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PRIMER NOTE

Development of EST-SSR markers for *Taxillus nigrans* (Loranthaceae) in southwestern China using next-generation sequencing¹

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- Premise of the study: We developed transcriptome microsatellite markers (simple sequence repeats) for Taxillus nigrans (Loranthaceae) to survey the genetic diversity and population structure of this species.
- Methods and Results: We used Illumina HiSeq data to reconstruct the transcriptome of *T. nigrans* by de novo assembly and used the transcriptome to develop a set of simple sequence repeat markers. Overall, 40 primer pairs were designed and tested; 19 of them amplified successfully and demonstrated polymorphisms. Two loci that detected null alleles were eliminated, and the remaining 17, which were subjected to further analyses, yielded two to 21 alleles per locus.
- *Conclusions:* The markers will serve as a basis for studies to assess the extent and pattern of distribution of genetic variation in *T. nigrans*, and they may also be useful in conservation genetic, ecological, and evolutionary studies of the genus *Taxillus*, a group of plant species of importance in Chinese traditional medicine.

Key words: Chinese traditional medicine; conservation; Loranthaceae; microsatellite marker; next-generation sequencing; *Taxillus nigrans*; transcriptome.

Taxillus nigrans (Hance) Danser (Loranthaceae) is a mistletoe species that is found attached to many canopy tree species in low mountains, hills, and river basins in subtropical areas of southwestern China at elevations of 300–1300 m. Flowering can occur throughout the year, and the fruiting period is mainly in November. The entire plant of this species can be used as raw material for Chinese traditional medicine (Jiang, 1998). However, because the range of the species has undergone rapid expansion mediated by birds in the urban area of Chengdu (Sichuan Province, China), it forms large groves on garden tree species and is sometimes harmful to its host trees, so that individuals of this species are often removed by gardeners. To date, apart from some basic taxonomic data on the species (Gong et al., 2004) and genome studies on other species of Taxillus Tiegh. (Rist et al., 2011; Wei et al., 2017), nearly all published research has focused on aspects relating to its medicinal value, for example, the extraction and identification of medicinal components and the optimization of extraction methods (Li et al., 2006, 2009; Zhang et al., 2016; Zhao et al., 2016). There is little information on the genetic diversity and population structure of the species. We are also interested in developing genetic approaches for identification of individuals and assignment testing, which will help in

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understanding how this species expands its distribution and jumps from host to host in urban areas as well as in the field.

Simple sequence repeat (SSR) markers, also known as microsatellites or short tandem repeats, are highly polymorphic and are therefore useful as molecular markers in population genetic studies (Zhang et al., 2012; Jiang et al., 2015). Transcriptome sequencing has proven to be a powerful and cost-effective tool that has greatly accelerated the process of discovering molecular markers, including single nucleotide polymorphisms (SNPs) and SSRs (Ashrafi et al., 2012; Qi et al., 2016). In this study, we sequenced and assembled the transcriptome of *T. nigrans* and developed a set of expressed sequence tag (EST)–SSR markers for population genetic studies of *T. nigrans*. We also tested the transferability of these markers in herbarium samples of *T. delavayi* (Tiegh.) Danser and five individuals of *Scurrula parasitica* L. (collected from the field), another Loranthaceae parasite that co-occurs with *T. nigrans*.

METHODS AND RESULTS

Approximately 10 μ g (400 ng/ μ L) of total RNA was extracted from fresh leaf material of one individual of *T. nigrans* using TRIzol Reagent (Invitrogen, Carlsbad, California, USA). Subsequently, mRNA was isolated using magnetic oligo (dT) beads (Illumina, San Diego, California, USA); it was then fragmented into short fragments using the Ambion RNA Fragmentation Kit (Ambion, Austin, Texas, USA) according to the manufacturer's protocols. First-strand cDNA synthesis was performed using reverse transcriptase (Invitrogen) with random primers, and second-strand cDNA was synthesized by RNase H and DNA Polymerase I (Invitrogen). Finally, the transcriptome was sequenced on an Illumina HiSeq 2000 system at Novogene (Beijing, China). Prior to the assembly, a stringent filtering process of raw sequencing reads was conducted. The number of

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Locus	Primer sequences $(5'-3')$	Rep	Repeat motif	Allele size (bp)	$T_{\rm a}$ (°C) ^a	Fluorescent dye	Genbank accession no.	r	Protein ^b	$Organism^c$	E-valı
#TR7149	F: GGCAAAATCAACCGAGAAGA		(CT) ₁₂	164	60	6-FAM	KY412965	0.0156	NEN1-like	Populus euphratica	3×10
			Ć		09 (0
#1K11504	F: CCTCAGGAACCTGCAAGAAG R: CGACACAGGACAGCTGTAA		(AAU) ₁₆	C17	00 00	НЕХ	KY412900	0700.0	w D repeat-containing protein RUP2	Elaets guineensis	8×10
TR24412			(CT) ₂₁	122	09	6-FAM	KY412967	0.0747	Predicted gene, 39330	Oryza sativa	4×10
	R: AAGCGAACTCGAATCACTGC				60						
TR47466	F: AGTCCTTCGTTCCCGATACC		$(AT)_{24}$	231	09	TAMRA	KY421968	0.3990	Unknown	Vigna angularis	2×10-
#TR51334			$(AG)_{26}$	206	9 9 (TAMRA	KY421969	0.0008	Transmembrane protein,	Medicago truncatula	1×10^{-1}
TR56117	R: CTGCACTCTTCCATACGGCT F: TCTTTCCATTCAGCGACTC		TC)	166	09 09	TAMRA	KV421970	0 1183	putative	Zizinkus ininka	2~10
1110/11			1015	001	61	CONTRACT	0/017171	C011.0	FOCIO1411000	nontal condition	01/7
TR59209			(TC) ₁₅	157	09	6-FAM	KY421971	0.1826	LOC107268204	Cephus cinctus	2×10
#TR83979	K: AGGAATCGAACAGGAGGGTC F: CCTCCGTCTCTCTCTCTCTCTC		(CT),,	2.45	00	HEX	KY421972	0.0748	At3602290	Orvza sativa	2×10
			77/	2	09						i I
TR85804			(AG) ₃₃	219	09	HEX	KY421973	0.0705	CARUB_v10002273mg	Capsella rubella	1×10^{-1}
TR87965	K: CCTTGCTAATTCCACCACCA F: TGGAGATCTTGGCTTCGTTC		(AG) ₁₄	216	09	6-FAM	KY421974	0.1080	DDB1- and CUL4-	Theobroma cacao	2×10
			+		09				associated factor 13		
#TR88317			$(TAT)_{15}$	129	09	6-FAM	KY421975	0.2512	Restricted Tev	Nicotiana	1×10
	R: TTGCAGGAACAGGTATGGCT				60				Movement 1-like	tomento siform is	
TR90181			(TA) ₂₅	217	59	HEX	KY421976	-0.0708	LOC107270001	Cephus cinctus	1×10
					59						
#TR91417	F: AGAGGAATTGGCATCGTCAG R: TCCAACTCACACTTGCCTCA		$(GA)_{26}$	213	09 09	6-FAM	KY421977	0.1064	LOC105638199	Jatropha curcas	2×10
#TR97121			(AG) ₁₅	204	09	HEX	KY421978	0.0685	Transcription factor	Vigna angularis	3×10-
			Ē		99 (00000	bHLH35		
I K98083	F: IGGCTACCCTOTOTICTCCC		$(C1)_{15}$	CC7	00	HEX	K Y 4 2 I 9 / 9	0.0839	LUC10439/400	Nelumbo nucifera	01X7
TR105177			(GA)20	217	909 09	TAMRA	KY421980	0.0798	LOC103319601	Prunus mume	1×10^{-1}
			67 /		09						
TR120023	F: CTTGATCTTCTGGTGCGGTT		$(GA)_{14}$	161	09	TAMRA	KY421981	0.1191	LOC104727032	Camelina sativa	4×10
:					09						
*#TR85478	F: GTCGTCATGGACTCTTCGCT		(TC) ₅	228	88	TAMRA	KY421982	QN	EUTSA_v10007584mg	Eutrema salsugineum	5×10
	••		Ő	to	00 (, ,
*1K8/192	F: CCTTTGGAGGGGTTCAACTT D: mcccccssscmmmscmcsscc		(GCG)4	271	00	HEX	KY421983	0.4202	LOC1039865/6	Musa acuminata	0.11
	JIN II VOODOT V		1.		8						

 0^{-12} 0^{-15} 0^{-22}

0-13

 0^{-21}

 0^{-25}

6-0 0-4 Ξ

6-0

Characteristics of 19 polymorphic microsatellite loci developed for Taxillus nigrans.

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TABLE 1.

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6-0

0-8

 0^{-21}

0-8 0-8 0^{-15} 0^{-26}

 0^{-21}

4-0

lue^d

Note: ND = not done; r = null allele frequency; T_a = annealing temperature, ^aThe annealing temperature for each primer is listed, and the final annealing temperature for each PCR reaction is given as the average annealing temperature of the adopted primer pair. ^bInformation from BLAST analysis on the protein most closely matching the EST. ^cOrganism from which the BLAST match was obtained. ^d E-value associated with the BLAST match.

* Null alleles (r > 0.4).

Primers successfully amplified for Taxillus delavayi.

TABLE 2. Genetic properties of 17 newly developed polymorphic microsatellite loci in three populations of *Taxillus nigrans*. Loci exhibiting null alleles are not included.^a

	Sic	huan University (n	= 100)		Tazishan ($n = 30$))		Huanhuaxi ($n = 3$	30)
Locus	A	$H_{\rm o}$	H _e	A	$H_{\rm o}$	H _e	A	$H_{\rm o}$	$H_{\rm e}$
TR7149	7	0.717	0.815	5	0.900	0.728	10	0.967	0.844
TR11564	5	0.667	0.781	4	0.767	0.672	5	0.667	0.727
TR24412	6	0.551	0.628	4	0.633	0.691	7	0.621	0.722
TR47466	6	0.333	0.453	2	0.034	0.034	4	0.367	0.476
TR51334	11	0.525	0.776	2	0.966	0.499	5	0.967	0.577
TR56117	9	0.583	0.745	7	0.724	0.737	8	0.567	0.787
TR59209	10	0.626	0.789	6	0.310	0.596	6	0.643	0.786
TR83979	17	0.737	0.859	8	0.931	0.829	10	0.828	0.757
TR85804	18	0.808	0.876	9	0.633	0.799	17	0.833	0.898
TR87965	7	0.646	0.786	5	0.586	0.703	6	0.667	0.764
TR88317	11	0.347	0.714	5	0.607	0.702	4	0.517	0.644
TR90181	14	1.000	0.786	7	0.963	0.747	5	1.000	0.621
TR91417	10	0.717	0.809	6	0.400	0.665	7	0.700	0.749
TR97121	2	0.380	0.476	2	0.500	0.408	2	0.400	0.464
TR98683	14	0.690	0.860	10	0.833	0.815	15	0.933	0.813
TR105177	20	0.764	0.893	8	0.733	0.807	10	0.931	0.835
TR120023	21	0.802	0.885	6	0.633	0.776	6	0.552	0.797

Note: A = number of alleles sampled; H_e = expected heterozygosity; H_o = observed heterozygosity; n = number of individuals sampled. ^aVoucher and locality information are provided in Appendix 1.

low-quality (Q \leq 3) bases in a single read was restricted to less than 50%, and paired reads were discarded if the number of unknown nucleotide bases in either of the paired reads exceeded 3% following the sequencing company's protocol (Novogene). After removing the adapter sequences and ambiguous reads, the clean reads obtained were de novo assembled using Trinity (release 2013-02-25; Grabherr et al., 2011) with default settings. The final assembly was composed of 299,147 unigenes and had an N50 size of 1056 bp. Raw transcriptome read data were deposited in the National Center for Biotechnology Information (NCBI) Short Read Archive (accession no. SRP105083).

SSRs were detected using the Perl script MISA (Thiel et al., 2003) with a motif size of one to six nucleotides and thresholds of eight, four, four, three, three, and three repeat units for mono-, di-, tri-, tetra-, penta-, and hexanucleotide SSRs, respectively. We selected 83,954 microsatellite loci and used the primer design software package Primer3 version 2.3.6 (Untergasser et al., 2012) to design primer sets. Following random browsing across the output files of these primer sets, 40 markers were selected based on length (19–20 bp), GC

TABLE 3. Fragment sizes detected in cross-amplification tests of the 19 newly developed microsatellite markers in *Taxillus delavayi* and *Scurrula parasitica*.^a

Locus	Taxillus delavayi $(n = 2)$	Scurrula parasitica $(n = 5)$
TR7149	167	152-163
TR11564	192	193
TR24412	_	124
TR47466		272
TR51334	182	174–182
TR56117	_	155–185
TR59209	_	125–143
TR83979	244	177–211
TR85804	_	179–255
TR87965	_	197
TR88317	100	100-130
TR90181	_	205-207
TR91417	196	196–204
TR97121	332	353
TR98683	_	244-260
TR105177	_	189
TR120023	_	152
TR85478	229	229
TR87192	_	269

Note: — = amplification failed; n = number of individuals sampled. ^aVoucher and locality information are provided in Appendix 1.

content (40–65%) of the primers, and annealing temperatures (59–61°C) of the primer sets. Nineteen of the 40 tested markers were selected based on PCR success rate and degree of polymorphism (difference in band length), and these were used to genotype individual *Taxillus* plants (Table 1).

Genomic DNA was extracted from the silica-dried leaves of 160 individuals from three populations of T. nigrans, two individuals of T. delavayi, and five individuals of S. parasitica (Appendix 1) using the cetyltrimethylammonium bromide (CTAB) method of Doyle and Doyle (1987). PCR reactions were performed in 25-µL volumes containing 12.5 µL 2× PCR buffer, 300.0 µM each dNTP, 0.3 µM each primer, 1.25 unit Taq DNA polymerase (Vazyme Biotech, Nanjing, China), and ca. 50 ng of genomic DNA. The cycling conditions were as follows: 95°C for 5 min, followed by 35 cycles of 95°C for 45 s, 55°C for 30 s, and 72°C for 45 s; the reactions were completed by a final elongation step at 72°C for 10 min. The PCR products were checked on 1% agarose gels to confirm PCR success and then sent to TsingKe (Chengdu, China) for microsatellite genotyping. Primer pairs were synthesized with the forward primer of each pair 5' end-labeled with either 6-FAM, TAMRA, or HEX (Applied Biosystems, Foster City, California, USA), and amplicons were analyzed on an ABI PRISM 3100 genetic analyzer. The microsatellite genotype at each locus for each individual was determined using GeneMarker (SoftGenetics, State College, Pennsylvania, USA). Allele sizes at each locus were then scored and checked for possible genotyping errors, such as stuttering, large allele dropouts, or null alleles, using CERVUS (Dakin and Avise, 2004). In total, null alleles (null allele frequency [r] >0.4) were detected at two loci (Table 1). These loci were eliminated, and the remaining 17 microsatellite loci were subjected to further analyses (Table 2).

These 17 microsatellite loci were highly polymorphic, with two to 21 alleles per locus. We used GenAlEx version 6 (Peakall and Smouse, 2006) to calculate the number of alleles and the observed and expected heterozygosity at each locus (Table 2). When using GIMLET version 1.3.3 (Valière, 2002), a minimum of two loci and six loci are needed to estimate, respectively, the unbiased probability that a genotype is shared by two individuals (P_{ID}) in a population, and the probability that a genotype is shared by two siblings ($P_{ID(sib)}$).

In the cross-species transferability test, eight of the 19 loci were successfully genotyped in two individuals of *T. delavayi* taken from herbarium specimens (Table 3). In contrast, all polymorphic loci were successfully amplified in *S. parasitica* (Table 3). The difference in success between *T. delavayi* and *S. parasitica* may have been due to a higher proportion of degraded DNA from *T. delavayi* herbarium specimens.

CONCLUSIONS

We developed and amplified a set of polymorphic EST-SSR markers for *T. nigrans*. These new SSR markers will serve as a basis for studies assessing the genetic diversity and population structure of *T. nigrans*. Our research will be useful for conservation genetic, ecological, and evolutionary studies of the genus *Taxillus*, a group of plant species of importance in Chinese traditional medicine. We plan to use these markers to explain the rapid demographic expansion and host specificity of *T. nigrans* in urban areas in southwestern China.

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Species	Ν	Population code	Locality	Geographic coordinates	Voucher specimen accession no. ^a
Taxillus nigrans (Hance) Danser	100	SCU	Sichuan University, Sichuan	30°37′48″N, 104°4′48″E	SZ-00545040, SZ-00545041, SZ-00545042, SZ-00545043, SZ-00545044
T. nigrans	30	TZT	Tazishan, Sichuan	30°38'7"N, 104°7'15"E	SZ-00545045, SZ-00545046, SZ-00545047
T. nigrans	30	НН	Huanhuaxi, Sichuan	30°39′28″N, 104°1′55″E	SZ-00545049, SZ-00545049, SZ-00545050
T. delavayi (Tiegh.) Danser T. delavayi Scurrula parasitica L.	1 1 5	Individual Individual TZS	Maerkang, Sichuan Muli, Sichuan Tazishan, Sichuan	31°54′46″N, 102°11′24″E 27°55′55″N, 101°16′43″E 30°38′7″N, 104°7′15″E	SZ-00280020 SZ-00280006 SZ-00545051

APPENDIX 1. Voucher specimen information for Loranthaceae used in this study.

Note: *N* = number of individuals sampled.

^aAll voucher specimens are deposited at the herbarium of Sichuan University (SZ), Sichuan, China.