

Rapid Communication

Transcriptome-Scale Developing of Simple Sequence Repeat Markers in Coffee Arabica

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Coffee is an important beverage crop in the world and the main commercially cultivated species are *Coffea arabica* (arabica) and *C. canephora* (robusta). *C. arabica* is a dominant coffee specie with a high potential for genetic improvement and it normally starts fruiting three years after planting, thereby significantly extending its breeding process. Thus, molecular marker assisted selection would efficiently accelerate the process. In the present study, we conducted a large-scale development of Simple Sequence Repeat (SSR) markers according to a high quality 454-pyrosequencing database. 1032 SSR loci were identified from 929 unigenes (6.66% of 13951). Mononucleotides (500, 48.45%), trinucleotides (411, 39.83%) and dinucleotides (98, 9.49%) were the main SSR types. The most abundant SSR motif was A/T (490, 47.57%), followed by AAG/CTT (126, 12.23%), ACG/CGT (88, 8.54%), ACT/AGT (70, 6.79%) and AG/CT (61, 5.92%). A total of 115 pairs of reported SSR primers were utilized for the SSR validation and two matched our results. Our work will expand the number of SSR loci and benefit relevant studies by applying these loci in *C. arabica*.

Keywords: *Coffea arabica*; Transcriptome; Simple sequence repeat markers**Abbreviations**

AFLP: Amplified Fragment Length Polymorphism; SSR: Simple Sequence Repeat; SNPs: Single Nucleotide Polymorphisms; MISA: MicroSatellite; COG: Clusters of Orthologous Groups; EST: Expressing Sequence Tag

Introduction

Coffee is an important beverage crop in the world and globally, the two main species of coffee that are commercially produced are *Coffea arabica* (arabica) and *C. canephora* (robusta). *C. arabica*, the dominant coffee species, contains a high potential for genetic improvement and normally starts fruiting about three years after being planted, which has significantly extended the crop breeding process [1]. Thus, molecular marker assisted selection would efficiently accelerate the process [2]. To date, molecular markers have been successfully utilized in germplasm evaluation of coffee, such as Amplified Fragment Length Polymorphism (AFLP) and Simple Sequence Repeat (SSR) [3,4]. But the markers employed in these studies are still on small amounts. In recent years, the fast development of next generation sequencing technology makes it possible for large scale marker-based germplasm evaluation [5]. A recent study has reported 1444 Single Nucleotide Polymorphisms (SNPs) associated with caffeine content by a draft genome sequence of *C. arabica* [1]. Although this genome data is still not released with the published study, transcriptome data in previous studies makes it possible for large scale SSR marker developing [6,7]. In this study, we conducted a large-scale development of SSR markers based on a high quality 454-pyrosequencing database [6]. This work has expanded the number of SSR loci which could complement available information and enhance future research efforts on the utilization of these loci in *C. arabica*.

Materials and Methods**Identification and characterization of SSRs**

De novo assembly of 13,951 unigenes of *C. arabica* CIFC H147/1 from a previous study [6] was utilized for SSR detection by using the MicroSatellite (MISA) identification tool with default criteria [8]. The maximal number of bases interrupting two SSR motifs in a compound microsatellite was 20. All the SSR-contained unigenes were searched against the Clusters of Orthologous Groups (COG) data by BLASTx set and then classified into COG categories with a cutoff Expected value (E-value) of $1e^{-5}$ [9].

SSR validation with previous study

Primers from previous studies were downloaded to validate the SSRs in the present study [5,10]. These primers were transformed into FASTA format and each primer considered as a single sequence. Then, comparison was conducted by BLASTn-short procedure to search all the unigenes with SSR loci [11]. Each pair of primers that matched the same sequence with a sequence similarity over 95% was selected for SSR loci comparison. If these SSRs were the same with our result, they would be highlighted and also a reliable proof for our work.

Results**Identification of SSRs**

A total of 13,951 unigenes from the previous study were utilized for SSR loci screening. As a result, we found 1,032 SSR sites in 929 unigenes with a frequency of 1 SSR per 8.53 kb sequences (Table 1). Among these, 87 unigenes had more than one SSR loci. 79, 6 and 4 sequences contained 2, 3 and 4 SSR loci, respectively. We separately counted the number of unigenes that contained SSR loci and used

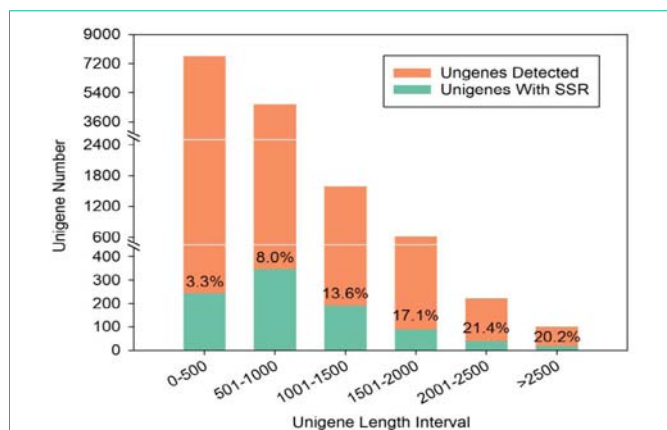


Figure 1: Length distribution of unigenes for SSR detection (orange) and with SSR loci (green). The percentages in bar represent the ratio of unigenes with SSR to unigenes for SSR detection.

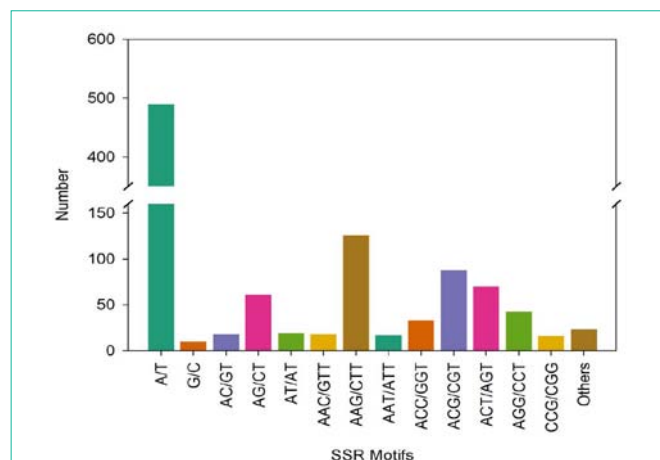


Figure 2: Numbers of SSR motifs.

Table 1: Results for SSR detection.

Item	Count
Total unigenes	13,951
Total size of unigenes (bp)	8,808,436
SSR loci	1,032
Unigenes contain SSR	929
Sequences containing more than 1 SSR	87
Mononucleotide	500
Dinucleotide	98
Trinucleotide	411
Tetranucleotide	15
Pentanucleotide	5
Hexanucleotide	3

for SSR detection at different length intervals (Figure 1). 3.3%, 8.0%, 13.6%, 17.1%, 21.4% and 20.2% of unigenes contained SSR loci at 6 different length intervals, respectively.

Mononucleotides (500, 48.45%), trinucleotides (411, 39.83%) and dinucleotides (98, 9.49%) were the main types of the 1,032 SSR loci (Table 1). Meanwhile, only 15 tetranucleotides, 5 pentanucleotides and 3 hexanucleotides were observed. Among all the identified SSR motifs, the most abundant SSR motif was A/T (490, 47.57%), followed by AAG/CTT (126, 12.23%), ACG/CGT (88, 8.54%), ACT/AGT (70, 6.79%) and AG/CT (61, 5.92%) (Figure 2).

COG annotation

The 929 unigenes with SSR loci were classified into 25 COG categories (Figure 3). Among these, function unknown (of those poorly characterized and labeled in green, 456) was the most abundant category. And general function prediction only (of those poorly characterized and labeled in green, 62), posttranslational modification, protein turnover, chaperones (of cellular processes and signaling that were labeled in blue, 54), signal transduction mechanisms (of cellular processes and signaling that were labeled in blue, 50) and translation, ribosomal structure and biogenesis (of information storage and processing that were labeled in yellow, 47)

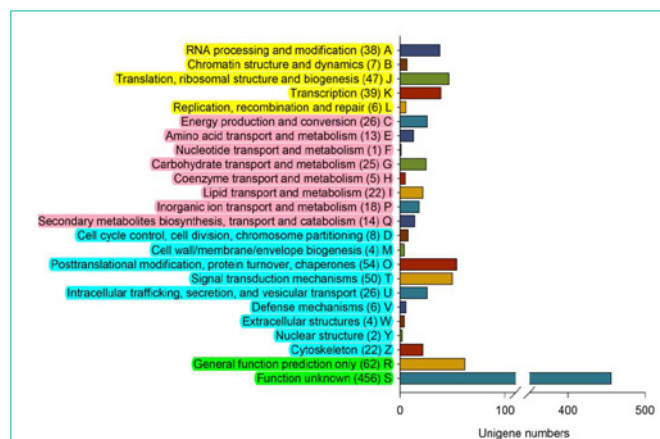


Figure 3: COG classification of the unigenes that containing SSRs and Arabic numerals represent unigenes numbers. The functional classifications were listed on the left of y-axis, among which consist of the four functional categories were labeled in yellow (information storage and processing), pink (metabolism), blue (cellular processes and signaling) and green (poorly characterized), respectively.

were at middle amount.

SSR validation

A total of 115 pairs of published primers were used for SSR validation [3,10]. As a result, two pairs matched unigenes in the current study (Table 2). They shared the same SSR loci but with different sizes of PCR amplification products in C1FC H147/1, respectively.

Discussion

In plant research works, more SSR markers are necessary for the construction of high-density linkage map and application in breeding processes. In this research, we successfully conducted a large-scale development of SSR markers according to previous transcriptome database [6]. The 1,032 Expressed Sequence Tag (EST)-SSR loci have significantly expanded SSRs for *C. arabica*. Only 6.66% of all the unigenes contained SSR loci, which is relatively lower than other plants, such as 12.29% in kenaf and 7.25% in ramie [12,13]. This observation might have resulted due to the sequence length. The

Table 2: Validated SSR loci with previous study [10].

SSR Loci	Units	Present Study						Previous Study	
		Name	Unigenes	Length	Start	Stop	Size	Name	Size
AG	6	Contig20847.505.ag.6	Contig20847	581	505	516	140	SSR03	142,148
AAAGG	3	Contig21585.776.aaagg.3	Contig21585	1160	776	790	135	SSR06	126

sequence lengths of 11758 unigenes were not over 1000 bp, among which, less than 8% unigenes contained SSR loci. Meanwhile, more than 13% of those unigenes contained SSR loci with sequence lengths over 1000 bp (Figure 1). The result indicated that long unigenes were very possible to contain more SSR loci than short unigenes, which might be also responsibility for rare appearance of tetranucleotides, pentanucleotides and hexanucleotides. Besides, the motifs with high GC region were also infrequent, such as G/C and CCG/CGG. We speculated that a regular successive G/C array might not reach over 30 bp in ESTs of *C. arabica* [13].

We further classified the COG categories of the 929 unigenes with SSR loci and found that more than half of the unigenes were characterized as function unknown and general function prediction only, which were relatively lower than the previous study in ramie [13]. The poorly annotated unigenes have restricted the functional characterization of these EST-SSR loci. Therefore, we collected a series of reported SSR markers to validated them. Interestingly, our SSR loci didn't match any of the 103 SSR loci from a previous EST-derived SSR marker development [3]. We just found two (2) pair of SSR markers consistent with our result (Table 2). SSR03 has been reported as a polymorphic SSR loci and SSR06 without polymorphism [10]. However, SSR06 actually has different amplification size in C1FC 147/1, which means it should be considered as a polymorphic SSR loci. In fact, there are still a lot of reported SSR markers that has been utilized for validating our result. More works would be needed in future study, especially PCR amplification. This study has supported a framework and guidance for further study.

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