ASSESSMENT OF THE SUITABILITY OF MOLECULAR SCoT MARKERS FOR GENETIC ANALYSIS OF COFFEE SPECIES

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Abstract

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Start codon targeted polymorphism (SCoT), a novel and gene-targeted marker, has recently become the marker of choice in genetic diversity studies. In the present study, 31 SCoT primers were tested for their suitability in the genetic analysis of 21 coffee genotypes representing 18 species. A total of 647 distinct PCR amplified fragments were produced with a mean of 20.9 fragments per primer and 80.80% of which were polymorphic. The polymorphic information content of SCoT primers ranged from 0.16 to 0.86, with a mean value of 0.63. Resolving power ranged from 6.19 to 28.29, with a mean value of 20.2. Species-specific unique PCR amplified fragments were identified for 16 species, which could be used as genetic fingerprints. The genetic similarity among various coffee species calculated using the Dice similarity coefficient ranged between 0.60 and 0.89. The dendrogram constructed using the unweighted pair group of arithmetic means (UPGMA) clustered the 21 coffee genotypes into two major groups. The study revealed that Coffea jenkinsii, an indigenous species from India, showed the highest similarity with C. arabica, which is of Ethiopian origin. The results proved the suitability of SCoT markers in genetic analysis of coffee genotypes.

Keywords: Coffea species, DNA-based markers, genetic diversity, species-specific fragments, start codon targeted polymorphism (SCoT).

INTRODUCTION

The genus Coffea L. belongs to the family Rubiaceae that consists of more than 124 species (Razafinarivo et al., 2013). Most coffee species naturally occur in tropical forests of Africa, Madagascar and Mascarene islands, while some of the species are found in the Indian subcontinent, Southeast Asia and tropical Australasia (Davis et al., 2006, 2010, 2011; Davis, 2011). However, all the coffee species are not commercially cultivated except two main species Coffea arabica L. and C. canephora Pierre ex A. Froehner, which provides the global commodity for consumption (Mishra, 2019). Coffee is grown in about 10.2 million hectares of land spanning over 80 countries and is the second most important commodity in the world trade market after petroleum. In 2018, the total global production of green coffee was about 173.09 million bags (60 kg capacity) with an export earning of over US$ 30.1 billion during 2020 (ICO report 2020). More than 125 million people in the world derive their income directly or indirectly from the produce of this crop (Mishra, 2019).

Due to the prodigious amount of economic importance, intensive research has been carried out on evolutionary and genetic aspects of various coffee species by using conventional and molecular analysis of both nuclear and plastid genomes (Davis et al., 2007; Maurin et al., 2007; Tesfaye et al., 2007; Anthony et al., 2010; Davis et al., 2011; Nowak et al.,
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About 100 mg of tissue from the young leaves was ground to a fine powder using liquid nitrogen and transferred to 1.5 ml microcentrifuge tubes containing 750 µL of preheated extraction buffer (2% CTAB (w / v), 100 mM Tris-HCl (pH 8.0), 25 mM EDTA, 2 M NaCl and 0.1% beta-mercaptoethanol). The tubes containing the lysates were incubated in a hot water bath at 65°C for 45 min and then centrifuged at 6000 rpm for 15 min. The supernatant was transferred to a fresh tube, and equal volumes of chloroform: isooamy alcohol (24:1) was added. The samples were gently mixed by inverting the tubes and centrifuged at 6000 rpm for 15 min. The clear upper phase was transferred into a fresh 1.5 ml microcentrifuge tube and DNA was precipitated by adding to 0.7 volume of ice-cold isopropanol and incubated at -20°C for 15 min. The samples were centrifuged at 8000 rpm for 15 min, and the DNA pellet was washed

Plant material
All the 21 genotypes representing 18 coffee species were grown in the nursery of Tissue Culture and Plant Biotechnology Division, Mysore. The details of coffee genotypes, along with their codes and country of origin, are provided in Table 1.

Isolation of genomic DNA
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with 75% (v/v) ethanol for 10 min following which the tubes were centrifuged at 8000 rpm for 5 min, and the ethanol was discarded. The DNA pellet was dried at room temperature until traces of ethanol was removed and dissolved in 100 μl of TE buffer (10 mM–Tris and 1 mM–EDTA). RNase treatment was carried out by adding 5 μL of RNase enzyme solution and incubating at 37°C. The RNase was deactivated by heating the samples to 70°C. The quality of DNA was accessed by separating it on 0.8% agarose gel stained with ethidium bromide (0.5 µg / ml) and quantified using a UV spectrophotometer at 260 nm and 280 nm. The DNA samples were diluted to a working concentration of 10 ng/μl using sterilised distilled water and stored at -20°C for future use.

**SCoT marker analysis**

Thirty-one SCoT primers developed by Collard & Mackill (2009) (Table 2) were used for assessing their suitability in the genetic analysis of coffee species. The integrity and reproducibility of the bands amplified by SCoT primers were validated by conducting the PCR amplification at least two times. PCR amplification was carried out in a total reaction volume of 20 μL using Bio-Rad Thermal cycler S1000. PCR reaction mixtures contained 5.0 μL of template DNA (10 ng/μL), 3.0 μL of 3 μM primer, 2.0 μL of 2 mM dNTPs, 2.0 μL of 10X Taq buffer, 2.0 μL of 25 mM MgCl₂ (all from Thermo Fisher Scientific, Waltham, USA), and 0.5 μL of 3 units/μL Taq DNA polymerase enzyme (GeNei). Standard PCR cycling parameters were used, which includes an initial denaturation step of 5 min at 94°C, followed by 30 cycles of 94°C for 30 s, primer annealing at 48°C for 2 min, primer extension at 72°C for 2 min, and final extension of 15 min at 72°C. The PCR amplified products were mixed with 5 μL Bromophenol blue dye (99.5% deionised formamide, 10 mM EDTA pH 8, 0.05% Bromophenol blue, xylene-cyanol dye solution, 1μL pure, sterile water) and separated on 1.5% agarose gel (SeaKem, Rockland USA) containing 0.5 µg ethidium bromide/ml in 1 × TBE (Tris-HCl, Boric acid, EDTA) buffer. After electrophoresis, the gels were visualised and documented using Gel Doc System (BioRad) with a Multi Analyst software programme.

| Species | Species code | Place of origin/distribution | Conservation status |
|---------|--------------|------------------------------|---------------------|
| *C. arabica* L. ‘Kents’ | S-1 | Ethiopia/India | Vulnerable |
| *C. canephora* Pierre ex A. Froehner, cv. S. 274 | S-2 | West Tropical Africa/India | Least concern |
| *C. canephora* var. *ugandae* (Cramer) A.Chev. | S-3 | West Tropical Africa | Least concern |
| *C. canephora* var. *quillon* Philipp. | S-4 | West Tropical Africa | Least concern |
| *C. congensis* var. *froehneri* Pierre ex De Wild. | S-5 | West Central Africa Congo | Least concern |
| *C. eugenioides* S.Moore | S-6 | West Tropical Africa | Least concern |
| *C. zanguebariae* Lour. | S-7 | East Tropical Africa | Vulnerable |
| *C. racemosa* Lour. | S-8 | Southern Tropical Africa | Near threatened |
| *C. kapakata* (A. Chev.) Bridson | S-9 | West Angola | Vulnerable |
| *C. stenophylla* G.Don | S-10 | West Tropical Africa | Least concern |
| *C. salvatrix* Swynn.,& Philipson | S-11 | East Tropical Africa | Near threatened |
| *C. abeokutae* P. J. S. Cramer | S-12 | West Tropical Africa | Least concern |
| *C. liberica* Hiern | S-13 | West Tropical Africa | Least concern |
| *C. dewevrei* De Wild. & T.Durand | S-14 | Democratic Republic of Congo | Vulnerable |
| *C. arnoldiana* De Wild. | S-15 | West Central Africa | Least concern |
| *C. dewevrei* var. *excelsa* (A.Chev.) A.Chev. | S-16 | West Central Africa | Least concern |
| *C. bengalensis* Roxb.ex Schult. | S-17 | India | Endangered |
| *C. travancorensis* Wight & Arn. | S-18 | India | Endangered |
| *C. khasiana* (Korth.) Hook.f. | S-19 | India | Endangered |
| *C. wightiana* Wall. ex Wight & Arn. | S-20 | India | Endangered |
| *C. jenkinsii* Hook.f. | S-21 | India | Endangered |
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estimated as PIC = (1–p_i^2)/n, where n is the number of band positions analysed in all the species, p_i is the frequency of the banding pattern. The resolving power of a primer is Rp = \sqrt{I_b}, where I_b (band informativeness) takes the value of 1 – [2x (0.5–p)] and p is the ratio of samples sharing the band. A pairwise similarity matrix was constructed using the Dice similarity coefficient (Sneath & Sokal, 1973). The relationship between the species was displayed as a dendrogram constructed using NTsys 2.10e software (Rohlf, 1995) based on the Unweighted Pair Group Method using Arithmetic averages (UPGMA) with a bootstrap value of 1000.

The multivariate analysis by employing GenAIEX 6.5 software (Peakall & Smouse 2012) was performed to define relationships between 21 coffee genotypes employing Principal Coordinate Analysis (PCoA) using the Dice similarity coefficient.

RESULTS

In the present study, highly polymorphic fingerprinting patterns were obtained by using 31 SCoT primers in 21 genotypes belonging to 18 different coffee species (Figs 1–2).

The 31 primers produced 647 distinct scoreable fragments with the number of amplified fragments ranging from 7 (SCoT-10) to 30 (SCoT-4 and SCoT-17) with a mean of 20.87 fragments per primer (Table 3). Out of the total 647 amplified fragments, 544 were polymorphic, with a mean of 18 polymorphic fragments per primer. The amplified product size generated using different primers ranged from 300bp to 4kb. The per cent polymorphism ranged from 15% to a maximum of 100% with a mean of 80.80%.

Among 31 primers screened, six primers showed 100% polymorphism. To determine the informative and discriminatory capacity of each primer, polymorphic information content (PIC) and resolving power (RP) were calculated (Table 3). The PIC or DI was estimated as PIC = (1–p_i^2)/n, where n is the number of band positions analysed in all the species, p_i is the frequency of the banding pattern. The resolving power of a primer is Rp = \sqrt{I_b}, where I_b (band informativeness) takes the value of 1 – [2x (0.5–p)] and p is the ratio of samples sharing the band. A pairwise similarity matrix was constructed using the Dice similarity coefficient (Sneath & Sokal, 1973). The relationship between the species was displayed as a dendrogram constructed using NTsys 2.10e software (Rohlf, 1995) based on the Unweighted Pair Group Method using Arithmetic averages (UPGMA) with a bootstrap value of 1000.

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occurred in 90–100% frequency, whereas 0.1 to 4.99% of fragments occurred in 1% frequency across the genotypes (Table 4).

Thirty-one SCoT primers generated 62 unique fragments in 18 coffee species out of which five Indian species (C. bengalensis, C. travancorensis, C. wightiana, C. khasiana and C. jenkinsii) generated 18 unique fragments (Table 5). Among the Indian species, the maximum number of unique fragments was generated by C. bengalensis (7) followed by C. travancorensis, C. wightiana (4 each), and the least number of unique fragments were observed in C. khasiana (3). However, C. arabica, C. canephora var. ugandae, C. eugeniodes, C. stenophylla and C. jenkinsii failed to produce any unique fragments (Table 5). The genetic similarity calculated between different species ranged from 0.60 (C. abeokutae and C. khasiana) to 0.89 (C. canephora cv. S.274 and C. canephora var. ugandae, C. canephora var. quillon and C. canephora var. ugandae). Among the indigenous species, C. bengalensis showed the highest similarity of 0.78 with C. travancorensis, and C. wightiana, whereas the least genetic similarity of 0.66 was observed between C. bengalensis and C. jenkinsii.

Based on the SCoT marker data analysis, a strict consensus tree was constructed using the UPGMA clustering algorithm, and all the 21 coffee genotypes were grouped into two major clusters (Fig. 3). The first major cluster consisted of only two species C. dewevrei and C. khasiana. The second major cluster was divided into four minor clusters. The first minor cluster brought together C. bengalensis, C. wightiana and C. travancorensis. In the second minor cluster, C. abeokutae, C. arnoldiana and C. dewevrei, var. excelsa were grouped together. The third minor clus-
Table 3. Evaluation of 31 SCoT primers in genetic analysis of coffee species

| Primers | Total bands | Size range (bp) | Number of bands in each species (mean) | No. of polymorphic bands | Percentage of polymorphism | Resolution power | Polymorphic information content |
|---------|-------------|----------------|----------------------------------------|---------------------------|----------------------------|-----------------|-------------------------------|
| SCoT 1  | 18          | 450–3000       | 2–10 (6.43)                            | 17                        | 94.44                      | 12.86           | 0.81                          |
| SCoT 2  | 29          | 450–3000       | 9–16 (12.76)                           | 28                        | 96.55                      | 25.52           | 0.72                          |
| SCoT 3  | 21          | 500–3000       | 5–15 (9.38)                            | 19                        | 90.47                      | 18.76           | 0.71                          |
| SCoT 4  | 30          | 450–2500       | 10–19 (14.14)                          | 28                        | 93.33                      | 28.29           | 0.69                          |
| SCoT 5  | 28          | 500–3000       | 4–17 (11.95)                           | 28                        | 100.00                     | 23.90           | 0.74                          |
| SCoT 6  | 20          | 500–4000       | 8–16 (11.95)                           | 17                        | 85.00                      | 23.90           | 0.53                          |
| SCoT 7  | 19          | 700–2500       | 4–11 (7.00)                            | 17                        | 89.47                      | 14.00           | 0.76                          |
| SCoT 8  | 27          | 700–3000       | 6–12 (08.43)                           | 27                        | 100.00                     | 16.86           | 0.86                          |
| SCoT 9  | 19          | 800–3000       | 4–12 (7.67)                            | 15                        | 78.94                      | 15.33           | 0.73                          |
| SCoT 10 | 7           | 1400–3000      | 2–6 (3.10)                             | 5                         | 71.42                      | 6.19            | 0.66                          |
| SCoT 11 | 16          | 450–3500       | 8–13 (11.76)                           | 8                         | 50.00                      | 23.62           | 0.30                          |
| SCoT 12 | 20          | 700–3000       | 8–16 (12.67)                           | 16                        | 80.00                      | 25.33           | 0.49                          |
| SCoT 13 | 13          | 500–3000       | 11–12 (13.33)                          | 2                         | 15.38                      | 22.67           | 0.16                          |
| SCoT 14 | 11          | 500–2500       | 7–10 (8.14)                            | 4                         | 36.36                      | 16.29           | 0.29                          |
| SCoT 15 | 22          | 900–2500       | 7–17 (11.81)                           | 18                        | 81.81                      | 23.62           | 0.59                          |
| SCoT 16 | 30          | 380–2800       | 9–17 (13.33)                           | 28                        | 93.33                      | 26.67           | 0.71                          |
| SCoT 17 | 19          | 500–3000       | 10–16 (12.62)                          | 11                        | 57.89                      | 25.24           | 0.42                          |
| SCoT 18 | 28          | 550–2200       | 9–19 (12.71)                           | 24                        | 85.71                      | 25.43           | 0.67                          |
| SCoT 19 | 20          | 400–2500       | 5–14 (9.86)                            | 16                        | 76.19                      | 19.71           | 0.63                          |
| SCoT 20 | 26          | 100–2500       | 4–17 (9.38)                            | 26                        | 100.00                     | 18.76           | 0.80                          |
| SCoT 21 | 19          | 500–2500       | 3–12 (8.09)                            | 18                        | 94.73                      | 16.19           | 0.73                          |
| SCoT 22 | 27          | 300–3500       | 9–19 (13.62)                           | 21                        | 77.77                      | 27.24           | 0.61                          |
| SCoT 23 | 15          | 550–2800       | 10–14 (11.90)                          | 6                         | 40.00                      | 23.81           | 0.27                          |
| SCoT 24 | 22          | 300–2000       | 7–2 (9.43)                             | 21                        | 95.45                      | 18.86           | 0.68                          |
| SCoT 25 | 17          | 550–2500       | 5–9 (6.86)                             | 14                        | 82.35                      | 13.71           | 0.71                          |
| SCoT 26 | 17          | 520–2500       | 4–11 (8.67)                            | 17                        | 100.00                     | 17.33           | 0.66                          |
| SCoT 27 | 21          | 700–2800       | 3–11 (6.57)                            | 21                        | 100.00                     | 13.14           | 0.86                          |
| SCoT 28 | 20          | 550–2500       | 9–13 (11.43)                           | 15                        | 75.00                      | 22.86           | 0.56                          |
| SCoT 29 | 21          | 500–3000       | 5–13 (9.91)                            | 19                        | 90.47                      | 19.81           | 0.68                          |
| SCoT 30 | 22          | 550–2800       | 3–10 (6.86)                            | 22                        | 100.00                     | 13.71           | 0.84                          |
| SCoT 31 | 22          | 550–3000       | 11–18 (13.33)                          | 16                        | 72.72                      | 26.67           | 0.53                          |
| Total   | 647         | 201–427        | 313.09                                 | 544                       | 2504.78                    | 626.3           | 19.40                         |
| Mean    | 20.87       | 6.5–14.0       | 10.10                                  | 18                        | 80.80                      | 20.20           | 0.63                          |

DISCUSSION

Characterisation and evaluation of genetic diversity among coffee species are essential to understand the value of germplasm collections and their usefulness in breeding programmes. Previously, the genetic diversity of coffee species has been studied by using RAPD, ISSR and SRAP markers (Mishra et al., 2011a, b). However, with the rapid development...
Table 4. Details of species-specific fragments amplified by SCoT markers in different coffee species

| Coffee species                          | SCoT marker                                      |
|----------------------------------------|--------------------------------------------------|
| C. arabica L. ‘Kents’                  | –                                                |
| C. canephora Pierre ex A.Froehner, cv. S. 274 | SCoT-28-3500                                    |
| C. canephora var. ugandae (Cramer) A.Chev. | –                                                |
| C. canephora var. quilllon Philipp.    | SCoT-18-720, SCoT-19-1400, SCoT-19-600           |
| C. congensis var. froehneri Pierre ex De Wild. | SCoT-9-2900, SCoT-34-1000                      |
| C. eugenioides S.Moore                 | –                                                |
| C. zanguebariae Lour.                  | SCoT-1-900, SCoT-2-680, SCoT-11-1400, SCoT-14-1200, SCoT-17-1300, SCoT-19-550, SCoT-28-480, SCoT-34-600, SCoT-39-2200, SCoT-62-710 |
| C. racemosa Lour.                      | SCoT-6-720, SCoT-22-1500, SCoT-36-2500           |
| C. kapakata (A. Chev.) Bridson         | SCoT-7-1000, SCoT-9-1900                         |
| C. stenophylla G.Don                   | –                                                |
| C. salvatrix Swynn.& Philipson         | SCoT-5-750, SCoT-11-900, SCoT-31-1200            |
| C. abeokutae P. J. S. Cramer           | SCoT-20-1600, SCoT-61-1000                       |
| C. liberica Hiern                      | SCoT-2-600, SCoT-4-300, SCoT-6-800, SCoT-31-2500, SCoT-31-600, SCoT-62-1000, SCoT-63-2800 |
| C. dewevrei De Wild. & T.Durand        | SCoT-6-750, SCoT-7-900, SCoT-7-800, SCoT-18-1400, SCoT-20-800 |
| C. arnoldiana De Wild.                | SCoT-3-600, SCoT-17-390, SCoT-34-1200, SCoT-48-650 |
| Coffea dewevrei var. excelsa (A.Chev.) A.Chev. | SCoT-20-650, SCoT-21-1400                      |
| C. bengalensis Roxb.ex Schult.         | SCoT-2-400, SCoT-7-1200, SCoT-8-3100, SCoT-17-500, SCoT-20-750, SCoT-20-710, SCoT-21-1150 |
| C. travancorensis Wight & Arn.        | SCoT-19-2400, SCoT-21-1200, SCoT-22-1300, SCoT-62-2500 |
| C. khasiana (Korth.) Hook.f.           | SCoT-3-680, SCoT-15-700, SCoT-62-550             |
| C. wightiana Wall. ex Wight & Arn.    | SCoT-2-680, SCoT-5-1000, SCoT-5-1100, SCoT-61-1400 |
| C. jenkinsii Hook.f.                   | –                                                |

in the field of genomics, new functional markers targeting the genes and promoters are available and used in the genetic analysis of many plant species with better resolution and reproducibility (POCZAI et al., 2013). SCoT marker assay preferentially targets the coding sequences in the genome, because the primers are based on the short conserved region surrounding the ATG translation start codon (COLLARD & MACKILL, 2009; XIONG et al., 2009). Therefore, the polymorphism revealed by the SCoT marker is directly related to the diversity at the gene level, which could be possibly involved in phenotypic trait variation (ANDERSEN & LUBBERSTEDT, 2003). Although a large amount of data was obtained with SCoT markers for various plant species, the genetic diversity and breeding value of the coffee gene pool using the SCoT marker is yet to be ascertained. The present study is the first report on the use of SCoT markers to estimate the breeding value of the coffee gene pool.

The data generated in the present study by screening 21 coffee genotypes with 31 SCoT revealed a mean of 80.80% polymorphic fragments. The percentage polymorphism obtained using SCoT markers was low as compared to SRAP and ISSR marker analysis, wherein 96.12 and 93.06% polymorphism was obtained, respectively (MISHRA, 2011b). The lower percentage of mean polymorphism could be due to the differential working principle of the SCoT marker system, wherein only the functional domains of genes were targeted. The functional domains of genes comprise a smaller portion of the total genome. They are more conserved, unlike non-functional regions, resulting in a lower polymorphic percentage of the marker system. Nevertheless, the polymorphism exhibited by the SCoT marker system is more critical as it is directly linked with gene sequence, which in turn governs phenotypic traits. However, the mean resolving power (Rp) detected by SCoT (20.2) markers was much higher than in previous studies employing SRAP (9.74) and ISSR (8.64) (MISHRA et al., 2011b). The higher Rp of SCoT can be related to the amplification of both dominant and co-dominant fragments in the marker system (POCZAI et al., 2013). Similarly, the PIC values, which are used as allele
Table 5. Frequency class of PCR amplified bands generated by analysing 21 coffee genotypes using SCoT assay

| Primers | Frequency class (%) | Total |
|---------|--------------------|-------|
|         | 0.1–4.99 | 5.0–9.99 | 10.0–29.99 | 30.0–49.99 | 50.0–69.99 | 70.0–89.99 | 90.0–100 |       |
| SCoT 1  | 1  | 2  | 43  | 28  | 22  | 18  | 21  | 135   |
| SCoT 2  | 4  | 6  | 12  | 56  | 63  | 86  | 41  | 268   |
| SCoT 3  | 2  | 6  | 21  | 28  | 60  | 18  | 62  | 197   |
| SCoT 4  | 1  | 6  | 44  | 35  | 64  | 67  | 80  | 297   |
| SCoT 5  | 3  | 2  | 27  | 31  | 114  | 34  | 40  | 251   |
| SCoT 6  | 3  | 2  | 6  | 15  | 25  | 117  | 83  | 251   |
| SCoT 7  | 4  | 0  | 27  | 14  | 23  | 18  | 61  | 147   |
| SCoT 8  | 1  | 6  | 61  | 51  | 25  | 33  | 0  | 177   |
| SCoT 9  | 2  | 4  | 34  | 25  | 12  | 0  | 84  | 161   |
| SCoT 10 | 0  | 2  | 10  | 0  | 11  | 0  | 42  | 65    |
| SCoT 11 | 2  | 4  | 0  | 0  | 0  | 17  | 225  | 248   |
| SCoT 13 | 0  | 6  | 11  | 20  | 36  | 51  | 142  | 266   |
| SCoT 14 | 1  | 0  | 10  | 0  | 17  | 210  | 238  |       |
| SCoT 15 | 1  | 2  | 3  | 0  | 0  | 18  | 147  | 171   |
| SCoT 16 | 0  | 8  | 11  | 33  | 37  | 18  | 141  | 248   |
| SCoT 17 | 3  | 8  | 31  | 35  | 75  | 48  | 80  | 280   |
| SCoT 18 | 2  | 0  | 17  | 10  | 13  | 15  | 208  | 265   |
| SCoT 19 | 4  | 6  | 28  | 9  | 23  | 93  | 104  | 267   |
| SCoT 20 | 5  | 0  | 19  | 16  | 14  | 48  | 105  | 207   |
| SCoT 21 | 3  | 4  | 45  | 34  | 40  | 51  | 20  | 197   |
| SCoT 22 | 2  | 0  | 31  | 32  | 12  | 52  | 41  | 170   |
| SCoT 28 | 2  | 4  | 41  | 14  | 39  | 0  | 186  | 286   |
| SCoT 30 | 0  | 4  | 0  | 9  | 13  | 35  | 189  | 250   |
| SCoT 31 | 3  | 8  | 21  | 7  | 13  | 66  | 80  | 198   |
| SCoT 34 | 3  | 4  | 28  | 0  | 12  | 34  | 63  | 144   |
| SCoT 36 | 1  | 2  | 7  | 39  | 28  | 66  | 39  | 182   |
| SCoT 39 | 1  | 8  | 22  | 33  | 58  | 16  | 0  | 138   |
| SCoT 48 | 1  | 0  | 21  | 33  | 24  | 18  | 143  | 240   |
| SCoT 61 | 2  | 4  | 22  | 19  | 64  | 17  | 80  | 208   |
| SCoT 62 | 4  | 10  | 29  | 26  | 39  | 16  | 20  | 144   |
| SCoT 63 | 1  | 0  | 36  | 0  | 50  | 48  | 145  | 280   |
| Total   | 62  | 118 | 708  | 662  | 1009  | 1135  | 2882  | 6576   |

diversity and frequency among different coffee species, varied from 0.163 to 0.864 with a mean of 0.625 and was lower than the mean PIC value obtained using ISSR (0.81) and SRAP (0.81) marker analysis (Mishra et al., 2011b). The PIC value obtained in our study was much higher than the PIC value (0.33) obtained in Cocos nucifera L. using SCoT marker analysis (Rajes et al., 2015).

The clustering pattern of 21 coffee genotypes in UPGMA dendrogram is largely in congruence with the results of PCoA analysis. In both dendrogram and PCoA plot, the Indian wild coffee species formed a separate group (I). African wild coffee species were arranged in II and III groups. In contrast, the cultivated species C. arabica and C. canephora were placed along with C. eugenioides and C. congensis in group IV (Fig. 4). Both C. canephora and C. eugenioides were considered as progenitors of C. arabica, and similarly, C. congensis is close to C. canephora and used in the breeding programme (Mishra, 2019). Hence it is quite apparent that all the three species are grouped together. The results indicated that SCoT markers could be efficiently used to identify polymorphism between coffee species.

In the present study, 31 SCoT primers amplified 62 unique fragments in 15 species, with a maximum number of 10 unique fragments generated by C. zanguebariae and followed by C. liberica and C. bengalensis. However, no unique fragments were generated in C. arabica, C. congensis, C. steno-
Fig. 3. Dendrogram generated using the unweighted pair group method with arithmetic average analysis (UPGMA) showing relationships among different coffee species using SCoT data. The numbers at the nodes indicate the bootstrap value. Coffee species listed in Table 1.

Fig. 4. Principal Coordinate Analysis (PCoA) of SCoT marker data obtained by evaluation of coffee species. Coffee species listed in Table 1.
Assessment of the suitability of molecular SCoT markers for genetic analysis of coffee species

phylla and C. jenkinsii, which could be attributed to the insufficient number of SCoT primers used. These unique fragments could be used as diagnostic fingerprinting tools to discriminate species. A close look at the fingerprinting pattern generated by different SCoT primers revealed that SCoT 20, SCoT 19, SCoT 7, SCoT 62 and SCoT 2 are more informative markers as they amplified the maximum number of unique fragments across the species (Table 5). Since single accession per species was used in the present study, the study could facilitate in selecting the highly polymorphic SCoT primers suitable for genetic analysis of coffee species. In future research, more number of plants belonging to each species could be included for assessing the total genetic variability in the coffee gene pool.

In the present study, the pairwise similarity coefficient between different coffee species ranged from 0.60 to 0.89. The highest similarity of 0.89 was obtained between C. canephora cv. S.274 and C. canephora var. ugandae, and C. canephora var. quillon and C. canephora var. ugandae using SCoT marker. However, in a previous study, the pairwise similarity coefficient between different coffee species ranged from 0.11-0.90 using SRAP and 0.27-0.89 using ISSR markers in coffee (Mishra et al., 2011b). The highest similarity (0.90) was obtained between C. canephora and C. canephora var. laurantii using SRAP and (0.89) between C. canephora and C. congensis using ISSR assay. The genotypes with the highest coefficient of similarity using the SCoT, SRAP and ISSR assays had a common origin (West Tropical Africa). In the present study, the lowest genetic similarity was observed between C. abeokutae and C. khasiana based on the SCoT marker data analysis. In the previous study, the lowest genetic similarity was established between C. bengalensis and C. liberica using the SRAP marker and between C. wightiana and C. congensis using ISSR marker analysis (Mishra et al., 2011b). In this study, a narrow range of similarity coefficient was observed between five Indian species, which coincided with the previous research employing SRAP and ISSR markers (Mishra et al., 2011b). Further, Indian species, C. bengalensis and C. travancorensis were closely grouped in the UPGMA dendrogram, which has been confirmed in the previous study. However, C. jenkinsii, an indigenous Indian species, showed the highest similarity to C. arabica, originated from Ethiopia and widely cultivated in India. The maximum similarity observed between C. jenkinsii and C. arabica ‘Kents’ could be due to the close similarity at the functional loci revealed by SCoT marker analysis.

CONCLUSIONS

The extent of the genetic variability of coffee species at the functional level was successfully tested for the first time using SCoT markers. The markers with their high resolving power successfully evaluated the genotypes and generated a high level of polymorphism. The assay successfully differentiated coffee species based on functional diversity and geographical origin. The species-specific amplicons produced by the SCoT marker assay could be used as fingerprints of species identification. Since the SCoT assay targets functional domains, the sequence information obtained by sequencing differential fragments generated across the species can be used in selecting the right parental material in the coffee improvement and conservation programmes.

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AUTHOR’S CONTRIBUTION

The corresponding author designed and collected the material for the experiment and wrote the manuscript. AKH and PJ carried out the experiments and participated in manuscript preparation. All authors read and approved the manuscript.

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MOLEKULINIŲ SCOT ŽYMENŲ TINKAMUMO COFFEE RŪŠIŲ GENETINEI ANALIZEI ĮVERTINIMAS

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Santrauka

Pastaruoju metu genetinės įvairovės tyrimuose vis plačiau naudojamas naujas startinio genų kodono (SCoT) gentinių žymenų metodas. Šiame darbe buvo išbandytas 31 SCoT genetinio pradmens efektyvumas, identifikuojant 18 kavos rūšių 21 genotipą. Buvo paruošti 647 genetiniai fragmentai, vidutiniškai 20,9 vienam pradmeniui, iš kurių 80,8% buvo polimorfiški. Nustatytas SCoT pradmenų polimorfizmas kito nuo 0,16 iki 0,86, o vidutinė vertė buvo 0,62. Skiriamoji pradmenų geba svyravo nuo 6,19 iki 28,29, vidutinė vertė – 20,2. Šešiolikoje kavos rūšių genotipų buvo nustatyti specifiniai DNR fragmentai, kurie gali būti naudojami kaip genetiniai „pirštų atspaudai“ rūšių identifikavimui. Kavos rūšių genetinis panašumas apskaičiuotas, naudojant Dice panašumo koeficientą, buvo 0,60–0,89. Dendrograma sudaryta remiantis porų grupavimo pagal aritmetinius vidurkius su vie nodais svoriais metodu (UPGMA) parodė, kad visi kavos genotipai buvo suskirstyti į dvi pagrindines grupes. Statistinė tirtų genotipų vizualizacija parodė, kad Indijos vietinė rūsis Coffea jenkensii genetiškai artima etiopinės kilmės C. arabica. Gauti rezultatai patvirtino molekulinį SCoT žymenų metodo tinkamumą tolesnei kavos rūšių genetinei analizei.