## **Notas Científicas**

# Microsatellites for genetic studies and breeding programs in common bean

Tatiana de Campos<sup>(1)</sup>, Luciana Lasry Benchimol<sup>(2)</sup>, Sérgio Augusto Moraes Carbonell<sup>(2)</sup>, Alisson Fernando Chioratto<sup>(2)</sup>, Eduardo Fernandes Formighieri<sup>(3)</sup> and Anete Pereira de Souza<sup>(1)</sup>

(¹¹)Universidade Estadual de Campinas (Unicamp), Centro de Biologia Molecular e Engenharia Genética, Caixa Postal 6.010, CEP 13083-970 Campinas, SP, Brazil. E-mail: tatyuni@unicamp.br, anete@unicamp.br (²¹)Instituto Agronômico, Centro de Pesquisa e Desenvolvimento de Recursos Genéticos Vegetais, Fazenda Santa Elisa, Caixa Postal 28, CEP 13001-970 Campinas, SP, Brazil. E-mail: llasry@iac.sp.gov.br, carbonel@iac.sp.gov.br, afchiorato@iac.sp.gov.br (³¹)Unicamp, Dep. de Genética e Evolução, Caixa Postal 6.109, CEP 13083-970 Campinas, SP, Brazil. E-mail: eduformi@lge.ibi.unicamp.br

Abstract – Twenty microsatelitte loci were identified and characterized in common bean. Microsatellites were tested in 14 genotypes. The allele number ranged from 1 to 3, and the polymorphism information content (PIC) was between 0.14 and 0.65. These polymorphic markers are available to be used for breeding programs.

Index terms: Phaseolus vulgaris, SSR, molecular markers.

# Microssatélites para estudos genéticos e programas de melhoramento em feijoeiro

Resumo – Vinte locos de marcadores microssatélites foram identificados e caracterizados em feijoeiro. Os microssatélites foram testados em 14 genótipos. O número de alelos variou entre 1 e 3, e o conteúdo informativo de polimorfismo (PIC) entre 0,14 e 0,65. Esses marcadores polimórficos estão disponíveis para serem usados em programas de melhoramento.

Termos para indexação: Phaseolus vulgaris, SSR, marcadores moleculares.

Common bean (*Phaseolus vulgaris* L.) is the principal leguminosae used for human nourishment, mainly in South America and Africa, where it represents an important source of protein. Brazil detains the most important productivity and consumption of common bean in the world. However, this crop has not reached high productivity (Silva, 2000). Common bean occurred in two centers of origin in South and Central America, comprising two major gene pools, Andean or large-seeded bean type, and Mesoamerican or small-seeded type (Gepts & Debouck, 1991).

Microsatellites or simple sequence repeats (SSR) are polymerase chain reaction (PCR) based markers developed, for a wide range of plant species, around short segments of DNA, in which a specific motif of one to six nucleotides is repeated in tandem and distributed over the euchromatic part of the genome (Morgante & Olivieri, 1993; Powell et al., 1996). In common bean there are about 200 available SSR markers (Yu et al., 2000; Gaitán-Solís et al., 2002;

Métais et al., 2002; Yaish & Pérez de la Vega, 2003; Buso et al., 2006), a small number when compared with other crops, like soybean (Cregan et al., 1999).

Microsatellite markers are powerful tools but their development is expensive and labour intensive. Consequently, many researchers have tried to use primer pairs developed for one species in another (Cipriani et al., 1999), attributing knowledge as transferability or cross-species amplification.

The variety IAC-UNA was used to construct a microsatellite enriched library for two dinucleotide repeat sequences (CT and GT). This enrichment was based on the procedure described by Billotte et al. (1999). IAC-UNA is a black-seeded variety, developed by Agronomic Institute (IAC, Campinas, SP, Brazil), resistant to anthracnose and susceptible to bean rust, *Fusarium* and angular leaf spot.

The genomic DNA was extracted from leaf tissue using a CTAB extraction method, as described by Hoisington et al. (1994). The extracted DNA was

digested with *RsaI* restriction enzime, and the digested fragments were linked to *RsaI* adapters. The library was enriched for dinucleotide sequences using (CT)<sub>8</sub> and (GT)<sub>8</sub> biotinylated microsatellite primers with labelled probes that were bound to Streptavidine MagneSphere Paramagnetic Particles as described by manufacturer.

Selected fragments were amplified by PCR using primer sequences complementary to the adapters, and then, attached to the vector pGEM-T (Promega). Plasmids were introduced into XL-1 Blue cells. Transformed cells were cultivated on agar plates containing 100 µg mL<sup>-1</sup> of ampicilin and 50 µg mL<sup>-1</sup> of X-galactosidase. Single white colonies were transferred onto microplates for long-term storage at 80°C. Sequencing of the inserts was performed using the ABI 377 Big Dye Terminator and 20 sequences containing microsatellites were selected to have primer pairs designed.

Primer designing was performed using Primer Select with the following conditions: amplification DNA size from 150 pb to 350 pb; GC content between 40–60%; temperature annealing (*Ta*) between 45 and 60°C; primer length between 18 and 22 pb; no hairpins or dimmers.

Twenty primers were designed for microsatellite loci, and were selected for characterization using 14 accessions from the IAC Germplasm Bank, including Andean (A) and Mesoamerican (M) gene pools. Total DNA was extracted from the following accessions: 'Sanilac' (M), 'Baetão' (M), 'Red Kidney' (A), 'Jamapa' (M), 'Flor de Mayo' (M), 'Tu' (M), 'Carioca Comum' (M), 'Jabola [CB]' (A), 'Fradinho Cruzeiro' (*Vigna* spp.), '87-JP-12' (A), 'BAT-93' (M), 'Jalo EEP-558' (A), 'IAC-UNA' (M), 'CAL-143' (A).

PCR reactions were carried out in a total volume of 25 μL containing 50 ng of template DNA, 0,8 μM of forward and reverse primer, 100 μM of each dNTP, 1,5 mM MgCl<sub>2</sub>, 10 mM Tris-HCl, 50 mM KCl, and 0.5 U *Taq* DNA Polymerase (Invitrogen). Reactions were performed using the following conditions: 1 min at 94°C; then, 30 cycles of [1 min 94°C, 1 min at specific Ta, 1 min at 72°C], followed by 5 min at 72°C. Amplification products were checked by electrophoresis on 3% agarose gels, and then loaded on 6% w/v denatured polyacrylamide gels using a 10 bp ladder (Invitrogen) as a size standard. After each run, gels were silver stained according to Creste et al. (2001).

The polymorphism information content (PIC) value was calculated by the following formula:

$$PIC = 1 \text{--} \sum_{i=1}^{n} \, f_{i}^{\ 2} \text{--} \sum_{i=l}^{n-1} \sum_{j=i+l}^{n} 2 \, f_{i}^{\ 2} \, f_{j}^{\ 2}$$

in which n is the number of alleles;  $f_i$  and  $f_j$  are the frequencies of the  $i^{th}$  and  $j^{th}$  alleles, respectively (Botstein et al., 1980). Polymorphisms were observed among the amplified alelles.

Fifteen loci were polymorphic, four were monomorphic and one was unable to amplify among the twenty analyzed microsatellites (Table 1). Thirteen primer pairs were perfect dinucleotide motifs, one was imperfect (FJ 20), and six presented compound motifs. Two loci could separate Andean from Mesoamerican accessions; showing two different alleles, each one associated to a genetic pool (FJ 05 and FJ 17). These SSRs would be tested in other genotypes, considering that they can be a tool to separate in domestication centers.

The number of alleles ranged from one to three, with an average of 2.07 alleles per locus. Yu et al. (2000) found an allele range that varied from 2 to 10, for a 24 polymorphic SSR evaluation in 12 genotypes. PIC values ranged from 0.14 to 0.65. The highest PIC (0.65) was found in FJ 14, that presented the greatest repetition motif, (GA)<sub>10</sub>, and the most elevated number of alleles (3). Métais et al. (2002) published a range of 0.12 to 0.72 for PIC values, with an average of 0.44, when evaluating 15 polymorphic SSRs in 45 different bean lines of nine different quality types.

The primer pairs were also used to amplify one genotype of Vigna spp., accession 'Fradinho Cruzeiro', and one genotype of *Phaseolus lunatus*, accession 87-JP-12, using the same conditions optimized for *Phaseolus vulgaris* accessions. The efficiency of heterologous amplification was 100%. However, it would be necessary a resequencing of the amplification products to check if they were really related to the original sequence from which primer pairs were developed. Even so, this shows a considerable level of sequence, conservation within the primer regions flanking microsatellite loci. These results suggest that the new microsatellites reported in this paper could be used for synteny studies, establishing the conservation of genes between species such as Phaseolus lunatus and Vigna spp.

In common bean, most of the molecular markers used for genetic studies and breeding programs are RAPD

**Table 1.** Primer sequences and characteristics of 20 common bean (*Phaseolus vulgaris*) microsatellite markers, obtained from variety IAC-UNA, tested on 14 accessions, comprising Mesoamerican and Andean gene pools.

Locus/ GeneBank accession	Primers $(5' \rightarrow 3')$	Repeat motif	Na <sup>(1)</sup>	Ta <sup>(2)</sup> (°C)	Size of cloned alelle (size range) in bp	PIC <sup>(3</sup>
FJ 01/	GTCGCCGCTACTTCTTTGTT	(AC) <sub>7</sub>	2	60	270 (265-270)	0.51
DQ469376	TTTTAATGTTGTGGGAGTGATG	_ ()/			()	
F.J. 02/	GGTCCACAATCAAGCAGTCA	$(AC)_{10} (AT)_5$	2	51,4	251 (258-260)	0.52
DQ469377 -	TATGGAACCTGATAGCAAGTG	_ (110)[0 (111)]	_	01,.	201 (200 200)	0.02
FJ 03/	TTCGCGAGCAAGCAACTA	(GT) <sub>6</sub>	1	45	178 (177)	0.00
DQ469378	TGAATGTTTTAAATGCGTTGAA	_ (31)6	-		170 (177)	0.00
FJ 04/	ATAGATGAAGGATTGGGAGAG	(AG) <sub>8</sub>	1	45	216 (218)	0.00
DO469379	GGGAAATTGAAGAGGAGATAC	_ (110)8	•		210 (210)	0.00
FJ 05/	AAGAAACAGAAACAATAAAAAC	(CT) <sub>2</sub> (CA) <sub>6</sub>	2	60	212 (220-222)	0.48
DQ469380 -	TTTCCATTTATTTTCAGTCACA	_ (01)2 (011)6	_	00	212 (220 222)	0.10
FJ 06/	TTGGAACACCGTGGAATGGA	(GT) <sub>7</sub>	0	60	152 (-) <sup>(4)</sup>	0.00
DO469381	GAGGCTTTAGACGTTGGAGACA	_ (31)/	Ü		10-()	0.00
FJ 07/	GAAAACGCGAAACAACCGA	(CA) <sub>8</sub>	2	60	290 (283-297)	0.52
DQ469382	ATGTCTCCAAATCCCAAGTG	_ (012)8	_			
FJ 08/	ATGGTCATGGTATCAGTTCA	(CA) <sub>6</sub> (TA) <sub>3</sub>	1	60	195 (195)	0.00
DO469383	TCTTTTCCATAGTATTCTCTTG	_ (011)6 (111)3	1	00	155 (155)	0.00
FJ 09/	ACCTTAGATAGTGCTTGTTAGAG	(TG) <sub>6</sub>	1	45	155 (153)	0.00
DQ469384 -	CATGACACCTAGGGCAAA	_ (10)6	•		100 (100)	0.00
FJ 10/	AGGGGAGTTGTGTTCTTAC	(TG) <sub>6</sub>	2	45	209 (207-209)	0.14
DO469385	ATACGTACGAGTGACTGGAGA	_ (10)6	_	15	203 (207 203)	0.1 1
FJ 11/	AAAAGGATCAAAGAGGAGAAAT	(CA) <sub>5</sub>	2	60	297 (308-310)	0.25
DO469386	GGGCAAGTAAAGCTAAACGAG	_ (011)5	_	00	257 (300 310)	0.20
F.J. 12/	TATCAGCCTAGTTATTTTCAAG	(CA) <sub>7</sub>	2	60	256 (255-258)	0.25
DQ469387	CATACTTTCTTATTTTCTGGA	_ (0/1)/	_	00	230 (233 230)	0.20
F.J.13/	TCGATGCAGGATTGGATT	(AC) <sub>9</sub>	2	60	266 (267-269)	0.48
DQ469388 -	CAGGTTGATTGTGATAGGTTAC	_ (110)9	_	00	200 (201 205)	0.10
F.J 14/	TTCATGGCAAGGTAAGTAAATA	(AG) <sub>10</sub>	3	60	148 (145-155)	0.65
DQ469389 -	TGAATGAACACAACAACAA	_ (/10)10	5	00	110 (115 155)	0.05
FJ 15/	AGAATGGAGGGAAAAGCAAAAG	(ATGAG) <sub>4</sub>	2	60	191 (200-205)	0.14
DQ469390 -	CCGAAGTCCAAGATTAGAAGCC	$-\frac{(\mathrm{FIG})_{4}}{(\mathrm{GT})_{3}}$	_	00	191 (200 203)	0.1
FJ 16/	TGGTGCTACAACAAAAGAGAAT	(TA) <sub>6</sub> (TG) <sub>7</sub>	2	60	284 (280-300)	0.20
DQ469391	TAGGCATGTGGGTAGGTCAG	_ (111)6(10)/	_	00	20. (200 200)	0.20
FJ 17/	TCCCGATTTATAGTTCTCATTT	(TG) <sub>8</sub> (TA) <sub>3</sub>	2	60	222 (220-230)	0.48
DQ469392	AGGGACCTCCTTCATCTC	_ (10)8(111)3	_	00	222 (220 250)	0.10
FJ 18/	CATTGAGATTTGAGGTTTCGTT	(TG) <sub>5</sub>	2	60	224 (224-230)	0.48
DQ469393	AGGTATTTCCATCGTGCTTTTC	_ (10)	-	00	22 (22 : 233)	0.10
FJ 19/	ATGTTAGTGCCTTATTTCTCT	(CA) <sub>7</sub>	2	60	205 (210-222)	0.51
DQ469394	AAGGTAGGGTTGGGATTGT	_ (0/1)/	_	00	203 (210 222)	0.51
FJ 20/	TTGGAACACCGTGGAATGGA	(AG) <sub>3</sub> AA	2	60	251 (250-263)	0.51
DQ469395 -	GAGGCTTTAGACGTTGGAGACA	$-\frac{(AG)_3}{(AG)_3}$	-	00	231 (230 203)	0.51

 $<sup>{\ }^{(1)}</sup>Number\ of\ alleles.\ {\ }^{(2)}Temperature\ annealing.\ {\ }^{(3)}Polymorphism\ information\ content.\ {\ }^{(4)}No\ amplification.$ 

markers, which are dominant, and not reproducible. SSR are codominant, more polymorphic and stable. These new informative microsatellites are an available tool in common bean research. They consist in important source of polymorphism which can be used in breeding programs or in genetic studies, as genetic and QTL mapping, marker-assisted selection and germplasm characterization in common bean.

## Acknowledgements

To Fapesp, for financial support and fellowships; to CNPq, for fellowships.

#### References

BILLOTTE, N.; LAGODA, P.J.L.; RISTERUCCI, A.M.; BAURENS, C. Microsatellite-enriched libraries: applied

methodology for the development of SSR markers in tropical crops. **Fruits**, v.54, p.277-288, 1999.

BOTSTEIN, D.; WHITE, R.L.; SKOLNICK, M.; DAVIS, R.W. Construction of a genetic linkage map in man using restriction fragment length polymorphisms. **American Journal of Human Genetics**, v. 32, p.314-331, 1980.

BUSO, G.S.C.; AMARAL, Z.P.S.; BRONDANI, R.P.V.; FERREIRA, M.E. Microsatellite markers for the common bean *Phaseolus vulgaris*. **Molecular Ecology Notes**, v.6, p.252-254, 2006.

CIPRIANI, G.; LOT, G.; HUANG, W.G.; MATARAZZO, M.T.; PETERLUNGER, E.; TESTOLIN, R. AC/GT and AG/CT microsatellite repeats in peach [*Prunus persica* (L.) Batsch]: isolation, characterisation and cross-species amplification in *Prunus*. **Theoretical and Applied Genetics**, v.99, p.65-72, 1999.

CREGAN, P.B.; JARVIK, T.; BUSH, A.L.; SHOEMAKER, R.C.; LARK, K.G.; KAHLER, A.L.; KAYA, N.; VAN TOAI, T.T.; LOHNES, D.G.; CHUNG, J.; SPECHT, J.E. An integrated genetic linkage map of the soybean genome. **Crop Science**, v.39, p.1464-1490, 1999.

CRESTE, S.; TULMANN-NETO, A.; FIGUEIRA, A. Detection of single sequence repeat polymorphisms in denaturing polyacrylamide sequencing gels by silver staining. **Plant Molecular Biology Reporter**, v.19, p.299-306, 2001.

GAITÁN-SOLÍS, E.; DUQUE, M.C.; EDWARDS, K.J.; TOHME, J. Microsatellite repeats in common bean (*Phaseolus vulgaris*): isolation, characterization and cross-species amplification in *Phaseolus* ssp. **Crop Science**, v.42, p.2128-2136, 2002.

GEPTS, P.; DEBOUCK, D. Origin, domestication, and evolution of the common bean (*Phaseolus vulgaris* L.). In: SCHOONHOVEN, A. van; VOYSEST, O. (Ed.). **Common beans**: research for crop improvement. Cali: CIAT, 1991. p.7-53.

HOISINGTON, D.; KHAIRALLAH, M.; GONZÁLEZ-DE-LEÓN, D. **Laboratory protocols**: CIMMYT applied molecular genetics laboratory. 2<sup>nd</sup> ed. Mexico: CIMMYT, 1994. 88p.

MÉTAIS, I.; HAMON, B.; JALOUZOT, R.; PELTIER, D. Structure and level of genetic diversity in various bean types evidenced with microsatellite markers isolated from a genomic enriched library. **Theoretical and Applied Genetics**, v.104, p.1346-1352, 2002.

MORGANTE, M.; OLIVIERI, A.M. PCR-amplified microsatellites as markers in plant genetics. **Plant Journal**, v.3, p.175-182, 1993.

POWELL, W.; MACHRAY, G.C.; PROVAN, J. Polymorphism revealed by simple sequence repeats. **Trends in Plant Science**, v.1, p.215-222, 1996.

SILVA, M.V. da. **Identificação de marcador RAPD ligado ao alelo co-7 de resistência do feijão ao agente causal da antracnose**. 2000. 41p. Dissertação (Mestrado) - Universidade Federal de Lavras, Lavras

YAISH, M.W.F.; PÉREZ DE LA VEGA, M. Isolation of (GA)<sub>n</sub> microsatellite sequences and description of a predicted MADS-box sequence isolated from common bean (*Phaseolus vulgaris* L.). **Genetics and Molecular Biology**, v.26, p.337-342, 2003.

YU, K.; PARK, S.J.; POYSA, V.; GEPTS, P. Integration of simple sequence repeat (SSR) markers into a molecular linkage map of common bean (*Phaseolus vulgaris*). **Journal of Heredity**, v.91, p.429-434, 2000.

Received on October 19, 2006 and accepted on February 14, 2007