Natural selection in common bean microsatellite alleles and identification of QTLs for grain yield

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Abstract Natural selection acts to select better adapted individuals or alleles in segregating population and help plant breeding. The objective of this work was to verify the effect of natural selection on microsatellite alleles as indicators of better adaptation and identification of quantitative trait loci (QTLs) for grain yield. This study evaluated 107 progenies from the F_8 and 107 from the F_{24} generation derived from crossing Carioca MG and ESAL 686 lines, carried out by the bulk method, and evaluated in three different seasons: winter 2001; rainy 2001 and dry 2002. It was utilized 22 polymorphic markers and the natural selection acted in all of them. The frequency of the alleles of the parent Carioca MG, the most adapted, was increased in all of the 22 loci in F_8 and 19 loci in F_{24} . Selection affected each locus with different intensities in different generations. All of the selected alleles can be important for breeding program. QTLs were identified in generation F_8 and F_{24} at varied magnitudes. The best marker PVttc002 explained 11.76% of variation in grain yield. However, an elevated interaction between QTLs and the environments was observed, showing the great difficulty in assisted selection.

Keywords: bulk method, fitness coefficient, Phaseolus vulgaris, SSR markers

INTRODUCTION

In the segregant population of common bean (*Phaseolus vulgaris*), advanced by bulk method, the natural selection maintains the individuals in the population with favorable characteristics for the environment in which they are found, contributing to their better adaptation. Grain yield, just as the majority of economically important characteristics, is affected by natural selection. Considering this trait one of the breeding methods that is most affected by natural selection is the bulk which maintains the most adapted individuals (Allard, 1999).

Natural selection can offer benefits to those who wish to improve plant traits such as yield and disease resistance, but in some cases such as growth habit and size of seeds, it can be disadvantageous, and artificial selection is necessary in these cases (Allard, 1999).

Molecular markers affected by natural selection can be an important alternative for selection of better adapted alleles (Allard, 1999). These markers can be identified from alteration in their allele frequencies, with the prevailing allele in advanced generations explaining the best adaptation, as observed by Rodrigues and Santos (2006).

Molecular markers can also be used as a tool to aid in the selection process of quantitative traits in which environmental influence is greater, such as grain yield. Better contribution of these markers is expected because of the difficulties involved in the selection process (Silva and Vencovsky, 2002). However, the results observed are not always the most promising (Blair et al. 2006; Rodrigues et al. 2007; Torga et al. 2010). The objective of this work was to identify microsatellite markers (SSR) selected by natural selection in segregating bean populations advanced by the bulk method and to identify markers related to grain yield.

MATERIALS AND METHODS

A segregant population that came from the crossing Carioca MG and ESAL 686 genitors was used. Carioca MG is a variety with small cream-colored beans with brown stripes, undetermined type II growth habit and carries the *Co-2* allele, which confers resistance to some types of the *Colletotrichum lindemuthianum* fungus, which causes anthracnose. ESAL 686 has determined type I growth habit, it has big and yellow beans, short cycles (80 days) and is resistant to angular leaf spot (*Pseudocercospora griseola*).

Beginning with the crossing of Carioca MG and ESAL 686 lines, the segregant F_2 population was carried forward in bulk until generation F_8 , from which a sample of 107 plants was taken (Silva et al. 2004). The rest of the population continued to be carried forward until F_{24} , at which time another 107 plant samples were taken. The segregant population was advanced by the bulk method. At each generation, after harvest, a sample of around 2000 bean seeds was stored in a cold chamber to be used for obtaining the next generation.

For this study, two parents and 107 progenies descended from each generation $F_{8:11}$ and $F_{24:27}$, were also utilized by Silva et al. (2004), Rodrigues and Santos (2006) and Rodrigues et al. (2007). Fifteen beans from each progeny were planted in a tray for DNA extraction as in Rodrigues and Santos (2006). Polymorphic chain reactions (PCR) were carried out with 135 pairs of microsatellite markers, whose sequences are published by Blair et al. (2003), Caixeta et al. (2005) and Hanai et al. (2010). The PCR reaction began with DNA denaturation at 95°C for 2 min followed by 32 cycles of denaturation at 94°C for 20 sec, annealing at 42 to 64°C (depending on the primer) for 20 sec, and elongation at 72°C for 20 sec, with a final elongation at 72°C for 10 min. After amplification the reaction products were separated by polyacrylamide gel (6 to 8%) electrophoresis, stained with silver nitrate and photographed with a digital camera (Benbouza et al. 2006).

Genotyping of F_8 and F_{24} progenies was carried out with twenty-two loci that had polymorphism in the genitors and an equal mixture of DNA from F_{24} progenies. For each microsatellite locus, the genotypic proportions of both generations were compared by means of the x^2 test, considering A^1 the allele from the Carioca MG genitor and A^2 from the ESAL 686 genitor. Thus, at the j^{th} segregant population, with j=1 corresponding to F_8 and j=2 corresponding to F_{24} , the j^{th} genotypes occur, with j=1 corresponding to A^1A^1 , j=1 corresponding to A^1A^2 , and j=1 corresponding to A^1A^2 . With the number of the j^{th} genotype observed in the j^{th} generation represented by n_{ij} , the corresponding number expected is given by

$$e_{i\,j}=(n_{i\,|}n_{\cdot\,j\,})/n_{\cdot\,\cdot\,}$$
 in which

$$n_{i.} = \sum_{j=1}^{2} n_{ij}$$
, $n_{.j} = \sum_{i=1}^{3} n_{ij}$ and $n_{..} = \sum_{i,j} n_{ij}$

$$\chi^2 = \sum \frac{(n_{ij} - e_{ij})^2}{e_{ij}}$$

 x^2 was estimated with two degrees of freedom, that is, the product of the number of populations minus one by the number of genotypes per progeny minus one.

Considering that the natural crossing rate of beans in the region is approximately T = 0.005 (Pereira Filho and Cavariani, 1984) and the corresponding self-fertilization rate is S = 1-T = 0.995, the genotypic frequencies for each primer pair (locus) were estimated.

Thus, with A^1 and A^2 the alleles of each locus, genotypic frequencies estimated in generations n and n + 1 are given by the following expressions (Allard, 1999; Rodrigues and Santos, 2006):

Freq
$$(A^1A^1) = f_1^{(n+1)} = S[f_1^{(n)} + 0.25f_2^{(n)}] + T[f_1^{(n)} 0.5f_2^{(n)}]^2$$

Freq
$$(A^1A^2) = f_2^{(n+1)} = S[0.5f_2^{(n)}] + 2T[f_1^{(n)} + 0.5f_2^{(n)}][f_3^{(n)} + 0.5f_2^{(n)}]$$

Freq
$$(A^2A^2) = f_3^{(n+1)} = S[f_3^{(n)} + 0.25f_2^{(n)}] + T[f_3^{(n)} + 0.5f_2^{(n)}]^2$$

Considering the relative adaptability coefficient of genotype A^1A^1 as ω_1 of genotype A^2A^2 as ω_3 and of genotype A^1A^2 as $\omega_2 = 1$, the accumulated relative adaptability coefficients of F_2 to F_8 and F_8 to F_{24} were estimated by means of the expressions (Allard and Hansche, 1964):

$$\omega_1 = \frac{O_1[0,5SH + 2T(P + 0,5H)(R + 0,5H)]}{O_2[S(P + 0,25H) + T(P + 0,5H)^2]}$$

$$\omega_3 = \frac{O_3[0,5SH + 2T(P + 0,5H)(R + 0,5H)]}{O_2[S(R + 0,25H) + T(R + 0,5H)^2]}$$

In which: P and O_1 are the proportions of A^1A^1 in generations n and n+1, respectively; H and O_2 are the proportions of A^1A^2 in generations n and n+1; and R and O_3 are the proportions of A^2A^2 in generations n and n+1.

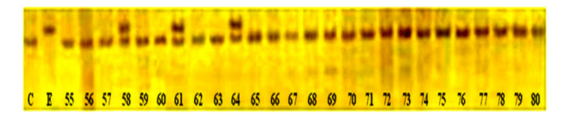


Fig. 1 Pattern of microsatellite bands amplified by the AJ416395 primer in poliacrilamyde gel; column with the letter C is Carioca MG parent and with the letter E is ESAL 686 parent; following columns are progenies 55 to 80 from F_{24: 27} generation and they are indicated by numbers.

The genotypic proportions in generation n are expected to be the same as those in generation F_7 if there is no natural selection, taking into consideration the rates of 0.005 crossing and 0.995 self-fertilization. While in F_{23} , these are the proportions expected beginning from the proportions observed at F_8 , also taking into consideration the absence of natural selection and the rates of crossing and self-fertilization.

The progenies were also evaluated for grain yield by Silva et al. (2004) in three seasons: winter 2001 ($F_{8:9}$ and $F_{24:25}$), planted in August in Ijaci using a simple 18 x 18 square lattice design, and plot with one meter row with 0.5 m spacing, with a density of 15 seeds; the rainy season in 2001 ($F_{8:10}$ and $F_{24:26}$), planted in November 2001 and the dry season of 2002 ($F_{8:11}$ and $F_{24:27}$), planted in March 2002, the latter two both in Lavras using a triple 18 x 18 square lattice design, and plot with two meter rows with 0.5 m spacing and a density of 15 seeds per meter. In all seasons, grain weight per plot was evaluated.

Simple and multiple linear regression analyses were carried out with the genotyping results, with model selection by the backward method for each of the 22 markers, with average grain yield data at each generation per season and also the mean grain yield of all seasons. The analyses were set up using SAS 8.0 version software.

RESULTS AND DISCUSSION

Among the 135 primer pairs utilized, 22 had polymorphism between the parents and the bulk of generation F_{24} , as shown by the PVatt003 and AJ416395 primers (Figure 1 and Figure 2). Thirteen of the primers chosen were mapped on seven different chromosomes and for the rest the position of the genome is unknown.

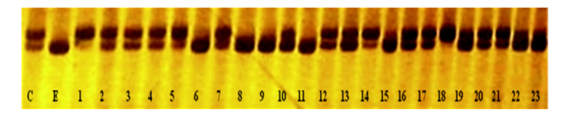


Fig. 2 Pattern of microsatellite bands amplified by the Bmd-16 primer in poliacrilamyde gel; column with the letter C is Carioca MG parent and with the letter E is ESAL 686 parent; following columns are progenies 1 to 23 from F_{8:11} generation and they are indicated by numbers.

In the absence of natural selection, the expected proportions, considering T = 0.005 and S = 0.995 and according to the expressions in Allard (1999) are 0.4938 A^1A^1 , 0.0124 A^1A^2 , and 0.4938 A^2A^2 for F_8 , and for F_{24} they are 0.4975 A^1A^1 , 0.0050 A^1A^2 ,and 0.4975 A^2A^2 , considering homozygous parents. The expected proportions in situations in which one of the parent has a locus in heterozygosis for F_8 are 0.244382 A^1A^1 , 0.011237 A^1A^2 ,and 0.744382 A^2A^2 and for F_{24} are 0.248134 A^1A^1 , 0.003731 A^1A^2 , and 0.748134 A^2A^2 . Considering these expected proportions for the genotypes at each generation, the differences between them are not statistically significant. This therefore shows that natural selection acted on all of the microsatellite loci (Table 1). It is noted that the effect of natural selection occurred in both the first self-fertilized generation up to F_8 and the more advanced generations, since they differed in 19 of the 22 loci.

For the three loci that were not affected by natural selection from F_8 to F_{24} , the markers amplified by primers BMd-33, PVatt003 and PvM15, natural selection was accentuated in the first segregant generations, favoring selection of genotype A^1A^1 , identical to the best adapted parent. Since the primers are mapped in seven different bean chromosomes, it can be inferred that natural selection took place throughout the genome.

An interesting result was the heterozygosis found in the Carioca MG parent for the three primers utilized (BMd-15, BMd-16, and PvM13). In addition to showing that the line (cultivar) were not totally pure, it was observed that even after several generations of self-fertilization, the cultivar maintain genetic variability through heterozygosis (Allard and Hansche, 1964). It is also noted that heterozygosis maintained a much greater frequency than expected for the three loci and the homozygote genotype with the Carioca-derived allele remained superior for two of the markers, especially at F₂₄ (Table 3). For the PvM13 marker, the great majority of the progenies remained in heterozygosis, clearly indicating the adaptive advantage of this genotype or, more precisely, the region of the genome marked by this locus (Table 1).

These observations show the superiority of the Carioca parent in relation to adaptation and also that heterozygosis can be maintained at a high frequency, even after a large number of successive self-fertilizations in a segregant population, when this genotype is favourable.

Since both populations originated from a biparental crossing, and the majority of genitor loci are homozygotes, it can be deduced that the allele frequencies in the segregant loci were 0.5 in generation F_2 . In the case of heterozygote loci in one parent, a frequency of 0.75 can be deduced for the allele that is found in both parents and 0.25 for the allele found in only one parent, as in the case of Carioca MG. The frequencies should remain unaltered in the absence of natural selection. Note that in F_8 (Table 2), in all of the microsatellite loci there was an increase in frequency of the allele coming from the Carioca MG parent in relation to what was expected, even for the loci in which Carioca MG had two

alleles, indicating that natural selection favored the plants that carried these alleles. In locus PvM 13, the allele from the ESAL 686 genitor was more common, although this frequency was much less than expected.

In the F_{24} population (Table 2), all 22 loci were affected by natural selection. In 21 of the loci, the Carioca MG genitor alleles appeared more frequently. For the PvM13 primer, the ESAL 686 parent allele was slightly more common, as observed in F_8 , because of the predominance of the heterozygote resulting in allele frequencies around 0.5 (Table 1 and Table 3). Comparing populations F_8 and F_{24} , their allele frequencies were different in 21 loci, except for PvM13, showing that natural selection was active not only until generation F_8 , but also from F_8 to F_{24} , maintaining the Carioca genitor allele in greater amounts. However, apparently natural selection for loci Pvtaaa003 and PVttc002 was more intense until F_8 and reversed direction from F_8 to F_{24} , while the Carioca allele became even more predominant in this late generation. In this case the first marker did not exhibit heterozygosis and the apparent inverse selection effect took place in the homozygotes, while in the latter, selection effect mainly increased heterozygote frequency. However, in both, the ESAL 686 allele genotype was favoured in more advanced generations.

Table 1. Number of genotypes observed for the amplified microsatellite fragments at F_8 and F_{24} and a comparison of the two populations by means of x^2 .

Primers		F ₈			F ₂₄		x²
11111010	A¹A¹	A^1A^2	A^2A^2	A¹A¹	A^1A^2	A^2A^2	×
BMd-1	63	12	32	60	39	8	28.7672***
BMb-2	80	0	27	105	1	1	28.5212***
BMb-9	46	41	20	87	17	3	35.1353***
BMd-20	66	0	41	95	0	12	21.0915***
BMd-26	67	0	40	105	0	2	42.7762***
BMd-33	103	2	2	106	0	1	2.3764 ^{ns}
BMd-45	59	40	3	73	25	19	16.5826***
Pvtaaa003	96	0	11	69	0	38	19.1158***
PVttc001	88	13	6	103	2	2	11.4133***
PVttc002	86	16	5	68	27	12	7.8002**
PVct002	97	1	9	104	2	1	6.9770**
PVatt003	84	12	11	94	5	13	2.8518 ^{ns}
AJ416395	84	34	9	101	6	0	30.1621***
AJ416402	61	28	18	80	8	20	13.7767***
AF293023.2	88	14	5	104	2	1	12.9999***
AZ301561.1	67	6	34	79	1	27	5.361*
PvM15	91	2	14	98	2	7	2.5926 ^{ns}
PvM21	66	10	31	82	1	24	9.9842**
PvM30	90	7	10	93	14	0	12.3825***
BMd-15 ^b	44	28	35	53	50	4	31.6811***
BMd-16 ^b	50	34	23	74	33	0	27.6606***
PvM13 ^b	2	103	4	0	91	16	32.0000***

ns: P = 0.95; *: $P \le 0.05$; **: $P \le 0.01$; ***: $P \le 0.001$; b: heterozygote loci in the Carioca parent.

Changes in allele frequency could take place because of the sampling effect; however, in the present study each generation came from 2,000 bean plants from the previous generation, minimizing this effect. Nonetheless, when 107 progenies were taken as samples for the two populations, the sampling effect became a little more important since allele frequency oscillation of 0.03 or less could have taken place by chance with a 0.05 probability (Rodrigues and Santos, 2006). Thus the majority of oscillations were mainly due to the effect of natural selection. These effects were specific for each locus and generation, even including reverse selection as was also observed in various monogenetic traits as well as in isozyme loci (Allard and Hansche, 1964; Allard, 1999). In general, these authors verified that natural selection increased the frequency of the alleles that favour adaptation to each environment. In relation to reverse selection seen in loci Pvtaaa003 and PVttc002, it could have occurred due to selection dependent on allele frequency when the adaptability of the genotype that is less common increases, as observed in Allard (1999). This phenomenon is considered important in maintaining variability in the population. An equivalent effect also occurred and corresponded to maintenance of excess heterozygotes in some loci.

Table 2. Allelic frequency estimates for F₈ and F₂₄ generations.

	F ₈		F ₂₄		
Primers	Carioca MG allele	Esal 686 allele	Carioca MG allele	Esal 686 allele	
BMd-1	0.6449	0.3551	0.743	0.257	
BMb-2	0.7477	0.2523	0.986	0.014	
BMb-9	0.6215	0.3785	0.8925	0.1075	
BMd-20	0.6168	0.3832	0.8879	0.1121	
BMd-26	0.6262	0.3738	0.9813	0.0187	
BMd-33	0.972	0.028	0.9907	0.0093	
BMd-45	0.7383	0.215	0.7991	0.2944	
Pvtaaa003	0.8972	0.1028	0.6449	0.3551	
PVttc001	0.8832	0.1168	0.972	0.028	
PVttc002	0.8785	0.1215	0.7617	0.2383	
PVct002	0.9112	0.0888	0.9813	0.0187	
PVatt003	0.8411	0.1589	0.9019	0.1449	
AJ416395	0.9439	0.243	0.972	0.028	
AJ416402	0.7009	0.2991	0.785	0.2243	
AF293023.2	0.8879	0.1121	0.9813	0.0187	
AZ301561.1	0.6542	0.3458	0.743	0.257	
PvM15	0.8598	0.1402	0.9252	0.0748	
PvM21	0.6636	0.3364	0.771	0.229	
PvM30	0.8738	0.1262	0.9346	0.0654	
BMd-15 ^b	0.5421	0.4579	0.729	0.271	
BMd-16 ^b	0.6262	0.3738	0.8458	0.1542	
PvM13 ^b	0.4907	0.5093	0.4252	0.5748	

^b: Heterozygote loci in the Carioca MG parent.

The predominance of natural selection favoring alleles of the Carioca MG parent agrees with the fact that the majority of the area cultivated with common bean in Brazil is planted with this type and indicates not just its better acceptance but also its better adaptability. High productivity of the Carioca also extends to various other countries because of its greater tolerance to acid soils, especially in the case of Brazil (Singh, 1992).

The relative fitness coefficients (ω_1) were estimated for the homozygote genotypes related to each primer pair for better understanding the effect of natural selection. For example, considering "A¹" to be the allele from the Carioca MG parent and "A²" the allele from the ESAL 686 parent, ω_2 was estimated for A¹A¹ and ω_3 for A²A². The relative fitness coefficient of the A¹A² heterozygote, ω_2 , was considered to be equal to 1.0 (Allard and Hansche, 1964). Estimates of ω_1 and ω_3 equal to 1.0 indicate that natural selection did not take place for these genotypes, that is, they had the same reproductive efficiency as the heterozygote (Allard and Hansche, 1964). Estimates of ω_1 and ω_3 less than 1.0 indicate that natural selection took place, reducing the frequency of these genotypes in relation to the heterozygote, which was more adapted and inversely, estimates larger than 1.0 indicate that selection increased the frequency of the homozygotes in relation to the heterozygote, which in this case is less adapted (Allard, 1999).

Table 3. Estimate of the accumulated relative fitness coefficients ω_1 and ω_3 , per locus, in the F₈ and F₂₄ generations of a *P. vulgaris* cultivar Carioca MG and ESAL 686 cross.

	F ₈		F:	24
Primers	ω ₁	ω3	ω ₁	ω3
BMd-1	0.1327	0.0674	0.0054	0.0034
BMb-2	-	-	0.3719	0.0165
BMb-9	0.0284	0.0123	0.0181	0.0029
BMd-20	-	-	-	-
BMd-26	-	-	-	-
BMd-33	1.3018	0.0253	-	-
BMd-45	0.0373	0.0019	0.0103	0.0125
Pvtaaa003	-	-	-	-
PVttc001	0.0896	0.0080	0.1824	0.0165
PVttc002	0.1359	0.0079	0.0089	0.0073
PVct002	2.4519	0.2275	0.1842	0.0083
PVatt003	0.1769	0.0232	0.0666	0.0429
AJ416395	0.0624	0.0067	0.0596	0.0000
AJ416402	0.0551	0.0162	0.0354	0.0413
AF293023.2	0.1589	0.0090	0.1842	0.0083
AZ301561.1	0.2823	0.1432	0.2798	0.4458
PvM15	1.1501	0.1769	0.1736	0.0578
PvM21	0.1668	0.0784	0.2905	0.3963
PvM30	0.3250	0.0361	0.0235	0.0000
BMd-15	0.0397	0.0316	0.0038	0.0013
BMd-16	0.0372	0.0171	0.0079	0.0000
PvM13b	0.0005	0.0010	0.0000	0.0029
ω ₁ s/heter	0.3685	0.0494	0.1059	0.05912

Accumulated Relative fitness coefficients for A^1A^1 (ω_1) from F_7 to F_8 (Table 3) varied from 0.0005 to 2.4519 and for A^2A^2 (ω_3) varied from 0.0010 to 0.227. The value zero occurs when no locus of the evaluated genotype is observed, as in the case of the PvM13 primer ($A^1A^1 = 0$ and $\omega_1 = 0$).

It is important to remember that these accumulated coefficients refer to the effect of natural selection on the homozygotes from generation F_2 to F_8 , during six generations. It is noted that there is a larger variation of ω_1 when compared with ω_3 , which shows a larger oscillation of the relative fitness coefficients in the different loci, with values higher than 1. These values higher than one show that for the BMd-33, PVct002 and PvM15 loci, the homozygote with the highest value adapted better than the heterozygote; however, for the rest of the loci, the ω_1 and ω_3 estimates were less than 1.0, showing that heterozygote combination was superior.

Average ω_1 accumulated fitness coefficient estimates (0.368468) were higher than ω_3 (0.0494349), showing that selection was more intense on the homozygote for the allele from the ESAL 686 line (A^2A^2) . These estimates, just as those obtained for each locus, confirm better adaptation for the majority of the homozygotes for alleles from the Carioca MG parent.

Table 4. Simple linear regression analysis for the grain yield in common bean, considering each marker separately in each season per generation, and the joint analysis of seasons in F₈ and F₂₄ generations.

Generations	Season	Primers	Pr > F	R² (%) adj
Generation F ₈		BMd-1	0.0063	6
	Winter	BMd-9	0.0068	5.88
		AF293023.2	0.0485	2.74
	Rainy	PvM30	0.024	3.85
	Dry	Pvtaaa003	0.0314	3.42
		PvM30	0.0417	2.97
	Joint	PvM30	0.0189	4.23
	Winter	-	-	-
		Pvtaaa003	0.011	5.1
	Rainy	AJ416402	0.0159	4.51
		BMd-45	0.0012	8.71
		PVttc002	0.0002	11.76
Generation F ₂₄		BMd-20	0.0422	2.96
		Pvtaaa003	0.0446	2.87
	Dry	AJ416395	0.0249	3.79
		PVttc002	0.0331	3.34
		PvM21	0.0193	4.2
	Joint	Pvtaaa003	0.0018	7.99
		PVttc002	0.0218	4

Due to the absence of heterozygote individuals, it was not possible to estimate accumulated fitness coefficient ω_1 and ω_3 for the following loci: BMb-2, BMd-20, BMd-26, and Pvtaaa003.

For the accumulated relative fitness coefficients in F_{23} to F_{24} , there was a variation from zero to 0.3719378 for ω_1 and a variation from zero to 0.4458374 for relative fitness coefficient ω_3 (Table 3). In this case, ω_1 and ω_3 estimates include the effect of natural selection on the homozygotes from generation F_8 to F_{24} , during 16 generations, 2.67 times the number of generations when compared to the estimates for F_8 .

Taking note that a greater amplitude of ω_3 estimates (0.4458374) when compared to those of ω_1 (0.3719378), imply greater oscillation of the relative fitness coefficients to A^2A^2 in the different loci.

Comparing ω_1 and ω_3 averages of F_{24} and F_8 , the same effect of natural selection is observed though at a lower intensity, probably because the more extreme genotypic frequencies in F_{24} than in F_8 . This reduced intensity can also be explained by reverse selection in some loci and maintenance of the intense heterozygosis in other loci; natural selection depending on low genotypic frequency may even have taken place.

Considering both generations, the effect of natural selection was more accentuated in the first segregant generations, as Allard (1999) observed, also because the genotypic frequencies were more similar due to the greater frequency of unfavourable alleles.

Also in F_{24} , due to the absence of heterozygotes, it wasn't possible to estimate the accumulated relative fitness coefficients for the following primers: BMd-20, BMd-26, BMd-33, and Pvtaaa003.

Despite not having detected heterozygotes in 4 loci of F_8 and 4 loci of F_{24} , the expected heterozygote frequency without natural selection is 0.0124 at F_8 and 0.005 at F_{24} , starting with pure and contrasting parents. When one of the parents is a heterozygote, the expected heterozygote frequency is 0.011237 at F_8 and 0.003731 at F_{24} . Such frequencies happen because the species mainly reproduces by self-fertilization in the environmental conditions in which the populations were evaluated. Therefore the frequencies of loci in which heterozygotes were observed in a sample of 107 plants were still higher with loci BMd-1, BMb-9, BMd-45, PVttc002, BMd-15, BMd-16, and PvM13 and especially at F_{24} , again showing the superiority of heterozygote adaptation in the genomic regions around these loci (Table 1).

According to Allard and Hansche (1964), the effect of natural selection favouring maintenance of heterozygotes contributed to retain genetic variability in the population. The populations utilized in this work were evaluated for grain yield in the progenies at various generations, and showed a far larger gain with natural selection than the artificial selection (Gonçalves et al. 2001; Corte et al. 2002; Silva et al. 2004). Therefore the increase in grain yield due to natural selection even after many generations of self-fertilization is the result of the better adaptive value of the loci in heterozygosis for this trait. Consequently, it can be inferred that the high number of microsatellite heterozygote loci at advanced generations of self-fertilization must also refer to regions of the genome that contribute to better adaptation, especially the alleles of the Carioca MG parent. So the microsatellite fragments chosen by natural selection can be utilized to perform assisted selection, as suggested by Allard (1999) and Rodrigues and Santos (2006). Thus homozygote genotypes chosen by natural selection could contribute to better adaptability in lines to be selected.

Table 5. Multiple regression analysis with model selection through the backward method for grain yield in common bean in each season per generation and joint analysis of seasons in F_8 and F_{24} generations.

Generations	Season	Primer	P > 0,05	R² (%)
	Winter	BMd-1+ BMd-9	0.0016	11.63
F ₈	Rainy	BMd-26 + BMd-45	0.015	7.76
	Dry	PvM15 + PVctt002	0.0196	7.28
	Joint	0	0	0
	Winter	0	0	0
	Rainy	BMd-26 + AJ416402 +	< 0001	30.74
F ₂₄		BMd-45+ PVttc002		
	Dry	AJ416395+PVttc002	0.0008	14.95
		PvM21		
	Joint	Pvtaaa003+ BMd-45	0.0005	15.87
		AJ416395		

Therefore it was verified among the SSR fragments if some are associated to grain yield. The results of simple and multiple linear regression analysis utilizing the backward model for the grain yield are on Table 4 and Table 5, respectively.

The adjusted determination coefficient values (R^2) obtained by the markers considering simple linear regression were low, varying from 2.74% to 11.76%, indicating that these markers explain a reduced part of the phenotypic variation (Table 4). Note that in the backward method, some markers identified by linear regression weren't identified in multiple regression and vice versa. However, this result was expected since the method only selects the minimum number of markers needed to explain the phenotypic variation. With this procedure, the amount of phenotypic variation explained was equally low. This fact is common; as stated in Bernardo (2008), 72% of the QTLs mapped in various studies for quantitative traits such as grain yield explained less than 10% of phenotypic variation, corroborating the present result. It is important to remember that the markers used were selected by natural selection during the generations of bulk populations and their efficiency can be reduced to identify QTLs. However, the deviation from the expected segregation is not always prejudicial (Hackett and Broadfoot, 2003).

QTLs interacted heavily with the environment, as can be seen when there is a differential expression of QTLs in relation to environmental changes, or when there is no QTL expression in some of the environments evaluated (Campbell et al. 2003; Torga et al. 2010). This interaction can be verified by observing that in different harvests in the two generations, different markers were found that explained some of the variation of grain yield. It was also observed that in the winter evaluation of the F_{24} progenies, there was no marker that could explain part of grain yield. Also, this interaction may explain the results of joint multiple regression analysis of F_8 at which no marker identified QTL and of F_{24} three markers together explained 15.84% of grain yield variation (Table 5). It is again necessary to remember that this apparent interaction could also be the selection effect that altered the frequency of the markers in both generations.

Table 6. Average common bean yield in Kg ha⁻¹ of each genotypic class for the significant markers using the backward method in the three seasons in each generation evaluated, and in the joint analysis.

Generations	Primers		Average		
		Carioca MG	Heterozygote	Esal-686	Seasons
	BMd-1	4917	4188	4402	Winter
	BMd-9	4885	4702	4169	Winter
Generation F ₈	BMd-45	1279	1291	1775	Rainy
	BMd-26	1389	-	1206	Rainy
	PvM15	2290	2323	2597	Dry
	PVttc002	2361	1577	2089	Dry
	BMd-26	1881	-	1526	Rainy
	AJ416402	1922	1880	1687	Rainy
	AJ416402	3156	3298	3150	Joint
	BMd-45	1816	1784	2193	Rainy
	BMd-45	3137	3130	3305	Joint
	PVttc002	1999	1622	1744	Running
Generation F ₂₄	PVttc002	2710	2565	2534	Dry
	AJ416395	2678	2333	-	Dry
	AJ416395	3174	3013	-	Joint
	PvM21	2708	2582	2518	Dry
	Pvtaaa003	3242	-	3026	Joint

The fact that QTLs interacted with the environment could also be explained by the variation in R² estimates for the same marker in different evaluations, as happened for the PVttc002 marker with R2 values varying from 0 to 11.76%. The magnitudes and the instability of QTL expression detected confirmed the difficulty of assisted selection since it is generally done for the purpose of utilizing the lines in several environments.

Of the markers affected by natural selection and chosen by the backward method, in each season and at each generation, some contributed to increased yield and others reduced expression of this trait (Table 6). For example, BMd-1 and BMd-9 together contributed 11.63% toward increase grain yield in the winter at generation F₈ and BMd-45 and PvM-15 reduced yield in the rainy and dry seasons, respectively.

At generation F₂₄, in winter, no marker was identified, but in the rainy season four markers, BMd-26, AJ416402, BMd-45 and PVttc002 were chosen that individually explained a small part of the yield variation, but together explained 30.74% of the variation, with three of them increasing yield and BMd-45 reducing it. It is worth stressing the high level of interaction among QTLs and environments, since these same primers did not explain the same amount of variation in any other season. However, in the joint analysis two other markers along with BMd-45 explained a smaller amount of yield variability. The high level of interaction for the grain yield QTLs with the environment has been detected many times including in common bean (Pereira et al. 2007; Rodrigues et al. 2007; Torga et al. 2010).

Therefore, natural selection acted in all of the 22 microsatellite loci and the frequency of the best adapted parental alleles increased in 22 of the loci in F₈ and 19 loci in F₂₄. The intensity of natural selection was different for each locus and each generation. Besides, there are markers influenced by natural selection that explain the effect of QTLs of grain yield, although the expression of the QTLs is generally unstable and interacts strongly with the environments, seasons and generations, and could have been increased by natural selection.

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