Selection strategies for marker-assisted background selection with chromosome-wise SSR multiplexes in pseudo-backcross programs for grapevine breeding

E. HERZOG¹⁾, R. TÖPFER²⁾, L. HAUSMANN²⁾, R. EIBACH²⁾ and M. FRISCH¹⁾

¹⁾ Institute of Agronomy and Plant Breeding II, Justus Liebig University, Giessen, Germany

²⁾ Julius Kühn Institut (JKI), Federal Research Centre for Cultivated Plants, Institute for Grapevine Breeding Geilweilerhof, Siebeldingen, Germany

Summary

Organizing SSR markers located on one chromosome into PCR multiplexes has the potential to reduce the costs of marker analysis. The optimal selection strategies for such chromosome-wise multiplexes have not yet been investigated. We investigated with computer simulations three different selection strategies for gene introgression with a pseudo-backcross scheme and a marker density of one marker every 10 cM. Selecting individuals with the highest number of chromosomes carrying *V. vinifera* alleles at all background marker loci reduced the number of required multiplexes by 7.24-7.87 % in generations pBC_4 - pBC_6 for population sizes $n_t = 150-300$ individuals per pseudo-backcross generation.

K e y w o r d s : gene introgression, simulation study, multiplex PCR, microsatellite, marker-assisted selection.

Introduction

American and Asian Vitis species carrying resistance genes against mildew disease have been employed in interspecific breeding programs (DI GASPERO and CATTONARO 2010, TÖPFER et al. 2011). Along this line only one example has been described for a systematic development of introgression lines as described by the pioneering work of Alain Bouquet (PAUQUET et al. 2001). This work turns out to be very time consuming as well as space and labor demanding. The development of molecular markers for early selection of seedlings with traits of agronomic interest is therefore of particular value (EIBACH et al. 2007). Simple sequence repeats (SSRs) are useful genetic markers, as they are abundant in the genome, highly polymorphic and transferable between *V. vinifera* and related species (SALMASO *et al.* 2008, VEZZULLI et al. 2008, BLANC et al. 2012). However, they have the disadvantage of low throughput compared to single nucleotide polymorphisms (SNPs), which limits their use in large-scale breeding programs. Organizing SSRs into PCR multiplexes considerably reduces the costs of marker analysis (MERDINOGLU et al. 2005). PATOCCHI et al. (2009) suggested that organizing SSR markers located on one linkage group in one multiplex is applicable and advantageous. Efficient selection strategies for *V. vinifera* genome recovery in pseudo-backcross programs for gene introgression with such chromosome-wise multiplexes have not yet been investigated. The objective of our study was to compare with computer simulations different selection strategies for gene introgression with chromosome-wise multiplexes.

Material and Methods

Computer simulations were carried out assuming no interference in crossover formation. Each simulation was run 10,000 times in order to reduce sampling effects and to obtain stable results with small standard error.

The genetic model for grapevine consisted of 19 chromosomes (2 x 40 cM, 7 x 60 cM, 5 x 80 cM, 5 x 100 cM). This corresponded to a total genome length of 1400 cM. The marker for the dominant target gene was assumed to be a gene-based marker and was located on a 100 cM chromosome at 61 cM from the telomere. Background markers were equidistantly spaced with one marker every 10 cM, the first and last marker of each chromosome being placed on the telomeres.

A pseudo-backcross scheme with changing *V. vinifera* parents in every generation was investigated up to generation pBC_6 . The goal was to recover as much *V. vinifera* genome as possible, irrespective from which of the parents. For chromosome-wise multiplexes it was assumed that one multiplex included genotyping all background marker loci located on one chromosome. This resulted in multiplexes comprising 5 to 11 SSRs. In advanced pseudo-backcross generations only those chromosomes were analyzed which did not yet carry *V. vinifera* alleles at all background marker loci.

The donor was heterozygous for the desired allele at the target locus. The *V. vinifera* parents could be distinguished from the donor at all marker loci. Initially, the donor and the first *V. vinifera* parent were crossed to produce n_{F1} F₁ individuals. From this F₁ population, one individual that carried the donor allele at the target locus was selected as parent for generation pBC₁. This individual was crossed to the second *V. vinifera* parent to produce n_1 pBC₁ individuals. From this pBC₁ population, one best individual was selected with the selection strategies described below (see

Correspondence to: Dr. M. FRISCH, Institute of Agronomy and Plant Breeding II, Justus Liebig University, Heinrich-Buff-Ring 26-32, 35392 Giessen, Germany. E-mail: matthias.frisch@uni-giessen.de

also Fig. 1), and crossed to the next *V. vinifera* parent. This procedure was repeated for t = 6 pseudo-backcross generations with constant population sizes $n_{\text{F1}} = n_{\text{t}} = 50, 100, 150, 200, 250, 300.$

For all selection strategies, carriers of the donor allele at the target locus were pre-selected in the first selection step. These individuals were then subjected to one of three genome-wide background selection strategies. For Strategy 1, a selection index $i = \sum_{m} x_m$ was created, where summation is over background markers and x_m is the number of *V. vinifera* alleles at the *m*th marker. One individual with the highest value of *i* was selected in the second selection step and crossed to the next *V. vinifera* parent of generation pBC_{t+1}. The individual with the highest proportion of *V. vinifera* alleles at background marker loci was thus selected as parent for the next generation pBC_{t+1}.

For Strategy 2, a selection index $j = \sum_{c xc}$ was created, where summation is over chromosomes and $x_c = 1$ if a chromosome carries *V. vinifera* alleles at all background marker loci. All individuals with the highest value of *j* were selected in the second selection step. For these individuals, the value of *i* was determined as described for Strategy 1 in the third selection step. One individual with the highest value of *i* was selected and crossed to the next *V. vinifera* parent of generation pBC_{t+1}. The best individual with the highest number of chromosomes carrying *V. vinifera* alleles at all background marker loci was thus selected as parent for the next generation pBC_{t+1}.

For Strategy 3, a selection index $k = \sum_{cxc}$ was created, where summation is over chromosomes and xc = length of chromosome c in M if a chromosome carries V. vinifera alleles at all background marker loci. All individuals with the highest value of k were selected in the second selection step. For these individuals, the value of i was determined as described for Strategy 1 in the third selection step. One individual with the highest value of i was selected and crossed to the next V. vinifera parent of generation pBC_{t+1}. The best individual with the highest cumulative length of chromosomes carrying V. vinifera alleles at all background marker loci was thus selected as parent for the next generation pBC_{t+1} .

To quantify the success of the respective pseudo-backcross programs, the 10th percentile (Q_{10}) , the arithmetic mean (\bar{x}) and the standard deviation (s_x) of the frequency distribution of *V. vinifera* genome in percentage, the average number of chromosomes carrying *V. vinifera* alleles at all background marker loci, and the average number and length of donor fragments were determined in every backcross generation for the selected individuals. In addition, the required number of chromosome-wise multiplexes for the respective pseudo-backcross programs, and the number of individuals n_i selected for evaluation with selection index *i* were determined in every backcross generation.

Results and Discussion

Up to generation pBC₃, Strategy 2 required a higher resource input than Strategy 1 for recovering equivalent levels of *V. vinifera* genome (Fig. 2). Strategy 3 was always inferior to Strategy 2. A $Q_{10} \ge 98$ % required $n_t = 100$ individuals per generation, 1753 multiplexes and three pseudo-backcross generations with Strategy 1 (Fig. 1). With Strategy 2, a $Q_{10} \ge 98\%$ required $n_t = 200$ individuals and 3162 multiplexes. With Strategy 3, a $Q_{10} \ge 98$ % required $n_t = 250$ individuals and 3947 multiplexes, or an additional pseudo-backcross generation.

In generations pBC₁-pBC₃, pre-selection for chromosomes carrying *V. vinifera* alleles at all background marker loci considerably reduced the number of individuals n_i from which the parent for the next generation was selected (Tab. 1). With Strategy 2 only $n_i = 1.8$, 2.4, 10.5 individuals were evaluated for selection index *i*. For Strategy 1, n_i were all individuals carrying the target gene (Tab. 1). These individuals have on average more, but shorter donor fragments than those selected with Strategy 2, and the probability that an individual with a higher overall proportion of *V. vinifera* genome is selected is higher than for Strategy 2.

Strategy 1	Strategy 2	Strategy 3						
Selection of best individual with								
highest proportion of background markers for <i>V. vinifera</i> parent	highest number of chromosomes carrying <i>V. vinifera</i> alleles at all background markers	highest cumulative length of chromosomes carrying <i>V. vinifera</i> alleles at all background markers						
Number of plants and multiplexes for selection up to pBC_3 at $Q_{10} \ge 98$ %								
<i>n₁</i> = 100 plants 1753 multiplexes	n, = 200 plants 3162 multiplexes	n, = 250 plants 3947 multiplexes						

Fig. 1: Strategies and effort required to get $Q_{10} \ge 98$ % of *V. vinifera* genome in generation pBC₃. Number of plants per generation and number of chromosome-wise multiplexes are indicated for each strategy.

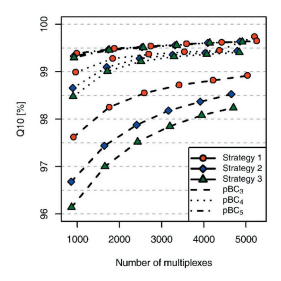


Fig. 2: Tenth percentile Q_{10} values of *V. vinifera* genome recovered in generations pBC₃-pBC₅ for Strategies 1, 2, 3 with constant population sizes $n_t = 50, 100, 150, 200, 250, 300$ individuals per pseudo-backcross generation plotted against the required number of chromosome-wise multiplexes.

In generations pBC₄-pBC₆, the differences in *V. vini-fera* genome between the selection strategies disappeared (Fig. 2 and Tab. 1). For population sizes of $n_t \ge 150$ individuals per pseudo-backcross generation, the differences in Q_{10} values were < 0.1% between Strategy 1 and 2 (Fig. 2). Moreover, Strategy 1 then required more chromosome-

wise multiplexes for equivalent levels of V. vinifera genome than Strategy 2 (Tab. 2). For $n_t = 150-300$ individuals per pseudo-backcross generation, 7.24-7.87 % of multiplexes were saved in generations pBC₄-pBC₆ with Strategy 2 compared to Strategy 1 (Tab. 2). In generation pBC₄, all non-carrier chromosomes carried V. vinifera alleles at all background marker loci on average, and selection focused on individuals with the shortest donor segment around the target gene for all three strategies (Tab. 1). The number of required multiplexes was then increasing at approximately the same rate for Strategy 1 and Strategy 2. However, Strategy 2 was more efficient than Strategy 1 in selecting for chromosomes which carried V. vinifera alleles at all background marker loci in generations pBC1-pBC3. While for Strategy 2, 9.3, 15.6, and 18.0 chromosomes on average carried *V. vinifera* alleles at all background marker loci, only 8.1, 14.5, 17.6 chromosomes carried V. vinifera alleles at all background marker loci with Strategy 1. These early savings resulted in an overall saving of chromosome-wise multiplexes in advanced pseudo-backcross generations.

Increasing the number of individuals per pseudo-backcross generation from $n_t = 150$ to $n_t = 300$ resulted in an additional *V. vinifera* genome recovery of 1.2-0.7 % for Strategy 1 in generations pBC₁-pBC₃ (data for pBC₁ and pBC₂ not shown, for pBC₃ see Fig. 2). In contrast, increasing the number of individuals per pseudo-backcross generation beyond $n_t = 150$ -200 had little effect on *V. vinifera* genome recovery in advanced backcross generations for both Strategy 1 and Strategy 2 (Tab. 2, see also Fig. 2).

Table 1

Recovered level of *V. vinifera* genome $(Q_{10}, \overline{x}, s_x)$ in percentage, required number of chromosome-wise multiplexes (CM), average number of chromosomes carrying *V. vinifera* alleles at all background marker loci (CCV), number of individuals evaluated for selection index *i*, number (\overline{x}, s_x) and length (\overline{x}, s_x) of donor fragments (cM) in generations pBC₁-pBC₆ for Strategies 1, 2, 3 with constant population size $n_t = 150$ individuals per pseudo-backcross generation

V. vinifera genome								No. donor		Length donor	
Strategy pBC _t		(%)		СМ	CM CCV		fragments		fragments (cM)		
	-	Q_{10}	\overline{x}	S_x				\overline{x}	S_x	\overline{x}	S_x
1	pBC_1	82.24	84.59	1.92	1425	8.1	75.0	12.9	2.2	33.55	26.22
	pBC ₂	94.90	96.24	1.04	2246	14.5	74.9	5.8	1.8	18.14	17.62
	pBC ₃	98.55	99.14	0.43	2585	17.6	75.0	2.2	1.1	10.91	9.71
	pBC_4	99.37	99.64	0.20	2692	18.4	75.0	1.4	0.6	7.05	5.08
	pBC ₅	99.54	99.75	0.15	2740	18.7	74.9	1.3	0.5	5.51	3.79
	pBC ₆	99.64	99.80	0.13	2761	18.9	74.9	1.2	0.4	4.91	3.26
2	pBC ₁	79.10	82.75	2.86	1425	9.3	1.8	11.5	1.7	41.96	29.21
	pBC_2	92.89	95.07	1.66	2152	15.6	2.4	4.7	1.5	29.40	26.45
	pBC ₃	97.88	98.82	0.69	2407	18.0	10.5	1.8	0.9	18.69	18.39
	pBC ₄	99.28	99.60	0.24	2481	18.3	54.9	1.4	0.6	8.25	6.30
	pBC₅	99.52	99.74	0.16	2534	18.7	47.9	1.2	0.5	5.84	4.02
	pBC ₆	99.63	99.79	0.13	2558	18.8	61.9	1.2	0.4	5.06	3.34
3	pBC ₁	78.29	82.19	3.06	1425	9.2	1.1	11.6	1.7	43.04	28.95
	pBC_2	92.02	94.47	1.87	2163	15.5	1.2	4.8	1.5	32.23	27.28
	pBC ₃	97.52	98.63	0.80	2428	18.0	8.9	1.8	0.9	20.95	20.68
	pBC_4	99.22	99.57	0.27	2506	18.3	55.3	1.4	0.6	8.67	6.88
	pBC ₅	99.51	99.74	0.16	2560	18.6	48.2	1.2	0.5	5.90	4.03
	pBC ₆	99.62	99.79	0.13	2586	18.8	61.4	1.2	0.4	5.10	3.35

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Table 2

Recovered level of *V. vinifera* genome (Q_{10}) and percentage of saved chromosome-wise multiplexes (CM (%)) for Strategy 2 compared to Strategy 1 in generations pBC₄-pBC₆ with constant population sizes $n_i = 150, 200, 250, 300$

Generation	$n_t = 150$		$n_t = 200$		$n_t = 250$		$n_t = 300$	
	Q_{10}	CM (%)						
pBC_4	99.28	7.84	99.36	7.83	99.40	7.63	99.44	7.87
pBC ₅	99.52	7.52	99.56	7.50	99.61	7.33	99.64	7.52
pBC ₆	99.63	7.35	99.67	7.37	99.68	7.24	99.68	7.47

We conclude that if SSR markers are analyzed as chromosome-wise multiplexes, selecting for individuals with the highest proportion of *V. vinifera* genome at background marker loci is the most efficient selection strategy for short gene introgression programs of up to three pseudo-backcross generations. For such short gene introgression programs, population sizes of $n_t \ge 300$ individuals per pseudobackcross generation maximize *V. vinifera* genome recovery. For gene introgression programs of four to six pseudobackcross generations, pre-selecting individuals with the highest number of chromosomes carrying *V. vinifera* alleles at all background marker loci has the potential to considerably reduce the number of required SSR multiplexes. For these longer gene introgression programs, population sizes of $n_t = 150-200$ individuals are sufficient.

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