Characterization and Transferable Utility of Microsatellite Markers in the Wild and Cultivated Arachis Species

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Abstract

Microsatellite or simple sequence repeat (SSR) is one of the most widely distributed molecular markers that have been widely utilized to assess genetic diversity and genetic mapping for important traits in plants. However, the understanding of microsatellite characteristics in Arachis species and the currently available amount of high-quality SSR markers remain limited. In this study, we identified 16,435 genome survey sequences SSRs (GSS-SSRs) and 40,199 expressed sequence tag SSRs (EST-SSRs) in Arachis hypogaea and its wild relative species using the publicly available sequence data. The GSS-SSRs had a density of 159.9 – 239.8 SSRs/Mb for wild Arachis and 1,015.8 SSRs/Mb for cultivated Arachis, whereas the EST-SSRs had the density of 173.5 – 384.4 SSRs/Mb and 250.9 SSRs/Mb for wild and cultivated Arachis, respectively. The trinucleotide SSRs were predominant across Arachis species, except that the dinucleotide accounted for most in A. hypogaea GSSs. From Arachis GSS-SSR and EST-SSR sequences, we developed 2,589 novel SSR markers that showed a high polymorphism in six diverse Arachis accessions. A genetic linkage map that contained 540 novel SSR loci and 105 anchor SSR loci was constructed by case of a recombinant inbred lines F6 population. A subset of 82 randomly selected SSR markers were used to screen 39 wild and 22 cultivated Arachis accessions, which revealed a high transferability of the novel SSRs across Arachis species. Our results provided informative clues to investigate microsatellite patterns across A. hypogaea and its wild relative species and potentially facilitate the germplasm evaluation and gene mapping in Arachis species.

Introduction

Cultivated peanut (Arachis hypogaea L.) is an important oil crop in the world for its direct consumption in the food industry and edible oil in cooking. It is widely cultivated in more than 100 countries with a global annual production of 45.7 Mt over an area of 25.4 Mha (http://faostat.fao.org/faostat/collections?subset=argriculture 2013). Peanut belongs to the genus Arachis containing at least 80 species that are divided into nine taxonomic sections based on
morphological variation, geographical distribution and cross-compatibility [1]. All wild Arachis species are diploid (2n = 2x = 18, 2n = 2x = 20) except A. monticola and certain species in section Rhizomatosae, whereas peanut is an allotetraploid species (AABB, 2n = 4x = 40) harboring A and B genomes [1]. Cytology and molecular studies have indicated that cultivated peanut results from a single hybridization event between the wild diploid species A. duranensis (AA, 2n = 2x = 20) and A. ipaënsis (BB, 2n = 2x = 20) followed by spontaneous chromosome duplication [2, 3]. The studies of genetic diversity in the Arachis genus have revealed that cultivated peanut possesses a narrow genetic base compared with wild Arachis species [4–7] perhaps due to the bottleneck effect in domestication and little natural gene exchange between wild Arachis species and cultivated peanut. Microsatellites, or simple sequence repeats (SSRs), defined as 1–6 nucleotides tandem repeats in the genomes, are mainly caused by replication slippage, leading to addition or removal of repeat motifs in plant genomes. Thus, the certain number and length of SSRs were probable to reflect the evolutionary history in particular species and its relatives [8].

Since no reference genome was currently available in any of Arachis species, the SSRs were still the preferable markers in assessing genetic diversity, genetic mapping and marker-assisted selection of important traits as SSRs are multi-allelic, easily detectable by PCR, abundantly distributed in genomes and codominantly inherited [9]. Although the Arachis community has made large efforts to develop microsatellite markers [10–18], there was little microsatellite characterization using a large number of sequences in the wild and cultivated Arachis species [18] and the understanding of microsatellite characteristics in Arachis species remains ambiguous. Furthermore, despite a large number of QTLs identified for important traits [19–25], the publicly available SSR markers remains insufficient for studies in gene fine-mapping and genome wide association mapping.

Currently, a total of 65,111 genome survey sequences (GSSs) and 281,115 expressed sequence tags (ESTs) of Arachis species are publicly available in NCBI (www.ncbi.nlm.nih.gov) [accessed 19 December 2014]. In this study, GSSs and ESTs in Arachis species were utilized to perform microsatellite characterization and marker development. The main objectives of this study were: (a) to characterize and compare the frequency, type of microsatellites in the assembled GSSs and ESTs of Arachis species; (b) to develop a set of novel SSR markers and validate the reliability and polymorphisms using six A. hypogaea accessions; (c) to genetically map the novel SSRs on the linkage groups of A. hypogaea by linkage analysis; (d) to test the utility of the novel SSR markers across the A. hypogaea and its wild relative species.

Materials and Methods

Plant materials and DNA isolation

A set of six diverse peanut accessions (Fuchuan Dahuasheng, ICG6375, Xuhua 13, Zhonghua 6, Zhonghua 10 and ICG12625) were used to screen the polymorphisms of SSR markers. These varieties have been used as parents for three established mapping populations. Fuchuan Dahuasheng (A. hypogaea var. hirsuta), Xuhua 13 (A. hypogaea var. hypogaea) and Zhonghua 10 (A. hypogaea var. vulgaris) are cultivars with large pods and seeds. Zhonghua 6 (A. hypogaea var. vulgaris) is a cultivar with small pods and seeds. ICG6375 (A. hypogaea var. vulgaris) and ICG12625 (A. hypogaea var. aequatoriana) are groundnut varieties received from the International Crop Research Institute for the Semi-arid Tropics. ICG6375 has small pods and seeds and ICG12625 has high plant height and dark purple testa. A recombinant inbred line (RIL) F6 population (n = 140) derived from the cross by Zhonghua 10 and ICG12625 was used to construct genetic linkage map. Finally, a subset of SSR markers evenly distributed on the linkage map were selected to evaluate the SSR transferability in the different Arachis species, containing 39 wild Arachis accessions representing five type genomes (A, B, P, E, and AB genome) and
22 cultivated peanut accessions. Detailed information for the 61 *Arachis* accessions was listed in S1 Table.

Genomic DNA was extracted from young leaves of these above mentioned accessions using a modified cetyltrimethyl ammonium bromide (CTAB) method. The integrity and quality of the DNA was evaluated on a 1% agarose gel by comparison with uncut lambda DNA.

**Source of sequences and SSR identification**

The GSSs and ESTs of *Arachis* species were downloaded in FASTA format from GenBank and subsequently used for SSR mining and marker development. These sequences downloaded were utilized to identify and select the SSR-containing sequences using SPUTNIK software (http://espressosoftware.com/pages/sputnik.jsp). The criteria for SSR selection were set at six repeats for dinucleotides and four repeats for tri-, tetra- and pentanucleotides. To remove the redundant SSR-containing sequences, the repetitive sequences were masked using RepeatMasker (http://www.repeatmasker.org) and then the masked sequences were assembled using CAP3 software (http://pbil.univ-lyon1.fr/cap.php) with overlap length cutoff of 40 nucleotides and overlap percent identity cutoff of 95. After removing redundancy, the resulting consensus sequences of contigs and singletons were again mined for SSRs.

**Development of SSR marker and polymorphism detection**

The above mentioned non-redundant SSR-containing sequences were employed to design PCR primers using Primer3 software [26]. The primer length was between 18 and 23 nucleotides with an optimum size of 20 nucleotides. The melting temperatures ranged from 50 to 70°C with an optimum temperature of 55°C. The GC content varied from 30% to 70% with an optimum GC content of 50%. The predicted PCR products ranged from 100 to 400 bp. After filtering out the SSRs with identical primer sequence to the publicly released SSRs in *Arachis* species, the novel SSR markers were identified in the present study and designated as “AGGS”, representing *Arachis* genus GenBank sequence.

The novel SSRs were tested for the reliability and polymorphism using a set of six diverse peanut accessions mentioned above. PCR amplifications were performed in a volume of 10 μl including 50 ng genomic DNA, 1 × Taq buffer, 2 mM MgCl₂, 0.2 mM dNTPs, 0.2 μM each primer and 0.25 U Taq DNA polymerase. Then the amplification was conducted by the ‘touch-down’ method, with the following thermal profile: initial denaturation at 94°C for 5 min; ten cycles of 30 s at 94°C, 30 s at 65°C with a 1°C decrease in annealing temperature per cycle and 45 s at 72°C; 30 cycles of 30 s at 94°C, 30 s at 55°C and 45 s at 72°C and a final extension at 72°C for 10 min. The PCR products were visualized on 6% polyacrylamide gel followed by silver staining. The fragment sizes of PCR products were estimated by comparison with a 50 bp DNA ladder.

**Linkage analysis and map construction**

The recombinant inbred lines (RIL) F₆ population containing 140 individuals derived from the cross between Zhonghua 10 and ICG12625 were developed in our laboratory. The polymorphic SSR markers between Zhonghua 10 and ICG12625 were used to construct a genetic linkage map. For each SSR marker, the set of the amplified SSR alleles (or fragments) were classified into a single-locus SSR, if they exhibited an obvious co-dominant pattern across the whole population and the observed heterozygosity rate was less than 5% as described previously [27, 28]. These multi-locus SSRs were named using the suffixes “-1”, “-2” and “-n” after the SSR name, respectively. For example, the locus AGGS0010-3 meant the third locus amplified by the SSR marker AGGS0010. The marker loci were grouped at logarithm of odds (LOD) 4.0 by JoinMap 3.0 [29]. The Kosambi mapping function was used to convert the
recombination frequency into genetic distance [30]. Pearson’s Chi square test was performed to evaluate the goodness of fit to the expected 1:1 segregation ratio for each marker. In order to assign the novel SSR markers to specific linkage groups, a set of 105 public SSRs that were previously evenly mapped on A1-A10 and B1-B10 [24] were used as anchor markers.

**Evaluation of the SSR markers across *Arachis* species**

To test the transferability of the novel SSR markers in different *Arachis* species and evaluate the utility in genetic diversity and phylogenetic analysis, a set of 82 SSR markers that contained 19 SSRs originated from the A-genome species, 22 SSRs from the B-genome species and 41 SSRs from the AB-genome species were selected to genotype 39 wild *Arachis* accessions and 22 cultivated peanut accessions. The allele richness for SSR markers were estimated for the wild and cultivated *Arachis* accessions using the PowerMarker v3.51 package [31]. The Nei’s distances [32] between all pairs of accessions were calculated and a phylogenetic tree was constructed with the Neighbor-joining (N-J) algorithm to depict the genetic relationship and differentiation between the wild and cultivated *Arachis* accessions [33].

**Results**

**Identification and characterization of microsatellites from GSSs in *Arachis* species**

A total of 65,111 GSSs available of *Arachis* spp. were obtained in the GenBank database (accessed 19 December 2014), including 44,761 GSSs from *A. duranensis* (AA genome), 3,276 GSSs from *A. batizocoi* (BB genome) and 17,074 GSSs from *A. hypogaea* (AABB genome), respectively. Through searching the di-, tri-, tetra-, and pentanucleotide SSR repeats in GSS sequence, a total of 4,401, 377 and 7,251 SSR-containing sequences were identified in *A. duranensis*, *A. batizocoi* and *A. hypogaea*, respectively (Table 1). All these SSR containing sequences were aligned with each other to remove the redundant GSSs, resulting in a total of 445 contigs and 10,823 singletons. Interestingly, a large fraction of singletons were identified in either cultivated *Arachis* species (6,888/7,251; *A. hypogaea*) or the two wild *Arachis* species (i.e., 3,574/4,401 for *A. duranensis* and 361/377 for *A. batizocoi*), indicating that the public data is informative and of little redundancy. By integrating the contigs and singletons, we finally identified 4,826, 454 and 11,155 unique GSS-SSRs with an overall frequency of 159.9, 239.8 and 1,015.8 SSRs/Mb in *A. duranensis*, *A. batizocoi* and *A. hypogaea*, respectively (Table 1).

Of the total GSS-SSRs identified in the three species, the di- and trinucleotide repeat motifs accounted for nearly 90% (Fig 1, S2 Table). The trinucleotide repeat motifs were the most abundant repeat types in *A. duranensis* and *A. batizocoi*, while the dinucleotide repeat motifs were the most abundant in *A. hypogaea*. Tetra- and pentanucleotide repeat motifs both had a low frequency (0.6~8.9%) in all the three species. In dinucleotide repeat motifs, the most abundant repeat motif were (AG/CT)hom *A. batizocoi* (23.6%) and *A. hypogaea* (45.0%), while (AT/AT)hom (14.4%) in *A. duranensis* (Fig 1, S2 Table). All ten possible combinations of trinucleotide repeat motifs were observed in the three species except in *A. batizocoi*. Among these trinucleotide repeat motifs, (AAG/CTT)hom was the most common motif in *A. duranensis*, *A. batizocoi* and *A. hypogaea*, with the frequency of 22.2%, 14.8% and 6.2%, respectively (Fig 1, S2 Table).

**Identification and characterization of microsatellites from ESTs in *Arachis* species**

In the GenBank database 281,115 ESTs were acquired in *Arachis* spp., which contained 400 ESTs from *A. appressipila* (PP genome), 35,292 ESTs from *A. duranensis* (AA genome), 6,264
| Sequences type | Species          | Genome | No. of sequences surveyed | Length of sequences searched (bp) | No. of sequences containing repeats | No. of singletons | No. of contigs | No. of sequences containing repeats after removing redundant sequences | No. of SSRs after removing redundant sequences | Overall density (SSRs/Mb) |
|---------------|------------------|--------|---------------------------|----------------------------------|----------------------------------|-------------------|----------------|-------------------------------------------------|-----------------------------------------------|--------------------------|
| GSS           | A. duranensis    | A      | 44,761                    | 30,185,265                       | 4,401                            | 3,574             | 263           | 3                                               | 4,072                          | 4,826                    | 159.9                       |
|               | A. batizocoi     | B      | 3,276                     | 1,892,928                        | 377                              | 361               | 8             | 2                                               | 377                            | 454                      | 239.8                       |
|               | A. hypogaea      | AB     | 17,074                    | 10,981,790                       | 7,251                            | 6,888             | 174           | 2                                               | 7,194                          | 11,155                   | 1,015.8                     |
| EST           | A. appressipila  | P      | 400                       | 216,494                          | 62                               | 56                | 3             | 2                                               | 60                             | 69                       | 318.7                       |
|               | A. duranensis    | A      | 35,292                    | 17,417,205                       | 5,980                            | 2,870             | 832           | 4                                               | 4,605                          | 5,857                    | 336.3                       |
|               | A. stenosperma   | A      | 6,264                     | 4,213,240                        | 868                              | 689               | 67            | 3                                               | 805                            | 969                      | 230.0                       |
|               | A. ipaënsis      | B      | 32,787                    | 14,420,631                       | 5,694                            | 2,883             | 764           | 4                                               | 4,423                          | 5,543                    | 384.4                       |
|               | A. magna         | B      | 750                       | 236,255                          | 77                               | 10                | 12            | 5                                               | 35                             | 41                       | 173.5                       |
|               | A. hypogaea      | AB     | 205,622                   | 110,480,671                      | 33,286                           | 13,795            | 3,145         | 6                                               | 21,524                         | 27,720                   | 250.9                       |

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ESTs from *A. stenosperma* (AA genome), 32,787 ESTs from *A. ipaensis* (BB genome), 750 ESTs from *A. magna* (BB genome) and 205,622 ESTs from *A. hypogaea* (AABB genome). A total of 62, 5,980, 868, 5,694, 77 and 33,286 SSR-containing sequences were identified from ESTs in *A. appressipila*, *A. duranensis*, *A. stenosperma*, *A. ipaensis*, *A. magna* and *A. hypogaea*, respectively. After removing the redundant sequences, 60, 4,605, 805, 4,423, 35 and 21,524 unique SSR-containing sequences were obtained from ESTs in *A. appressipila*, *A. duranensis*, *A. stenosperma*, *A. ipaensis*, *A. magna* and *A. hypogaea*, respectively. Eventually, 69, 5,857, 969, 5,543, 41 and 27,720 EST-SSRs were identified from these unique SSR-containing sequences in *A. appressipila*, *A. duranensis*, *A. stenosperma*, *A. ipaensis*, *A. magna* and *A. hypogaea*, respectively, with an overall frequency of 318.7, 336.3, 230.0, 384.4, 173.5 and 250.9 SSRs/Mb (Table 1).

Among these EST-SSRs identified, the trinucleotide repeat was the most abundant repeat type in all six *Arachis* species and had a high frequency of 73.82% on average with a range from 69.4% in *A. ipaensis* to 79.7% in *A. appressipila* (Fig 1, S3 Table). The dinucleotide repeat motif occurred at a median frequency of 20.3% on average with a range from 14.6% in *A. magna* to 24.3% in *A. ipaensis*. Tetra- and pentanucleotide repeat motifs were observed with a low frequency (0~4.9%) in the six *Arachis* species. Among the dinucleotide repeats, (AG/CT)_n motif was observed with the highest frequency in the six *Arachis* species except *A. magna*, in which only one dinucleotide type (AT/AT)_n was observed. All ten possible combinations of trinucleotide repeat motifs were identified in the six *Arachis* species except *A. appressipila* and *A. magna*, possibly due to the significantly less available ESTs. Among these trinucleotide repeats, the most abundant motif was (AAG/CTT)_n for *A. appressipila*, *A. duranensis*, *A. ipaensis* and *A. hypogaea*, while (AAT/ATT)_n for *A. stenosperma* and *A. magna* (Fig 1, S3 Table).

**Development of SSR markers and polymorphism analysis**

After filtering out the SSRs with identical primer sequence to the publicly released SSRs in *Arachis* species, we totally identified 2,589 novel SSR markers in the present study (S4 Table). The set of novel SSRs from *Arachis* species were tested for reliability and polymorphism by amplification in six *A. hypogaea* accessions. Among 2,589 novel SSRs, there were 2,207 SSRs (85.2%) that the primers enable to amplify one or more clear fragments and 925 SSRs (35.7%) showing polymorphism in the six *A. hypogaea* accessions. The dinucleotide repeat (39.7%) and the (AG/CT)_n motif (42.0%) accounted for the largest fraction of the novel SSRs, and these SSRs also exhibited the highest level of polymorphism in *A. hypogaea* (S5 Table). To further distinguish the characteristic of SSR markers, the whole set of developed SSRs were classified into two major classes based on the length of SSR. The ‘Class I’ type of SSRs were defined as the ones with the length of SSR more than 20 bp, and the ‘Class II’ type of SSRs as the ones with the length of SSR less than 20 bp but more than 12 bp [34]. The ‘Class I’ type of SSRs contained 959 SSRs and the ‘Class II’ type of SSRs contained 1,630 SSRs. Notably, there were 47.8% of SSRs as ‘Class I’ type while 15.2% of SSRs as ‘Class II’ type that were polymorphic in the six accessions, respectively (Fig 2).

**Construction of the genetic linkage map**

Among the 925 polymorphic SSRs in six diverse *A. hypogaea* accessions, a subset of 559 SSR markers that were polymorphic between Zhonhua 10 and ICG12625 were employed to genotype a derived RIL population for allocating the specific positions to the novel SSRs by genetic linkage map. According to segregation patterns of SSRs, the 559 polymorphic SSRs totally identified 580 SSR loci in the RIL population. There were one SSR with three loci, 19 SSRs with two loci for each SSR and 539 SSRs with a single-locus for each SSR. With the set of 580 novel
SSR loci and 105 anchor SSR markers, a genetic linkage map consisting of 20 linkage groups was constructed (Fig 3). The genetic map was covered by a total of 645 SSR loci with the total length of 1,711.47 cM and an average interval of 2.65 cM between flanking markers (S6 Table). The 20 linkage groups were designated as A1-A10 for the A subgenome and B1-B10 for the B subgenome by aligning anchor markers to a previously published map [24].

In the linkage map, there were totally 305 SSR loci for the A subgenome with 253 novel SSR loci and 52 anchor loci, while 340 SSR loci for the B subgenome with 287 novel SSR loci and 53 anchor loci (Table 2). All of the anchor SSRs were mapped to the originally reported linkage groups [24], indicating the reliability of linkage map constructed using these novel SSR markers. In the A subgenome, the number of novel SSR loci on the linkage groups varied from 41 (A3) to 15 (A8) with an average of 25. In the B subgenome, all the linkage groups contained more than 22 novel SSR loci, among which the B10 linkage group had the most novel SSR loci.

**Fig 1. Distribution of the motif and repeat type of microsatellites in the assembled GSSs and ESTs in *Arachis* species.** For the bar plot, the horizontal axis showed the abundances (%) of microsatellites with different motif types (the legends lay below the plot). For the pie plots, the portions with different colors showed the proportions (%) of microsatellites with different repeat types (the legends lay right of the plot). In the plot, only the repeat and motifs from dinucleotide to pentanucleotide SSRs were shown.

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In the linkage map, Chi square ($\chi^2$) analysis revealed that only 57 of 580 novel SSR loci exhibited a significant level of segregation distortion in the RIL population ($P<0.05$) (S6 Table), which further verified the reliability of the novel SSRs in present study.

The utility of novel SSR markers in *A. hypogaea* and its wild related species

To test the transferability of the novel SSR markers in different *Arachis* species, we randomly selected 82 SSRs from linkage map to genotype 22 *A. hypogaea* accessions and 39 of its wild relatives. Among these SSRs, there were 37 SSRs that successfully amplified all 61 tested *Arachis* accessions, including 9 SSRs (47.4%) derived from *A. duranensis*, 4 SSRs (18.2%) from *A. batizocoi* and 24 SSRs (58.5%) from *A. hypogaea* (Table 3). Notably, all 82 SSRs not only amplified clear fragments in each of 22 *A. hypogaea* accessions, but also exhibited a relatively high amplification rate in the wild *Arachis* accessions for AB genome (77.3~97.6%), B genome (81.8~85.4%) and E genome (81.8~85.4%). Comparatively, the SSRs derived from *A. batizocoi* had a relatively low amplification rate in the wild *Arachis* accessions of P genome (36.4%) and A genome (36.4%) (Table 3). These results revealed that the SSRs derived from *A. hypogaea* and *A. duranensis* had relatively higher transferability across the cultivated and wild *Arachis* species than those from *A. batizocoi*. Furthermore, it was found that the SSRs from *A. hypogaea* had a high level of polymorphism in all accessions with 7.6 alleles/SSR, followed by SSRs from *A. batizocoi* (5.1 alleles/SSR) and *A. duranensis* (4.6 alleles/SSR). All the SSRs had a similar level of polymorphism in wild *Arachis* accessions of P genome, E genome, A genome, B genome and AB genome, respectively, while the SSRs developed from *A. hypogaea* sequences had a significantly higher level of polymorphism in the *A. hypogaea* accessions than those SSRs developed from *A. duranensis* sequences and *A. batizocoi* sequences (Table 3). Additionally, the novel SSRs were used to evaluate the genetic relationship among the 61 wild and cultivated *Arachis* accessions. A phylogenetic dendrogram based on the N-J algorithm was constructed that clustered all the *Arachis* accessions into two major genetic groups basically in accordance to the *Arachis* taxonomy (Fig 4). The largest group consisted of all the *A. hypogaea* accessions and the majority of accessions from its diploid ancestor of A and B genomes. The diploid and tetrapolyploid wild *Arachis* accessions were clustered into a separate large group. Our results
indicated that the novel SSRs had a high transferability across the *Arachis* species and had the ability to assess genetic diversity and phylogenetic relationship between wild and cultivated *Arachis*.

**Discussion**

To date, there was little knowledge about microsatellite distributions in the wild and cultivated *Arachis* species. In this study, the microsatellite characteristics across the *Arachis* species were analyzed by using the publicly released GSS and EST sequences. We found that the GSSs had a SSR density of 159.9 and 239.8 SSRs/Mb for two diploid wild *Arachis* species (*A. duranensis* and *A. batizocoi*), but exhibited an extremely high density of 1,025.8 SSRs/Mb for the tetraploid *Arachis* (*A. hypogaea*). The microsatellite density in *Arachis* species was sharply higher than those in *Gossypium* species (41.2 to 49.1 SSRs/Mb) [35], but lower than those in *Brassica* crop species (420.6 to 496.8 SSRs/Mb) [36] except in *A. hypogaea*. The significantly higher SSR density of *A. hypogaea* relative to other *Arachis* species may be attributed to the lack and the
uneven genomic distribution of GSS sequences in the Arachis species. In the present study, the length of assessed sequences accounted for a small proportion of genome in each Arachis species, which might lead to the difficulties for accurately evaluating characteristics of microsatellite across the Arachis species. On the other hand, some of the sequences of GenBank database were obtained with the initial focus on mining and development of microsatellite markers in the previous studies, thus probably resulting in overestimating the numbers and density of SSRs in the present study, especially in A. hypogaea [11, 13, 16]. For ESTs, the tetrapolyploid Arachis species (A. hypogaea) had a SSR density of 250.9 SSRs/Mb, but its diploid wild ancestors (A. duranensis and A. ipaensis) had a relatively high SSR density of 336.3 and 384.4 SSRs/Mb. This result was congruent to the finding that the genetic diversity of cultivated peanut was lower than the wild diploid Arachis species [7], which may be attributed to the origin of

Table 2. Information of linkage group constructed using the novel SSRs in present study.

| Linkage Group | SSR loci | Anchor Loci | Novel Loci | Length (cM) |
|---------------|----------|-------------|------------|-------------|
| A1            | 31       | 6           | 25         | 87.59       |
| A2            | 25       | 5           | 20         | 66.56       |
| A3            | 46       | 5           | 41         | 114.18      |
| A4            | 25       | 5           | 20         | 68.00       |
| A5            | 31       | 5           | 26         | 89.82       |
| A6            | 25       | 5           | 20         | 95.07       |
| A7            | 31       | 5           | 26         | 91.62       |
| A8            | 20       | 5           | 15         | 90.73       |
| A9            | 46       | 6           | 40         | 104.91      |
| A10           | 25       | 5           | 20         | 60.39       |
| B1            | 27       | 5           | 22         | 57.39       |
| B2            | 32       | 7           | 25         | 105.40      |
| B3            | 37       | 5           | 32         | 106.16      |
| B4            | 36       | 5           | 31         | 121.80      |
| B5            | 36       | 5           | 31         | 104.51      |
| B6            | 40       | 5           | 35         | 64.35       |
| B7            | 30       | 5           | 25         | 76.76       |
| B8            | 29       | 5           | 24         | 63.61       |
| B9            | 28       | 5           | 23         | 60.47       |
| B10           | 45       | 6           | 39         | 82.15       |
| A subgenome   | 305      | 52          | 253        | 868.87      |
| B subgenome   | 340      | 53          | 287        | 842.60      |
| whole genome  | 645      | 105         | 540        | 1,711.47    |

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Table 3. The amplification of SSRs derived from A. duranensis, A. batizocoi and A. hypogaea in Arachis species.

| SSR resources | SSRs No. | Amplified SSRs (%) | alleles/SSR |
|---------------|----------|--------------------|-------------|
|               | I II III IV V VI VII | I II III IV V VI VII |
| A. duranensis | 19       | 9 (47.4) 10 (52.6) | 15 (78.9) 11 (57.9) 15 (78.9) 17 (89.5) 19 (100) | 4.6 3.1 2.8 4.3 2.6 2.3 2.4 |
| A. batizocoi  | 22       | 4 (18.2) 8 (36.4) | 18 (81.8) 8 (36.4) 18 (81.8) 17 (77.3) 22 (100) | 5.1 3.3 3 4.5 2.9 2.6 1.7 |
| A. hypogaea   | 41       | 24 (58.5) 30 (73.2) | 35 (85.4) 27 (65.9) 35 (85.4) 40 (97.6) 41 (100) | 7.6 2.8 3 3.3 2 4.9 |

I: all Arachis accessions; II: 5 wild Arachis accessions of P genome; III: 6 wild Arachis accessions of E genome; IV: 19 wild Arachis accessions of A genome; V: 6 wild Arachis accessions of B genome; VI: 3 wild Arachis accessions of AB genome; VII: 22 A. hypogaea accessions

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cultivated peanut in a single allopolyploid event between the wild diploid ancestors, *A. duranensis* and *A. ipaensis* [2, 3]. Despite the limitation of the current data, the results of this study provided an informative clue to characterize the differentiation of microsatellites across the *Arachis* species. With the future release of the reference genomes of the cultivated peanut and its two diploid ancestors, it would be possible to comprehensively characterize the evolutionary pattern of microsatellites between wild and cultivated *Arachis* species.

The composition of microsatellites varied across the *Arachis* species. In GSSs, the dinucleotide repeats dominated in *A. hypogaea* (76.0%), which is similar to Chinese spring wheat [8].

Fig 4. The phylogenetic analysis of the 61 cultivated and wild *Arachis* accessions in this study.

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rice and Arabidopsis [37], while the trinucleotide repeats were the predominant in A. duranensis and A. batizocoi (63.2% and 54.4%) as well as Brachypodium [38], bamboo [39] and Setaria italica [40]. These results revealed the different patterns of microsatellite motifs across the plant species, which may reflected the genomic footprints of speciation and evolution in these species. In ESTs, the trinucleotide repeats were predominant across all the Arachis species. This result was in accordance with previous ones on EST-SSR in peanut [18]. However, it was inconsistent with another study on GSS-SSR and EST-SSR, in which 3.5 Mbp cultivated peanut genome sequences and 29.3 Mbp A. duranensis genome sequences were used to identify microsatellites [17]. This may be attributed to the data from the different Arachis species used in these studies. The use of more Arachis species in present study would be more reliable to infer the pattern of microsatellites in Arachis genus, although the large amount of data and reference genome of the Arachis species were still needed to make a comprehensive evolutionary conclusion in Arachis genus. In the present study, the most frequent dinucleotide repeat motif was (AG/CT)n, which was consistent with that identified in the previous studies in peanut [10, 11, 14, 16, 18], while the dinucleotide motif (AT)n was enriched in Chinese spring wheat [8], Gossypium species [35] and Brassica crop species [36, 41, 42]. Additionally, the top two trinucleotide repeat motifs were (AAG/CTT)n and (AAT/ATT)n in this study, which was in accordance with those results of previous studies in peanut [10, 11, 14, 16, 18], Gossypium species [35] and Brassica crop species [36, 41, 42].

In Arachis, SSRs has served as an informative and manageable marker in germplasm evaluation, QTL analysis and marker-assisted selection for a long time. Extensive efforts had been made to develop SSR markers in Arachis from various resources [10–18], but the amount of SSR markers remain insufficient to map QTL in fine-scale or saturate the whole genome for genome wide association study (GWAS) [23–25, 43–45], due to the low genetic diversity in A. hypogaea. To this end, we developed novel 2,589 SSRs based on the publicly released Arachis sequences in present study. The majority of the novel SSRs were able to clearly amplify fragments in A. hypogaea, and 540 of these markers were successfully allocated to the specific genetic positions in linkage groups of A. hypogaea. These results would provide an informative resource to localize QTL and marker-assisted selection. Moreover, as the polyploidy nature of A. hypogaea, the SSRs probably amplify multiple alleles at homologous DNA sequences. That fact would lead to a substantial risk of allele calling bias in natural populations for GWAS in tetrapolyploid species [27, 28, 46]. Therefore, there is an urgent need to develop the single-locus SSR markers to facilitate their applications, as currently only a small proportion of the previously developed publicly available SSRs were single-locus markers [43–45]. In the present study, we totally identified 539 SSRs that were probably to be single-locus markers according to the segregation pattern in a biparental RIL population. These single-locus SSRs with the specific genetic positions gave the research community a useful resource to evaluate linkage disequilibrium in Arachis, although they were still needed to be validated by different mapping populations. Additionally, the novel SSRs in this study were highly transferable across Arachis species, especially for A. hypogaea, and would be informative to assess genetic diversity and phylogenetic relationship between wild and cultivated Arachis.

Supporting Information

S1 Table. Information for the 61 cultivated and wild Arachis accessions used in this study. (XLSX)

S2 Table. Number, repeat number and total repeat length of the di- to pentanucleotide repeats motifs of microsatellites in the assembled GSSs of Arachis species. (XLSX)
S3 Table. Number, repeat number and total repeat length of the di- to pentanucleotide repeats motifs of microsatellites in the assembled ESTs of *Arachis* species. (XLSX)

S4 Table. Detailed information for the novel SSRs developed in the present study. (XLSX)

S5 Table. Polymorphism rates of different repeat motifs in six *A. hypogaea* cultivars. (XLSX)

S6 Table. The detailed information of genetic positions and segregation distortion of the SSRs in the linkage map. (XLSX)

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Author Contributions

Conceived and designed the experiments: LH HJ. Performed the experiments: BW JZ WC YZ YC XZ. Analyzed the data: LH HL. Contributed reagents/materials/analysis tools: LH XR YC XZ. Wrote the paper: LH YL BL HJ.

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