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## **Microsatellite (SSR) Analysis on Genetic Diversity of (*Coffea canephora*) Germplasm in Kagera Region, Tanzania**

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### **Abstract**

*Coffea canephora* (Robusta) is one of the two important coffee species grown in Tanzania for commercial purpose. Robusta coffee contributes 40 - 50% of the total coffee production in Tanzania. However, the production of Robusta coffee in Tanzania has been hampered by coffee wilt disease. Despite the importance of Robusta coffee in Tanzania, its genome has not thoroughly researched. A study was conducted to investigate the molecular diversity of cultivated and wild coffee found in Kagera region in Tanzania. One hundred twenty four genotypes (124) of cultivated and wild coffee were analyzed by simple sequence repeat (SSR) marker techniques using 12 microsatellite markers. Genetic diversity, similarity or dissimilarity, genetic distances between individuals and genetic differentiation between populations was analyzed. Findings indicate high genetic variations among cultivated and wild coffee genotypes ranging between 20 and 83%. Polymorphism was 80% among SSR markers with 8 loci. Two distinctive genetic groups were identified. The first genetic group comprised four distinctive genetic groups one to four.

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The second genetic group consisted of four genetic groups, five to eight. Wild coffee genotypes had similarities to some of cultivated *C. canephora* in groups one to four implying that some cultivated *C. canephora* originates from wild coffee. Groups' five to eight comprise genotypes from cultivated *C. canephora*. Detailed study is needed to compare the identified eight (8) genetic groups of *C. canephora* in Kagera region in Tanzania with the already know groups worldwide.

**Keywords:** Genetic diversity of *Coffea canephora*; (SSR) Microsatellite.

## 1. Introduction

*Coffea canephora* is one of two important coffee species grown in the world. It accounts 35-40% of world production (International Coffee Organization. www.ico.org). It is self-incompatible and diploid; while *C. arabica* which accounts 60- 65% is tetraploid and self compatible [13]. *Coffea canephora* is found in low and middle altitudes areas in Africa, far East and parts of Southern America [22]. In Africa, *C. canephora* is found in Ivory Coast, Democratic Republic of Congo, Cameroon, Uganda, Angola, Ghana, Togo, Madagascar, Republic of Central Africa and Tanzania. In Far East this coffee *spp* is found in India, Indonesia, Vietnam, Philippines and Brazil in Southern America. In most countries, *C. canephora* is conserved in ex-situ collection plots. *Coffea canephora* is self-incompatible coffee *spp* in natural, cross-pollinated perennial plant, while wind and insects being are the main pollinators. Based on the nature of *C. canephora* and its wide geographical distribution genetic characterization are very important for both effective crop improvement and conservation purposes [26, 22]. Comprehensive conversation of genetic resources for *C. canephora* is very important due to threats of extinction of the species resulting from invasion of the most devastating coffee wilt disease and rapid extinction of natural forests caused by human activities and climates changes [20]. Assessment of genetic variability between and among *C. arabica* and *C. canephora* populations using different molecular markers' techniques has be conducted by several scientists [7, 19, 15, 10,8,23, 3, 18, 17, 20, 22 and 31]. Bertrand [13] reported the genetic diversity of *C. canephora* for the first time in 1986 when studied the wild and cultivated coffee genotypes from Western and Central Africa Republic. During this study two diversity groups were identified. These groups included Congolese group, which comprised genotypes from Central African Republic and Cameroon, and a Guinean group, which comprised genotypes from Ivory Coast [20]. Montagnon *and his colleagues* [7] who further investigated the diversity of *C. canephora* and wild coffee genotypes reported three groups namely Guinea, Congolese SG1 and Congolese SG2 with groups SG1 and SG2 being the sub-groups of the Congolese group. Dussert *and his colleagues* [23] using Restriction Fragment Length Polymorphism molecular markers (RFLP) reported five diversity groups of cultivated *C. canephora* and wild coffee genotypes, adding 2 Congolese groups B and C. Cubry *and his colleagues* [18] and Cubry *and his colleagues* [17] using microsatellites on *C. canephora* confirmed the previous findings of 5 diversity groups: SG1, SG2, B, C and Guinean. Musoli *and his colleagues* [20] while studying the diversity of wild coffee and cultivated *C. canephora* from Ugandan coffee discovered 7 distinctive genetic diversity groups of which 5 were related to the previous know groups and one unique group from Uganda coffee. Thomas [4], Nyange and Marandu [9] reported that some of wild and cultivated *C. canephora* were indigenous to Uganda and the Northern part of Tanzania in Kagera. In 16<sup>th</sup> century large seed *C. canephora* was introduced from Congo, and commercial cultivation started in Uganda and the Northern part of Tanzania region in Kagera region in 1900s [2, 1].

However, due to self-incompatibility of the species and the proximity of cultivated *C. canephora* to the forests might have led to the add mixtures between introduced *C. canephora* and wild coffee [20]. The previous studies conducted to investigate the diversity of cultivated *C. canephora* and wild coffee genotypes did not include coffee genotypes from the Northern part of Tanzania. On these contexts, therefore a study was conducted to determine the diversity and relatedness and unrelatedness of cultivated *C. canephora* and wild coffee genotypes found in Kagera region in Tanzania. This paper presents the results of a study of genetic diversity of Tanzanian cultivated *C. canephora* and wild coffee genotypes.

## 2. Materials and Methods

### 2.1 Samples collections

A total of one hundred and four (104) cultivated robusta coffee cultivars and twenty (20) genotypes of wild Robusta coffee were used in this study. One hundred and four cultivars of cultivated Robusta coffee collected from farmers' fields in Kagera region were selected randomly from 656 accessions in germplasm established at ARI-Maruku in Bukoba District. A total of twenty (20) trees of wild Robusta coffee sampled from Bushenyi (4 samples) and Minziro (16 samples) forests in Missenyi district were raised at Maruku (Table 1). One hundred and four (104) of cultivated *C. canephora* and twenty (20) wild coffee young leaves were harvested using Eppendorf tubes and immediately wrapped in the plastic bags. The samples were kept in the cool ice box and eventually stored at -20°C before DNA extraction.

### 2.2 Coffee DNA extraction

Extraction of DNA was done at Sokoine University of Agriculture (SUA) using a modified CTAB protocol for isolating DNA from plant tissues as described by Mahuku in [24] and Dellaporta *and his colleagues* in [25]. Samples were ground using pestles in the eppendorf tubes. Fifty milligrams (50 mg) of each leaf sample were put in the 1.5 ml eppendorf tube containing DNA extraction buffer prepared from 100 mM Tris-HCL (pH 8.0), 2.0 M NaCL, 20 mM EDTA (pH 8.0) 2% (w/v) CTAB, (1% w/v) PVP (PVP K10, MW10.000) and 0.5% (w/v) activated charcoal. Charcoals and PVP were used to bind the phenolic compounds and left pure DNA. Samples were centrifuged at 16000g for 10 minutes at room temperature and the supernatants were transferred to the new microfuge tubes. One millilitre (1 ml) of chloroform / isoamyl alcohol (24:1) was added to each supernatant sample and then centrifuged at 16000 g for 10 minutes at room temperature. The aqueous (upper) phases were transferred to the fresh eppendorf tubes and 0.45 ml of isopropanol was added to each sample and mixed by inversion and thereafter samples were incubated at 25°C for 1 hour and then centrifuged at 700 g for 10 minutes at room temperature to get DNA pellets. The supernatants were discarded to leave the DNA pellets. The pellets were washed by adding 1 ml of wash buffer made up of 15 mM ammonium acetate in 75% (v/v) ethanol to each tube and then centrifuged at 900 g for 10 minutes at room temperature. The supernatants were discarded and the pellets were air dried at room temperature for twenty minutes. Each pellet was dissolved in 50 µL of TE buffer prepared from (10 mM Tris-HCL (pH 8.0) and 1 mM EDTA (pH 8.0) and then centrifuged at 16000 g for 10 minutes at room temperature to remove some impurities and remain with pure supernatants. The supernatants which contained DNA were transferred to the new tubes and in forms of solutions DNA were stores at -20°C in

the deep freezer. The DNAs' quantities and qualities were assessed by using 1% agarose gel electrophoresis using the standard protocol established at molecular laboratory at Sokoine University of Agriculture (SUA).

**Table 1:** Genotypes of 104 cultivated Robusta and 20 wild coffee characterized by using microsatellite (SSR) markers

| Genotype code | Origin  | Genotype code | Origin  | Genotype code | Origin  | Genotype code | Origin          |
|---------------|---------|---------------|---------|---------------|---------|---------------|-----------------|
| 001 MI 1      | Misenyi | 087 ML12      | Muleba  | 192 ML1       | Muleba  | 337 MI21      | Misenyi         |
| 002 MI 2      | Misenyi | 091 KR23      | Karagwe | 193 ML2       | Muleba  | 342 MI20      | Misenyi         |
| 003 MI 3      | Misenyi | 092 KR24      | Karagwe | 194 ML3       | Muleba  | 344 MI19      | Misenyi         |
| 004 MI 4      | Misenyi | 108 BK4       | Bukoba  | 240 BK14      | Bukoba  | 346 MI11      | Misenyi         |
| 005 MI 5      | Misenyi | 109 BK5       | Bukoba  | 255 BK16      | Bukoba  | 347 MR10      | Bukoba          |
| 006 MI 6      | Misenyi | 112 BK6       | Bukoba  | 257 BK18      | Bukoba  | 348 (13/61)   | Bukoba          |
| 007 MI 7      | Misenyi | 113 BK 7      | Bukoba  | 259 BK19      | Bukoba  | 349 ML2       | Muleba          |
| 008 MI 8      | Misenyi | 114 BK4       | Bukoba  | 263 BK20      | Bukoba  | MSI           | Bukoba          |
| 009 MI 9      | Misenyi | 115 BK8       | Bukoba  | 268 BK21      | Bukoba  | FB1           | Busenyi forest  |
| 010 MI 10     | Misenyi | 117 BK        | Bukoba  | 269 BK22      | Bukoba  | FB2           | Busenyi forest  |
| 011 MI 11     | Misenyi | 118 ( 1/62)   | Bukoba  | 280 KR1       | Karagwe | FB3           | Bushenyi forest |
| 012 MI 12     | Misenyi | 120 M L13     | Muleba  | 283 KR2       | Karagwe | FB4           | Bushenyi forest |
| 020 MS5       | Bukoba  | 123 BK 10     | Bukoba  | 284 KR3       | Karagwe | FM5           | Minziro forest  |
| 023 KR20      | Karagwe | 125 BK11      | Bukoba  | 287 KR4       | Karagwe | FM6           | Minziro forest  |
| 025 KR19      | Karagwe | 127 ML12      | Muleba  | 288 KR5       | Karagwe | FM7           | Minziro forest  |
| 026 BK26      | Bukoba  | 131 MS1       | Bukoba  | 292 KR6       | Karagwe | FM8           | Minziro forest  |
| 030 KR18      | Karagwe | 139 MI11      | Misenyi | 293 KR7       | Karagwe | FM9           | Minziro forest  |
| 036 KR 12     | Karagwe | 142 BK13      | Karagwe | 294 KR8       | Karagwe | FM10          | Minziro forest  |
| 037 ML17      | Muleba  | 147 KR25      | Karagwe | 295 BK23      | Bukoba  | FM11          | Minziro forest  |
| 046 KR22      | Karagwe | 158 MI12      | Misenyi | 306 ML20      | Muleba  | FM12          | Minziro forest  |
| 047 MS2       | Bukoba  | 160 MI13      | Misenyi | 308 MI21      | Misenyi | FM13          | Minziro forest  |
| 049 KR12      | Karagwe | 162 MI14      | Misenyi | 310 MI25      | Misenyi | FM14          | Minziro forest  |
| 054 KR16      | Karagwe | 164 MI15      | Misenyi | 311 KR9       | Karagwe | FM15          | Minziro forest  |
| 055 KR15      | Karagwe | 165 MI16      | Misenyi | 312 KR10      | Karagwe | FM16          | Minziro forest  |
| 057 BK2       | Bukoba  | 167 MI17      | Misenyi | 315 KR11      | Karagwe | FM17          | Minziro forest  |
| 059 BK3       | Bukoba  | 170 ML10      | Muleba  | 316 ML22      | Muleba  | FM18          | Minziro forest  |
| 060 KR13      | Karagwe | 172 ML9       | Muleba  | 320 KR12      | Karagwe | FM19          | Minziro forest  |
| 062 KR14      | Karagwe | 175 ML8       | Muleba  | 323 ML24      | Muleba  | FM20          | Minziro forest  |
| 077 ML18      | Muleba  | 177 ML7       | Muleba  | 324 ML15      | Muleba  |               |                 |
| 079 ML17      | Muleba  | 179 ML6       | Muleba  | 330 MI24      | Misenyi |               |                 |
| 080 ML16      | Muleba  | 181 ML5       | Muleba  | 332 MI23      | Misenyi |               |                 |
| 086 ML15      | Muleba  | 185 ML4       | Muleba  | 333 MI22      | Misenyi |               |                 |

### 2.3 SSR genotyping

Fourteen polymorphic markers (SSRs) mapped on the *C. canephora* and *C. arabica* genomes were used to genotype the 124 genotypes collected from Bukoba, Karagwe, Missenyi and Muleba districts, and Bushenyi and Minziro forests in Kagera region in Tanzania (Table 9). Markers 394, 445, 501,364, 368, 384,355,456 DL 020 and 456 were designed from *C. canephora* clone 126 [14, 28]. Markers 774 and 782 were designed from *C. arabica* (Catura) [21]. Markers (DL013 and DL025) were designed for the study of sugar metabolism in coffee [28]. These markers were chosen because of their applications in *C. canephora* and related coffee species [29, 17, 20]. Markers were chosen based on linkage groups (n=11). In this study 10 out of 11 linkage groups which are located at a distance of at least 50 cM from each other (20) were used for the assessment of the diversity. The set of markers used were enough to allow differentiations of genetic diversity within the species [20, 17].

### 2.4 PCR amplification and visualization

Polymerase chain reaction (PCR) were performed in 20  $\mu$ L containing 13.4  $\mu$ l PCR water, 2  $\mu$ l PCR buffer, 1.2  $\mu$ l of  $MgCl_2$  (25 mM), 0.4  $\mu$ l of dNTP (10 mM), 1  $\mu$ L of primer forward (10  $\mu$ M), 1  $\mu$ L of primer reverse (10 $\mu$ M), Taq DNA polymerase 5  $\mu$ / $\mu$ L and 1  $\mu$ L of DNA. Polymerase chain reaction (PCR) amplifications were run in an Eppendorf PCR tubes in the PCR machine.

The amplification protocol consisted of an initial denaturation cycles of 5 minutes at 94°C followed by 30 cycles (45s at 94°C, 30s at  $T_M^C$  in accordance to the design of each primer (Table 4.1), 2 minutes at 72° C and the final elongation step at 72° C for 5 minutes. The 8  $\mu$ L of each PCR products together with molecular marker ladder were loaded into the wells on 2% solidified agarose gels in 1 x TBE (Tris Borate EDTA) buffer. Electrophoresis separation of DNA fragments was conducted at 120 V for 2.30 hours. Resulting DNA fragments were stained with 0.5  $\mu$ g / ml ethidium bromide in the bath of water for 30 to 60 minutes.

The gel images were visualized under UV light chamber and were retrieved by using digital camera CANON and eventually modified using Picasa 3 software. The DNA fragments from different loci sizes for each primer were scored as 0 and 1 represented absence and presence of bands respectively. The pair of alleles from each locus was scored as AA, Aa and aa to represent dominant, heterozygote and recessive alleles respectively.

### Data analysis

Data were entered in a computer using excel software package as binary matrices. Data were analyzed by using hierarchical structure genomic statistical package. Analysis of genotypic frequency, alleles frequency, polymorphic loci, Shannon index, HW test, allele numbers, observed heterozygosity. F-Statistic, fixed index effective allele numbers, expected heterozygote and genetic distance were performed using the POPGENE software version 1.31 using the following models:

- i. The index of genetic similarity (GS):  $GS = 2N_{ij} / (N_i + N_j)$ , where  $N_{ij}$  is the number of SSR alleles common to genotypes i and j while  $N_i$  and  $N_j$  are the total numbers of SSR alleles observed for genotype i and j respectively.

- ii. The mean number of alleles 'N':  $N = \sum_{i=1}^n (N_i/n)$ , where  $N_i$  is the number of alleles at  $i$ th allele, where  $N$  is the number of alleles at  $i$ th allele.
- iii. The effective allelic number 'N<sub>e</sub>':  $N_e = \sum_{i=1}^n N_{ei}/n = \sum_{i=1}^n (1/\sum q_j^2)/n$ , where  $N_e$  is the effective allelic number at  $i$ th allele, and  $q_j$  the frequency of the  $j$ th allele.
- iv. H<sub>o</sub> is the observed heterozygosity:  $H_o = \sum H_{oi}/n = \sum (1 - \sum_{j=1}^m q_{ij}^2)/n$ , where  $H_{oi}$  represents the observed heterozygosity of the  $i$ th allele and  $q_{ij}$  is the frequency of  $j$ th homozygous allele at  $i$ th allele.
- v. The expected heterozygosity, (index of gene diversity):  $H_e = \sum H_i/n = \sum (1 - \sum q_{ij}^2)/n$ , where  $H_i$  is the expected heterozygosity of the  $i$ th allele and  $q_{ij}$  is the frequency of the  $j$ th homozygous allele at  $i$ th allele.
- vi. Wright fix index, defined as inbreeding coefficient 'F':  $F = 1 - H_o/H_e$  ranges from -1 to 1. The value of F is 1 when the population is heterozygous. 'F<sub>it</sub>' 'F<sub>is</sub>' and 'F<sub>st</sub>' are Wright F- statistics parameters. F<sub>it</sub> and F<sub>is</sub> are defined as genetic deviation from Hardy- Weinberg expectation within and among genotypes respectively. Genotypes arrive at Hardy- equilibrium when F<sub>it</sub> and F<sub>is</sub> are 0. F<sub>st</sub>, ranging from 0 to 1, is an estimate of gene differentiation between genotypes, which represents genetic variation among genotypes [32]. F<sub>st</sub> is 0 if there was no genetic variation among genotypes.
- vii. Genetic distances 'D' between each pair of landraces was estimated by the modified Rogers distance as follows:  $D = 1/n \sum_{i=1}^n \sum_{j=1}^m 1/2(p_{ij}^X - p_{ij}^Y)^2$ , where  $p_{ij}^X$  and  $q_{ij}^Y$  are frequencies of  $i$ th allele at  $j$ th allele/locus in genotypes X and Y respectively. Molecular data were further analyzed by using the hierarchical cluster analysis method. Dendrogramme tree was drawn based on Nei's genetic distances using un-weighted pair-group method with arithmetic mean (UPGMA) generated by the nearest neighbour Jaccard similarities coefficients.

### 3. Results

#### 3.1 Amplification of PCR products of 104 cultivated *C. canephora* and 20 wild coffee

Figure 1 summarizes amplification products of genotyped 104 cultivated Robusta coffee and 20 wild coffee genotypes using markers designed from Robusta clone 126. Among twelve (12) markers tested, 10 showed amplification and produced clear bands that were scored (Figure 1).

#### 3.2 Genetic diversity of cultivated Robusta and wild coffee

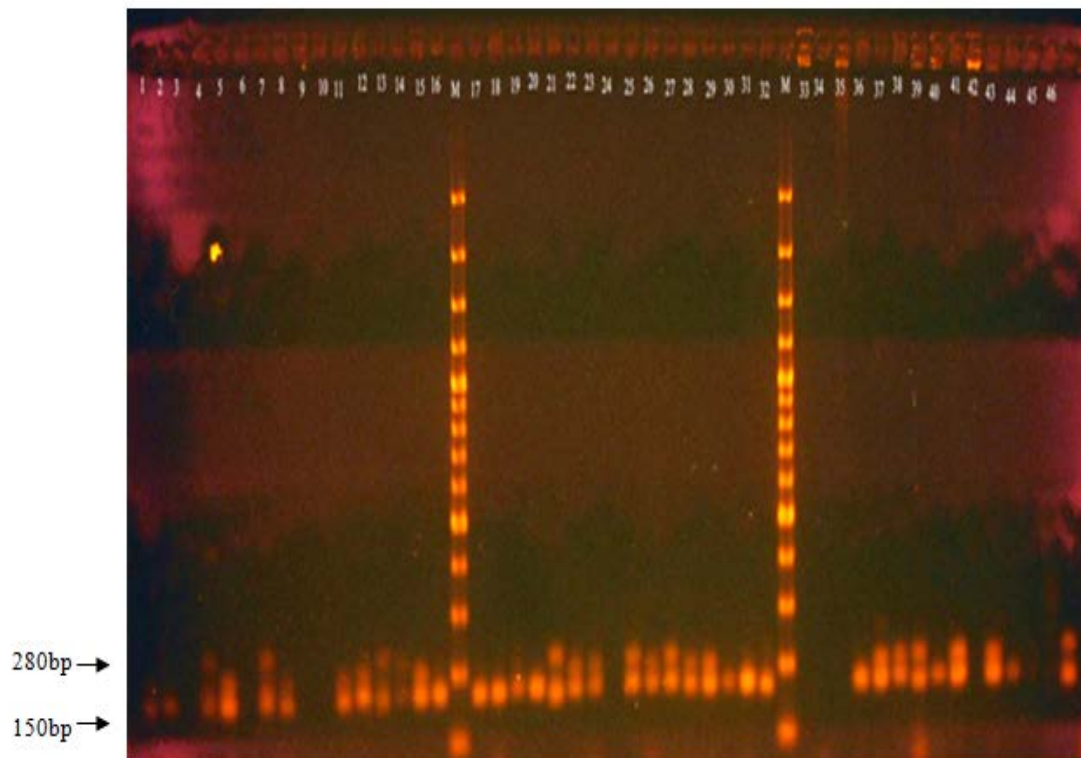
Figure 1 and Table 2 summarize results on the dissimilarity/or similarity of assessed cultivated Robusta coffee and wild coffee in Kagera region. A total number of alleles observed for tested primers were 248. Eight out of 10 amplified markers generated 8 polymorphic loci, and 80% polymorphisms were detected from 124 genotypes investigated showing the greatest allelic diversity of the populations. For each SSR locus, the number of alleles ranged from 1 to 2 corresponding to an average of 1.8. The effective allelic number ranged from 1.0 to 1.9995 with average of 1.4779. The average of observed heterozygosity (H<sub>o</sub>) was 0.12 ranging from 0.00 to 0.35. The average of expected heterozygosity was 0.28, ranging from 0.00 to 0.50. The overall mean for heterozygosity ranged between 0.00 and 0.18 with average of 0.05.

**Table 2:** SSR primers used for PCR analysis of 124 coffee genotypes

| SN  | EMBL.acc.no. | Marker Name | Primer Sequences (5' - 3')         | Allele size (bp) | TM <sup>c</sup> |              | Primer origin                  | Species origin        |
|-----|--------------|-------------|------------------------------------|------------------|-----------------|--------------|--------------------------------|-----------------------|
|     |              |             |                                    |                  | Designer        | PCR products |                                |                       |
| 1   | AM231563     | 394         | Forward. GCCGTCTCGTATCCCTCA        | 124              | 52.9            | 54.0         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. GAAGCCAGAAAAGTCAGTCACATAG |                  |                 | 53.8         |                                |                       |
| 2.  | AM231567     | 445         | Forward. CCACAGCTTGAATGACCAGA      | 275              | 52.1            | 53.3         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. AATTGACCAAGTAATCACCGACT   |                  |                 | 53.3         |                                |                       |
| 3.  | AM231576     | 501         | Forward. CACCACCATCTAATGCACCT      | 343              | 51.9            | 52.4         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. CTGCACCAGCTAATTCAAGC      |                  |                 | 52.4         |                                |                       |
| 4.  | AJ871899     | DL020       | Forward.TGCTCAAACCTTCTTGCT         | 250              | 42.5            | 42.5         | Leroy and his colleagues (28)  | <i>Coffea</i>         |
|     |              |             | Reverse. CGCCAACCTAATGTGT          |                  |                 | 42.5         |                                |                       |
| 5.  | AJ871892     | DLO13       | Forward. AGAGGGATGTCAGCATAA        | 267              | 44.1            | 44.2         | Leroy and his colleagues (28)  | <i>Coffea</i>         |
|     |              |             | Reverse. ATTTGTGTTTGGTAGATGTG      |                  |                 | 44.3         |                                |                       |
| 6.  | AJ871904     | DL025       | Forward.TTGTGAGAGTGGAGGA           | 197              | 42.0            | 44.0         | Leroy and his colleagues (28)  | <i>Coffea</i>         |
|     |              |             | Reverse. CCAAAGACAGTGCAGTAA        |                  |                 | 43.0         |                                |                       |
| 7.  | AM231556     | 364         | Forward. AGAAGAATGAAGACCAAACACA    | 90               | 50.5            | 50.4         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. TAACGCCTGCCATCG           |                  |                 | 48.3         |                                |                       |
| 8.  | AM231558     | 368         | Forward. CACATCTCCATCCATAACCATTT   | 160              | 54.2            | 54.5         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. TCCTACCTACTTGCTGTGCT      |                  |                 | 53.0         |                                |                       |
| 9.  | AM231560     | 384         | Forward. ACGCTATGACAAGGCAATGA      | 255              | 52.9            | 54.5         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. TGCAGTAGTTTACCCTTTATCC    |                  |                 | 54.0         |                                |                       |
| 10. | AM231552     | 355         | Forward. CTATGATGTCTTCCAACCTTCTAAC | 177              | 52.2            | 52.5         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. GGTCCAATTCTGTTTCAATTTTC   |                  |                 | 51.8         |                                |                       |
| 11. | AJ308774     | 774         | Forward. GCCACAAGTTTCGTGCTTTT      | 228              | 54.2            | 55.0         | Poncet and his colleagues (29) | <i>Coffea arabica</i> |
|     |              |             | Reverse. GGGTGTCCGGTGTAGGTGTATG    |                  |                 | 53.8         |                                |                       |
| 12. |              |             | Forward. AAAGGAAAATTGTTGGCTCTGA    | 114              | 54.4            | 53.0         | Poncet and his colleagues (30) | <i>Coffea arabica</i> |
| 13. | AJ308782     | 782         | Reverse. TCCACATACATTCCAGCA        |                  |                 | 53.4         |                                |                       |
| 14. | AM231568     | 456         | Forward.TGGTTGTTTTCTTCCATCAATC     | 297              | 53.0            | 53.0         | Poncet and his colleagues (29) | <i>Coffea</i>         |
|     |              |             | Reverse. TCCAGTTTCCACCTCT          |                  |                 | 52.5         |                                |                       |

Note: Source Cubry *And his colleagues* [17] and Musoli *And his colleagues* [20]

The results showed two distinctive main genetic groups among 104 cultivated robusta and 20 wild coffee genotypes (Fig. 1). The first genetic group consisted groups I, II, III and IV. The second genetic group composed groups V, VI, VII and VIII. The distances of each group from the two main groups varied significantly ( $P = 0 \leq 0.05$ ) (Fig. 1).



**Figure 2:** Gel electrophoresis of DNA of *C. canephora* genotypes, M is a 100bp Ladder, amplification product from SSR marker 355 with alleles ranging between 150bp - 280bp using accessions 1- 46 (Table 1).

#### **Genetic structure of cultivated and wild coffee populations of Tanzanian germplasm**

The results for the genetic structure of cultivated and wild coffee population varied from  $F_{st} = 0$  for loci DL 025 and 774 to  $F_{st} = 0.97$  for locus DL013. The overall mean of  $F_{st}$  for 104 cultivated *C. canephora* and 20 wild coffees was 0.81 indicating that 81% of genetic variations were observed among investigated genotypes and only 19% of genetic variations were noted within cultivated *C. canephora* and wild coffees populations.

The structured analysis revealed ten groups of cultivated *C. canephora* and wild coffee coffees in Kagera region of Tanzania with four distinctive populations of cultivated *C. canephora* (Fig. 1).

The wild coffee from Bushenyi forest was clearly identified as different group with admixture of very few individuals from cultivated *C. canephora* (Table 4.3). However, the results showed that some wild coffees from Bushenyi and Minziro forests were genetically closely related with some cultivated *C. canephora*.

#### **Genetic distances between cultivated *C. canephora* genotypes from Tanzanian germplasm**



Tables 4 and 5 present genetic distances of 14 selected cultivated *C. canephora* and coffee populations from Missenyi, Muleba, Bukoba, Karagwe and wild coffees from Bushenyi and Minziro forests. The genetic distances among 14 selected *C. canephora* ranged from the lowest value (0.00) between MI-4, MI-6 and MI-5 to the highest (0.64) between MI-13 and MI-2 (Table 4). Furthermore, the genetic distances between the genotypes of cultivated *C. canephora* and wild coffee varied greatly among the six origins.

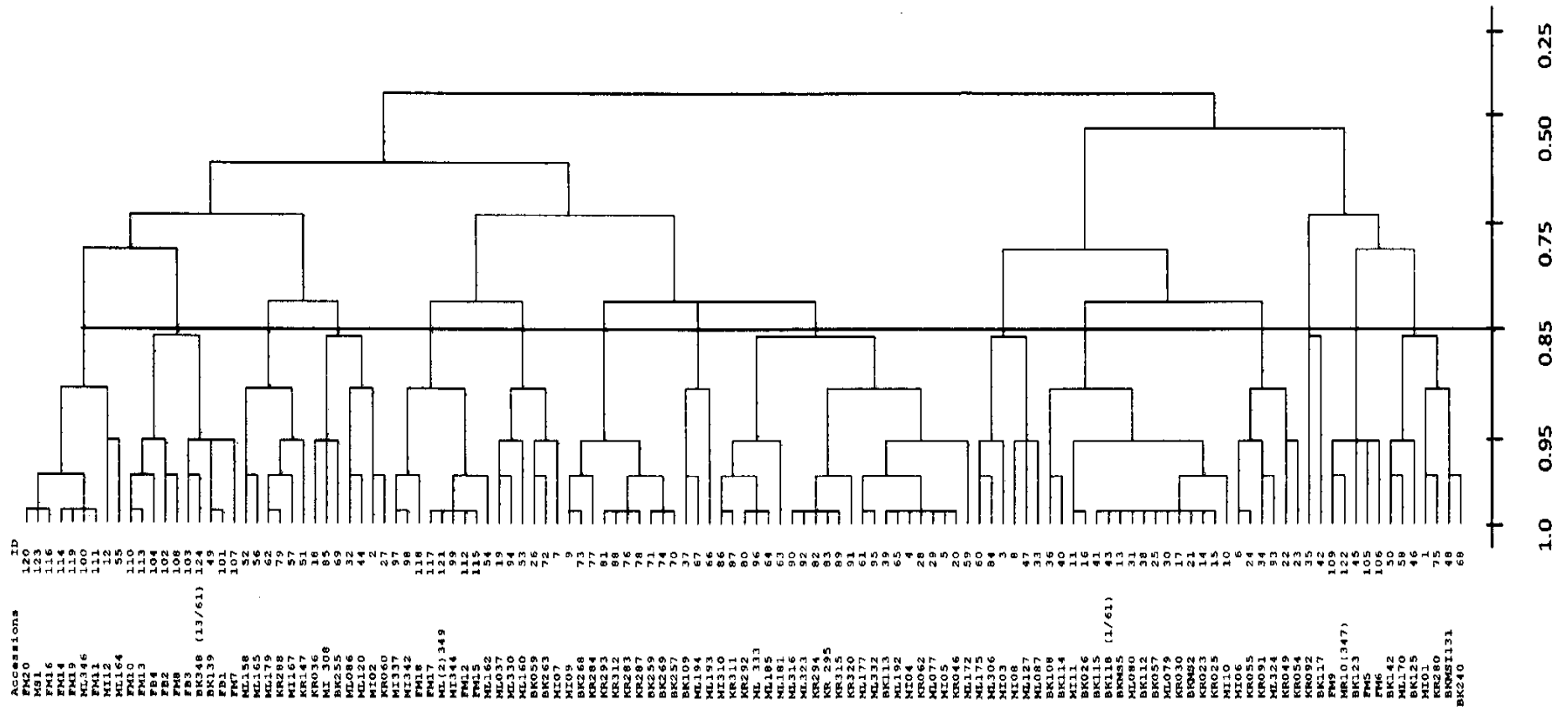
The lowest genetic distance was 0.0034 between the genotypes for cultivated *C. canephora* from Karagwe and Bukoba districts. The overall results of genetic distances of cultivated *C. canephora* from Missenyi, Bukoba, Karagwe and Muleba were genetically closely related (Table 5).

The highest genetic distance was observed between wild coffee from Bushenyi forest and cultivated *C. canephora* from Muleba (0.13), Karagwe (0.11) and Missenyi (0.10) districts. The results also showed that wild coffee from Minziro forest were closely related to cultivated *C. canephora* from Bukoba and Karagwe districts.

**Table 3:** Genetic variations of 104 cultivated *C. canephora* and 20 wild coffees based on 10 SSR loci in Tanzania

| Locus  | Simple Size | N <sub>a</sub> | N <sub>e</sub> | I      | H <sub>o</sub> | H <sub>e</sub> | Ne**   | H <sub>av</sub> | F <sub>st</sub> | Null <sub>m</sub> |
|--------|-------------|----------------|----------------|--------|----------------|----------------|--------|-----------------|-----------------|-------------------|
| 456    | 248         | 2              | 1.3402         | 0.4213 | 0.1532         | 0.2549         | 0.2539 | 0.0766          | 0.6982          | 0.1081            |
| 364    | 248         | 2              | 1.4018         | 0.4612 | 0.0242         | 0.2878         | 0.2866 | 0.0121          | 0.9578          | 0.0110            |
| 368    | 248         | 2              | 1.7763         | 0.6288 | 0.2581         | 0.4388         | 0.4370 | 0.1290          | 0.7048          | 0.1047            |
| 394    | 248         | 2              | 1.0411         | 0.0987 | 0.2442         | 0.0397         | 0.0395 | 0.0121          | 0.6938          | 0.1103            |
| 501    | 248         | 2              | 1.6103         | 0.5667 | 0.1210         | 0.3805         | 0.3790 | 0.0605          | 0.8404          | 0.0475            |
| 355    | 248         | 2              | 1.9995         | 0.6930 | 0.3548         | 0.5019         | 0.4999 | 0.0121          | 0.6451          | 0.1376            |
| 384    | 248         | 2              | 1.6205         | 0.5710 | 0.0645         | 0.3845         | 0.3829 | 0.0605          | 0.9158          | 0.0230            |
| DL013  | 248         | 2              | 1.9895         | 0.6905 | 0.0242         | 0.4994         | 0.4974 | 0.1774          | 0.9757          | 0.0062            |
| DL025  | 248         | 2              | 1.000          | 0.000  | 0.000          | 0.000          | 0.000  | 0.0324          | 0.000           | *****             |
| 774    | 248         | 2              | 1.000          | 0.000  | 0.000          | 0.000          | 0.000  | 0.0121          | 0.000           | *****             |
| Mean   | 248         | 1.8            | 1.4779         | 0.4131 | 0.1024         | 0.2787         | 0.2776 | 0.0512          | 0.8155          | 0.0565            |
| St.Dev |             | 0.42           | 0.3849         | 0.2775 | 0.1209         | 0.1997         | 0.1997 | 0.0605          |                 |                   |

N<sub>a</sub> = Observed number of alleles per locus, N<sub>e</sub> = Effective number of alleles {(Kimura and Crow (1964))}, I = Shannon's Information index {Lewontin (1972)}, H<sub>o</sub> = Observed heterozygosity computed using Levene (1949), H<sub>e</sub> = Expected heterozygosity computed using Levene (1949), Nei's = (1973) Expected heterozygosity, H<sub>av</sub> = Average heterozygosity and Null<sub>m</sub> = Estimated from F<sub>st</sub> = 0.25(1-F<sub>st</sub>)/F<sub>st</sub>. At P = ≤ 0.05)



**Figure 1:** The Cluster Dendrogramme representing the dissimilarity among cultivated *C. canephora* and wild coffee genotypes from Kagera region, Tanzania obtained by UPGMA method based on the weighted Jaccard index estimated from the polymorphism of 12 microsatellite

**Table 4:** Genetic distances between selected 14 cultivated *C. canephora* from Missenyi district

|       | MI-1   | MI-2   | MI-3   | MI-4   | MI-5   | MI-6   | MI-7   | MI-8   | MI-9   | MI-10  | MI-11  | MI-12  | MI-13  | MI-14 |
|-------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|-------|
| MI-1  | *****  |        |        |        |        |        |        |        |        |        |        |        |        |       |
| MI-2  | 0.3567 | *****  |        |        |        |        |        |        |        |        |        |        |        |       |
| MI-3  | 0.0256 | 0.2620 | *****  |        |        |        |        |        |        |        |        |        |        |       |
| MI-4  | 0.3495 | 0.2064 | 0.3239 | *****  |        |        |        |        |        |        |        |        |        |       |
| MI-5  | 0.3495 | 0.2064 | 0.3239 | 0.0000 | *****  |        |        |        |        |        |        |        |        |       |
| MI-6  | 0.3495 | 0.2064 | 0.3239 | 0.0000 | 0.0000 | *****  |        |        |        |        |        |        |        |       |
| MI-7  | 0.5816 | 0.2451 | 0.5559 | 0.0949 | 0.0949 | 0.0949 | *****  |        |        |        |        |        |        |       |
| MI-8  | 0.3567 | 0.5108 | 0.4051 | 0.0813 | 0.0813 | 0.0813 | 0.2451 | *****  |        |        |        |        |        |       |
| MI-9  | 0.2520 | 0.5722 | 0.3054 | 0.1808 | 0.1808 | 0.1808 | 0.2936 | 0.1369 | *****  |        |        |        |        |       |
| MI-10 | 0.3567 | 0.2231 | 0.2620 | 0.3495 | 0.3495 | 0.3495 | 0.3993 | 0.5108 | 0.2620 | *****  |        |        |        |       |
| MI-11 | 0.5166 | 0.2064 | 0.4039 | 0.1942 | 0.1942 | 0.1942 | 0.1638 | 0.3495 | 0.2498 | 0.0813 | *****  |        |        |       |
| MI-12 | 0.0256 | 0.2620 | 0.0541 | 0.3239 | 0.3239 | 0.3239 | 0.4606 | 0.4051 | 0.2364 | 0.2620 | 0.4039 | *****  |        |       |
| MI-13 | 0.3040 | 0.6405 | 0.2783 | 0.5592 | 0.5592 | 0.5592 | 0.5289 | 0.4581 | 0.3525 | 0.3040 | 0.2968 | 0.3247 | *****  |       |
| MI-14 | 0.1054 | 0.5106 | 0.1369 | 0.3495 | 0.3495 | 0.3495 | 0.5816 | 0.2231 | 0.1369 | 0.2231 | 0.3495 | 0.1280 | 0.1516 | ***** |

Genetic distance among 14 selected Robusta accessions (P = 0.05)

**Table 5:** Genetic distance of cultivated *C. canephora* and wild coffees based on geographical locations

| Population(ID) | MI     | ML     | BK     | KR     | FB     | FM    |
|----------------|--------|--------|--------|--------|--------|-------|
| <b>MI</b>      | *****  |        |        |        |        |       |
| <b>ML</b>      | 0.0345 | *****  |        |        |        |       |
| <b>BK</b>      | 0.0185 | 0.0055 | *****  |        |        |       |
| <b>KR</b>      | 0.0215 | 0.0063 | 0.0034 | *****  |        |       |
| <b>FB</b>      | 0.1006 | 0.1294 | 0.0948 | 0.1141 | *****  |       |
| <b>FM</b>      | 0.0785 | 0.1076 | 0.0783 | 0.0902 | 0.0076 | ***** |

Genetic distances among population of cultivated robusta and wild coffee species ( $P = 0.05$ ).

Abbreviations represent geographical locations where the materials were collected: MI-

Missenyi, ML= Muleba, BK = Bukoba, KR = Karagwe, FB = Bushenyi forest and FM = Minziro forest.

#### 4. Discussion

The analysis of genetic diversity of cultivated *C. canephora* and wild coffee revealed unrelated and relatedness of coffee species in Kagera region of Tanzania. The high polymorphism (80%), variations on F- statistics, expected heterozygosity; coefficient of similarity and genetic distances found in this study reflects a highest genetic diversity among 124 genotypes of both wild and cultivated *C. canephora*. The highest observed genetic diversity on this study provides evidence that SSR markers are adequate for assessing intra-specific and inter specific variations, and informative for detecting genetic diversity and relationships among cultivated *C. canephora* and their related wild coffee genomes. These findings are in agreement with those reported in the previous studies on the global diversity of *C. canephora* and wild coffee using SSR markers [18, 17, and 20]. In this study, two main groups of cultivated *C. canephora* with eight [8] subgroups had been identified of which four sub groups composed of mixtures of cultivated and wild coffee genotypes and the other four sub groups composed of only cultivated *C. canephora*. The first and second groups comprised 75 and 25 % of the total genotypes investigated, respectively indicating that high proportions of robusta coffee cultivars growing in Kagera region are mixtures of cultivated and wild coffees. Furthermore, two distinctive groups of wild coffee genotypes were observed from Bushenyi and Minziro forests. The SSR techniques showed that coffee genotypes found in Minziro and Bushenyi forests were closely related to cultivate *C. canephora* indicating that all investigated coffee genotypes could have a common genome. Moreover, observed genetic dissimilarity between cultivated *C. canephora* and some wild coffee in Minziro and Bushenyi imply that some coffee genotypes lack common genome. However, cultivated *C. canephora* and wild coffee species from Minziro are not easily distinguishable morphologically and they were either erect or bending types.

The wild genotypes from Bushenyi forest despite of being genetically related to cultivated *C. canephora*, but they have distinctive morphological characteristics. Individual coffee trees found in Bushenyi forest were short, with small branches, few branches, few berry clusters with small seeds, resistant to coffee leaf rust and coffee wilt disease. Similar relationships between wild coffee materials and Nganda - erect populations had been observed in Uganda [20] and Ivory Coast [16].

Reports by Musoli *and his colleagues* [20] and Montagnon *and his colleagues* [6] showed that cultivated *C. canephora* in most of African, *C. canephora* growing countries resulted from natural crossing of wild coffee materials and introduced genotypes from other regions or countries leading to a mixture of all genotypes. Genetic relationships between cultivated *C. canephora* observed in Bukoba, Karagwe, Missenyi and Muleba districts with wild coffee found in Bushenyi and Minziro forests support that hypothesis and indicate that the early established coffee bushes in Kagera region in Tanzania originated in Kagera region and Uganda [9,5,11]. From this study therefore, it was believed that cultivated coffee in Kagera region in Tanzania originated from wild coffee genotypes found in the region and Uganda [5, 4]. The presence of both Nganda and Erect coffee types in Uganda and Tanzania implies that both countries share the genetic pool of *C. canephora*. The variations of identified genetic groups between coffee species found in Tanzania and those of Uganda probably attributed to the origin sources of tested materials. Coffee genotypes used in this study represented the actual natural origin where there was no any kind of improvement while those used in Uganda probably could have been undergone through improvements. The genotypes of wild coffee probably have been cultivated in admixtures with introduced genotypes from Congolese sub groups SG2, SG1 B, C and Guinean leading to existence of spreading (Nganda, Erect and Semi erect cultivated types). The high diversity of cultivated *C. canephora* genotypes could be due to multiples origins of coffee species, resulting from successive natural hybridization of wild species and introduced Congolese genotypes [20]. According to TCB [27] cultivated *C. canephora* was introduced in Kagera region during the 16<sup>th</sup> century from Congo. Musoli *and his colleagues* [20] reported the presence of the genetic diversity of the Congolese *C. canephora* groups B, C, SG1, SG2 and Guinean in Ugandan *C. canephora* genotypes. Since Tanzania and Uganda share borders there is a high possibility of having Guineans and Congolese groups in Tanzanian cultivated *C. canephora*. The genomic molecular study on cultivated *C. canephora* and wild coffee genotypes revealed the diversity and the complex mixtures between wild and cultivated *C. canephora* in Kagera region in Tanzania. The value of expected dissimilarity coefficient within Tanzanian genotypes ranged from 0.08 to 0.83 with overall mean of 0.51. The mean value of gene diversity observed in this study was close to the previous values (0.55) observed within *C. canephora* [17]. Furthermore, the values of expected variations on this study were close to those reported by Musoli *and his colleagues* [20] who reported the diversity of wild and *C. canephora* in Uganda using 18 SSRs microsatellite markers ranged 0.48 to 0.59, and 0.47 to 0.68 for out crossing perennial plants [12].

The molecular analysis of wild coffee genotypes found in Bushenyi and Minziro forests revealed the genetic relatedness and distinctiveness to minority and majority of cultivated *C. canephora* found in Kagera region in Tanzania respectively. The existence of the similarity between cultivated *C. canephora* and wild genotypes mixture observed in this study is further supported by Cubry *and his colleagues* [16] who reported the highest diversity of the Guinean group which comprised a large number of natural populations from the forests of Ivory Coast and smallholder plantations. The overall findings with regards to wild coffee support the previous reports

that *C. canephora* is indigenous to Uganda and Tanzania [9, 20, 11].

## 5. Conclusion and Recommendations

In conclusion, this study revealed the richness of genetic diversity of cultivated *C. canephora* and wild coffee in Kagera region in Tanzania. The results obtained from this study will be useful in planning strong breeding programme for genetic conservation and improvement of *C. canephora* the second important coffee species in Tanzania. The observed genetic diversity in this study will be utilized in hybridization of coffee species to develop coffee varieties which are high yielding, resistant to diseases, with good qualities and which can grow well in various agro ecological environments. In addition, these findings will be the basis for exploring more information on existence of coffee species within the country, collect and conserve them for further uses in the breeding programme.

The identified eight (8) genetically diverse groups of cultivated *C. canephora* and two (2) genetic diversity groups of wild coffee should be thoroughly studied to compare their relationships with the genotypes found in the other countries which grow robusta coffee worldwide.

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