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PRIMER NOTE

CHLOROPLAST AND MITOCHONDRIAL MICROSATELLITES FOR *MILLETTIA PINNATA* (FABACEAE) AND CROSS-AMPLIFICATION IN RELATED SPECIES¹

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- *Premise of the study:* Chloroplast and mitochondrial microsatellites were identified to study the population genetics of *Millettia pinnata* (Fabaceae).
- *Methods and Results:* Based on publicly available plastid genome sequence data of *M. pinnata*, 42 primer pairs were developed, of which 17 displayed polymorphisms across 89 individuals from four populations. For chloroplast loci, two to six alleles were recovered and the unbiased haploid diversity per locus ranged from 0.391 to 0.857. For mitochondrial loci, two to four alleles were recovered and the unbiased haploid diversity ranged from 0.264 to 0.740. Sixteen of the 17 screened markers could be successfully amplified in the related species *M. pulchra*.
- *Conclusions:* The 17 microsatellite markers developed here exhibited variation in *M. pinnata* and 16 presented transferability in the related species *M. pulchra*, suggesting that these markers will be valuable for genetic studies across *M. pinnata* and its related species.

Key words: chloroplast microsatellite; cross-amplification; Fabaceae; *Millettia pinnata*; *Millettia pulchra*; mitochondrial microsatellite.

Millettia pinnata (L.) Panigrahi (syn. *Pongamia pinnata* (L.) Pierre; Fabaceae) is an arboreal legume of the subfamily Papilionoideae and, more specifically, of the tribe Millettieae. According to Scott et al. (2008), this species is widely distributed in the Indian subcontinent and Southeast Asia, extending to Polynesia and northern Australia. Historically, this plant has been used as a source of traditional medicine, animal fodder, green manure, timber, fish poison, and fuel in India and neighboring regions (Satyavati et al., 1987). Seeds of *M. pinnata* contain oils that are inedible but useful for biodiesel, and thus it has received increasing attention as a sustainable biofuel crop in the past decade.

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Population genetic studies on M. pinnata have used molecular biological methods including amplified fragment length polymorphisms (AFLPs), three endonuclease (TE)-AFLP (Sharma et al., 2011), inter-simple sequence repeats (ISSRs) (Sahoo et al., 2010; Sharma et al., 2014), and the chloroplast trnK/matK and the internal transcribed spacer (ITS) region of nuclear ribosomal DNA (Hu et al., 2000, 2002; Arpiwi et al., 2013). Huang et al. (2016) developed nuclear simple sequence repeat (SSR) markers for M. pinnata; however, there is still a lack of plastid (chloroplast and mitochondrial) SSR markers capable of detecting high levels of polymorphism in this species. The uniparentally inherited characteristics of chloroplast and mitochondria can supply information on phylogenetic relationships between individuals because their lineages are not disturbed by recombination (Soranzo et al., 1999). In general, for the maternally inherited chloroplast, the higher sequence variations of SSR loci are distributed throughout the noncoding regions and the flanking regions are conserved (Powell et al., 1995), which makes it possible to monitor the population structure affected by pollen flow and seed-mediated gene flow (Provan et al., 2001). The search for mitochondrial SSR (mtSSR) loci might also be informative, although preliminary studies on plant mitochondrial microsatellites have shown little intraspecific variability (Soranzo et al., 1999). Therefore, plastid SSR markers can be effective for analyzing genetic diversity, population structure, paternity inheritance, and germplasm resource identification (Provan et al., 2001).

Applications in Plant Sciences 2017 5(5): 1700034; http://www.bioone.org/loi/apps © 2017 Wang et al. Published by the Botanical Society of America. This is an open access article distributed under the terms of the Creative Commons Attribution License (CC-BY-NC-SA 4.0), which permits unrestricted noncommercial use and redistribution provided that the original author and source are credited and the new work is distributed under the same license as the original. In this study, we developed a set of novel SSRs based on publicly available chloroplast and mitochondrial genome sequence data of *M. pinnata* to assess the genetic variation and population genetic structure of this species. Furthermore, we tested the transferability of these markers in the related species *M. pulchra* (Benth.) Kurz.

METHODS AND RESULTS

In this study, the complete chloroplast and mitochondrial genome sequence data of *M. pinnata* were downloaded from the National Center for Biotechnology Information (NCBI) GenBank database (GenBank accession no. JN673818.2 and JN872550.1, respectively). The SSR loci distributed throughout the *M. pinnata* chloroplast and mitochondrial genomes were screened using MISA software (Thiel et al., 2003). The SSR motifs contained one to five nucleotides, five for dinucleotides, four for trinucleotides, and three for tetranucleotides and pentanucleotides. A total of 97 repeat motifs were identified in the chloroplast and mitochondrial genomes, among which the most frequent types were mononucleotides (69 [71.1%]) and dinucleotides (17 [17.5%]), while tri- (5 [5.1%]) and tetranucleotide (6 [6.2%]) motifs were rare. Forty-two loci were selected at random to design primers using Primer3 (Rozen and Skaletsky, 1999), with the optimum conditions set at length of 22 bp (20–26 bp), temperature of 55–60°C, and product size range of 100–500 bp.

Eighty-nine individuals of M. pinnata from four natural populations (Appendix 1) were used to evaluate polymorphism of the target microsatellite loci. Genomic DNA from silica-dried leaves was isolated using the advanced cetvltrimethylammonium bromide (CTAB) method (Doyle, 1991). PCR amplifications were performed in a final volume of 15 µL, containing 30 ng of genomic DNA, 1× PCR buffer (10 mM Tris-HCl [pH 8.4] and 1.5 mM MgCl₂; TransGen Biotech Co., Beijing, China), 0.2 mM dNTPs (Bocai Biotech Co., Shanghai, China), 0.5 µM of each primer (BGI Sequencing Co., Beijing, China), and 0.5 units EasyTaq DNA polymerase (TransGen Biotech Co.). PCR reactions were conducted in a Bio-Rad PTC-200 thermal cycler (Bio-Rad Laboratories, Hercules, California, USA) under the following conditions: initial denaturation at 94°C for 4 min; followed by 35 cycles of 94°C for 1 min, 45 s at the specific annealing temperature for each primer pair (Table 1), and 72°C for 60 s; and a final extension of 10 min at 72°C. PCR products were detected using 1.0% agarose gel electrophoresis to test the utility of the primers. Finally, among the 42 selected primer pairs, 40 were successfully amplified but products from only 17 primer pairs exhibited clear SSR polymorphisms. Six individuals from the related species M. pulchra were used to evaluate the transferability of these polymorphic markers applying the 17 screened primers. With the Quant-iT PicoGreen dsDNA Reagent and Kit (including the 35-400-bp Range DNA Ladder; Invitrogen, Carlsbad, California, USA), the Fragment Analyzer Automated CE System (Advanced Analytical Technologies [AATI], Ames, Iowa, USA) was applied to perform SSR genotyping. Raw data were exported, and the number of alleles and allele sizes per locus were called using PROSize software (version 2.0, AATI). All sequences of cpSSR and mtSSR loci were deposited in GenBank, and their accession numbers are presented in Table 1. Because it would be difficult to score the mononucleotide microsatellites consistently, we

TABLE 1. Characteristics of 17 novel microsatellite markers developed in Millettia pinnata.

Locus	Primer sequences $(5'-3')$	Repeat motif	$T_{\rm a}(^{\circ}{\rm C})$	ES (bp)	Position ^a	GenBank accession no
POSSRB38 ^b	F: TTAAGGAGGCCCCTAATGAAAT	$(T)_{10}(A)_{10}$	55	258	trnG-psal IGS	KY189098
	R: TTTTAGATACGGGCAGTAGGGA	()10 ()10			1	
POSSRB548 ^b	F: ATTAATCGGGGATACACGACAG	$(T)_{10}(A)_{13}$	55	191	ycf3 CDS	KY189099
	R: ATTCCGACAACTTCAGGAGAAA					
POSSR22 ^b	F: ATTGCAGGTTAACCCCCTTC	$(AT)_7$	55	190	rpl20 intron	KY189089
	R: AAAATCGGCGGAGAAGTTTT					
POSSR30 ^b	F: TCGTCGGTAAATTCAACGGT	$(AT)_7$	55	360	trnD intron	KY189092
	R: AGGGGATCCCTCTTGTTTTT					
POSSR44 ^b	F: GAATTTGTTTCTTCGTCTTTACAAAA	$(AT)_7$	55	173	trnH intron	KY189093
	R: GTGGATCAAGGCAGTGGATT					
POSSR81 ^b	F: GCTCCGTTCCATGTCTCATT	$(TA)_8$	55	275	<i>ycf1</i> intron	KY189096
	R: TGAAAAATCATCGCAAACCTC					
POSSRB253 ^b	F: AATTAGGCTCGATCAACTGGAA	(T) ₁₂	56	242	ycf2-ycf15 IGS	KY411953
	R: CTGTTCCTATCTAACGGAACGC					
POSSRB426 ^b	F: ATTCAGTATCCTGCCACGAAAT	$(A)_{10}$	56	175	ycf2-ycf15 IGS	KY411954
	R: CGACCTTGCATTTTTCTACCTC					
POSSRS6 ^b	F: CTGTTCCTATCTAACGGAACGC	(A) ₁₂	56	242	ycf2-ycf15 IGS	KY411955
	R: AATTAGGCTCGATCAACTGGAA					
POSSRS89 ^b	F: TTCGTGATTCCTTGGTAAATCC	$(A)_{10}$	56	203	atpF CDS	KY411956
	R: AGGCCATTTTATCGACATGAGT					
POSSRS217 ^b	F: GATTACGATAAATTGGATCGGC	$(T)_{11}$	56	168	rps7 intron	KY411957
	R: ATCTTTCGAGATCCACCCTACA					
POSSRS225 ^b	F: TTTCTTCCAACATAACAACCCC	(T) ₁₅	56	152	rps18 intron	KY411958
	R: AAAGCGAGTCGACCACTAGAAC					
POSSRB218 ^c	F: CGTTCCTTTTCTCTTGGACATC	(TATTA) ₅	55	168	atp9 intron	KY189090
	R: TGCATGGGTCTTCCTTTCTTAT					
POSSRB310 ^c	F: GAGTTTGAATTGACCCGGTTAG	$(CT)_6$	55	242	trnD-trnI IGS	KY189091
	R: CTAGGCAGCTATCGGAAAGAAA					
POSSR53 ^c	F: AGGCTGCTAGGAAGGAGGAC	$(TA)_6$	55	175	ccmC-rps3 IGS	KY189094
	R: GGCACCATTTCAGAGAGGAG					
POSSR61 ^c	F: CGGTGAAGTACCCCCTTACA	$(GA)_6$	55	242	nad1-atp1 IGS	KY189095
	R: GCTTCGGTGTAAGCACCAGT					
POSSR95 ^c	F: GTATGTACCCCGATTCGCAC	(CAAA) ₅	55	203	Rrn26-mttB IGS	KY189097
	R: GTCTAGGCTGATTTGGCAGG					

Note: A = number of alleles; CDS = coding sequence; ES = expected band size; IGS = intergenic space; T_a = annealing temperature.

^aPosition of each SSR is according to the NCBI sequence information of *M. pinnata* (GenBank accession no. JN673818.2 for the complete chloroplast genome and JN872550.1 for the complete mitochondrial genome).

^bLoci located in the chloroplast.

^cLoci located in the mitochondria.

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								Millettia pinnata	ı pinna	ta								Millettic	Millettia pulchra	
		POPH	POPH ($N = 26$)			POLS	OLS (N = 25)			POC	POCN (N = 14)			PON	PONS ($N = 24$)			MVSD	MVSD (N = 6)	
Locus	A	A_{e}	Ι	$h_{\rm unb}$	A	A_{e}	Ι	$h_{ m unb}$	A	$A_{ m e}$	Ι	$h_{\rm unb}$	A	$A_{ m e}$	Ι	$h_{ m unb}$	A	A_{e}	Ι	$h_{\rm unb}$
POSSRB38 ^b	4	3.159	1.209	0.711	ю	2.323	0.944	0.593	ю	2.882	1.079	0.703	4	3.032	1.190	0.699	ю	2.571	1.011	0.733
POSSRB548 ^b	5	4.225	1.487	0.794	9	4.496	1.614	0.810	С	2.579	1.004	0.659	5	3.945	1.452	0.779	б	2.000	0.868	0.600
POSSR22 ^b	4	2.840	1.185	0.674	С	1.947	0.779	0.507	0	1.690	0.598	0.440	0	1.600	0.562	0.391	0	1.800	0.637	0.533
POSSR30 ^b	4	3.314	1.266	0.726	б	2.462	0.974	0.620	0	1.960	0.683	0.527	б	1.646	0.675	0.409	0	2.000	0.693	0.600
POSSR44 ^b	б	2.828	1.068	0.673	0	1.771	0.627	0.453	0	1.690	0.598	0.440	б	2.743	1.051	0.663	б	2.000	0.868	0.600
POSSR81 ^b	4	2.704	1.170	0.655	4	3.307	1.290	0.727	0	1.849	0.652	0.495	4	2.969	1.200	0.692	0	1.800	0.637	0.533
POSSRB253 ^b	4	2.704	1.170	0.655	б	1.947	0.779	0.507	4	2.390	1.055	0.626	4	3.245	1.228	0.723	б	2.571	1.011	0.733
POSSRB426 ^b	2	3.347	1.345	0.729	4	2.376	1.085	0.603	4	2.882	1.195	0.703	9	5.434	1.738	0.851	0	1.800	0.637	0.533
POSSRS6 ^b	2	2.522	1.173	0.628	9	5.631	1.760	0.857	4	3.267	1.277	0.747	9	3.556	1.501	0.750	0	2.000	0.693	0.600
POSSRS89 ^b	4	3.414	1.307	0.735	б	2.828	1.068	0.673	4	3.379	1.272	0.758	4	2.717	1.179	0.659	б	2.571	1.011	0.733
POSSRS217 ^b	2	3.453	1.401	0.740	5	3.307	1.315	0.727	4	3.267	1.254	0.747	5	3.550	1.397	0.751	0	1.800	0.637	0.533
POSSRS225 ^b	4	3.073	1.220	0.702	5	3.342	1.356	0.730	S	3.500	1.390	0.769	5	3.789	1.471	0.768	0	1.471	0.500	0.400
POSSRB218 ^c	4	3.414	1.307	0.735	0	1.676	0.593	0.420	0	1.324	0.410	0.264	б	2.743	1.051	0.663	б	2.571	1.011	0.733
POSSRB310 ^c	б	2.770	1.058	0.665	б	1.900	0.807	0.493	0	1.960	0.683	0.527	б	2.323	0.960	0.594	0	1.800	0.637	0.533
POSSR53°	4	2.965	1.174	0.689	4	2.706	1.111	0.657	б	2.579	1.004	0.659	б	1.805	0.739	0.466	0	1.800	0.637	0.533
POSSR61 ^c	б	2.104	0.901	0.547	б	2.990	1.097	0.693	0	1.960	0.683	0.527	б	2.667	1.028	0.652				
POSSR95°	4	3.453	1.309	0.740	0	1.923	0.673	0.500	Э	2.085	0.892	0.560	4	3.097	1.238	0.707	ю	2.000	0.868	0.600
<i>Note:</i> — = not available; A = number of alleles per population;	ot avail	able; $A =$	number of	f alleles pe	r popu		= number	of effecti	ve alle	les per pot	oulation; h	unb = unbi	ased h	aploid div	A_e = number of effective alleles per population; h_{unb} = unbiased haploid diversity; I = Shannon's information index; N = number of	Shannon's	inform	ation inde	x; N = nun	nber of

Characterization of 17 novel microsatellite markers in populations of Millettia pinnata and M. pulchra.^a TABLE 2.

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individuals analyzed. ^a Voucher and locality information are provided in Appendix 1. ^b Loci located in the chloroplast. ^c Loci located in the mitochondria.

conducted direct sequencing of the PCR products using the BigDye Terminator v3.1 Cycle Sequencing Kit and ABI PRISM 3730 sequencer (Applied Biosystems, Waltham, Massachusetts, USA) with both forward and reverse primers, to verify the allelic variants tested in this study. Sequences were visualized and analyzed with the DNASTAR software package (DNASTAR, Madison, Wisconsin, USA).

The 17 selected primers exhibited high polymorphisms across 89 individuals of four *M. pinnata* populations. For each of these loci, the number of alleles per population, the number of effective alleles per population, Shannon's information index, and the unbiased haploid diversity (h_{unb}) of each microsatellite locus were calculated using GenAlEx version 6.5 (Peakall and Smouse, 2012). Among chloroplast loci, the number of alleles per locus per population varied from two to six, while h_{unb} ranged from 0.391 to 0.857. For mitochondrial loci, two to four alleles per locus per population were detected and h_{unb} ranged from 0.264 to 0.740 (Table 2). Subsequently, 16 of the 17 developed markers were successfully amplified in the related species *M. pulchra*, demonstrating their transferability (Table 2).

CONCLUSIONS

The 17 polymorphic SSR markers developed here proved useful in the evaluation of the genetic diversity of *M. pinnata*, and 16 showed high transferability within the related species *M. pulchra*. This set of novel polymorphic SSR markers will serve as a very useful tool for the genetic diversity analysis, clonal identification, and germplasm conservation of *M. pinnata* and its related species.

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APPENDIX 1. Sampling information for the populations of Millettia pinnata and M. pulchra used in this study.

Species	Population code	Collection locality	Geographic coordinates	Ν	Voucher no. ^a
Millettia pinnata (L.) Panigrahi	POPH	Perhentian Islands, Malaysia	5°54′34.12″N, 102°44′42.43″E	26	Liu20150926
	POLS	Laem Son, Thailand	6°56′21.10″N, 99°42′43.64″E	25	Huang20160701
	POCN	Chilaw, Sri Lanka	7°33′43.16″N, 79°48′5.96″E	14	Liu20161001
	PONS	Tainan, Taiwan, China	23°1'1.2"N, 120°7'1.20"E	24	He20121208
Millettia pulchra (Benth.) Kurz	MVSD	Heishiding, Guangdong, China	23°27′43.91″N, 111°54′32.37″E	6	Shi20161210

Note: N = number of individuals sampled.

^aAll voucher specimens are deposited at the herbarium of Sun Yat-sen University (SYS), Guangzhou, China.