

# Characterization and Development of EST-Derived SSR Markers in Sinowilsonia henryi (Hamamelidaceae)

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PRIMER NOTE

## CHARACTERIZATION AND DEVELOPMENT OF EST-DERIVED SSR MARKERS IN SINOWILSONIA HENRYI (HAMAMELIDACEAE)<sup>1</sup>

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- Premise of the study: Polymorphic microsatellite markers were developed to reveal the genetic diversity of extant populations
  and the mating system of Sinowilsonia henryi (Hamamelidaceae).
- *Methods and Results:* In this study, nuclear simple sequence repeat (SSR) markers were developed using the Illumina high-throughput sequencing technique (RNA-Seq). The de novo—assembled transcriptome generated a total of 64,694 unique sequences with an average length of 601 bp. A total of 2941 microsatellite loci were detected. Of the 121 tested loci, 13 loci were polymorphic and eight were monomorphic among 72 individuals representing three natural populations of the species. The number of alleles per locus ranged from one to four, and the observed and expected heterozygosity at population level were 0.00–1.00 and 0.10–0.66, respectively.
- Conclusions: The developed expressed sequence tag (EST)–SSRs will be useful for studying genetic diversity of S. henryi as well as assessing the mating system among Sinowilsonia species.

**Key words:** Hamamelidaceae; microsatellite; RNA-Seq; *Sinowilsonia henryi*.

The tree genus *Sinowilsonia* Hemsl. is a member of the Hamamelidaceae family and comprises only one species, *S. henryi* Hemsl. This species is narrowly distributed in the mountains of central China at an elevation of 600–1400 m (Zhang et al., 2003). Currently, the natural habitats of this species are severely deteriorated and fragmented, with population sizes ranging from as few as five individuals to approximately 50 flowering plants (Zhou et al., 2014). Thus, *S. henryi* has been listed as an endangered plant species in the China Plant Red Data Book (Fu and Jin, 1992).

Knowledge of genetic diversity and genetic structure of extant populations is essential to the formulation of effective conservation and management strategies for threatened species (Frankham et al., 2002). Due to their codominance, hypervariability, and reliable scorability, microsatellite markers have been widely used in population genetic studies (Selkoe and Toonen, 2006). However, microsatellite markers for *S. henryi* are currently not available. High-throughput RNA sequencing (RNA-Seq) is one of the most useful next-generation sequencing techniques for identifying microsatellites. In the current study, we developed and characterized 21 expressed sequence tag-simple sequence repeat (EST-SSR) markers for *S. henryi* using RNA-Seq.

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#### METHODS AND RESULTS

Total RNAs were isolated from young leaves using a cetyltrimethylammonium bromide (CTAB) procedure (Chang et al., 1993). The poly(A)+ RNA (mRNA) was purified with the RNA Clean-up Kit (Invitrogen, Carlsbad, California, USA) according to the manufacturer's instructions. The purified RNA was subsequently fragmented into small pieces (200 bp) by the fragmentation buffer. Then, the cleaved RNA fragments were used for first-strand cDNA synthesis using reverse transcriptase (Invitrogen) with random hexamer primers. Subsequently, second-strand cDNA was synthesized using RNase H and DNA polymerase I (Tiangen, Beijing, China). Illumina paired-end sequencing adapters were then ligated to the ends of the 3'-adenylated cDNA fragments. The cDNA library was sequenced by Shanghai Haiyu Biotechnology Co. Ltd. on the Illumina HiSeq 2000 instrument (Illumina, San Diego, California, USA). Before assembly, raw reads were filtered to remove those containing adapter or lowquality reads (more than 20% of nucleotides with Q-value ≤ 10) and reads containing poly N (>10% ambiguous base calls). Transcriptome assembly was performed using the Trinity package (version 2013-02-25) with the default parameters (Grabherr et al., 2011).

A total of 28.7 million 300-bp, clean, paired-end reads were obtained. All clean reads are available from the National Center for Biotechnology Information (NCBI) Short Read Archive (SRA) database (Bioproject accession no. PRJNA394173). De novo assembly of clean reads resulted in 64,694 unique sequences with an average length of 601 bp and an N50 length of 999 bp. The MIcroSAtellite identification tool (MISA; Thiel et al., 2003) was used to screen for the presence of microsatellites. The parameters used to identify microsatellites were seven repeats for di-, five for tri- and tetra-, four for penta-, and three for hexanucleotide repeats. Subsequently, SSR primers were designed with minimum GC content of 40% and an expected product size ranging from 100 to 280 bp using Primer3 (Rozen and Skaletsky, 1999).

A total of 8892 SSRs containing repeats from di- to pentanucleotides were identified from 64,694 unique sequences. Dinucleotides were the most abundant repeat type (5232), followed by trinucleotides (2198), hexanucleotides (1035), pentanucleotides (259), and tetranucleotides (168). The dinucleotide repeat (AG/CT)<sub>n</sub> (3646) was followed by (AT/AT)<sub>n</sub> (1192), (AC/GT)<sub>n</sub> (384), and (CG/CG)<sub>n</sub> (11). Among the trinucleotide repeat motifs, the most frequent SSR motif was

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Table 1. Frequency of repeat motifs in nonredundant Sinowilsonia henryi ESTs.

SSR motifs	No. of repeats											
	3	4	5	6	7	8	9	10	11	12	>12	Total
AC/GT	_	_	_	_	106	86	71	64	54	3	0	384
AG/CT	_	_	_	_	717	802	1268	742	111	6	0	3646
AT/TA	_	_	_	_	283	275	325	248	59	1	1	1192
CG/GC	_	_	_	_	5	4	1	1	0	0	0	11
AAC/GTT	_	_	59	17	13	6	0	0	0	0	0	95
AAG/CTT	_	_	300	200	164	3	0	0	0	0	0	667
AAT/ATT	_	_	147	103	61	3	0	0	0	0	0	314
ACC/GGT	_	_	111	47	33	3	0	0	0	0	0	194
ACG/CGT	_	_	35	14	10	4	0	0	1	0	0	63
ACT/AGT	_	_	19	9	2	1	0	0	0	0	0	31
AGC/CTG	_	_	139	94	64	4	0	0	0	0	0	301
AGG/CCT	_	_	81	44	32	7	0	0	0	0	0	164
ATC/ATG	_	_	136	57	54	5	0	0	0	0	0	252
CCG/CGG	_	_	73	31	9	4	0	0	0	0	0	117
Tetra-	_	_	146	22	0	0	0	0	0	0	0	168
Penta-	_	252	7	0	0	0	0	0	0	0	0	259
Hexa-	844	191	0	0	0	0	0	0	0	0	0	1035

*Note*: — = number of repeats not calculated.

AAG/CTT (667), followed by AAT/ATT (314), AGC/CTG (301), and ATC/ATG (252) (Table 1). Of the 8892 identified SSRs, 2941 (33%) were suitable for designing locus-specific primers (Appendix S1).

SSR loci with a minimum of 10 repeats for dinucleotides and seven for trinucleotides were selected for amplification. A total of 121 primer pairs were selected and used for further characterization. Eight individuals of *S. henryi* from Wuhan Botanical Garden, China, were collected to initially assess microsatellite polymorphism. Genomic DNA was isolated using the CTAB method (Doyle and Doyle, 1987). PCR reactions were performed in a 10-μL reaction mixture (final volume) containing approximately 50 ng of genomic DNA, 0.2 μM each of forward and reverse primer, 10 mM Tris-HCl (pH 8.4), 50 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 1.5 mM MgCl<sub>2</sub>, 0.2 mM dNTPs, and 1 unit *Taq* polymerase (Fermentas, Vilnius, Lithuania). The PCR cycling program included 5 min of initial denaturation at 94°C; followed by 35 cycles of 50 s at 94°C, 50 s at 56–60°C depending on the primer pair (Table 2), and 1 min at 72°C; followed by a final 10-min extension step at 72°C. The PCR products were separated on a 6% polyacrylamide denaturing gel of high resolution with silver stain. A 25-bp marker ladder (Promega Corporation, Madison, Wisconsin, USA) was used to identify the alleles.

Of the 121 primer pairs tested, 21 successfully amplified the target fragments (Table 1); of these, 13 loci were polymorphic (SH01-SH13), while eight were detected as monomorphic (SH14-SH21; Table 2). The level of genetic variability was estimated by genotyping 72 individuals of S. henryi from three wild populations (Appendix 1). For each locus, the number of alleles (A), observed heterozygosity  $(H_o)$ , and expected heterozygosity  $(H_e)$  were estimated using the program GENEPOP version 3.4 (Raymond and Rousset, 1995). Null alleles were detected at three loci (SH03, SH04, and SH07) using the program CERVUS 2.0 (Marshall et al., 1998). In the SNJ population, A ranged from one to three,  $H_e$  ranged from 0 to 0.60, and  $H_o$  ranged from 0 to 1.00. In the FS population, A ranged from one to three,  $H_e$  ranged from 0 to 0.66, and  $H_o$  ranged from 0 to 0.80. In the WD population, A ranged from one to four,  $H_e$  ranged from 0 to 0.63, and  $H_0$  ranged from 0 to 0.63. Three loci deviated from Hardy-Weinberg equilibrium after correction for multiple tests (Table 3). The observed departures from Hardy-Weinberg equilibrium may be due to null alleles. Significant linkage disequilibrium was observed in 10 pairs of loci before correction for multiple tests (P < 0.05). However, no loci were observed to be in linkage disequilibrium after correction for multiple tests (P < 0.0006). The sequences containing microsatellites were BLASTed against the NCBI nonredundant protein database using BLASTX with a threshold of E-value < 2.00E-5. Ten loci showed significant similarities to known proteins in the NCBI nonredundant protein database (Table 2).

### **CONCLUSIONS**

In the current study, a total of 2941 primer pairs were successfully designed based on transcriptome sequences. In total, 121

PCR primers of SSR loci were used for validation of amplification and polymorphism; of these, 13 revealed microsatellite polymorphism. To the best of our knowledge, this is the first study to develop microsatellites for *S. henryi*. These EST-derived SSRs could provide valuable tools for studying genetic diversity and assessing the mating system among *Sinowilsonia* species. In addition, because EST-derived SSRs may be associated with functional genes, the remaining untested 2820 SSRs and 21 loci developed in the current study may be useful for examining adaptive variation using genome scan methods.

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Table 2. Characterization of 21 EST-SSR primers developed in Sinowilsonia henryi.

Locus		Primer sequences (5'–3')	Repeat motif	Allele size range (bp)	T <sub>a</sub> (°C)	BLASTX top hit description	E-value	GenBank accession no.
SH01		TTTGACCCCGAAACAACAGC TGATACCGCTCAAGTCTCCC	$(CAT)_7$	209–213	58	_		MF503975
SH02	F:	CATCACCTTCTGCTGGAACG ACCCCGGAGCATATATCAGC	$(TC)_{10}$	223–231	58	Hypothetical protein EUGRSUZ_G03166	3E-40	MF503976
SH03	F:	CCACCTCGTTCTTCTCGTCT	$(ATC)_7$	210–213	58	Phospho- <i>N</i> -acetylmuramoyl- pentapeptide-transferase	2E-176	MF503977
SH04	F:	CCTGACGGTAAAGAGAAACGC GAGTCGGAGTCCCATTTTGC	$(TTC)_7$	258–275	60	NUDIX domain-containing	3E-165	MF503978
SH05	F:	GTCTTCGAACATGAGGCGTC TATGCTAGTGGTGGTGCTGT	(GCA) <sub>7</sub>	195–202	58	protein —		MF503979
SH06	F:	TAGTCGTCGGGCTCATCATC ATTGAAGGCGTTTGGATCCG	(GCC) <sub>7</sub>	148–158	58	_		MF503980
SH07	R: F:	TGGCTTCCCTCTCGTCTTTT TGACATGGAGGGTAGTGTGG	$(ATG)_7$	183–186	58	_		MF503981
SH08		TCACCTCTTCCATTGCCTTCT GAAGCTGGAGTTTGTGACGG	(GTT) <sub>8</sub>	214–225	58	_		MF503982
SH09		CTTCGGGGCCTATAGTTGGT GGGGTGTTGTCCATTGATACAG	(ACC) <sub>7</sub>	232–240	58	CBL-interacting protein	0	MF503983
SH10		CCAGCAGTTGAAGTTCAGGAG AACCAAATCAGGCTCGCTTT	(AGC) <sub>7</sub>	225–239	59	kinase 07 Pre-mRNA-splicing factor	0	MF503984
SH11		CCGCTGCCAGATGAAATTGA GGATTGCCATCATGCTGTTG	(TC) <sub>10</sub>	209–215	58	SYF1 Transmembrane protein 230	1E-61	MF503985
SH12		AGCAAATTTGGCCACTGGAG GGCATCCACAGTGTTGCTAG	(TC) <sub>10</sub>	154–156	58	_		MF503986
SH13	R:	ACTTCTGGGGCCATTTCCTT AAGGACGAGGATGAATGGGG	(GCG) <sub>7</sub>	265–268	56	_		MF503987
SH14	R:	CCCAATTCCCCTCGAGAAGT TCACCATCATCACCACCTTCA	(TTG) <sub>7</sub>	175	56	ABC transporter G family	0	MF510515
	R:	AGGCTCATGGGTTTACAGCT				member 5-like	U	
SH15	R:	AGCAAGAGGACCAACACTCT TGCTGCTTTTACTTCCCCTC	(AAG) <sub>7</sub>	200	58	_		MF510516
SH16		CCAAGAGACCCCACCAACTA AGACGTTGCCTCAGTCTTGT	(GCT) <sub>7</sub>	256	56	_		MF510517
SH17		TGGCTTCCAACCTCCTCAAA GGTGGGGTGGAAGAAGAGAG	$(ACA)_8$	250	56	_		MF510518
SH18		ACCCGCGATCATACTGACAA GGTCCGTCATCACTTCTCCT	$(CTG)_7$	165	56	DExH-box ATP-dependent RNA helicase	3E-35	MF510519
SH19		GAGCAAACCCACAATCCAGA GCTGCCATGGTGAAGAAACA	$(GAG)_7$	200	58	_		MF510520
SH20	F:	GGGTGGGGAGAATAGGGAAG AGAGGGAGAGAGGGTCACAA	$(CT)_{10}$	200	56	NADP-dependent malic enzyme	0	MF510521
SH21	F:	CCATATCCGCCGCCAATAAG GCTCAATTTGCTACCTTTGAAG	(GGT) <sub>7</sub>	275	58	Receptor-like protein 1, putative isoform 2	2E-35	MF510522

*Note*:  $T_a$  = annealing temperature.

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Table 3. Genetic diversity of 13 SSR loci in three populations of Sinowilsonia henryi.<sup>a</sup>

Locus	SNJ ( $N = 15$ )				FS $(N = 25)$	)	WD $(N = 32)$		
	A	$H_{\mathrm{e}}$	$H_{\rm o}$	A	$H_{\mathrm{e}}$	$H_{\rm o}$	A	$H_{\mathrm{e}}$	$H_{\rm o}$
SH01	2	0.18	0.13	2	0.30	0.28	3	0.47	0.41
SH02	3	0.56	0.88	3	0.52	0.24	3	0.59	0.63
SH03	2	0.50	0.50	1	0.00	0.00***	1	0.00	0.00***
SH04	2	0.31	0.38	3	0.63	0.32	3	0.48	0.38
SH05	3	0.60	1.00	3	0.25	0.28	2	0.42	0.41
SH06	3	0.51	0.75	3	0.37	0.32	3	0.41	0.38
SH07	1	0.00	0.00	2	0.42	0.20	1	0.00	0.00***
SH08	2	0.22	0.25	3	0.62	0.72	2	0.48	0.38
SH09	1	0.00	0.00***	2	0.08	0.08	1	0.00	0.00***
SH10	2	0.38	0.25	3	0.63	0.56	3	0.63	0.59
SH11	3	0.53	0.75	3	0.66	0.80	4	0.36	0.38
SH12	2	0.50	0.75	2	0.34	0.36	2	0.49	0.47
SH13	2	0.22	0.25	2	0.39	0.36	2	0.44	0.34
Average	2.15	0.35	0.45	2.46	0.40	0.35	2.31	0.37	0.44

Note: A = number of alleles;  $H_e =$  expected heterozygosity;  $H_o =$  observed heterozygosity; N = number of individuals sampled.

APPENDIX 1. List of vouchers of Sinowilsonia henryi used in this study.

Population code	N	Location	Voucher no.a	Geographic coordinates	Altitude (m)
SNJ	15	Shennongjia Mountain, Hubei Province	Q. G. Ye 1102	31°30′09″N, 110°24′03″E	1405
FS	25	Yangchashan, Fang County, Hubei Province	Q. G. Ye 1108	31°53′01″N, 110°27′53″E	1201
WD	32	Wudang Mountain, Hubei Province	Q. G. Ye 1109	32°40′59″N, 111°01′01″E	1035

*Note*: N = number of individuals sampled.

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<sup>&</sup>lt;sup>a</sup>Locality and voucher information are provided in Appendix 1.

<sup>\*\*\*</sup> Denotes significant departure from Hardy–Weinberg equilibrium after Bonferroni correction (P < 0.0006).

<sup>&</sup>lt;sup>a</sup>All vouchers are deposited at the Wuhan Botanical Garden Herbarium (HIB), Wuhan, Hubei Province, China.