EST-SSR markers identification based on RNA-sequencing and application in Schisandra chinensis

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Abstract

Background: Schisandra chinensis, a climbing woody vine, is the best-known and representative genus of the Schisandraceae family which is an important plant in Chinese herbal medicine; however, the application of molecular breeding is restricted by the few genetic markers for this species.

Results: In this study, we performed transcriptome sequencing of S. chinensis using the Illumina HiSeq platform to establish a library of expressed sequence tag-simple sequence repeat (EST-SSR) markers. A total of 59,786 unigenes were obtained and 6254 putative SSR sites were detected with a frequency of 10.46%. The predominant type of repeat motif was dinucleotide (35.71%), followed by trinucleotide (13.22%), hexanucleotide (0.50%), tetranucleotide (0.06%), and pentanucleotide (0.22%). We randomly selected 50 EST-SSR primer pairs and used 14 of these for genetic diversity analysis in 42 S. chinensis genotypes. All 42 accessions were successfully identified and formed four major clusters, indicating that the SSR markers can be used for genetic diversity analysis and genetic linkage map construction. In addition, using the polymorphic bands associated with 10 markers as DNA fingerprints, we generated a manual cultivar identification diagram that can distinguish between the 42 accessions, with different individuals identifiable based on polymorphic band patterns.

Conclusion: S. chinensis transcriptome data is an effective resource for developing SSR markers. These results can provide a basis for the identification of S. chinensis accessions and construction of genetic linkage maps as part of future selective breeding and conservation efforts for this valuable plant.

Background

Schisandra chinensis (Turcz.) Baill. belonging to family Schisandraceae is an important plant in Chinese herbal medicine [1–2]. While only S. sphenanthera and S. chinensis are listed in the Chinese Pharmacopoeia, in fact, most Schisandra species have medicinal value [3–4]; specifically, the fruit has multiple beneficial therapeutic and physiologic properties including adaptogenic, hepatoprotective, anticancer, antioxidant, and anti-inflammatory properties. These are mostly attributable to dibenzocyclooctadiene lignans, which are widely referred to as Schisandra lignans [5–6]. About 150 lignan derivatives with a dibenzocyclooctadiene skeleton have been identified in fruit extracts [7].
Due to habitat loss and excessive economic exploitation, the abundance of Schisandra species has decreased markedly in recent years, such that they are now endangered [8].

Molecular marker-assisted selective breeding can be applied to the genetic diversification of Schisandra varieties. This requires the identification of DNA markers and generation of linkage maps for target gene localization. However, only few types of marker have been used to date in Schisandra—namely, inter simple sequence repeats (ISSRs) [9-10], simple sequence repeats (SSRs) [11-12], amplified fragment length polymorphisms (AFLPs) [13-14], and randomly amplified polymorphic DNA (RAPD) [15]. In order to increase the collection of available markers, in the present study, we screened EST-SSR markers for Schisandra. Our results provide a foundation for research on germplasm resources and functional gene localization as well as for marker-assisted breeding of Schisandra.

Results

**SSR site distribution in S. chinensis**

We searched 59,786 unigene sequences in the transcriptome data and detected 6254 SSR sites in 4989 sequences. The SSR frequency was 10.46%. There were 897 unigene sequences containing two or more EST-SSR sites, and all sequences included a complex SSR site.

The types of EST-SSR detected in the transcriptome varied and their frequencies differed significantly (Table 2); mono-, di-, and trinucleotides were the most common, accounting for 60.06%, 31.61%, and 7.84%, respectively, of all SSRs. Tetra-, penta-, and hexanucleotides were rare, accounting for 0.49% of all SSR. There were 10 SSRs at most SSR sites (22.91%).

**Characteristics of EST-SSRs**

We analyzed di-, tri-, tetra-, penta-, and hexanucleotides; mononucleotides were excluded because of the potential for poor sequencing quality caused by homopolymerization [16]. In total, 82 types of motif were identified, with 8, 30, 25, 5, and 14 di-, tri-, tetra-, penta-, and hexanucleotide repeats, respectively (complementary sequences were considered one type of motif) (Table 3). The most highly represented EST-SSR type was dinucleotide (35.71%); TC/GA was the most common motif (48.46%) followed by CT/AG (39.52%), with other motif types constituting just 12.02% of all EST-SSR
dincleotides. There were 30 types of trinucleotide repeat motif among *S. chinensis* ESTs; the most frequent were GAA/TTC and AGA/TCT, which accounted for 16.59% (846) and 13.49% (688) of the total, respectively.

The frequency of different motifs in the EST datasets varied, with 20572 (79.39%) dinucleotides, 5100 (19.68%) trinucleotides, 206 (0.81%) tetranucleotides, 27 (0.1%) pentanucleotides, and six (0.02%) hexanucleotides (Fig. 1a). The number of motif repeats ranged from five to 12, with six being the most common number of repeats (Fig. 1b).

**Development *S. chinensis* EST-SSR primer pairs and detection of polymorphisms**

To obtain high quality SSR primers that could detect polymorphisms, we randomly selected 50 primer pairs to evaluate polymorphisms among four accessions of *S. chinensis* (Yanhong, Zaohong, Jinwuwei, and 12-(-2)-1). We identified 14 pairs of primer sets that were effective (Additional file 1), with a mean amplification rate of 28%.

**Discrimination between different *S. chinensis* genotypes using EST-SSR primer pairs**

In the genetic diversity analysis, the 14 EST-SSR primer pairs identified as described above could be used to differentiate between the 42 *S. chinensis* accessions. Using NTSYS-pc software to analyze genotype data, the accessions were classified into four groups at a similarity index of 0.63 (Fig. 2). A dendrogram revealed clear distinctions between the accessions, reflecting a high genetic diversity that can be exploited for *S. chinensis* identification based on a DNA fingerprint. The relatedness of the 42 accessions was supported by similarity coefficients ranging between 0.61 and 0.97. Group I, which comprised 28 varieties mostly originating in Jilin, was the largest group with four subgroups and a similarity coefficient of 0.682. Group II included four varieties; most were from Heilongjiang, with one accession from Jilin. ‘18-10-3’ and ‘162-1-4’ did not cluster with any of the groups and were designated as group IV, and the remaining accessions constituted group III. All 42 accessions were distinguishable based on the 14 EST-SSR markers and their clustering pattern was concordant with their distribution, indicating that EST-SSR data obtained by transcriptome analysis can reveal the genetic relatedness of *S. chinensis* germplasm resources.

**Accession identification**
There is a need for a simple, practical, and reliable method for identifying *S. chinensis* accessions. Of the 14 primer pairs that were tested, ten were required to clearly distinguish between the 42 accessions (Fig. 3). All accessions were initially identified based on different combinations of the 220-, 270-, and 280-bp bands amplified by primer pair no. 30 (Fig. 3). The smallest group contained only two strains—18-10-3 and 17-N1-N1—that were further distinguished based on a 240-bp band amplified by primer set no. 11. Likewise, all five of the other groups could be differentiated using the primers shown in Figure 3. Thus, all of the accessions could be identified using 10 pairs of primers for the construction of a manual cultivar identification diagram (MCID).

**Discussion**

*S. chinensis* (Turcz.) Baill. is a plant used to treat asthma and cough in traditional Chinese herbal medicine that is mainly distributed in northeastern China as well as in Korea, far eastern Russia, and northern Japan [17–19]. However, appreciation of the medicinal value of *S. chinensis* has led to its over-exploitation as well as habitat destruction, which has severely depleted natural *S. chinensis* sources. There is growing interest among herbalists and across the general population to preserve natural sources of important herbal medicine plants, including *S. chinensis* [8, 20–21]. Sustainable use of *S. chinensis* requires an understanding of its population genetic structure and diversity.

DNA markers are useful tools for genetic diversity analysis owing to their abundance, codominance, reproducibility, and high degree of polymorphism. A variety of markers including *rbcL*, internal transcribed spacers, AFLPs, ISSRs, and SSRs have been described in *S. chinensis* [11–13, 20, 22]. However, although DNA markers have been used to analyze genetic diversity in *Schisandrae*, there are no reports of DNA fingerprints or linkage maps for different genotypes. In the present work, transcriptome data for 42 *Schisandra* accessions were screened for EST-SSR markers. We found 6254 SSR sites in 59,786 unigenes, corresponding to a frequency of 10.46%. This value was influenced by the SSR search threshold and large number of single nucleotide repeats. In order to facilitate comparisons with other species, we analyzed the results in two groups: one including and the other excluding mononucleotide repeats. In the former context, the SSR frequency in *S. chinensis* was 10.46%, which is higher than onion (5.57%) [23] and Phoebe nanmu (9.90%) [24] but lower than oil
palm (11.26%) [25], Miscanthus (14.44%) [26], and eggplant (18.32%) [27]. Without considering mononucleotide repeats, the SSR distribution frequency was 5.79%, which is higher than corn (1.5%), barley (3.4%), rice (4.7%) [28], cucumber (4.03%) [29], Taxus chinensis (2.24%) [30], and Korean pine (4.24%) [31]; similar to peanut (6.63%) [32], pepper (7.83%) [33], and dal (7.6%) [34]; and lower than Rosa roxburghii (20.37%) [35], oak (27.61%) [36], and precocious trifoliate orange (26.88%) [37]. These differences may reflect actual variations in SSR abundance across species if we ignore the influence of the amount and source of EST data [35]. Thus, transcriptome SSRs are abundant in Schisandra, making these an ideal markers for genetic mapping.

Dinucleotides (37.83%) and trinucleotides (14.00%) were the main types of SSR in Schisandra; the two times higher frequency of the former compared to the latter is consistent with the trend observed in peanut [32], precocious trifoliate orange [37], and litchi [20], although there is also evidence that trinucleotides are the most common SSR type [20, 23, 26, 28, 31]. This difference may be related to the characteristics and quantity of EST-SSR and EST resources in plants. TC/GA was the most common dinucleotide repeat unit in S. chinensis, which is the same as pigeonpea [34] but in contrast to barley, wheat, corn, sorghum, rice [28], eggplant [27], peanut[32], Miscanthus [26], thorn pear [35], and T. chinensis [30]. GAA/TTC was the most abundant trinucleotide repeat motif, as reported for pansy [38].

Of the 50 EST-SSR primer pairs that we tested, 14 produced stable and polymorphic bands of the expected size, with an amplification efficiency of 28%; this is comparable to Morinda officinalis How (24%) [39] but higher than pigeon pea [34] and lower than onion (60%) [23], thorn pear (54.76%) [35], and T. chinensis (53.23%) [30]. Using these primer sets, the accessions could be divided into four distinct groups. To date, only nine EST-SSR markers have been identified to date for S. chinensis. In this study, we used 10 EST-SSR markers to generate an MCID identification map for different S. chinensis resources. The large number of SSR loci and polymorphic markers identified here based on transcriptome data can facilitate the identification of accessions and construction of genetic linkage maps in S. chinensis as part of broader efforts for the selective breeding and conservation of this valuable plant.

Conclusion
In this study, specific SSR markers were developed based on the transcriptome sequencing data of *S. chinensis*, and the frequency and distribution of SSR markers were analyzed. Using these markers, we successfully analyzed the genetic diversity and identified different accessions of *S. chinensis*. The results demonstrated that transcriptome sequencing is an effective method to identify molecular markers. Our work may lay a foundation for genetic diversity, genetic mapping, and marker-assisted selection in *S. chinensis*. It may facilitate *S. chinensis* breeding, as well as studies with other Schisandra plants with economic and medicinal value.

**Materials And Methods**

**Sample collection and total RNA extraction**

The plant materials were obtained from germplasm resource garden of Jilin Agriculture University (125°24′15″E, 43°48′5″N; Changchun Jilin Province, China) in July, 2019. These accessions originated from the wild and the information on 42 accessions is shown in Table 1, and there are no permissions were necessary to obtain branches of *Schisandra chinensis*. ‘Yanhong’ and ‘Jinwuwei’ were examined and approved by Jilin Provincial Variety Examination and Approval Committee in 2012 and 2016, respectively. Young leaves (0.5–1.0 g) were collected in July 2019 and ground into powder after freeze drying in liquid nitrogen. Total RNA was extracted using a modified version of the cetyltrimethylammonium bromide (CTAB) method [41]. RNA was visualized by 1.5% agarose gel electrophoresis.

**Transcriptome analysis**

Transcriptome data were obtained in 2018 by Illumina (San Diego, CA, USA) high-throughput deep sequencing. RNA was extracted with the CTAB method from six to eight *S. chinensis* seedlings at the true leaf stage and then reverse-transcribed into cDNA, which was sent to Biomarker Technologies (Beijing, China) for transcriptome sequencing. A total of 59,786 unigenes were assembled with the Trinity method [42] and used as background data for analysis.

**SSR site identification and primer design based on transcriptome data**

SSR loci of unigenes were analyzed using MISA software (https://omictools.com/misa-tool); the minimum number of mono-, di-, tri-, tetra-, penta-, and hexanucleotide repeats was 10, 6, 5, 5, 5, and
5, respectively. Primer 3 (http://bioinfo.ut.ee/primer3-0.4.0/) was used to design primers with the following features: (1) length between 18 and 23 bp; (2) annealing temperature of 55°C–65°C, with a difference of less than 2°C between forward and reverse primers; (3) amplification product size between 80 and 300 bp; and (4) GC content of 40%–60%. In total, 50 SSR primers were randomly selected and synthesized by Shenggong Biological Engineering (Shanghai, China).

**PCR amplification and data analysis**

The 50 EST-SSR primers were tested to identify those that were stable and could detect polymorphisms. The PCR reaction, which had a total volume of 16 µl, contained 8 µl of 2× Ex Taq Master Mix, 0.8 µl of each of primer, 5.4 µl ultrapure water, and about 20 ng DNA template. PCR amplification was carried out as previously described [10]. The products were separated on a 5% polyacrylamide gel; bands were visualized by silver staining. A clear band was assigned a value of 1 and a weak or no band at the same position was given a value of 0.

**Declarations**

**Abbreviations**

EST-SSR: Expressed Sequence Tag-Simple Sequence Repeat; ISSRs: Inter Simple Sequence Repeats; AFLPs: Amplified Fragment Length Polymorphisms; RAPD: Randomly Amplified Polymorphic DNA; CTAB: Cetyltrimethylammonium bromide; MCID: Manual Cultivar Identification Diagram.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

The raw data of RNA-seq are deposited in Sequence Read Archives Database (http://www.ncbi.nlm.nih.gov/bioproject/609148) under accession number PRJNA609148. Other dataset supporting the conclusions of this article are included within the article (and its additional file).

**Competing interests**
The authors declare that they have no conflict of interest.

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**Authors’ contributions**

SGL and AJ conceived and designed the research. SGL performed the research and wrote this manuscript. WZX, SD, ZSS and GJH analyzed the data. All authors read and approved the manuscript.

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Not Applicable

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Tables
Due to technical limitations, Tables 1 - 3 are only available for download from the Supplementary Files section.

Figures
Figure 1

Frequency distribution of EST-SSRs identified from S. chinensis ESTs, according to (a) motif type and (b) number of repeats.
Figure 2

Dendrogram of S. chinensis germplasms generated using the unweighted pair group method with arithmetic mean.
Figure 3

Manual cultivar identification diagram for S. chinensis accessions based on 10 pairs of EST-SSR primers. Numbers next to each horizontal line in the diagram indicate the size (in bp) of polymorphic bands used to differentiate the cultivars; (+) and (−) indicate the presence and absence of a polymorphic band, respectively.
Supplementary Files

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