Development of 12 Chloroplast Microsatellite Markers in Vigna unguiculata (Fabaceae) and Amplification in Phaseolus vulgaris

Authors: Pan, Lei, Li, Yi, Guo, Rui, Wu, Hua, Hu, Zhihui, et al.

Source: Applications in Plant Sciences, 2(3)

2

Published By: Botanical Society of America

URL: https://doi.org/10.3732/apps.1300075

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, Downloaded From: https://sfaging.bioone.org/fournals/Applications-in-Plant-Sciences on 26 Dec 2024
Terms of **Usel Stepel//sta**gin**g.sidene.ug/terms-of-us**eDI ESSES.

PRIMER NOTE

DEVELOPMENT OF 12 CHLOROPLAST MICROSATELLITE MARKERS IN *VIGNA UNGUICULATA* **(FABACEAE) AND AMPLIFICATION IN** *PHASEOLUS VULGARIS*¹

Lei Pan^{2,3}, Yi Li^{2,3}, Rui Guo^{2,3}, Hua Wu^{2,3}, Zhihui Hu^{2,3}, and Chanyou Chen^{2,3,4}

²School of Life Sciences, Jianghan University, Wuhan 430056, Hubei, People's Republic of China; and ³Hubei Province Engineering Research Center of Legume Plants, Jianghan University, Wuhan 430056, Hubei, People's Republic of China

- *Premise of the study: Vigna unguiculata* is an economically important legume, and the complexity of its variability and evolution needs to be further understood. Based on publicly available databases, we developed chloroplast microsatellite primers to investigate genetic diversity within *V. unguiculata* and its related species *Phaseolus vulgaris* .
- *Methods and Results:* Twelve polymorphic chloroplast microsatellite markers were developed and characterized in 62 *V. unguiculata* individuals. The number of alleles per locus varied between two and four, the unbiased haploid diversity per locus ranged from 0.123 to 0.497, and the polymorphism information content varied from 0.114 to 0.369. In cross-species amplifications, nine of these markers showed polymorphism in 29 *P. vulgaris* individuals.
- *Conclusions:* The newly developed chloroplast microsatellite markers exhibit variation in *V. unguiculata* as well as their transferability in *P. vulgaris* . These markers can be used to investigate genetic diversity and evolution in *V. unguiculata* and *P. vulgaris* .

Key words: chloroplast microsatellite; cross-amplification; Fabaceae; *Phaseolus vulgaris*; *Vigna unguiculata* .

Cowpea (*Vigna unguiculata* (L.) Walp.) ($2n = 2x = 22$), a legume crop of economic importance, is widely distributed in the arid and semiarid regions of Africa, Asia, Europe, Latin America, and some parts of the United States (Citadin et al., 2011). As a member of the legume family, it belongs to Phaseoleae, the same tribe as common bean (*Phaseolus vulgaris* L.). Compared to its close relatives and many other crop species, *V. unguiculata* shows a greater tolerance to drought and has the ability to fix nitrogen in poor soils (Muchero et al., 2009). Its grains are a major source of dietary protein for humans, and cowpea hay is fed to livestock as a nutritious fodder (Badiane et al., 2012). However, even though restriction fragment length polymorphism (RFLP), amplified fragment length polymorphism (AFLP), random amplified polymorphic DNA (RAPD), and simple sequence repeat (SSR) molecular makers have been developed for the cowpea nuclear genome, knowledge of variability and evolution in the chloroplast genome of *V. unguiculata* is limited at the molecular level (Provan et al., 2001; Xu et al., 2010).

 Chloroplast microsatellite, or chloroplast simple sequence repeat (cpSSR), markers can be used to detect DNA variability in the chloroplast genome. They have the same characteristics

 The authors thank Professor Zeng Chen (George Washington University, Washington, D.C., USA) for his suggestions and improvement to the English. This work was supported by the Wuhan Planning Project of Science and Technology (no. 2013021001010478, no. 201250499145-11) and by the Research Foundation for Talented Scholars of Jianghan University (no. 2012027), Hubei Province, People's Republic of China. 4 Author for correspondence: ccy@jhun.edu.cn

doi:10.3732/apps.1300075

as nuclear microsatellites, including a multiallelic and codominant nature. Moreover, cpSSR markers are found to be polymorphic and transferable among related species because the flanking regions of cpSSR loci are conserved. Of particular importance, cpSSR markers are maternally inherited in most angiosperms, which allow monitoring of influence on population structure by seed-mediated gene flow and pollen flow (Provan et al., 2001). Therefore, they are useful for analysis of population genetics, genetic diversity, paternity analysis, and germplasm resource identification (Provan et al., 2001). In this study, we developed 12 cpSSR markers for *V. unguiculata* and evaluated their transferability to a related legume species, *P. vulgaris* . These results will be helpful for the future exploration and germplasm conservation in both *V. unguiculata* and *P. vulgaris* , although chloroplast microsatellite diversity in *P. vulgaris* has been investigated (Angioi et al., 2009; Desiderio et al., 2013 .

METHODS AND RESULTS

 The complete chloroplast genome sequence of *V. unguiculata* was downloaded from GenBank (GenBank accession no. NC_018051). The cpSSR loci distributed throughout the *V. unguiculata* chloroplast genome were screened using SSRHunter 1.3 software (Li and Wan, 2005). SSRs were selected based on the length of the core repeat motif $(\geq 10$ nucleotides), for example, five units of dinucleotide repeat motifs, four units of trinucleotide repeat motifs, or three units of tetranucleotide repeat motifs. Primer pairs were designed based on the flanking regions of each SSR locus using Primer3 (Rozen and Skaletsky, 2000). The parameters of each primer were set using the following criteria: (1) primer size of 20–24 nucleotides in length; (2) GC content of 40–60%; (3) annealing temperature between $50-60^{\circ}\text{C}$; and (4) expected amplicon size of 100-300 bp. In total, 15 cpSSR primer pairs of *V. unguiculata* were designed and synthesized (Sangon, Shanghai, China). Twelve of them showed polymorphic bands

Applications in Plant Sciences 2014 2(3): 1300075; http://www.bioone.org/loi/apps © 2014 Pan et al. Published by the Botanical Society of America. This work is licensed under a Creative Commons Attribution License (CC-BY-NC-SA).

¹ Manuscript received 20 September 2013; revision accepted 5 December 2013.

Note: IGS = intergenic spacer; LSC = long single-copy region; SSC = short single-copy region; T_a = annealing temperature.

Position of each SSR in chloroplast complete genome of *Vigna unguiculata* (GenBank accession number: NC_018051).

in *V. unguiculata* accessions, two were monomorphic, and one primer pair gave no products. The 12 polymorphic markers were used in the following analysis.

 A total of 91 samples were used in this study, including 62 *V. unguiculata* accessions and 29 *P. vulgaris* accessions (Appendix 1). All the samples were collected from an agricultural field in Anshan (30.46°N, 113.94°E), Caidian District, Wuhan City, and preserved in Hubei Province Engineering Research Center of Legume Plants, Wuhan, China. Tender young leaves of each sample were collected and stored at −80°C until use. Total DNA was extracted from all the samples using the cetyltrimethylammonium bromide (CTAB) method (Doyle and Doyle, 1987). The yield and purity of the DNA were measured using a spectrophotometer SP-1910UVPC (Shanghai, China) at an A260/A280-nm wavelength.

 Characteristics of cpSSR markers were examined in both *V. unguiculata* and *P. vulgaris* . The same PCR conditions were applied in the two species. The PCR amplifications were performed in a 20- μ L reaction mixture containing 1× *Taq* buffer, 30 ng of genomic DNA, 1.5 mM MgCl₂, 200 μM dNTPs, 0.5 μM for each primer, and 0.5 U *Taq* polymerase (MBI Fermentas, Vilnius, Lithuania). The PCR conditions were as follows: an initial denaturation at 94°C for 5 min; followed by 35 cycles of 30 s at 94° C, 30 s at the locus-specific annealing temperature (Table 1), and 40 s at 72°C; and a final extension at 72°C for 5 min. The PCR products were separated using 6% denaturing polyacrylamide gels $(Ar: Bis = 19:1)$ and visualized with silver staining. Due to the nonrecombining nature of the chloroplast genome, each pair of chloroplast microsatellite primers was considered as a "locus" at a cpSSR site. Length variants of chloroplast microsatellites at each cpSSR site were treated as alleles. Alleles detected from polymorphic primer pairs were used to generate a chloroplast haplotype of each individual; multilocus haplotypes were obtained by combining alleles from all polymorphic loci. Based on the polymorphic cpSSR markers, the fragment size amplified from each locus was scored by referring to a 20-bp DNA ladder (TaKaRa Biotechnology Co., Dalian, China). The number of alleles (A) and unbiased haploid diversity index (h) per polymorphic locus were calculated using the software GenAlEx version 6.41 (Peakall and Smouse, 2006). To estimate the informativeness of each SSR marker, the polymorphism information content (PIC) was calculated using the formula described by Botstein et al. (1980) .

As shown in Table 2, the characteristics of the 12 polymorphic cpSSR loci are tested in 62 *V. unguiculata* samples. *A* ranged from two to four in *V. unguiculata* (average: 2.75), *h* ranged from 0.123 (VgcpSSR4) to 0.497 (VgcpSSR5) (average: 0.240), and PIC ranged from 0.114 (VgcpSSR4) to 0.369 (VgcpSSR5) (average: 0.211).

 The transferability of the 12 *V. unguiculata* cpSSR markers was assessed in a related species, *P. vulgaris*; parameters of genetic variation were evaluated in

29 *P. vulgaris* individuals (the *P. vulgaris* group) (Table 2). All of the 12 cpSSR markers were successfully amplified in the *P. vulgaris* group, and nine showed polymorphisms, with the exception of VgcpSSR7, VgcpSSR9, and VgcpSSR13, which were monomorphic markers. Therefore, it indicated that 75% of these markers can amplify polymorphic bands. In *P. vulgaris* , *A* ranged from one to two, with an average value of 1.75. For each cpSSR locus, *h* was between 0.000 (VgcpSSR7, VgcpSSR9, and VgcpSSR13) and 0.529 (VgcpSSR10 and VgcpSSR14) (average: 0.312). The PIC value varied between 0.183 (VgSSR3) and 0.374 (VgcpSSR2, VgcpSSR10, and VgcpSSR14) (average: 0.312).

CONCLUSIONS

 Twelve polymorphic cpSSR markers were developed in *V. unguiculata* and showed high transferability in *P. vulgaris.* Further

 TABLE 2. Characterization of the 12 cpSSR markers in *V. unguiculata* and their cross-species amplification in *P. vulgaris*.

Locus	V. unguiculata group			P. vulgaris group		
	А	h	PIC	A	h	PIC
VgcpSSR1	3	0.210	0.196	$\overline{2}$	0.323	0.262
VgcpSSR2	3	0.362	0.303	$\overline{2}$	0.516	0.374
VgcpSSR3	2	0.153	0.139	$\overline{2}$	0.212	0.183
VgcpSSR4	2	0.123	0.114	$\overline{2}$	0.380	0.298
VgcpSSR5	\overline{c}	0.497	0.369	$\overline{2}$	0.467	0.332
VgcpSSR7	\overline{c}	0.125	0.116	1	0.000	
VgcpSSR9	\overline{c}	0.151	0.138		0.000	
VgcpSSR10	4	0.256	0.237	$\overline{2}$	0.529	0.374
VgcpSSR11	3	0.202	0.185	$\overline{2}$	0.441	0.329
VgcpSSR12	4	0.270	0.255	$\overline{2}$	0.349	0.280
VgcpSSR13	3	0.154	0.146	1	0.000	
VgcpSSR14	3	0.383	0.328	$\overline{2}$	0.529	0.374
Average	2.75	0.240	0.211	1.75	0.312	0.312

Note: A = number of alleles for each locus; h = unbiased haploid diversity; PIC = polymorphism information content.

analyses indicated that the cpSSR markers of *V. unguiculata* could reveal a relatively high level of genetic diversity in both *V. unguiculata* and *P. vulgaris* germplasm. These markers can be used to investigate genetic diversity and evolution in *V. unguiculata* and *P. vulgaris* .

LITERATURE CITED

- ANGIOI, S. A., D. RAU, M. RODRIGUEZ, G. LOGOZZO, F. DESIDERIO, R. PAPA, AND G. ATTENE. 2009. Nuclear and chloroplast microsatellite diversity in *Phaseolus vulgaris* L. from Sardinia (Italy). *Molecular Breeding* 23 : $413 - 429$
- BADIANE, F. A., B. S. GOWDA, N. CISSÉ, D. DIOUF, O. SADIO, AND M. P. TIMKO. 2012. Genetic relationship of cowpea (Vigna unguiculata) varieties from Senegal based on SSR markers. *Genetics and Molecular Research* 11: 292-304.
- BOTSTEIN, D., R. L. WHITE, M. SKOLNICK, AND R. W. DAVIS. 1980. Construction of a genetic linkage map in man using restriction fragment length polymorphism. *American Journal of Human Genetics* 32: 314-331.
- CITADIN, C. T., A. B. IBRAHIM, AND F. J. ARAGÃO, 2011. Genetic engineering in cowpea (*Vigna unguiculata*): History, status and prospects. *GM Crops* 2: 144–149.
- DESIDERIO, F., E. BITOCCHI, E. BELLUCCI, D. RAU, M. RODRIGUEZ, G. ATTENE, R. PAPA, AND L. NANNI. 2013. Chloroplast microsatellite diversity in *Phaseolus vulgaris. Frontiers in Plant Science* 3: 312.
- DOYLE, J. J., AND J. L. DOYLE. 1987. A rapid DNA isolation procedure for small quantities of fresh leaf tissue. *Phytochemical Bulletin* 19: $11 - 15$.
- LI, Q., AND J. M. WAN. 2005. SSRHunter: Development of a local searching software for SSR sites. *Hereditas* 27: 808-810 (in Chinese) .
- MUCHERO, W., J. D. EHLERS, T. J. CLOSE, AND P. A. ROBERTS. 2009. Mapping QTL for drought stress-induced premature senescence and maturity in cowpea [*Vigna unguiculata* (L.) Walp.] . *Theoretical and* Applied Genetics 118: 849-863.
- PEAKALL, R., AND P. E. SMOUSE. 2006. GenAlEx 6: Genetic analysis in Excel. Population genetic software for teaching and research. *Molecular Ecology Resources 6: 288-295.*
- PROVAN, J., W. POWELL, AND P. M. HOLLINGSWORTH. 2001. Chloroplast microsatellites: New tools for studies in plant ecology and evolution. *Trends in Ecology & Evolution* 16: 142-147.
- ROZEN, S., AND H. SKALETSKY. 2000. Primer3 on the WWW for general users and for biologist programmers. *In* S. Misener and S. A. Krawetz [eds.], Methods in molecular biology, vol. 132: Bioinformatics methods and protocols, 365–386. Humana Press, Totowa, New Jersey, USA.
- XU, P., X. WU, B. WANG, Y. LIU, D. QIN, J. D. EHLERS, T. J. CLOSE, ET AL. 2010 . Development and polymorphism of *Vigna unguiculata* ssp. *unguiculata* microsatellite markers used for phylogenetic analysis in asparagus bean (*Vigna unguiculata* ssp. *sesquipedialis* (L.) Verdc.). *Molecular Breeding* 25: 675-684.

