High-throughput SSR marker development and its application in a centipedegrass (*Eremochloa ophiuroides* (Munro) Hack.) genetic diversity analysis

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**Abstract**

Centipedegrass (*Eremochloa ophiuroides* (Munro) Hack.) is a perennial, warm-season C4 grass species that shows great potential for use as a low-maintenance turfgrass species in tropical and subtropical regions. However, limited genetic and genomic information is available for this species, which has impeded systematic studies on the enhancement of its turf quality and resistance against biotic and abiotic stress. In this study, Illumina HiSeq high-throughput sequencing technology was performed to generate centipedegrass transcriptome sequences. A total of 352,513 assembled sequences were used to search for simple sequence repeat (SSR) loci, and 64,470 SSR loci were detected in 47,638 SSR containing sequences. The tri-nucleotides were the most frequent repeat motif, followed by di-nucleotides, tetra-nucleotides hexnucleotides, and pentanucleotides. A total of 48,061 primer pairs were successfully designed in the flanking sequences of the SSRs, and 100 sets of primers were randomly selected for the initial validation in four centipedegrass accessions. In total, 56 (56.0%) of the 100 primer pairs tested successfully amplified alleles from all four centipedegrass accessions, while 50 were identified as polymorphic markers and were then used to assess the level of genetic diversity among 43 centipedegrass core collections. The genetic diversity analysis exhibited that the number of alleles (Na) per locus ranged from 3 to 13, and the observed heterozygosity (Ho) ranged from 0.17 to 0.83. The polymorphism information content (PIC) value of the markers ranged from 0.15 to 0.78, and the genetic distances (coefficient Nei72) between the accessions varied from 0.07 to 0.48. The UPGMA-based dendrogram clustered all 43 core collections into two main groups and six subgroups, which further validated the effectiveness of these newly developed SSR markers. Hence, these newly developed SSR markers will be valuable and potentially useful for future genetic and genomic studies of *E. ophiuroides*. 

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**Citation:** Li J, Guo H, Wang Y, Zong J, Chen J, Li D, et al. (2018) High-throughput SSR marker development and its application in a centipedegrass (*Eremochloa ophiuroides* (Munro) Hack.) genetic diversity analysis. PLoS ONE 13(8): e0202605. https://doi.org/10.1371/journal.pone.0202605

**Editor:** Tzen-Yuh Chiang, National Cheng Kung University, TAIWAN

**Received:** May 7, 2018

**Accepted:** August 5, 2018

**Published:** August 22, 2018

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**Data Availability Statement:** All data files are available from the NCBI database (accession number(s) SRP134136).

**Funding:** This work was financially supported by Grants from the Scientific Facility Project of Jiangsu Province, China (Grant No. BM2015019-1), and Jiangsu Provincial Natural Science Foundation (Grant No. BK20171327). The funders had no role in study design, data collection and
Introduction

Simple sequence repeats (SSRs) are a valuable source of genetic markers because of their abundance, high rate of polymorphism, ubiquitous distribution throughout the genome, codominant inheritance, high extent of allelic diversity, and ease of assay by PCR [1,2]. Thus, SSRs are considered excellent molecular markers in studies of germplasm characterization, genetic diversity, and genetic mapping [3,4]. However, the traditional development of SSR markers has relied on the screening of genomic libraries using repetitive probes and the sequencing of positive clones, which is time-consuming and requires the use of specialized laboratory equipment [5]. However, recent advances in next-generation sequencing (NGS) technologies provide a cost-effective, convenient and reliable approach for sequence information acquisition in non-model species and greatly accelerated the development process for molecular markers [6,7]. RNA-Seq, which is based on NGS, is a high-throughput technology that has great advantages in obtaining a large amount of sequence data for SSR mining [8]. Using transcriptome data resources, rapid progress in the development of SSR loci has been made in many green plant species [9]. Recently, a large amount of SSR markers were developed in forage and turfgrass crops such as perennial ryegrass [10], Italian ryegrass [11], alfalfa [12], hemarthria [13], red clover [14], orchardgrass [15], sudangrass [16], common bermudagrass [17], creeping bentgrass [18], seashore paspalum [19], and zoysiagrass [20]. To date, there is not a large quantity of SSR markers developed in centipedegrass, with the exception of a recent report on EST-SSR development from a cold-stressed transcriptome of centipedegrass [21].

Centipedegrass (Eremochloa ophiuroides (Munro) Hack.) is an important turfgrass that belongs to warm-season (C4) perennial grass species and is distributed extensively in Southeast Asia and United States. E. ophiuroides is a native grass in South and Central China [22,23] and is known for its good adaptation to infertile soils and a range of climatic conditions [24,25]. E. ophiuroides has great potential as a low-maintenance turf and is often referred as ‘lazy man’s grass’ or ‘poor man’s grass’ because of its lower management requirements than those of most warm-season turfgrasses [26]. Centipedegrass usually presents a broad resistance to insect and disease infestations and shows excellent heat tolerance as well [26]. These outstanding characteristics make E. ophiuroides a popular turfgrass in tropical and subtropical regions. Moreover, centipedegrass is also used as a forage grass in Japan for its heavy grazing-tolerance [24].

Over the past two decades, a series of studies, including accession identification, an analysis of genetic diversity, and construction of genetic map in centipedegrass, was performed based on the use of limited universal molecular markers [27–31]. Some efforts in improving the lawn traits of centipedegrass, including traditional selection breeding [24,32,33], irradiation mutagenesis [34–36] and somatic variation, have also been carried out [25,37–39]. Despite progress in preliminary studies on genetic analysis and germplasm innovation, genetic and genomic information on this turfgrass species is still largely limited. Thus far, less than one hundred nucleotide sequences of DNA have been deposited in a public database (National Center for Biotechnology Information), which is markedly incomparable to that for other turfgrass plants, e.g., ryegrass, festuca, bermudagrass, or zoysiagrass. The paucity of available information on the nucleotide sequences has hindered its genetic and genomic studies, such as the large-scale development of molecular markers, the construction of high-density linkage maps, and gene discovery.

To further complement the genomic sequence information and the number of molecular markers, in this study we conducted large-scale SSR mining employing the RNA sequencing (RNA-Seq) data of centipedegrass leaf, stolon and spikes based on the high-throughput Illumina HiSeq 2000 platform. The resultant SSR sequences were characterized and validated
through the successful amplification of randomly selected target loci across a selection of four distinct *E. ophiuroides* accessions. The newly developed SSR markers were subsequently utilized to assess the genetic diversity level of core collections, including 43 centipedegrass accessions from diverse geographic origins. The datasets and results reported in the present study provide a public resource and information for future genetic studies and breeding programs in *E. ophiuroides*.

**Materials and methods**

**Plant materials and isolation of RNA and DNA**

Two *E. ophiuroides* accessions E092 and E092-1 were used for cDNA sequencing. The accession E092 is a wild-type with red-purple stolons and spike tissues during its developmental stages, and it was originally collected from Chongqing city in the Southwest China. The accession E092-1 is a natural mutant with uniform green stolon and spike tissues and was isolated from E092. Initially, four *E. ophiuroides* accessions including two purple-stolon accessions (E092 and E022) and two green-stolon ones (E092-1 and E039) were used for validating the SSR primers, and then, 43 accessions from the *E. ophiuroides* core collection were adopted to test the SSR markers and assess the genetic diversity level (Table 1). All these accessions of *E. ophiuroides* were maintained by the Main Warm-season Turfgrass Germplasm Resource Preserving Centre, Nanjing Botanical Garden Men. Sun Yat-Sen, Jiangsu Province and Chinese Academy of Sciences, Nanjing, China. All the plant materials were grown in plastic pots (13 cm top diameter × 10 cm bottom diameter × 11 cm depth), with a mix of soil, sand and peat at a ratio of 1:1:1, and were cultivated in a greenhouse under natural sunlight, with an average temperature of 30°C day/20°C night and a relative humidity of ~80%.

The total RNA, which was required for the transcriptome sequencing, was extracted from the stolons, leaves and inflorescences of accessions E092 and E092-1 using the Trizol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instructions. Prior to the RNA extraction, the tissue samples were frozen in liquid nitrogen and were homogenized by hand using a glass tissue grinder (DUALL 20, Kontes Glass Co.). The isolated RNA was treated with RNase-free DNase I (Ambion, Austin, TX, USA) to eliminate the potential genomic DNA. The RNA concentration and quality were determined by a NanoDrop 8000 spectrophotometer (NanoDrop, Wilmington, DE), and its integrity was confirmed using an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, Calif.). Samples with an RNA Integrity Number (RIN) ≥ 7, and 28S:18S ratio ≥ 1.5:1, and total amount ≥ 3 μg were considered acceptable.

The total genomic DNA was isolated from the young leaves of each plant using the EZgene™ SuperFast Plant Leaves DNA Kit (Biomiga, San Diego, CA, USA) following the manufacturer’s protocols for plant leaves with high levels of phenolic compounds and polysaccharides. The DNA was dissolved in 50 μL of sterile ultra-pure water, diluted to a final concentration of 30 ng/μL and stored at -20°C until the PCR analysis.

**cDNA library construction and Illumina sequencing**

Illumina sequencing was performed at the Decode Genomics Ltd., in Nanjing, China, following the manufacturer’s protocols (Illumina, San Diego, CA). Briefly, the poly (A) mRNA was purified from the total RNA using Sera-mag Magnetic Oligo (dT) beads from Illumina, and then, the mRNA-enriched RNAs were chemically fragmented into short sequences using the fragmentation solution (Ambion, USA). The double-stranded cDNA was synthesized using the Superscript Double-Stranded cDNA Synthesis Kit (Invitrogen, USA). Finally, the pair-end RNA-seq libraries were constructed using the Illumina Paired End Sample Prep kit and were subsequently sequenced on the Illumina HiSeqTM 2000 platform [40].
Table 1. The germplasm collections of E. ophiuroide used in this study.

| No. | Sample | Source                        |
|-----|--------|-------------------------------|
| 1   | E-004  | Fuzhou, Fujian, Eastern China |
| 2   | E-006  | Putuoshan, Zhejiang, Eastern China |
| 3   | E-013  | Yongtai, Fujian, Eastern China |
| 4   | E-015  | Lushan, Jiangxi, Central China |
| 5   | E-017  | Hangzhou, Zhejiang, Eastern China |
| 6   | E-019  | Hangzhou, Zhejiang, Eastern China |
| 7   | E-022  | Wuxi, Jiangsu, Eastern China |
| 8   | E-041  | Lingchuan, Guangxi, South China |
| 9   | E-042  | Taiping, Anhui, Eastern China |
| 10  | E-047  | Lianyungang, Jiangsu, Eastern China |
| 11  | E-055  | Guangzhou, Guangdong, South China |
| 12  | E-061  | Heyuan, Guangdong, South China |
| 13  | E-063  | Yingde, Guangdong, South China |
| 14  | E-065  | Yingde, Guangdong, South China |
| 15  | E-072  | Guilin, Guangxi, South China |
| 16  | E-074  | Yichang, Hubei, Central China |
| 17  | E-077  | Zhangjiagang, Hunan, Central China |
| 18  | E-078  | Yichang, Hubei, Central China |
| 19  | E-084  | Anshun, Guizhou, Southwestern China |
| 20  | E-087  | Guiyang, Guizhou, Southwestern China |
| 21  | E-091  | Chongqing, Southwestern China |
| 22  | E-092-1| Chongqing, Southwestern China |
| 23  | E-097  | Xinyang, Henan, Central China |
| 24  | E-098  | Xinyang, Henan, Central China |
| 25  | E-099  | Jinzhai, Anhui, Eastern China |
| 26  | E-102  | USA |
| 27  | E-112  | Xinjiang, Sichuan, Southwestern China |
| 28  | E-115  | Yiyang, Hunan, Central China |
| 29  | E-124  | Changsha, Hunan, Central China |
| 30  | E-131  | Yiyang, Hunan, Central China |
| 31  | E-134  | Guiyang, Guizhou, Southwestern China |
| 32  | E-135  | Guiyang, Guizhou, Southwestern China |
| 33  | E-141  | Feixi, Anhui, Eastern China |
| 34  | E-144  | Xinyang, Henan, Central China |
| 35  | E-145  | Xinyang, Henan, Central China |
| 36  | E-152  | Xinyang, Henan, Central China |
| 37  | E-154  | Liuhe, Jiangsu, Eastern China |
| 38  | E-155  | Lianyungang, Jiangsu, Eastern China |
| 39  | E-158  | USA |
| 40  | E-182  | Lanxi, Zhejiang, Eastern China |
| 41  | E-183  | Nanjing, Jiangsu, Eastern China |
| 42  | E-187  | Yixing, Jiangsu, Eastern China |
| 43  | E-188  | Jinzhai, Anhui, Eastern China |

[10.1371/journal.pone.0202605.t001](https://doi.org/10.1371/journal.pone.0202605.t001)

**Data filtering and de novo assembly of cDNA**

Prior to the transcriptome assembly, a stringent filtering criterion of an FPKM (fragments per kilobase of exon per million reads mapped) value of 1.0 in at least one sample or an FPKM
value of 0.5 in at least two samples was used to filter the raw sequencing reads. The clean reads were obtained from the raw sequencing reads by removing the adapter sequences, the reads with more than 10% unknown nucleotides, and the low-quality reads (> 50% bases with quality value Q ≤ 5 in a read). The de novo assembly of the transcriptome was accomplished using all the clean reads and the Trinity program (version trinityrnaseq_r20140717) using the de Bruijn graph method and default settings [41].

**Sequence data**

The assembled transcriptome sequences from the *E. ophiuroides* accessions E092 and E092-1 were deposited in the NCBI, with the BioProject accession PRJNA437781. They were de novo assembled from the raw sequencing data (7.5-fold coverage) of the accessions E092 and E092-1, which were deposited in the NCBI under the SRA accession number SRP134136 (SAMN08640788, SAMN08640789, SAMN08640790, SAMN08640791, SAMN08640792, SAMN08640793, SAMN08640794, SAMN08640795, and SAMN08640796).

**SSR motif detection and SSR marker development**

We employed the software package Genome-wide Microsatellite Analyzing Tool Package (GMATA2.1) (http://sourceforge.net/projects/gmata/?source=navbar) to mine the SSRs, perform the statistical analysis, and design primers from the identified SSR loci using constraints of more than five repeats and a motif length between 2 and 10 bp [42]. We followed the GMATA procedures described by Wang and Wang (2016) [42]. The primers were synthesized by Sangon Biotech Company (Shanghai, China).

Sequence IDs were given to the SSR-containing sequences used as templates for designing the primers (S1 Table). A marker name was assigned to each of the randomly selected SSR primer pairs for the PCR analysis. The marker name was comprised of the research center name “TJIB,” which stands for “Turfgrass Research Center of the Institute of Botany, Jiangsu Province and Chinese Academy of Sciences (JIB),” and the suffix showing the plant species *Eremochloa ophiuroides* (Eo) and the serial number.

**Genotyping the *E. ophiuroides* collections using the SSR markers**

A subset of 100 SSR primer pairs was randomly selected for validating the SSR locus amplification by polymerase chain reactions (PCR). The primer pairs that produced a reproducible and clear amplicon of the expected size were then used for assessing the genetic diversity among the centipedegrass accessions. PCR amplification was conducted in a 10 μL reaction volume containing 5 μL of 2 x TaqPCR MasterMix (TsingKe Biological Technology Co., Beijing, China), 1 μL of primer pair (10 μM), 1.5 μL of genomic DNA (30 ng) and 2.5 μL of ddH2O. The PCR conditions comprised an initial denaturing step (95˚C /3 min) followed by ten cycles of 94˚C /30 s, 55–60˚C /30 s, and 72˚C /30 s, and then 25 cycles of 94˚C /30 s, 55˚C /30 s, and 72˚C /30 s, and finally by an elongation step (72˚C /10 min). The PCR products were separated on 8.0% non-denaturing polyacrylamide gels and were visualized by 0.1% silver nitrate staining. The band sizes were determined by comparing them against a DNA ladder.

**Data analysis**

The genotyping data were used to determine the genetic relationships among the 43 *E. ophiuroides* core collections. The number of alleles (Na), the number of effective alleles (Ne), the observed heterozygosities (Ho), and the Shannon’s information index (I) were calculated using GenALEX software (version: 6.5) [43,44]. The polymorphic information content (PIC)
value of the alleles revealed by each primer pair was calculated by the formula: \( PIC = 1 - \sum (Pi)^2 \), where \( Pi \) is the frequency for the \( i \)th microsatellite allele. The genetic distances across the collections were calculated with the POPGENE software (version 1.31; https://www.ualberta.ca/~fyeh/popgene_download.html) [45]. Based on Nei's unbiased measures of genetic distance, a cluster analysis of the 43 collections was carried out using the unweighted pair-group method with arithmetic average (UPGMA), and the dendrogram was subjected to 1000 bootstraps using the MEGA4 [46].

Results

Illumina paired-end sequencing and characterization of the sequencing reads

In this study, to remove the highly similar or redundant sequences, we merged sequences with a sequence identity higher than 95% using the CD-HIT-EST algorithm [47]. A total of 390,247,286 clean reads were generated using the Illumina HiSeq2000 system, and 352,513 assembled sequences were used for further analysis after adaptor removal. The length of the assembled sequences varied from 200 bp to \( \geq 16 \) kb, with the average of approximately 735 bp, and the CG content was approximately 48% (Table 2).

SSR mining and characterization

The GMATA strategy was performed to search the SSR loci from the assembled transcriptome sequences. A total of 64,470 SSR loci were found in 47,638 SSR-containing transcriptome sequences. Of the SSR-containing transcriptome sequences, 11,725 contained more than one SSR locus (Table 2 and S1 Table).

Among all the SSRs detected, the most abundant repeat motifs were trinucleotides (33,614, 52.14%), followed by dinucleotides (28,783, 44.65%), tetranucleotides (1,399, 2.17%), hexanucleotides (331, 0.51%) and pentanucleotides (317, 0.49%) (Fig 1A, S2 Table). The highest frequency of the grouped SSR motif units was dimer AG/CT (5,774, 8.96%) followed by GA/TC (5,362, 8.32%), TG/CA (4,746, 7.36%), AC/TG (4,353, 6.75%), GCC/GGC (3,674, 5.70%) and CGC/GCG (3,613, 5.60%) (Fig 1B, S3 Table). The SSR length of 15 bp (17,296, 26.83%) was the most abundant, followed by an SSR length of 10 bp (13,212, 20.49) and then of 18 bp (9,563, 14.83) (Fig 1C, S4 Table).

Development and validation of the SSR markers

A total of 48,061 non-redundant primer pairs were designed using Primer 3.0 software based on the criteria of the melting temperature, CG content, lack of secondary structure and length
of amplification bands (S5 Table). The expected length of the target bands was between 100 bp and 400 bp.

Initially, a subset of 100 primer pairs was randomly selected for validating the SSR locus amplification. A total of 81 (81.0%) of the primer pairs tested successfully generated amplification products in at least one of the four *E. ophiuroides* accessions (E022, E039, E092, E092-1), and 56% of the primer pairs successfully amplified alleles from all four centipedegrass accessions. Of the 81 that amplified, 31 (38.27%) were monomorphic, and the other 50 (61.73%) were polymorphic between the four accessions and were selected for further evaluating the genetic diversity in the *E. ophiuroides* core collections (Table 3).

**Evaluation of the genetic diversity within the *E. ophiuroides* core collections**

The fifty polymorphic SSR markers were further used to assess the genetic diversity of the 43 *E. ophiuroides* core collections from different geographic locations. A total of 420 alleles were detected in 43 collections, wherein 285 alleles were determined to be collection-specific and 135 alleles were generally detected in multiple collections. The Na amplified per SSR locus ranged from 3 to 13, with an average of 8.40; the Ne ranged from 1.20 to 5.27, with an average of 3.01; the Ho ranged from 0.17 to 0.83, with the average of 0.64. The PIC value ranged from 0.15 to 0.78, with an average of 0.58. The I ranged from 0.31 to 1.76, with an average of 1.17 (Table 4).

The genetic distances, with the Nei72 coefficient, among the collections ranged from 0.07 to 0.48 (S6 Table). The largest genetic distance was observed between E188 from Jinzhai, Anhui, Eastern China and E102 from the USA. The UPGMA-based dendrogram placed the 43 accessions into two main groups, which were further subdivided into six distinct clusters (Fig 2). Cluster I comprised 21 collections from Eastern (E004, E006, E047, E042, E017, E019, E013), South (E065, E015, E063, E041, E072, E055), Southwestern (E092-1, E087, E084) and Central China (E022, E077, E078, E097, E098). Cluster II comprised only two accessions from South (E061) and Central China (E074). Cluster III included three accessions from Southwestern (E091, E112) and Eastern China (E099), and one accession from the USA (E102). Cluster IV consisted of four accessions from Central (E115, E124), eastern (E141), and Southwest China (E134). In addition to one collection from Southwestern China (E135), cluster V consisted mainly of the accessions from Central (E131, E144, E145) and Eastern China (E187, E188, E182, E183). Cluster VI contained four collections consisting of one collection from Central China (E152), two from Eastern China (E155, E154) and another one from the USA (E158).

**Discussion**

Simple sequence repeats (SSRs) are widely used as molecular markers in plant genetic studies due to their abundance, reproducibility, high allelic variation at each locus and simplicity to analyze using conventional PCR amplification. Recently developed next generation sequencing (NGS) platforms, such as Roche’s 454 GS FLX, Illumina’s GenomeAnalyzer (GA) and ABI’s SOLiD, offer opportunities for high-throughput and inexpensive genome sequencing and rapid marker development. In the present study, SSR markers for the diploid species *E. ophiuroides* were developed based on de novo assembled cDNA by the Illumina sequencing system from two phenotypically distinct accessions E092 and E092-1, which were maintained at the China National Germplasm Resources Nursery for warm-season turfgrasses in the Institute of Botany, Jiangsu Province and Chinese Academy of Sciences. The reliability of the newly developed SSR primers was tested via a PCR analysis on the core germplasm collections of centipedegrass, including 41 accessions from China and 2 accessions from the USA.
Table 3. Screening of SSR primer pairs by PCR in four *E. ophiuroides* accessions.

| ID          | Marker Name   | SSR                  | Forward primer (5'-3') | Reverse primer (5'-3') | Product size (bp) | PCR amplification in |
|------------|--------------|----------------------|------------------------|------------------------|-------------------|----------------------|
| TR80_e0_g1_i1 | TJB.Eo_001 | (CTC)12              | AGTTCATTTGCCAGCTCAT    | TAAACTGCGTGAGCAGGAAAAC | 195               | + - - -             |
| TR993_e0_g1_i1 | TJB.Eo_002 | (ACA)16              | TGTCATCAGGAGAAGAGCA   | GGTTGCGTGATGAGTTG     | 141               | + + - +             |
| TR1158_e0_g1_i1 | TJB.Eo_003 | (GT)11               | CCAGATCGCTGATGATCC    | GGGCAGCTGATGAGTCA     | 227               | + - - +             |
| TR1654_e0_g1_i1 | TJB.Eo_004* | (CAA)13              | GTGTTGTGTGGCTGGTGG    | TGTTGCTGATGAGACGAC    | 197               | + + + +             |
| TR1713_e0_g1_i1 | TJB.Eo_005 | (TCT)10              | ACCACAGCAGGAGAAGGAC   | ATGTTCTGCTGATGATCC    | 276               | + + + +             |
| TR2338_e0_g1_i1 | TJB.Eo_006* | (GCC)5               | GGTGCCTGTTCCGCTATCT   | CTGCTGCTCCGCTCCGTCCTG | 264               | + + + +             |
| TR2644_e0_g1_i1 | TJB.Eo_007 | (ACA)5               | AGCAGCCGAGATGAGCTGG   | CAGGGGCGTGATGAGTTG    | 207               | + + + +             |
| TR2941_e0_g1_i1 | TJB.Eo_008* | (ACA)5               | AGCAGGAGAGCAAGGACGAGA | ATGTTCTGCTCCGCTCCGTCCTG | 274               | + + + +             |
| TR3535_e0_g2_i1 | TJB.Eo_010* | (CTG)13              | GGTGCCTGAGATGAGATGAGA | GGTGCCTGAGACTGACGAGG | 183               | + + + +             |
| TR5354_e0_g2_i1 | TJB.Eo_011 | (GCC)5               | GGTGCCTGTTGAGATGAGATG | GGTGCCTGAGACTGACGAGG | 173               | + + + +             |
| TR6729_e0_g1_i1 | TJB.Eo_013* | (CTG)13              | GGTGCCTGTTGAGATGAGATG | GGTGCCTGAGACTGACGAGG | 173               | + + + +             |
| TR7108_e0_g1_i1 | TJB.Eo_014* | (CAT)7ttg(CTC)5      | GCCCAATCTATCTACATCA   | GATGAGGAGCAAGGACGAGGGA | 208               | - + + +             |
| TR7160_e0_g1_i1 | TJB.Eo_015 | (TG)11               | GGTGCCCTCTGTCGTAGTGG   | TTGGCAGCTGTTATGATAG  | 227               | - - - -             |
| TR7421_e0_g1_i1 | TJB.Eo_016* | (AGA)11              | AGGAGGAAGCAAGCAAGCAA  | GCTGACCTGTTGGCTGCTAG  | 107               | + + + +             |
| TR8081_e0_g1_i1 | TJB.Eo_017* | (GAG)10              | AGGAGGAAGCAAGCAAGCAA  | GCTGACCTGTTGGCTGCTAG  | 107               | + - + +             |
| TR9123_e0_g1_i1 | TJB.Eo_018 | (GTC)5tt(GTC)7       | CGTTGCTCTTCCGCTTGGT   | GAGGCTAACCATCTAAGCAGG | 103               | - + - +             |
| TR9124_e1_g1_i1 | TJB.Eo_019 | (TG)8                | TGTCATCCAGAGATGAGC    | GCCCATGGAAAAACCTACACT | 148               | - - - -             |
| TR9278_e0_g2_i1 | TJB.Eo_020* | (GCC)11              | AGGAGGAAGCAAGCAAGCAA  | GCTGACCTGTTGGCTGCTAG  | 264               | + + + +             |
| TR10202_e0_g1_i1 | TJB.Eo_021 | (A)14                | TTGGGCTCGCCGATATAGAG  | CAGGGGCGTGCTCTGCTGCT  | 123               | + + + +             |
| TR11518_e0_g1_i1 | TJB.Eo_022* | (CA)11               | AAACAAATTGCCAGAGATC   | GTGTTGCTGGTCTCCAGCAG  | 132               | + + + +             |
| TR12898_e0_g1_i1 | TJB.Eo_023* | (CAG)5               | CAGGAAGGAAGCAAGCAAAG  | GCTGCTGATGAGATGAGT   | 184               | + + + +             |
| TR13811_e0_g1_i1 | TJB.Eo_024 | (TCA)6               | ATTTTCCCTGCAAAGCCCTG  | GGGCAGCTGTTGTGATGTT   | 166               | + + + +             |
| TR13845_e0_g1_i1 | TJB.Eo_025* | (A)12                | ATCTCTGACATGGAGAAGCC  | AGGAGGAAGCAAGCAAGCAA  | 118               | + + + +             |
| TR13927_e0_g2_i1 | TJB.Eo_026 | (ACA)5               | CAAAGCGAGATGAGATGGCG  | GGTGTGCTGCGGAGATGTTG  | 279               | + + + +             |
| TR14515_e0_g1_i1 | TJB.Eo_027 | (TG)6                | GGTGTGCTGTTATGAGGGC   | TGTTGCTGCTGATGAGTGG   | 215               | - - - -             |
| TR14777_e0_g1_i1 | TJB.Eo_028* | (CTG)10              | AGTTCATTTGCCAGCTCAT   | TAAACTGCGTGAGCAGGAAAAC | 195               | + - - -             |

(Continued)
| ID            | Marker Name | SSR       | Forward primer (5’-3’) | Reverse primer (5’-3’) | Product size (bp) | PCR amplification in E092-1 | E092 | E022 | E039 |
|---------------|-------------|-----------|------------------------|------------------------|-------------------|-----------------------------|-------|------|------|
| TR15964_c0_g1_i1 | TJB_Eo_029* | (CAA)5    | AACAAGGCAACAGGACAGAG   | ATTTTCGACCGGATAGTTTT  | 182               | +                            | +     | +    | +    |
| TR16265_c0_g1_i1 | TJB_Eo_030  | (ATC)7    | CGCATATGTTGAGGAGTGTGA | CGACAATGCGGTGTGATA    | 254               | -                            | -     | -    | -    |
| TR16281_c0_g1_i1 | TJB_Eo_031* | (CAC)13   | GACGACGATGAGATAGAGA    | CTAGTGTTGAGGATAGAGAG | 154               | +                            | +     | +    | +    |
| TR16284_c1_g1_i1 | TJB_Eo_032  | (ACA)16   | CAAAATAGGCAATGAGGCA    | TCTGAGCTTCTCTGAGAGAG | 109               | -                            | -     | -    | -    |
| TR16410_c1_g1_i1 | TJB_Eo_033  | (GTT)12   | AGCTTATGCTCTCTGAGAG    | CAATATCCGAGACAGTGGTT | 155               | +                            | -     | -    | -    |
| TR17043_c1_g1_i1 | TJB_Eo_034  | (TCA)10   | CATCTGCTATCATCAGGAG    | GCTGGCTTGGGACTAGTGCG | 204               | -                            | -     | -    | -    |
| TR17216_c0_g1_i1 | TJB_Eo_035  | (CAA)19   | ATTTGGCAATGAGATAGAGA   | TGCTGAGCTTCTCTGAGAG   | 261               | +                            | +     | +    | +    |
| TR17372_c0_g1_i1 | TJB_Eo_036  | (TTG)10   | GATGGTGGTGGGAGGAGGAG   | ACAAATAGGCAATGAGGCA   | 254               | +                            | +     | +    | +    |
| TR17509_c1_g1_i1 | TJB_Eo_037  | (TGG)5    | GCAACGATGAGAGCAGAAG    | GCTGGCTTGGGACTAGTGCG | 185               | -                            | -     | -    | -    |
| TR18313_c0_g1_i1 | TJB_Eo_038  | (GCT)6gg  | GCTGCTGCTGCTGCTGCTGCT | TACTGCTCAGGACAGGATTT  | 236               | +                            | +     | +    | +    |
| TR18539_c0_g2_i1 | TJB_Eo_039  | (AAC)10   | CCAGGAAAGAACAGGACAG    | GCTGGAGCTTCTCTGAGAG   | 103               | -                            | -     | -    | -    |
| TR21460_c3_g4_i1 | TJB_Eo_040  | (CGT)5    | CACCTGCTCAAGACACAC     | TCAAGGCAACAGAAGAGGAG  | 165               | +                            | +     | +    | +    |
| TR21890_c0_g1_i1 | TJB_Eo_041  | (GCT)5    | AGACGACCTCAAGGACTCT    | GAGCTTCTCTGAGAGAAGAG  | 192               | +                            | +     | +    | +    |
| TR23033_c0_g1_i1 | TJB_Eo_042  | (GTT)10   | AGCTTATGCTCTCTGAGAG    | CAAGGCAGCTACCATCAAGA  | 108               | -                            | -     | -    | -    |
| TR23780_c0_g1_i1 | TJB_Eo_043  | (ACC)6as(CAG)5(CAA)5 | CGGAAATGCAACACCAAACTA | ACAAGCTGACCTGAGACT   | 141               | -                            | -     | -    | -    |
| TR23987_c0_g1_i1 | TJB_Eo_044* | (TTG)11   | CTGTTTCAAGCGGAGCTTCCC  | GAGCAGTATACGAGCAGAAA  | 185               | -                            | +     | +    | +    |
| TR26689_c0_g1_i1 | TJB_Eo_045  | (TCT)10   | ACAGTCTGCTCCACACATC    | GCAGAGCTGACACAGAAGAG  | 275               | +                            | +     | +    | +    |
| TR26876_c2_g2_i9 | TJB_Eo_046* | (A)11     | AAAATCAACCAGACGACAC    | ATGCTGCTCTGAGAAGAG    | 166               | -                            | +     | +    | +    |
| TR27357_c0_g3_i1 | TJB_Eo_047  | (ACA)10   | GCAACATATGCTTGCTAGAG   | CAAAGCGCAATGCAATGTA   | 237               | +                            | +     | +    | +    |
| TR27781_c0_g1_i1 | TJB_Eo_048  | (CAG7)(CAA)5 | TTACAAGCTGCTGCTGAGGAGT | GTCTGCTGCTGAGATAGGGAT | 172               | +                            | +     | +    | +    |
| TR28391_c0_g1_i1 | TJB_Eo_049  | (GT)6     | ACATGAAAGCTGCTGCTGAGG  | AAAAATAGGCAATGAGGCA   | 110               | +                            | +     | +    | +    |
| TR28656_c0_g1_i1 | TJB_Eo_050  | (TATG)11  | GCAATTGCTATGCTGAGAGAG  | GCTGGCTTGGGACTAGTGCG  | 170               | -                            | -     | -    | -    |
| TR29033_c0_g3_i1 | TJB_Eo_051* | (TTG)11   | TGTATGCTGCGGAGATGGG    | GCTGGCAACCTGCTAGAGAG  | 108               | +                            | +     | +    | +    |
| TR29155_c1_g3_i1 | TJB_Eo_052* | (CAA)5    | GCAACACACACACACAAACAC  | GCAAGCTGATGCTGAGGAT   | 258               | +                            | +     | +    | +    |
| TR29597_c0_g1_i1 | TJB_Eo_053  | (GTT)6    | GCTGATGACACACGAGACAG   | TCAAGATGCTGAGGCAGCA   | 202               | +                            | +     | +    | +    |
| TR29883_c0_g1_i4 | TJB_Eo_054* | (GGA)10   | CTTAGGCAACACCCCATCCT   | GTGAGCTGCTGAGCAGAGAG  | 161               | +                            | +     | +    | +    |
| TR30407_c0_g1_i1 | TJB_Eo_055* | (AG)12    | GCTGCTGCTGACTACACACA   | TCTGGCTCTCTCTGAGAG    | 136               | +                            | +     | +    | +    |
| TR30845_c0_g1_i1 | TJB_Eo_056* | (ATG)10   | ACCACACGACAGTGAGCAGC   | CATTGCTCTTCTCTGAGAG   | 202               | +                            | +     | +    | +    |
| TR31759_c0_g1_i1 | TJB_Eo_057  | (ATC)11   | TGGCTGAGAAACGTTGACA    | TGCTGCTGCTGCTGAGGATG | 192               | +                            | +     | +    | +    |
| TR32331_c0_g1_i1 | TJB_Eo_058  | (GAG)6    | GAAAGAAGGAGAGTGAGCAA   | ACTGACTACACTCCCAACAC  | 191               | +                            | +     | +    | +    |

(Continued)
| ID          | Marker Name            | SSR | Forward primer (5’-3’) | Reverse primer (5’-3’) | Product size (bp) | PCR amplification in E092-1 | E092 | E022 | E039 |
|-------------|------------------------|-----|------------------------|------------------------|------------------|-----------------------------|-------|------|------|
| TR33540_c0_g1_i1 | TJIB.Eo_059  | (ATG)10 | TGTTCAAGTGATGCAAGACGATC | AGATCGTGGTGTTGCAAGATC | 244              | -                        | -    | -    | -    |
| TR33948_c0_g1_i1 | TJIB.Eo_060  | (TGT)5(TGC)7 | GCTGTATTGGATGTCAGCATC | GCTGTAAGTGTCAGCATC | 250              | +                        | -    | -    | -    |
| TR34533_c0_g3_i1 | TJIB.Eo_061  | (CTG)12 | TCAAGCTGTGGTGTAACGC | GATGCAAGAGGATGTTCTAG | 276              | +                        | +    | +    | +    |
| TR34630_c0_g1_i3 | TJIB.Eo_062  | (CCG)5  | GATGCTGAGGAGCATTTTGG | AGATGCAAGAGGATGTTCTA | 237              | +                        | +    | +    | +    |
| TR35413_c0_g1_i1 | TJIB.Eo_063  | (TGT)8  | TTATGATGAGTAAAAAGGACGAAG | TTTATGATGAGTAAAAAGGACGAAG | 181             | +                        | +    | +    | +    |
| TR39034_c2_g2_i1 | TJIB.Eo_064  | (TGA)12 | CTATGCTGATGCTCACTGCTAA | CGGCTCTGCTCAGGCAATC | 134              | -                        | -    | -    | -    |
| TR39482_c0_g1_i1 | TJIB.Eo_065  | (CCT)7  | TGCTGATGCTGCTCGAGCAG | AGATGCAAGAGGATGTTCTA | 142              | +                        | +    | +    | +    |
| TR39866_c0_g1_i2 | TJIB.Eo_066  | (ATC)14 | GGTGTTGTGGATGATGTTCTA | AGATGCAAGAGGATGTTCTA | 198              | -                        | -    | -    | -    |
| TR42732_c0_g1_i1 | TJIB.Eo_067  | (GA)6   | GAGATCTACCGAGCACGCAAG | AGATGCAAGAGGATGTTCTA | 124              | +                        | +    | +    | +    |
| TR43258_c0_g1_i1 | TJIB.Eo_068  | (ATC)14 | AAGCTGCTCAGATGCTGTTCTA | AGATGCAAGAGGATGTTCTA | 268              | -                        | -    | -    | -    |
| TR44815_c0_g2_i1 | TJIB.Eo_070  | (CAC)10 | AGGCAAGAGGATGTTCTAAGG | AGATGCAAGAGGATGTTCTA | 198              | -                        | -    | -    | -    |
| TR45627_c0_g1_i1 | TJIB.Eo_071  | (GA)6   | GAGATCTACCGAGCACGCAAG | AGATGCAAGAGGATGTTCTA | 124              | +                        | +    | +    | +    |
| TR46377_c0_g1_i1 | TJIB.Eo_072  | (GAA)10 | GAGATCTACCGAGCACGCAAG | AGATGCAAGAGGATGTTCTA | 229              | +                        | +    | +    | +    |
| TR46548_c0_g1_i2 | TJIB.Eo_073  | (CCG)11 | TACAGCAATCCTCCAGCTCCT | GAGAAGATGCTCCTCCAGCTCCT | 278              | +                        | +    | +    | +    |
| TR46887_c0_g3_i1 | TJIB.Eo_074  | (CTG)11 | CTTTGAGAGGAGGCTTATTTG | TGACATGAGTGAGATGCTC | 115              | +                        | +    | +    | +    |
| TR46947_c0_g1_i1 | TJIB.Eo_075  | (CCT)11 | TCTCCAGAAGATGCTGTTCTA | GCTGACTCCTCCAGCTCCT | 210              | +                        | +    | +    | +    |
| TR47176_c0_g1_i1 | TJIB.Eo_076  | (AAC)10 | CTTTGCTCAGACTCAATGTT | GCTGACTCGTGGATGCTC | 179              | -                        | -    | -    | -    |
| TR47761_c0_g2_i1 | TJIB.Eo_077  | (GTT)14 | CCTTCTGAGGCTCCTCCAGCTC | GAGAAGATGCTCCTCCAGCTC | 135              | +                        | +    | +    | +    |
| TR48071_c1_g1_i2 | TJIB.Eo_078  | (ACC)10 | CAGTGAGTCTCCACCTCCCTCTC | AGATGCAAGAGGATGTTCTA | 235              | +                        | +    | +    | +    |
| TR49631_c0_g1_i1 | TJIB.Eo_079  | (AAC)7  | AAATCATTGAGTGGTGGAGCAG | TGCTGCTCCTAAGATGATGCTC | 211              | -                        | -    | -    | -    |
| TR51210_c0_g2_i1 | TJIB.Eo_080  | (ATC)14 | AAGGCCTTCTCCAGCTCACC | ATATGAGTGGGCTCCTGTTT | 195              | +                        | +    | +    | +    |
| TR51244_c7_g2_i1 | TJIB.Eo_081  | (CGG)12 | TGCTGTTGAAAAATCCACAC | TCTTCTGCGTGGGATGCTC | 259              | -                        | -    | -    | -    |
| TR51733_c0_g4_i1 | TJIB.Eo_082  | (TCT)13 | AGAAGACATAGGGCCACAC | GCTGACTCCTCAATGTTCTGAT | 156              | +                        | +    | +    | +    |
| TR51940_c0_g2_i1 | TJIB.Eo_083  | (CTT)15 | TGCGAATCGAGTGGTGTTCTT | CCAGCTGCTTCCTCTACTG | 272              | -                        | +    | +    | -    |
| TR52704_c4_g2_i1 | TJIB.Eo_084  | (CTC)10 | GGGTTATGAGTAATGGAAGT | GCTGACTCCTCCCTCCTCCTG | 144              | +                        | +    | +    | +    |
| ID           | Marker Name | SSR                          | Forward primer (5'-3') | Reverse primer (5'-3') | Product size (bp) | PCR amplification in |
|--------------|-------------|------------------------------|------------------------|------------------------|------------------|----------------------|
| TR53512_c1_g1_i5 | TJB_E0_085  | (A)12                        | CACCTTCAAAACCTGCCATT   | CACAACACACACTACGGGAG  | 258              | + + + +              |
| TR53516_c0_g1_i1 | TJB_E0_086" | (TGA)10                      | GGGCATATGCCTGGCAAT     | AAGCATAGAAGCGACGCAAT  | 139              | - + + +               |
| TR53584_c1_g1_i1 | TJB_E0_087" | (ACA)6acagcgtcgc(CAA)6       | TGTGCACTATTGGTACAGCC   | GGAATGCTGAGAAGAGAGA   | 255              | + + + +               |
| TR53641_c0_g1_i1 | TJB_E0_088  | (GTT)10                      | GCCCTTAGAAATGAGACCGT   | CACAACCCAAAATCATACGA  | 144              | - + + -               |
| TR53936_c0_g1_i1 | TJB_E0_089" | (CAA)5                       | CCTCTGTCGTGATGATGCA    | ACTGCAACACACAGGTCG    | 209              | + + + +               |
| TR54047_c0_g1_i1 | TJB_E0_090" | (CTAC)18                     | GGCCACACTGCTTTAACCAT   | TCACGGTTAAATCTTTGGAC  | 246              | + + + +               |
| TR54725_c0_g4_i1 | TJB_E0_091" | (TCA)5                       | CACACTCCGAGGTGACTT     | AGGAGTGCCCACAATGACA   | 218              | + + + +               |
| TR54851_c2_g5_i1 | TJB_E0_092" | (TCC)10                      | CGGCCATCTTTGTCGCA      | GAGAGGAGGAGGAGGACG    | 228              | + + + +               |
| TR56883_c0_g2_i1 | TJB_E0_093  | (TTG)10                      | TACACTGCTGAGCAAATCG    | CAAGCGATGTCGAAAGCAA   | 225              | + - + -               |
| TR57592_c0_g1_i1 | TJB_E0_094  | (CAA)10                      | CAGAATGCGCCGATTTGCT    | TCTGATACCCGATGCTGCG   | 229              | - - - -               |
| TR58461_c0_g2_i1 | TJB_E0_095  | (GAA)12                      | AACTCGAAGCGAAACGCTC    | CTGCGAGCCCTGATGCTGA   | 213              | - - - -               |
| TR59379_c0_g2_i2 | TJB_E0_096  | (CAT)5                       | GTTGTCGATACGACGCGC     | TAGCGTGACAAATGACG     | 138              | - - + +               |
| TR59407_c0_g1_i1 | TJB_E0_097" | (CCT)10                      | CATAACACGACACATGCGG    | GAGGAGGAGGAGAAGAAGA   | 264              | + + + +               |
| TR60466_c0_g1_i1 | TJB_E0_098  | (CAG)6                       | TACGGCTGACGATATGCTAC   | TGTGCTGATTGGATGCTG    | 108              | + + + +               |
| TR60586_c0_g1_i1 | TJB_E0_099  | (TGC)5                       | TGGACACCTGGTGGTGGGAG   | CGCAAGAGACGATGACG     | 247              | - - + -               |
| TR61247_c0_g4_i1 | TJB_E0_100  | (GTT)17                      | ACAGATGCTCGTGGGTTTT    | ATTTGAGTCGCGTCCAGACT  | 238              | + - - -               |

* These marker loci showed polymorphic amplifications in the four accessions of *E. ophiuroides*.

[https://doi.org/10.1371/journal.pone.0202605.t003](https://doi.org/10.1371/journal.pone.0202605.t003)
Centipedegrass is an important warm-season grass species that is widely used for turf in the southeastern United States and in the south of China. Because of its superiorities over other turf-grasses, which mainly include its low maintenance requirements and good adaptation to poor soil, centipedegrass shows great potential for commercial application in the turf industry. However, a lack of genomic information has hampered critical research on augmenting marker assisted breeding programs for this species. Hence, the development of an effective marker system to assess genetic diversity in centipedegrass collections facilitates the maintenance of germplasm and cultivar improvement. Prior to our study, 30 SSR primer pairs were developed in *E. ophiuroides* using the traditional method [29]. Recently, approximately 3,500 SSR primer pairs were successfully designed from 5,839 identified SSR loci based on Illumina paired-end sequencing reads [21]. In the present study, a total of 48,061 SSR primer pairs were developed from assembled centipedegrass transcriptome sequences using an NGS-based RNA-seq technique. These SSRs were mined from the transcriptomes, meaning in the gene coding regions only. However, the gene-coding regions should have fewer SSRs than those of the intergenic regions [48]. Therefore, the SSRs developed here are only part of the SSRs in the genome of centipedegrass. Even so, these results represent an important complementation to the molecular toolbox of centipedegrass, which might contribute to promoting genetic and genomic research on centipedegrass.

The motif type and abundance of SSRs are the main characteristics of microsatellites. In this study, the most abundant motif was that of tri-nucleotides (52.14%) followed by di-nucleotides (44.65%), tetra-nucleotides (2.17%), hex-nucleotides (331, 0.51%) and penta-nucleotides (0.49%). This result is consistent with previous findings, showing that tri-nucleotides are the most common type in centipedegrass [21], and is also in accord to the findings in many other plant species, such as alfalfa [49], Dysosma [50], peanut [51], common bean [52], and...
Table 4. Polymorphism statistics of novel SSR markers in 43 *E. ophiuroides* core collections.

| ID     | Marker name   | Na  | Ne  | Ho  | I   | PIC |
|--------|---------------|-----|-----|-----|-----|-----|
| TR1654_c2_g1_i1 | TJIB.Eo_004 | 3   | 1.88 | 0.48 | 0.66 | 0.36 |
| TR2338_c1_g1_i1 | TJIB.Eo_006 | 10  | 3.52 | 0.73 | 1.41 | 0.67 |
| TR2941_c0_g1_i2 | TJIB.Eo_008 | 8   | 5.16 | 0.83 | 1.75 | 0.78 |
| TR4044_c0_g1_i1 | TJIB.Eo_009 | 7   | 1.40 | 0.29 | 0.52 | 0.26 |
| TR5354_c0_g2_i1 | TJIB.Eo_010 | 11  | 4.07 | 0.77 | 1.65 | 0.72 |
| TR6729_c0_g1_i1 | TJIB.Eo_013 | 10  | 2.34 | 0.59 | 1.02 | 0.49 |
| TR7108_c0_g1_i1 | TJIB.Eo_014 | 7   | 2.59 | 0.63 | 1.01 | 0.54 |
| TR7421_c0_g1_i1 | TJIB.Eo_016 | 5   | 1.20 | 0.17 | 0.31 | 0.15 |
| TR8741_c0_g1_i1 | TJIB.Eo_017 | 8   | 3.82 | 0.76 | 1.44 | 0.69 |
| TR9728_c0_g2_i1 | TJIB.Eo_020 | 8   | 5.27 | 0.83 | 1.71 | 0.78 |
| TR11518_c0_g1_i1 | TJIB.Eo_022 | 7   | 2.56 | 0.62 | 1.14 | 0.56 |
| TR12898_c0_g1_i1 | TJIB.Eo_023 | 10  | 5.24 | 0.83 | 1.76 | 0.78 |
| TR13845_c0_g1_i1 | TJIB.Eo_025 | 6   | 1.90 | 0.48 | 0.83 | 0.42 |
| TR14777_c0_g1_i1 | TJIB.Eo_028 | 5   | 1.57 | 0.38 | 0.55 | 0.30 |
| TR15964_c0_g1_i1 | TJIB.Eo_029 | 10  | 2.48 | 0.61 | 1.11 | 0.55 |
| TR16281_c0_g1_i1 | TJIB.Eo_031 | 12  | 3.36 | 0.72 | 1.40 | 0.65 |
| TR18313_c0_g1_i1 | TJIB.Eo_038 | 11  | 3.36 | 0.72 | 1.34 | 0.65 |
| TR21460_c3_g4_i1 | TJIB.Eo_040 | 10  | 3.30 | 0.71 | 1.36 | 0.65 |
| TR23987_c0_g1_i1 | TJIB.Eo_044 | 9   | 3.38 | 0.72 | 1.31 | 0.65 |
| TR26876_c2_g2_i9 | TJIB.Eo_046 | 11  | 3.04 | 0.69 | 1.22 | 0.62 |
| TR27357_c0_g3_i1 | TJIB.Eo_047 | 9   | 4.31 | 0.79 | 1.52 | 0.73 |
| TR29033_c0_g3_i1 | TJIB.Eo_051 | 7   | 3.74 | 0.75 | 1.41 | 0.69 |
| TR29155_c1_g3_i3 | TJIB.Eo_052 | 13  | 3.31 | 0.71 | 1.26 | 0.64 |
| TR29833_c0_g1_i4 | TJIB.Eo_054 | 9   | 2.25 | 0.57 | 1.09 | 0.51 |
| TR30407_c0_g1_i1 | TJIB.Eo_055 | 11  | 4.68 | 0.81 | 1.65 | 0.76 |
| TR30845_c0_g1_i1 | TJIB.Eo_056 | 9   | 3.01 | 0.68 | 1.28 | 0.62 |
| TR33948_c0_g1_i1 | TJIB.Eo_060 | 11  | 2.40 | 0.60 | 0.99 | 0.49 |
| TR34333_c0_g3_i1 | TJIB.Eo_061 | 10  | 4.50 | 0.80 | 1.60 | 0.74 |
| TR34630_c0_g1_i3 | TJIB.Eo_062 | 7   | 2.04 | 0.52 | 0.81 | 0.42 |
| TR35413_c0_g1_i1 | TJIB.Eo_063 | 9   | 1.60 | 0.39 | 0.56 | 0.31 |
| TR39482_c0_g1_i1 | TJIB.Eo_065 | 8   | 2.98 | 0.68 | 1.25 | 0.60 |
| TR39754_c0_g1_i1 | TJIB.Eo_066 | 10  | 3.47 | 0.73 | 1.35 | 0.66 |
| TR42732_c0_g1_i1 | TJIB.Eo_068 | 8   | 2.14 | 0.55 | 0.97 | 0.48 |
| TR45627_c0_g1_i1 | TJIB.Eo_071 | 10  | 2.62 | 0.63 | 1.03 | 0.55 |
| TR46377_c0_g1_i1 | TJIB.Eo_072 | 10  | 2.29 | 0.58 | 0.98 | 0.48 |
| TR46548_c0_g1_i2 | TJIB.Eo_073 | 11  | 1.94 | 0.50 | 0.75 | 0.39 |
| TR46887_c0_g3_i1 | TJIB.Eo_074 | 9   | 2.86 | 0.67 | 1.21 | 0.59 |
| TR46947_c0_g1_i1 | TJIB.Eo_075 | 8   | 2.46 | 0.61 | 1.04 | 0.52 |
| TR47761_c0_g2_i1 | TJIB.Eo_077 | 5   | 1.91 | 0.49 | 0.74 | 0.38 |
| TR48071_c1_g1_i2 | TJIB.Eo_078 | 8   | 2.65 | 0.64 | 1.13 | 0.57 |
| TR51210_c0_g2_i1 | TJIB.Eo_080 | 9   | 2.43 | 0.60 | 1.21 | 0.55 |
| TR51733_c0_g4_i1 | TJIB.Eo_082 | 8   | 3.72 | 0.75 | 1.35 | 0.68 |
| TR52704_c4_g2_i1 | TJIB.Eo_084 | 5   | 3.07 | 0.69 | 1.23 | 0.61 |
| TR53516_c0_g1_i1 | TJIB.Eo_086 | 9   | 4.37 | 0.79 | 1.62 | 0.74 |
| TR53584_c1_g1_i1 | TJIB.Eo_087 | 8   | 5.01 | 0.82 | 1.66 | 0.77 |
| TR53936_c0_g1_i1 | TJIB.Eo_089 | 11  | 2.70 | 0.65 | 1.15 | 0.57 |
| TR54047_c0_g1_i1 | TJIB.Eo_090 | 4   | 3.16 | 0.70 | 1.25 | 0.63 |

(Continued)
chrysanthemum [53]. This result indicates that tri-nucleotide repeats are dominant over the other motif types in plant cDNA sequences, which might be explained by the fact that the variations of tri-nucleotide repeats will not affect the open reading frames and only lead to the adding or subtracting of amino acid repeats [54]. AG/CT was the most common motif unit of the grouped SSRs, representing 8.96% of the grouped SSR motif units in the present study, which is common in the genomes of many plant species [49, 55, 56].

In the present investigation, the 100 randomly selected primer pairs were used to check the effectiveness of the actual amplification of all the developed markers. Of these selected primer pairs, 81 (81.0%) of the primers successfully produced amplification in at least one of the four *E. ophiuroides* accessions, and 56% successfully amplified alleles from all four centipedegrass accessions, while 19 primers showed no amplification. The unsuccessful amplification indicated that transcriptome-based markers may not work for genomic DNA, which is mainly attributed to the fact that the primer originated from an erroneously assembled transcript or that the primer sequence contained exon-exon junctions. Out of these 81 primers, 50 (61.73%) showed polymorphic patterns, which was lower than the previously reported 87.78% polymorphic genic-SSRs across 14 centipedegrass accessions [21]. In addition to the higher proportion of monomorphic markers, the initial use of a lower quantity of centipedegrass accessions (only four accessions used) for the polymorphism screening was the main reason for the lower proportion of polymorphic markers detected in our study. For the further genetic diversity analysis among the 43 core accessions of *E. ophiuroides*, the Na detected by the 50 SSR markers varied from three to thirteen, with an average of 8.40 alleles per locus, while the Ne per locus was from 1.20 to 5.27, with an average of 3.01. This discrepancy can be explained by the fact that quite a few of the alleles identified were collection-common. At the same time, the average Na (8.4) and Ne (3.01) and the Ho and the I detected in the present study were higher than those in a previous study by Wang et al [21], which was because more centipedegrass accessions were used in the present study and because the 43 accessions derived from a core collection of centipedegrass germplasms were preliminarily constructed through a systematic work, including the initial collection of germplasms and the subsequent identification and screening [57]. Moreover, the PIC value is commonly used as a measure of polymorphism for a marker locus and is determined by both the number of alleles and their frequency distribution within the population. According to the classification basis for the marker loci informativeness level proposed by Botstein et al (1980) [58], all the *E. ophiuroides* SSRs validated in the present study indicated a moderate level (0.50 > PIC > 0.25) to a high level of informativeness (PIC > 0.5), with an average PIC value of 0.58, with the exception of TJIB.Eo_016 (PIC = 0.15), which was considered a poor marker in these respects. In addition, 70.0% (35 out of 50) of the markers showed a high informativeness level and worked as resolving power markers.

Currently, NGS-based RNA-seq has become one of the most efficient ways for developing genic SSR markers in both model and non-model plants, and many RNA-seq derived SSR

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**Table 4. (Continued)**

| ID            | Marker name | Na | Ne    | Ho    | I     | PIC  |
|---------------|-------------|----|-------|-------|-------|------|
| TR54725_c0_g4_i1 | TJIB.Eo_091 | 5  | 2.62  | 0.63  | 1.02  | 0.54 |
| TR54851_c2_g5_i1 | TJIB.Eo_092 | 6  | 3.07  | 0.69  | 1.17  | 0.61 |
| TR59407_c0_g1_i1 | TJIB.Eo_097 | 5  | 1.92  | 0.49  | 0.97  | 0.36 |
| **Total**     |             | 420| –     | –     | –     | –    |
| **Mean**      |             | 8.40| 3.01  | 0.64  | 1.17  | 0.58 |

Na, number of alleles; Ne, number of effective alleles; Ho, observed heterozygosity; I, Shannon's information index; PIC, polymorphism information content.

https://doi.org/10.1371/journal.pone.0202605.t004
markers have been widely utilized in genetic diversity analysis [59,60], chromosome mapping [61,62] and gene-based association studies [63–65]. In the present study, the usefulness of the 50 newly developed polymorphic EST-SSR markers for the evaluation of genetic diversity among centipedegrass accessions was clearly demonstrated. Based on the analysis results, the 50 markers divided the 43 E. ophiuroides accessions into two main groups and into six subgroups using the UPGMA cluster analysis. Although the resulting dendrogram could not sufficiently cluster the accessions based on their geographical origins, it precisely demonstrated the
effectiveness of these SSR markers in the *E. ophiuroides* genetic analysis. It is reasonable that the genetic relationship of the 43 *E. ophiuroides* accessions did not exactly correspond to their geographical positions due to their complex genetic background and evolution history. Therefore, these large-scale developed SSR markers from centipedegrass transcriptome are valuable tools for further genetic and genomic analyses of centipedegrass accessions. Moreover, Wang et al (2017) [21] reported that centipedegrass EST-SSRs were applicable for six Poaceae relatives and that a higher cross-species transferability of the SSRs was detected in C4 plant maize, sorghum, and sultan grass than in C3 plant wheat, rice, and barley. A good proportion of the centipedegrass genic SSRs mined in this study also functioned in two other Poaceae C4 members, zoysiagrass and bermudagrass (S1 Fig). Both findings proved the cross-species transferability of the centipedegrass SSR markers to other genetically closed species and reflected the close relationship of the C4 plants. Furthermore, the identification of the EST-SSR within the sequences provides a future opportunity to mine the expressed sequences for significant physical and functional association with turf traits of interest in marker-assisted breeding in *E. ophiuroides* and other closely related turf species.

Conclusions

The present work represents a substantial advance in the identification of a large number of informative SSR loci in *E. ophiuroides* by high-throughput RNA sequencing technology based on the Illumina HiSeq 2000 platform. A total of 64,470 SSR loci were identified from the assembled transcriptome of *E. ophiuroides*. Among them, the trinucleotide SSRs were the most dominant repeat motif (52.14%), followed by dinucleotides (28,783, 44.65%), tetranucleotides (1,399, 2.17%), hexanucleotides (331, 0.51%) and pentanucleotides (317, 0.49%). In total, 48,061 primer pairs were successfully designed from these identified SSR loci, and a subset of the 100 primer pairs was randomly selected and preliminarily tested in two different types of centipedegrass accessions. The PCR analysis revealed that 81.00% of the primer pairs successfully worked in at least one of the four *E. ophiuroides* accessions, and that 56% of the primer pairs successfully amplified alleles from all four accessions. Among the primers tested, 50 (61.73%) of the primers generated polymorphic bands and were further applied to assess the genetic diversity level of 43 centipedegrass core collections. In total, 420 alleles were detected by these newly developed SSR markers in the 43 collections, of which 285 were collection-specific alleles and 135 were multi-accession alleles. The PIC values ranged from 0.15 to 0.78, with an average of 0.58, and the I ranged from 0.31 to 1.76 with an average of 1.17. The 43 *E. ophiuroides* core collections were successfully clustered into two main groups and six subgroups based on the UPGMA cluster analysis. Thus, in the present study, the newly developed polymorphic SSR markers successfully shed light on the levels of the molecular diversity inherited in the core collections of centipedegrass deposited in the Turfgrass Research Center of Institute of Botany, Jiangsu Province and Chinese Academy of Sciences (JIB). The SSR primers and sequence information developed in the present study will be useful and robust resources for future genetic and genomic studies, such as genetic map construction and comparative genomic analyses, and molecular marker-assisted breeding in centipedegrass and its related species.

Supporting information

S1 Fig. Examples of PCR amplification of centipedegrass SSR markers in bermudagrass (*Cynodon dactylon* L). M: DNA ladder marker; C1: *Cynodon* accession 118M; C2: *Cynodon* accession 118. The captital letter A, B, . . .,N represents SSR marker, respectively. (PPTX)
S1 Table. Statistics of *E. ophiuroides* SSR mining.
(XLSX)

S2 Table. Statistics of SSR motif (mer) types of *E. ophiuroides*.
(XLSX)

S3 Table. Statistics of grouped SSR motif units of *E. ophiuroides*.
(XLSX)

S4 Table. Statistics of SSR length of *E. ophiuroides*.
(XLSX)

S5 Table. SSR markers developed from the de novo transcriptome sequence of *E. ophiuroides*.
(XLSX)

S6 Table. Nei’s unbiased measures of genetic distance between *E. ophiuroides* core collections.
(XLS)

**Acknowledgments**

The authors would like to thank anonymous reviewers for their constructive comments. We also thank Ms. Ai-gui Guo and Miss Jingjing Ma for help in maintaining experimental materials. We also acknowledge the helpful guidance of Dr. Weicong Qi in the processes of data analysis.

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**References**

1. Gupta PK, Varshney RK, Sharma PC, Ramesh B. Molecular markers and their applications in wheat breeding. Plant Breed. 1999; 118: 369–390.
2. Kuleung C, Baenziger PS, Dweikat I. Transferability of SSR markers among wheat, rye, and triticale. Theor Appl Genet. 2004; 108: 1147–1150. https://doi.org/10.1007/s00122-003-1532-5 PMID: 15067402

3. Powell W, Morgante M, Andre C, Hanafey M, Vogel J, Tingeys S, et al. The comparison of RFLP, RAPD, AFLP and SSR (microsatellite) markers for germplasm analysis. Mol Breed. 1996; 2: 225–238.

4. Varshney RK, Sigmund R, Börner A, Korzun V, Stein N, Sorrells ME, et al. Interspecific transferability and comparative mapping of barley EST-SSR markers in wheat, rye and rice. Plant Sci. 2005; 168: 195–202.

5. Iniguezly FL, Voort AV, Osborn TC. Development of a set of public SSR markers derived from genomic sequence of a rapid cycling Brassica oleracea L. genotype. Theor Appl Genet. 2008; 117: 977–985. https://doi.org/10.1007/s00122-008-0837-9 PMID: 18651126

6. Yang H, Tao Y, Zheng Z, Li C, Sweetingham M, Howieson J. Application of next-generation sequencing for rapid marker development in molecular plant breeding: a case study on anthracnose disease resistance in Lupinus angustifolius L. BMC Genomics 2012; 13: 318. https://doi.org/10.1186/1471-2164-13-318 PMID: 22805587

7. Zheng Y, Guo H, Zang G, Liu J. Genetic linkage maps of centipedegrass Eremochloa ophiuroides (Munro) Hack based on sequence-related amplified polymorphism and expressed sequence tag-simple sequence repeat markers. Sci Hortic. 2013; 156: 86–92.

8. Fu BD, He SP. Transcriptome Analysis of Silver Carp Hypophthalmichthys molitrix by Paired-End RNA Sequencing. DNA Res. 2012; 19: 131–142. https://doi.org/10.1093/dnares/dsr046 PMID: 2279088

9. Hodel RGJ, Gitzendanner MA, Germain-Aubrey CC, Liu X, Crowl AA, Sun M, et al. A new resource for the development of SSR markers: millions of loci from a thousand plant transcriptomes. Appl Plant Sci. 2016; 4: 1600024.

10. Studer B, Asp T, Frei U, Hentrup S, Meally H, Guillard A, et al. Expressed sequence tag-derived microsatellite markers of perennial ryegrass (Lolium perenne L.). Mol Breed. 2008; 21: 533–548.

11. Hirata M, Cai H, Inoue M, Yuyama N, Miura Y, Komatsu T, et al. Development of simple sequence repeat (SSR) markers and construction of an SSR-based linkage map in Italian ryegrass (Lolium multiflorum Lam.). Theor Appl Genet. 2006; 113: 270–279. https://doi.org/10.1007/s00122-006-0292-4 PMID: 16791693

12. Yang SS, Tu ZJ, Cheung F, Xu WW, Lamb JH, et al. Using RNA-seq for gene identification, polymorphism detection and transcript profiling in two alfalfa genotypes with divergent cell wall composition in stems. BMC Genomics. 2011: 12: 199. https://doi.org/10.1186/1471-2164-12-199 PMID: 21504589

13. Huang X, Yan HD, Zhang XQ, Zhang J, Frazier TP, Huang DJ, et al. De novo transcriptome analysis and molecular marker development of two Hemarthria species. Front Plant Sci. 2016; 7: 496. https://doi.org/10.3389/fpls.2016.00496 PMID: 27148320

14. Yates SA, Swain MT, Hegarty MJ, Chernukin I, Lowe M, Allison GG, et al. De novo assembly of red clover transcriptome based on RNA-seq data provides insight into drought response, gene discovery and marker identification. BMC Genomics. 2014; 15: 453. https://doi.org/10.1186/1471-2164-15-453 PMID: 24912738

15. Bushman BS, Larson SR, Tuna M, West MS, Hernandez AG, Vullaganti D, et al. Orchardgrass (Dactylis glomerata L.) EST and SSR marker development, annotation, and transferability. Theor Appl Genet. 2011; 123: 119–129. https://doi.org/10.1007/s00122-011-1571-2 PMID: 21465196

16. Zhu Y, Wang X, Huang L, Lin C, Zhang X, Xu W, et al. Transcriptomic identification of drought-related genes and SSR markers in Sudan grass based on RNA-Seq. Front Plant Sci. 2017; 8: 687. https://doi.org/10.3389/fpls.2017.00687 PMID: 28523007

17. Guo Y, Wu Y, Anderson JA, Moss JQ, Zhu L, Fu J. SSR marker development, linkage mapping, and QTL analysis for establishment rate in common bermudagrass. Plant Genome. 2017; 10: 1–11.

18. Kubik C, Honig J, Bonos SA. Characterization of 215 simple sequence repeat markers in creeping bentgrass (Agrostis tifoliana L.). Mol Ecol Resour. 2011; 11: 872–876. https://doi.org/10.1111/j.1755-0998.2011.03006.x PMID: 21843299

19. Jia X, Deng Y, Sun X, Liang L, Ye X. Characterization of the global transcriptome using illumina sequencing and novel microsatellite marker information in seashore paspalum. Genes Genom. 2015; 37: 77–86.

20. Tsuruta SI, Hashiguchi M, Ebina M, Matsuo T, Yamamoto T, Kobayashi M, et al. Development and characterization of simple sequence repeat markers in zoysia japonica steud. Grassl Sci. 2005; 51: 249–257.
21. Wang P, Yang L, Zhang E, Qin Z, Wang H, Liao Y, et al. Characterization and development of EST-SSR markers from a cold-stressed transcriptome of centipede grass by illumina paired-end sequencing. Plant Mol Biol Rep. 2017; 35: 215–223.

22. Hanna WW, Burton WG. Cytology, reproductive behavior and fertility characteristics of centipede grass. Crop Sci. 1978; 18: 835–837.

23. Watson L, Dallwitz MJ. The grass genera of the world. CAB International, Wallingford, UK. 1992.

24. Islam M, Hirata M. Centipede grass (Eremochloa ophiuroides [Munro] Hack.): Growth behavior and multipurpose usages. Grassl Sci. 2005; 51: 183–190.

25. Liu M, Yang J, Lu S, Guo Z, Lin X, Wu H. Somatic embryogenesis and plant regeneration in centipede grass (Eremochloa ophiuroides [Munro] Hack.). In Vitro Cell Dev-Pl. 2008; 44: 100–104.

26. Hanna WW. Centipede grass–Diversity and vulnerability. Crop Sci. 1995; 35: 332–334.

27. Xuan J, Guo H, Liu J, Chen S. Initial identification of cold tolerance in the Eremochloa ophiuroides germplasm resource. Acta Pratacult Sin. 2003; 12: 110–114 (in Chinese with English abstract).

28. Millalewis SR, Kimball JA, Zuleta MC, Harrisshultz KR, Schwartz BM, Hanna WW. Use of sequence-related amplified polymorphism (SRAP) markers for comparing levels of genetic diversity in centipede grass (Eremochloa ophiuroides [Munro] Hack.) germplasm. Genet Resour Crop Ev. 2012; 59: 1517–1526.

29. Harrisshultz KR, Millalewis SR, Zuleta MC, Schwartz BM, Hanna WW, Brady JA. Development of simple sequence repeat markers and the analysis of genetic diversity and ploidy level in a centipede grass collection. Crop Sci. 2012; 52: 383–392.

30. Guo H, Guo A, Zong J, Yuan X, Zhang F, Liu J. Identification analysis of eight centipede grass materials by SRAP molecular markers. Acta Agrestia Sin. 2014; 22: 203–207 (in Chinese with English abstract).

31. Luo J, Liu M, Zhang C. Transgenic centipede grass (Eremochloa ophiuroides [Munro] Hack.) Overexpressing-Adenosylmethionine Decarboxylase (SAMDC) gene for improved cold tolerance through involvement of H_2O_2 and no signaling. Front Plant Sci. 2017; 8: 1655. https://doi.org/10.3389/fpls.2017.01655 PMID: 29018465

32. Reuter JA, Spacek D, Snyder MP. High-throughput sequencing technologies. Mol Cell. 2015; 58: 586–597. https://doi.org/10.1016/j.molcel.2015.05.004 PMID: 26000844

33. Grabherr MG, Haas BJ, Yassour M, Levin JZ, Thompson DA, Amit I, et al. Full-length transcriptome assembly from ma-seq data without a reference genome. Nat Biotechnol. 2011; 29: 644–652. https://doi.org/10.1038/nbt.1883 PMID: 21572440

34. Wang X, Wang L. GMATA: an integrated software package for genome-scale SSR mining, marker development and viewing. Front Plant Sci. 2016; 7: 1350. https://doi.org/10.3389/fpls.2016.01350 PMID: 27679641

35. Peakall R, Smouse PE. GenAIEx 6.5: genetic analysis in Excel. Population genetic software for teaching and research–an update. Bioinformatics. 2012; 28: 2537–2539. https://doi.org/10.1093/bioinformatics/bts460 PMID: 22820204
44. Peakall R, Smouse PE. GENALEX 6: genetic analysis in Excel. Population genetic software for teaching and research. Mol Ecol Notes. 2006; 6: 288–295.

45. Yeh FC, Yang R, Boyle T. POPGENE Version 1.31. University of Alberta, Edmonton, Canada. 1999; Available form: http://www.ualberta.ca/~fyeh/.

46. Tamura K, Dudley J, Nei M, Kumar S. MEGA4: Molecular evolutionary genetics analysis (MEGA) software version 4.0. Mol Biol Evol. 2007; 24: 1596–1599. https://doi.org/10.1093/molbev/msm092 PMID: 17488738

47. Li W, Godzik A. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. Bioinformatics. 2006; 22: 1658–1659. https://doi.org/10.1093/bioinformatics/btl158 PMID: 16731699

48. Wang X, Yang S, Chen Y, Zhang S, Zhao Q, Li M, et al. Comparative genome-wide characterization leading to simple sequence repeat marker development for Nicotiana. BMC Genomics. 2018; 19: 500. https://doi.org/10.1186/s12864-018-4878-4 PMID: 29945549

49. Liu Z, Chen T, Ma L, Zhao Z, Nan Z, et al. Global transcriptome sequencing using the illumina platform and the development of EST-SSR markers in autotetraploid alfalfa. PloS ONE. 2013; 8: e83549. https://doi.org/10.1371/journal.pone.0083549 PMID: 24349529

50. Guo R, Mao YR, Cai JR, Wang JY, Wu J, Qiu YX. Characterization and cross-species transferability of EST–SSR markers developed from the transcriptome of Dysosma versipellis (Berberidaceae) and their application to population genetic studies. Mol Breed. 2014; 34: 1733–1746.

51. Zhou X, Dong Y, Zhao J, Huang L, Ren X, Chen Y, et al. Genomic survey sequencing for development and validation of single-locus SSR markers in peanut (Arachis hypogaea L.). BMC Genomics. 2016; 17: 420. https://doi.org/10.1186/s12864-016-2743-x PMID: 27251557

52. Kalavacharya V, Liu Z, Meyers BC, Thimmapuram J, Melmaiee K. Identification and analysis of common bean (Phaseolus vulgaris L.) transcriptomes by massively parallel pyrosequencing. BMC Plant Biol. 2011; 11: 135. https://doi.org/10.1186/1471-2229-11-135 PMID: 21985325

53. Morgante M, Hanafey M, Powell W. Microsatellites are preferentially associated with nonrepetitive DNA in plant genomes. Nat Genet. 2002; 30: 194–200. https://doi.org/10.1038/ng822 PMID: 11799393

54. He X, Zheng J, Zhou J, He K, Shi S, Wang B. Characterization and comparison of est-ssrs in salix, populus, and eucalyptus. Tree Genet Genomes. 2015; 11: 820.

55. Chen H, Wang L, Wang S, Liu C, Blair MW, Cheng X. Transcriptome sequencing of mung bean (Vigna radiate L.) genes and the identification of EST-SSR markers. PloS ONE. 2015; 10: e0120273. https://doi.org/10.1371/journal.pone.0120273 PMID: 25830701

56. Zheng YQ. Detection of genetic diversity, construction of genetic linkage map and mapping QTL in centipedegrass (Eremochloa ophiuroides). Ph. D thesis, Nanjing Agricultural University, Nanjing, China. 2009.

57. Botstein D, White RL, Skolnick M, Davis RW. Construction of genetic linkage map in man using restriction fragment length polymorphisms. Am J Hum Genet. 1980; 32: 314–331. PMID: 6247908

58. Ding Y, Zhang J, Lu Y, Lin E, Lou L, Tong Z. Development of EST-SSR markers and analysis of genetic diversity in natural populations of endemic and endangered plant Phoebe chekiangensis. Biochem Syst Ecol. 2015; 63: 183–189.

59. Zhang J, Liu T, Rui F. Development of EST-SSR markers derived from transcriptome of Saccharina japonica, and their application in genetic diversity analysis. J Appl Phycol. 2018; 30: 2101–2109.

60. Du FK, Xu F, Qu H, Feng S, Tang J, Wu R. Exploiting the transcriptome of Euphrates Poplar, Populus euphratica (Salicaceae) to develop and characterize new EST-SSR markers and construct an EST-SSR database. PloS ONE. 2013; 8: e61337. https://doi.org/10.1371/journal.pone.0061337 PMID: 23593466

61. Yang T, Fang L, Zhang X, Hu J, Bao S, Hao J, et al. High-throughput development of SSR markers from Pea (Lisum sativum L.) based on next generation sequencing of a purified Chinese commercial variety. PloS ONE. 2015; 10: e0139775. https://doi.org/10.1371/journal.pone.0139775 PMID: 26440522

62. Xiao Y, Zhou L, Wei X, Mason AS, Yang Y, Ma Z, et al. Exploiting transcriptome data for the development and characterization of gene-based SSR markers related to cold tolerance in oil palm (Elaeis guineensis). BMC Plant Biol. 2014; 14: 384. https://doi.org/10.1186/s12870-014-0384-2 PMID: 25526814

63. Nie G, Tang L, Zhang Y, Huang L, Ma X, Cao X, et al. Development of SSR markers based on transcriptome sequencing and association analysis with drought tolerance in perennial grass Miscanthus from China. Front Plant Sci. 2017; 8: 801. https://doi.org/10.3389/fpls.2017.00801 PMID: 28559912

64. Xie Y, Sun X, Ren J, Fan J, Lou Y, Fu J, et al. Genetic diversity and association mapping of cadmium tolerance in bermudagrass [Cynodon dactylon (L.) Pers.]. Plant Soil. 2015; 390: 307–321.