gene analogues (Rgenes-RFLPs), arranged in 10 LGs corresponding to the haploid chromosome number of *T. cacao* (Fig. 3, Supplementary Table S5). Among the 1262 markers, 65% were gene-based markers, including SNPs, SSRs and RFLPs.

Table 2. Nucleotide substitution types of the 1536 selected in silico SNPs

| Types | Number of SNPs | Percentage | Percentage |
|-------|----------------|------------|------------|
| A → C | 126            | 8          | Transversion 32 |
| A → T | 128            | 8          |            |
| C → G | 112            | 7          |            |
| T → C | 126            | 8          |            |
| T → G | 612            | 40         | Transition 68 |
| A → G | 432            | 29         |            |
Figure 1. Genetic map constructed from an F1 progeny of 264 individuals (located in CNRA, Côte d'Ivoire) belonging to the UPA402 × UF676 cross. This map consists of 1043 markers of a known DNA sequence (461 SNPs, 524 SSRs, and 58 RFLPs), spanning 752 cM. The average distance between two markers is 0.7 cM. The new markers added to this map are printed in red.
The total map length was 733.6 cM, i.e. slightly shorter than previously constructed maps (782.8 cM for Pugh et al.\textsuperscript{8} and 779.2 cM for Fouet et al.\textsuperscript{9}). The average distance between adjacent markers on this map was 0.6 cM, and thus shorter than the 1.3 cM of the map of Fouet et al.\textsuperscript{9}

The number of mapped loci varied substantially between LGs on the consensus map; from 63 in LG10 to 201 in LG1. The average distance between two markers in the different LGs ranged from 0.4 cM in LG1 to 0.9 cM in LG10.

In total, 844 new markers (681 SNP markers and 163 SSR defined in scaffolds) were mapped. These new markers were well distributed over all chromosomes allowing to fill some gaps in the previous maps, for example on chromosome 10.

### 4. Discussion

A large set of EST-SNP markers was generated and mapped in \textit{T. cacao}. New SSR markers were added to these SNPs, providing an efficient tool for high-throughput genotyping of cocoa populations.

SSR markers are multiallelic and well adapted for fine analysis of population diversity structure.\textsuperscript{40–43} In \textit{T. cacao}, an average number of 5.8 alleles per SSR was observed by Looor Solorzano\textsuperscript{14} after genotyping a collection of genetic resources of various genetic origins, and with a maximum of 15 alleles revealed by one SSR (mTcCIR322). This is not the case for SNPs that are only biallelic, but a higher number of SNP markers (several thousands) can be easily revealed at once using high-throughput technologies.

We used our new SNP and SSR markers to construct a very high-density genetic map. Sixty-five per cent of the markers were from within genes and the average distance between adjacent markers was 0.6 cM.

Several chromosome regions include markers with skewed segregations, particularly on LG 1, LG 3, LG 4, and LG 6. The region on LG 4 includes the locus for self-incompatibility previously identified by Crouzillat et al.\textsuperscript{5} The gameto-sporeophytic incompatibility system existing in \textit{T. cacao}\textsuperscript{45,46} could possibly explain the segregation distortion on this LG 4 region. Other factors which could explain segregation distortion, such as chromosome rearrangements in banana\textsuperscript{47} which are responsible for highly skewed marker segregations, have not been reported in \textit{T. cacao}.

This high-density genetic map can be used as a major tool for efficient genome-wide association studies (GWASs) in \textit{T. cacao} populations. This method, first applied in human and animal genetics,\textsuperscript{48–51} was also found to be highly effective for studying the determinism of useful traits in plants,\textsuperscript{52–56} particularly in cocoa with the analysis of some recent hybrid populations.\textsuperscript{8,57,58} GWAS is an alternative to QTL analyses in cross progenies for the purpose of studying genetic control of phenotypic traits in cocoa.

GWASs can be carried out on unrelated genetic resources such as wild or cultivated populations or germplasm collections. Large cocoa germplasm collections are maintained in many countries and characterized for useful traits. Two international cocoa collections are hosted at the International Cocoa Genebank, Trinidad (ICGT, preserving 2300 accessions),\textsuperscript{59} and at the Centro Agronomico Tropical de Investigacion y Ensenanza, Turrialba, Costa Rica (CATIE, preserving 1150 accessions).\textsuperscript{60} The markers identified here will now certainly facilitate such

| LG   | Length (cM) | Total number of markers | Average distance between markers (cM) | SNP | RFLP | Genomic SSR | SSR from scaffold | EST-SSR |
|------|-------------|-------------------------|--------------------------------------|-----|------|-------------|-------------------|---------|
| LG1  | 90.3        | 150                     | 0.6                                  | 69  | 8    | 33          | 22                | 18      |
| LG2  | 97.5        | 126                     | 0.8                                  | 55  | 6    | 26          | 28                | 11      |
| LG3  | 74.4        | 126                     | 0.6                                  | 57  | 6    | 33          | 18                | 14      |
| LG4  | 75.6        | 120                     | 0.6                                  | 61  | 11   | 27          | 14                | 8       |
| LG5  | 77.4        | 121                     | 0.6                                  | 55  | 6    | 34          | 15                | 11      |
| LG6  | 62.7        | 73                      | 0.9                                  | 28  | 2    | 19          | 12                | 12      |
| LG7  | 48.9        | 51                      | 1.0                                  | 11  | 6    | 16          | 16                | 2       |
| LG8  | 60.3        | 64                      | 0.9                                  | 30  | 1    | 17          | 5                 | 11      |
| LG9  | 103.2       | 154                     | 0.7                                  | 78  | 6    | 34          | 17                | 20      |
| LG10 | 61.3        | 54                      | 1.1                                  | 17  | 6    | 12          | 16                | 3       |
| Total| 751.7       | 1043                    | 0.7                                  | 461 | 58   | 251         | 163               | 110     |

SNP, single-nucleotide polymorphism; RFLP, restriction fragment polymorphism; SSR, simple sequence repeat; EST, expressed sequence tag.
Figure 2. Genetic map constructed from an F2 progeny of 132 individuals (located at CEPLAC, Brazil) obtained by selfing of a single (Scavina 6 × ICS1) selected hybrid. This map consists of 531 SNP markers, spanning 754 cM. The average distance between two markers is 1.4 cM.
Figure 3. Consensus map of (UPA402 × UF676) and F2 progenies. Markers segregating in both progenies are indicated in black, those segregating only in (UPA402 × UF676) are printed in green (previously mapped markers) and blue (newly mapped markers). Markers segregating only in the F2 progeny are printed in pink. This consensus map consists of 1262 markers of a known DNA sequence, and it has a length of 734 cM. The average distance between two markers is 0.6 cM. Among the 1262 markers, 810 correspond to markers defined in expressed genes. Significant skewed segregations are indicated by asterisks (*P < 0.05, **P < 0.01) or dots (F2 population).
GWASs, providing added-value to this wide characterization work, thus boosting knowledge on the genetic determinants of useful cocoa traits.

Another benefit of this large set of mapped markers is the possible integration of molecular information in conventional cocoa breeding schemes using marker-assisted selection (MAS).

In cocoa, few MAS experiments are currently underway.3,61 The efficiency of MAS in selecting P. palmivora-resistant cocoa plants has been reported by Lanaud et al.3

Until now, MAS studies have mainly been focused on traits controlled by a small number of genes, using only markers close to QTLs. However, such methods are of limited use for traits that are determined by a large number of genes of small effects.

Table 4. Distribution of SNP markers in the LGs of the F2 map (SCA6 × C2ICS1) selfing

| LG  | Length (cM) | Number of SNP markers | Average distance between markers (cM) |
|-----|-------------|-----------------------|--------------------------------------|
| LG1 | 98.6        | 104                   | 0.9                                  |
| LG2 | 102.8       | 74                    | 1.4                                  |
| LG3 | 78.7        | 73                    | 1.1                                  |
| LG4 | 69.1        | 49                    | 1.4                                  |
| LG5 | 85.4        | 56                    | 1.5                                  |
| LG6 | 71.0        | 28                    | 2.5                                  |
| LG7 | 48.7        | 19                    | 2.6                                  |
| LG8 | 43.3        | 28                    | 1.5                                  |
| LG9 | 104.9       | 78                    | 1.3                                  |
| LG10| 51.5        | 22                    | 2.3                                  |
| Total| 753.9       | 531                   | 1.4                                  |

SNP, single-nucleotide.

Table 5. Distribution of each marker type in the LGs of the consensus genetic map

| LG  | Length (cM) | Total number of markers | Average distance between markers (cM) | SNP | RFLP | Genomic SSR | SSR from scaffold | EST-SSR |
|-----|-------------|-------------------------|--------------------------------------|-----|------|-------------|-------------------|---------|
| LG1 | 77.1        | 201                     | 0.4                                  | 120 | 8    | 33          | 22                | 18      |
| LG2 | 101.1       | 156                     | 0.6                                  | 85  | 6    | 26          | 28                | 11      |
| LG3 | 76.9        | 162                     | 0.5                                  | 91  | 6    | 33          | 18                | 14      |
| LG4 | 64.2        | 135                     | 0.5                                  | 75  | 11   | 27          | 14                | 8       |
| LG5 | 78.1        | 147                     | 0.5                                  | 81  | 6    | 34          | 15                | 11      |
| LG6 | 64          | 81                      | 0.8                                  | 36  | 2    | 19          | 12                | 12      |
| LG7 | 52.6        | 62                      | 0.8                                  | 22  | 6    | 16          | 16                | 2       |
| LG8 | 59.2        | 73                      | 0.8                                  | 39  | 1    | 17          | 5                 | 11      |
| LG9 | 100.9       | 182                     | 0.6                                  | 106 | 6    | 33          | 17                | 20      |
| LG10| 59.5        | 63                      | 0.9                                  | 26  | 6    | 12          | 16                | 3       |
| Total| 733.6       | 1262                    | 0.6                                  | 681 | 58   | 250         | 163               | 110     |

SNP, single nucleotide polymorphism; RFLP, restriction fragment polymorphism; SSR, simple sequence repeat; EST, expressed sequence tag.
integration of both genetic and genomic (functional and structural) data. Overall, about 300 QTLs or marker/trait associations have already been identified in \textit{T. cacao}. High-throughput genotyping associated with a high marker density will facilitate fine mapping of genes involved in trait variation (with GWAS or classical QTL analyses conducted on large progenies), thus allowing to refine the QTL position in the genome, while facilitating the search for candidate genes in corresponding genome sequences. Several functional studies have already been conducted in cocoa, focused mainly on genes generally expressed in specific physiological conditions or metabolisms.\(^{73}\) It will be now possible to focus more specifically on the expression of genes directly responsible for trait variation after candidate gene validation.

Analysing genome evolution during domestication processes or adaptation to climate change can also help us to identify key genes underlying adaptive traits.\(^{74}\) Loss of diversity generally occurs during genome evolution, and some genes are selectively involved in natural selection or domestication. A large set of SNPs defined in expressed genes, such as those identified in this study, provides a key tool for identifying selection signatures or adaptive substitutions, and then highlighting candidate genes potentially involved in the adaptation\(^{60}\) or domestication processes and their corresponding molecular functions.\(^{75}\) All SNPs reported in this paper were identified in orthologous genes or gene families, thus facilitating comparative genomic approaches, and benefiting from gene knowledge accumulated in other species to accelerate cocoa breeding.

5. Availability

Information on the consensus linkage map, molecular markers, and primers are available in the Map Study ‘SSR_SNP_consortium_map’ of the cocoa module of TropGeneDB database (http://tropgene.db.cirad.fr).

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