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Quantitative trait loci from the host genetic background modulate the durability of a resistance gene: a rational basis for sustainable resistance breeding in plants.

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Running title:

QTLs for durability of plant resistance to pathogen

Abstract:

The combination of major resistance genes with quantitative resistance factors is hypothesized as a promising breeding strategy to preserve the durability of resistant cultivar, as recently observed in different pathosystems. Using the pepper (*Capsicum annuum*)/Potato virus Y (PVY, genus *Potyvirus*) pathosystem, we aimed at identifying plant genetic factors directly affecting the frequency of virus adaptation to the major resistance gene *pvr2*³ and at comparing them with genetic factors affecting quantitative resistance. The resistance breakdown frequency was a highly heritable trait ($h^2=0.87$). Four loci including additive quantitative trait loci (QTLs) and epistatic interactions explained together 70% of the variance of *pvr2*³ breakdown frequency. Three of the four QTLs controlling *pvr2*³ breakdown frequency were also involved in quantitative resistance, strongly suggesting that QTLs controlling quantitative resistance have a pleiotropic effect on the durability of the major resistance gene. With the first mapping of QTLs directly affecting resistance durability, this study provides a rationale for sustainable resistance breeding. Surprisingly, a genetic trade-off was observed between the durability of PVY resistance controlled by *pvr2*³ and the spectrum of the resistance against different potyviruses. This trade-off seemed to have been resolved by the combination of minor-effect durability QTLs under long term farmer selection.

Keywords: resistance durability, quantitative trait locus, resistance breakdown, *Capsicum spp.*, PVY, eukaryotic translation initiation factor 4E, potyviruses, resistance spectrum

INTRODUCTION

The use of resistant cultivars provides efficient, economical and environmentally friendly crop protection against plant pathogens, and many cultivars carrying major resistance genes successfully protected cultivated crops against their main pathogens. When available in genetic resources, major resistance genes are very attractive for breeding because of their simple inheritance that makes rapid and cheap their introgression through backcrosses to high-yielding but susceptible cultivars, and because they are expected to confer a nearly-complete resistance against the targeted pathogen. These resistances involve a gene-for-gene interaction and are usually race-specific. The widespread deployment of such resistance genes in many elite cultivars imposes a strong selection pressure on the pathogen population leading to the appearance and/or the increase in frequency of pathogen variants overcoming the resistance (or resistance breaking (RB) variants). This adaptation of the target pathogen makes the resistance ineffective, losing its economical interest. In order to preserve the durability of cultivar resistance, management strategies were developed to handle the adaptation of the pathogen populations and limit the appearance of RB variants. When several major resistance genes are available to control a given pathogen species, pyramiding of the distinct genes into a same cultivar, alternation of resistance genes across cultivation cycles or association of cultivars with distinct resistance genes during the same cultivation cycle were proposed (Wolfe, 1985; Mundt *et al.*, 2002; Pink, 2002; McDonald and Linde, 2002).

More recently, another strategy was suggested from experimental data showing that the durability of a major resistance gene was dependent on the plant genetic background in which it was introgressed. In 3 distinct pathosystems (plant-virus/fungus/nematode) the emergence and/or frequency of the pathogen variants breaking down the major resistance gene remained low or did not occur in the cultivars which combined the major resistance gene in a quantitatively resistant genetic background, but increased when the resistance gene was introgressed into a plant genotype with a susceptible genetic background (Palloix *et al.*, 2009; Brun *et al.*, 2010; Fournet *et al.*, 2013). These results raise the hypothesis that the selective value of allelic variation at a resistance locus depends on the genetic background. It addresses the question of how genetic background impacts the effect of mutations and evolution (Chandler *et al.*, 2013). Considering plant breeding practices it suggests that combining qualitative and quantitative resistance increases the durability of plant protection. This also raises a paradox, since the increase in resistance efficiency is expected to increase the selection pressure on the pathogen population and thus the risk to select adapted pathogen genotypes (Poland *et al.* 2009). However, none of the pathosystem studies cited above formally demonstrated that the increase in durability resulted from the action of quantitative resistance factors, since the durability of the major resistance was compared between pairs of genotypes with fully different genetic backgrounds. Consequently the decrease in resistance breakdown frequency might be due to any other genes from those genetic backgrounds. Further on, Quenouille *et al.* (2013) showed that the breakdown frequency of the major resistance allele *pvr2*³ controlling *Potato virus Y* (PVY) was correlated to the quantitative resistance level in a set of pepper genotypes carrying *pvr2*³ and differing in their genetic background. This suggested that the breakdown frequency of the major allele and the quantitative resistance shared a common genetic basis. When very few major resistance genes are available, this would open the possibility to preserve their durability. Moreover, it would provide tools for breeders to construct resistant cultivars with enhanced durability.

In our study, we (i) examined whether resistance breakdown is a heritable trait, (ii) described the genetic architecture underlying the genetic background which affect the breakdown frequency of a major resistance gene and (iii) discussed how breeders can exploit these results to preserve the durability of major resistance genes. Using the pepper (*Capsicum annuum*)/*Potato virus Y* (PVY) system, our strategy consisted in (i) detecting QTLs affecting the breakdown frequency of the *pvr2*³ major resistance allele, since breakdown frequency is a key determinant of resistance durability and

(ii) comparative mapping between these QTLs and QTLs affecting the quantitative resistance to PVY.

MATERIALS AND METHODS

Mapping population

A segregating doubled-haploid (DH) population comprising 350 lines was obtained from the F₁ hybrid between two *Capsicum annuum* inbred lines: ‘Perennial’ carrying the PVY major resistance allele *pvr2*³ in a partially resistant genetic background is an inbred line issued from an Indian landrace and ‘Yolo Wonder’ carrying the PVY susceptibility allele *pvr2*⁺. All DH lines were genotyped with a tetra-primer ARMS-PCR (amplification refractory mutation system – polymerase chain reaction) marker which targets SNP (single nucleotide polymorphism) signatures differentiating *pvr2*⁺ from *pvr2*³ (Rubio *et al.*, 2008). For phenotyping, we used the 153 DH lines carrying the *pvr2*³ allele but segregating for the genetic background.

Virus material

For evaluation of RB frequencies, we used a PVY chimera between PVY clones SON41p and LYE84.2 (Montarry *et al.*, 2011). This PVY chimera, named “CI chimera”, differs from SON41p only by the CI cistron, which was replaced by that of LYE84.2. The rest of the genome and more particularly the region directly involved in *pvr2*³ RB (the VPg cistron) are strictly identical between the two PVY clones. This “CI chimera” was chosen because of its higher ability to break the *pvr2*³ resistance down compared to the parental clones (Montarry *et al.*, 2011). However, the breakdown capacity of the “CI chimera” was shown to be highly correlated with that of SON41p when inoculated to a large range of pepper genotypes (Quenouille *et al.*, 2013). The “CI chimera” and SON41p were not infectious *per se* in plants carrying *pvr2*³, i.e. in these plants, the “CI chimera” showed very weak accumulation detected through RT-PCR but not ELISA and suggesting that the RB mutation occurred in the resistant plant (Montarry *et al.*, 2011) whereas mutants of the “CI chimera” possessing single non-synonymous substitutions in the VPg cistron could be detected at high concentration in systemically infected leaves. In addition, the CI chimera showed the same distributions of RB mutations when inoculated on plants carrying *pvr2*³ when compared to SON41p. The level of quantitative resistance controlled by the plant genetic background was evaluated using a single-nucleotide mutant of the “CI chimera” carrying an amino acid substitution at position 119 (aspartic acid to asparagine) of the VPg (named “CI chimera VPg-N”) which conferred the breakdown capacity towards the *pvr2*³ resistance gene and was shown to be the most frequent in isolates from field as well as from experimental evolution experiments (Ayme *et al.*, 2006; Palloix *et al.*, 2009). This mutant enables the pathogen to overcome the resistance conferred by *pvr2*³ and to reveal the quantitative resistance due to the genetic background.

Measure of resistance breakdown frequency

The breakdown frequency of *pvr2*³ was tested after inoculation of the PVY “CI chimera” to the 153 DH lines carrying the *pvr2*³ allele. Throughout the test, plants were grown in a climate-controlled room at 20-22 °C, 12h light /day. Inoculum was obtained after virus propagation in *Nicotiana tabacum* cv. Xanthi plants. Fifty grams of Xanthi leaves showing PVY symptoms were crushed in 200 mL of 0.03M phosphate buffer (pH 7.0) supplemented with 2% (w:v) diethyldithiocarbamate, 4.5 g of active charcoal and 4.5 g of Carborundum. Thirty pepper seedlings with two expanded cotyledons (2-3 weeks after sowing) per genotype were inoculated mechanically on their cotyledons. Thirty-eight days post-inoculation (dpi), plants were submitted to virus detection by DAS-ELISA as described by Ayme *et al.* (2006). In these conditions, in each plant infected systemically, the virus population was shown to be composed of one or several VPg mutants

carrying a *pvr2*³ RB mutation (Montarry *et al.*, 2011). For each pepper line, the RB frequency of *pvr2*³ was assessed by the ratio of the number of systemically infected plants over the total number of inoculated plants. Two independent tests of 30 plants per DH line were performed.

Measure of quantitative resistance

To evaluate the level of the quantitative resistance due to the genetic background, we used the “CI chimera VPg-N” PVY mutant which overcomes the resistance conferred by *pvr2*³ (infects all the genotypes carrying *pvr2*³) and reveals the quantitative resistance due to the genetic background. Two different traits were assessed : the area under the disease progress curve (AUDPC) and the virus accumulation. The PVY “CI chimera VPg-N”, was mechanically inoculated on 20 plants of each DH lines carrying *pvr2*³. The plants were grown in a climate-controlled room as previously described. The AUDPC assessment combined both intensity of symptoms and latency period: symptom intensity was evaluated, on the 20 plants, in a 3-class scale (1= no symptom, 2 = weak mosaic and 3 = severe mosaic/ necrosis) at 14, 21, 28 and 35 dpi. AUDPC was calculated using the formula:

$$\text{AUDPC} = \sum_{i=1}^{n-1} \left(\frac{y_i + y_{i+1}}{2} \right) (t_{i+1} - t_i)$$

where “n” is the number of symptom assessments, “y” the symptom intensity and “t” the time in dpi.

At 36 dpi, the virus accumulation was independently evaluated on 10 individual plants per DH line by semi-quantitative DAS-ELISA, using a dilution range of extracts of PVY infected plants, and expressed relatively to a common reference sample incorporated in each ELISA plate, as in Ayme *et al.* (2006).

Genotyping of molecular markers and linkage map

The 350 DH progeny was an extension of the former ‘PY’ mapping population (Lefebvre *et al.*, 2002). It was genotyped with 236 molecular markers: 108 single-nucleotide polymorphisms (SNPs, Nicolai *et al.*, 2012), 29 SNPs (Jung *et al.*, 2010), 47 simple sequence repeats (SSRs, Lee *et al.*, 2004; Nagy *et al.*, 2007; Minamiyama *et al.*, 2006), 33 amplified fragment-length polymorphisms (AFLPs, Lefebvre *et al.*, 2002), one single-strand conformation polymorphism (SSCP-PM5, Fazari *et al.*, 2012), two known genes and 16 new markers (supplementary table I) corresponding to previously mapped restriction fragment length polymorphisms (RFLPs) and/or conserved ortholog set (COS II) sequences and designed using the Sol Genomics network (www.solgenomics.net). The two known genes were *pvr2* (the major resistance locus to PVY) genotyped using the tetra-ARMS marker as in (Rubio *et al.*, 2008), and *pvr6* coding for the eIF(iso)4E (eukaryotic initiation factor iso4E) protein genotyped with a new marker targeting a 82 nucleotide deletion differentiating alleles *pvr6*⁺ and *pvr6* (Ruffel *et al.*, 2006) (supplementary data I). The DH progeny was also genotyped for two phenotypic markers, the presence/absence of pungency in the fruits (locus C) and the erected/pendant fruit position (locus *up*).

The genetic linkage map was constructed using the Mapmaker software version 3.0b (Lander *et al.*, 1987) with a LOD score threshold of 3.0, a maximum recombination fraction of 0.3 and distances between markers were calculated using the Kosambi mapping function. For each linkage group, marker order was verified with the “ripple” command and markers were conserved only if the LOD value was greater than 2.0. Three out of the 236 markers remained unlinked and 33 were deleted because they had no unique placement at LOD 2.0. The resulting map contained 200 markers distributed in 13 linkage groups (LG), which were assigned to the 12 pepper chromosomes through alignment of markers with previous pepper maps (Lefebvre *et al.*, 2002; Wu *et al.*, 2009)

(supplementary data II). The total length of the map was 2457.7 cM with an average length interval between markers of 12.3 ± 12.4 cM. The increased length of this map compared to former pepper maps was mainly attributed to the larger size of the progeny (350 individuals) that allowed to declare significant a recombination rate up to 0.45 cM (slightly more than 50 Kosambi cM), and also to the assignment of the two linkage groups to the single P7 chromosome, resulting in longer chromosomes (supplementary data II).

Statistical analyses

Descriptive statistics were performed with the R software (<http://www.r-project.org/>). For each phenotypic trait, narrow-sense heritability (h^2) was estimated using the formula $h^2 = \sigma_G^2 / (\sigma_G^2 + \sigma_E^2/n)$, where σ_G^2 corresponds to the genotypic variance, σ_E^2 to the environment (error) variance and n to the number of replicates.

For AUDPC, mean distributions were close to a normal distribution and we analyzed non-transformed data for QTL detection. Means of virus accumulation (VA) were non-normally distributed. Thus, for QTL detection, these data were log transformed [$\ln(\text{VA}) + 1$] to approximate a normal distribution. Resistance breakdown frequency was a non-normally distributed trait with an excess of low values, and no transformation was efficient to improve the distribution.

QTL analysis by regression interval mapping

For a non-normal distribution, QTL detection can be performed with a standard regression-based interval mapping approach, which is robust against deviations from normality when the threshold value is calculated by permutations (Rebai, 1997). QTL analyses were performed using iterative composite interval mapping (iQTLm) methods implemented in the MCQTL v.5.2.4 (Jourjon *et al.*, 2005) software. MCQTL performed QTL mapping using a linear regression model and the iQTLm scan method allows a multiple QTL model to be determined automatically (Charcosset *et al.*, 2001). The QTL significance was determined using the MCQTL test, which is equal to $-\log_{10}$ of the Fisher test Pvalue as described in the MCQTL v.5 reference manual (<http://carlit.toulouse.inra.fr/MCQTL/>). The significance threshold of the MCQTL test was estimated after performing 1000 permutation tests for each trait to reach a genome-wide type-I error rate of 0.05 and 0.1 for QTLs and cofactors, respectively. This resulted in a threshold of 3.4 for [$\log(\text{VA}) + 1$] and of 3.5 for RB and AUDPC. The cofactor threshold was fixed at 3.2 for the three traits. Cofactor selection was performed by forward selection of markers. The QTL confidence intervals (CI) were defined as a 2 unit drop-off of the MCQTL test around the maximum MCQTL score to obtain approximately a 95% confidence interval : $\text{CI} = [\max(\text{MCQTL test}) - 2; \max(\text{MCQTL test}) + 2]$. Epistasis between QTLs and the genetic background was tested using the iterative method implemented in MCQTL (threshold of 3.5). The part of phenotypic variation explained by additive and/or epistatic effects of each QTL (R^2 coefficients) and the global R^2 were estimated using the full QTL model in MCQTL.

QTL analysis by maximum likelihood interval mapping

For AUDPC and virus accumulation ($\log(\text{VA}) + 1$), QTL analyses were also performed using multiple QTL mapping (MQM) methods implemented in the R/qrtl package of the R (Broman *et al.*, 2003) software. We did not use this method for the RB trait because data were non-normally distributed. MQM is a maximum likelihood-based interval mapping method composed of three steps. First, missing data are “augmented”, i.e. all genotype possibilities are modeled and their probability calculated from neighboring marker information. In the second step, significant markers are detected with the “mqmscan” function without cofactor, which corresponds to a simple interval

mapping (SIM) with a backward elimination. In the last step, QTLs are detected using markers detected in step two as cofactors, which corresponds to a composite interval mapping, and a backward elimination is applied to remove the non-significant cofactors and to establish the final QTL model. Significance LOD thresholds were estimated for each trait, with a genome-wide type-I error rate of 0.05, after performing 1000 permutation tests. The LOD threshold was estimated at 2.6 for the two traits. For each QTL, the CI was defined as a 2-LOD drop-off around the maximum LOD score to obtain approximately a 95% confidence interval. R^2 coefficients were calculated with the “fitqtl” function of R/qtl.

RESULTS

Characterization of the phenotypic traits

To evaluate the frequency of *pvr2³* resistance breakdown, the “CI Chimera”, which is not infectious *per se* toward *pvr2³* resistant plants, was inoculated on the set of 153 DH carrying *pvr2³*. In these plants, only RB mutants of the “CI Chimera” possessing single non-synonymous substitutions in the factor (VPg) conferring virulence towards the *pvr2³* allele were detected in previous studies (Ayme *et al.*, 2006; Palloix *et al.*, 2009; Montarry *et al.*, 2011.). In our tests, mosaic symptoms appeared in 100% of the susceptible (*pvr2⁺*) control Yolo Wonder 2 weeks after inoculation, but mosaic or necrotic symptoms appeared later (4 to 5 weeks after inoculation) in some of the plants of some DH lines. Thirty-eight days after inoculation (dpi), the ELISA revealed an important phenotypic variability between the 153 DH lines, with frequencies of systemic infection, corresponding to RB frequencies, varying from 0% to 93.2% with a mean (\pm standard deviation) of 14.7% (\pm 23.8). The correlation between the two independent tests was high ($\rho_{\text{Spearman}}=0.75$; $p<10^{-7}$, supplementary data 3), and a heritability of 0.87 was estimated. The distribution of the trait was strongly skewed towards low values, with 70 DH lines displaying no resistance breakdown and 83 HD lines displaying a RB frequency between 1.7% and 93.2% (figure 1).

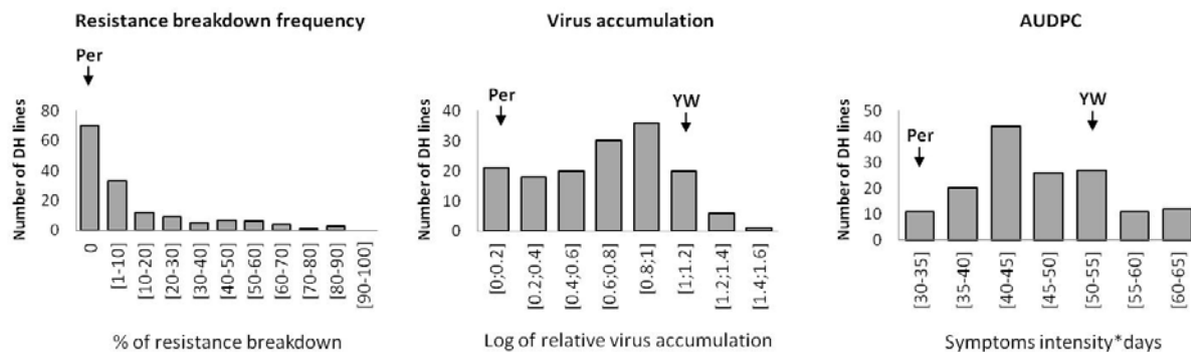


Figure 1: Histograms represent distributions of DH lines into resistance breakdown frequency, virus accumulation (ratio concentration of the sample/concentration of the control) and AUDPC classes. Arrows indicate the position of each parent in the phenotypic distribution. Per : Perennial, YW : Yolo Wonder.

The quantitative resistance was assessed using a *pvr2³*-breaking mutant of the “CI Chimera”, the “CI Chimera VPg-N”. With this virus, the *pvr2³* resistant lines displayed symptoms as soon as 14 dpi and 100% of the plants were infected at 35 dpi as attested by DAS-ELISA. Highly significant variations for virus accumulation (VA) and AUDPC were observed between DH lines with a heritability of 0.64 for $\text{Ln}(\text{VA}+1)$ and 0.98 for AUDPC. For statistical and QTL analyses, VA values were log transformed $[\text{Ln}(\text{VA}+1)]$ to approximate a normal distribution. The $\text{Ln}(\text{VA}+1)$ varied from 0 to 1.43 with a mean equal to 0.67 ± 0.35 and AUDPC varied from 29.75 to 63 with a mean equal to 46.6 ± 7.9 (figure 1).

Significant correlations ($p < 0.05$) were observed between the three traits with Spearman's coefficients equal to 0.40, 0.33 and 0.32 for RB/VA, RB/AUDPC and VA/AUDPC, respectively.

Mapping QTLs for the quantitative resistance traits

Distributions of the quantitative resistance traits AUDPC and $\log(VA+1)$ were close to normality. For these two traits, QTL detection was performed using iQTLm and MQM methods. Two significant QTLs for VA and three for AUDPC were detected by both iQTLm and MQM methods. These QTLs were named VA-3, VA-6, A-1, A-3 and A-9, according to the trait considered and the chromosome location. Hereafter, we will detail only the results from iQTLm because results from both methods are similar. The position, significance and effect of each QTL are detailed in table 1 and figure 2.

VA-3 and VA-6 explained 34.5% and 15.7% of the VA variation, respectively. Epistasis tests did not reveal any interaction between QTLs and the genetic background. The final model combining the effects of the two significant QTLs explained 43.9% of VA phenotypic variation corresponding to 69% of the trait heritability ($h^2=0.64$).

A-1, A-3 and A-9 explained from 14.69%, 15.5% and 16.1% of the AUDPC variation. Epistasis tests did not reveal any interaction between these QTLs and the genetic background. The part of the AUDPC phenotypic variation explained by A-1, A-3 and A-9 was equal to 34.1% corresponding to 35% of the trait heritability and a large part of the variability remained unexplained ($h^2=0.98$).

Table 1: QTL detected with the iterative QTL mapping (iQTLm) method.

Virus	Trait	QTL	Location ^a	Closest marker	CI ^b	Direction ^c	MCQTL test ^d	Variation explained		Heritability	
								locus	trait		
Wild-type PVY clone (CI Chimera)	Resistance breakdown	<i>QTLs with additive effect :</i>									
		Rb-1	1@87	Gpms_178	42-100	Per	5.2	12.8	68.9	0.87	
		Rb-3	3@43.1	Pvr6	41 - 49	YW	17.4	39.8			
		Rb-5	5@57.1	TG437	12-103	Per	3.6	8.6			
		<i>QTLs with epistatic effect :</i>									
		eRb-1-3	1@87 - 3@43.1	Gpms_178 / Pvr6			7.4	18.9			
eRb-3-5	3@43.1 - 5@57.1	Pvr6 / TG437			6.6	16.9					
eRb-3-6	3@43.1 - 6@188.3	Pvr6 / SNP11391			4.2	10.6					
Mutant PVY clone (CI Chimera VPg-N)	Virus accumulation	<i>QTLs with additive effect :</i>									
		Va-3	3@49.1	SNP23714	35-54	YW	14.6	34.5	43.9	0.64	
		Va-6	6@190.3	SNP11391	158 - 202	Per	6.3	15.7			
<i>QTLs with additive effect :</i>											
Mutant PVY clone (CI Chimera VPg-N)	AUDPC	A-1	1@87	Gpms_178	77-99	Per	6.3	16.1	34.1	0.98	
		A-3	3@40.8	SNPISO_2	16 - 52	YW	6.2	15.5			
		A-9	9@132	SSCP_MP5	117 - 149	YW	5.9	14.9			

^a QTL location are coded in the following way : Chromosome @ position on the chromosome in cM. For example, the QTL RB-1 is localized on the chromosome 1 at 87 cM (1@87).

^b The confidence interval was defined as a 2-MCQTL test drop-off around the maximum MCQTL test score.

^c Indicates the parent which decreases the value of the trait.

^d The MCQTL test significance threshold was set at $P=0.05$ and is equal to 3.4 for Virus accumulation and 3.5 for resistance breakdown and AUDPC.

Mapping QTLs for the frequency of *pvr2*³-resistance breakdown

Because of the skewed distribution of the RB frequency, QTL detection was performed with the standard regression-based interval mapping approach (Haley and Knott, 1992; Rebaï, 1997) using the iQTLm (Charcosset *et al.*, 2001) method implemented in MCQTL software (Jourjon *et al.*, 2005). Three significant QTLs, explaining 12.8%, 39.8% and 8.6% of the *pvr2*³ RB frequency variation, were detected on chromosomes 1, 3 and 5, respectively. QTLs were named RB-1, RB-3

and RB-5 according to the trait and the chromosome location. Table 1 and figure 2 detail position, significance and effect of each QTL. Epistasis tests revealed that RB-3 interacts with three distinct positions in the pepper genome, on chromosomes 1, 5 and 6. The positions on chromosomes 1 and 5 interacting with RB-3 are included in the CIs of RB-1 and RB-5, respectively, suggesting interaction effects between RB-1 and RB-3 and between RB-3 and RB-5 (figure 2). The position on chromosome 6 that interacts with RB-3 did not show any additive effect and will be further named RB-6. The final model combining additive and epistatic effects of the significant QTLs explained 68.9% of the variation of *pvr2*³ RB frequency, which corresponds to a great part (79%) of the trait heritability ($h^2=0.87$). Because of the skewed segregation of the RB trait, the QTL effects (as well as heritability) may have been overestimated. Indeed, the residues of the final model did not fit a normal distribution (Shapiro-Wilk normality test $W=0.87$, $p<10^{-7}$). The analysis was reproduced after removing the 70 lines with a RB value of 0. The same additive QTLs were detected and the final model including these 3 additive QTLs explained 48% of the phenotypic variance (instead of 52% with the whole population) with a normal distribution of residues ($W=0.98$, $p=0.55$). The interaction effects were not significant anymore. This very probably resulted from the fact that removing the 70 individuals with RB=0 also removed most of the individuals carrying the Yolo Wonder allele at RB-3, resulting in highly unbalanced allele frequencies at this locus (71/12), reduced population size ($n=83$) and decreased statistical power.

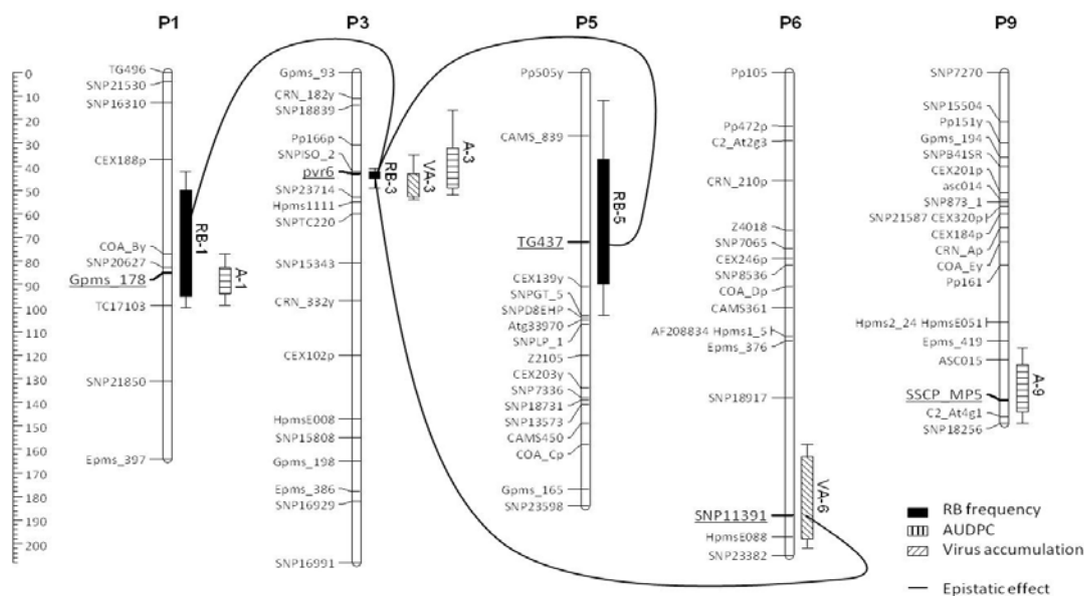


Figure 2: Map positions on Pepper genetic map of the significant quantitative trait loci (QTLs) for *pvr2*³ resistance breakdown (RB) frequency, the area under the disease progress curve (AUDPC) and the virus accumulation (VA). On the left, the scale indicates map distances calculated by the Kosambi mapping function. For each QTL, the length of the block represents the confidence interval (CI) of the QTL using a MCQTL test unit fall of 1 and the length of the vertical bar represents the CI of the QTL using a MCQTL test unit fall of 2. For each QTL, the closest marker is underlined. Lines indicate epistatic effects between loci.

Looking for co-localization between QTLs, the confidence interval of A-1, proved fully included in the confidence interval of RB-1, as was the confidence interval of RB-3 within those of A-3 and VA-3. The major QTL affecting the RB frequency of *pvr2*³ (RB-3) co-localizes with the major QTLs VA-3 and A-3 affecting VA and AUDPC, respectively. The QTL RB-1 also co-localizes with QTL A-1. The RB-6 locus displaying interaction effect with RB-3 also located at the LOD peak of VA-6, indicating that RB-6 colocalizes with VA-6 (figure 2).

Effect of parental alleles on the RB frequency

All genotypes carrying the Yolo Wonder allele at QTL RB-3 showed a very low resistance breakdown frequency with a mean equal to 0.31% whereas, genotype carrying the Perennial allele at QTL RB-3 showed a much higher resistance breakdown frequency with a mean equal to 29.63%. The phenotypic values of the different allele combination between QTL RB-3 and epistatic QTLs are presented in Table 2. For genotypes carrying the Yolo Wonder allele at QTL RB-3, the resistance breakdown frequency was independent of the alleles at the interacting QTLs (mean of resistance breakdown frequency varies between 0.17% and 0.37%), so that these QTL effects were not significant in these genotypes. Contrarily, in genotypes carrying the Perennial allele at QTL RB-3, the RB frequency strongly depended on the alleles present at the interacting QTLs. The Yolo Wonder alleles at QTLs RB-1, RB-5 or RB-6 increased the RB frequency when combined with the Perennial allele at QTL RB-3, whereas the Perennial alleles at these QTLs strongly decreased the RB frequency (Table 2). Altogether, the combination of the Perennial alleles at the three epistatic QTLs (RB-1, RB-5 and RB-6) with the Perennial allele at RB-3 decreased the RB frequency from 69.7% to 5.3% (table 2). The high effect of these QTLs in genotypes carrying the Perennial allele at RB3, contrarily to those carrying the Yolo Wonder allele at this QTL probably caused the significant interaction (scale effect).

Table 2: Effect of QTLs interaction on the resistance breakdown frequency of *pvr2³*.

		Rb-1 : Gpms_178 (1@85)		Rb-5 : TG437 (5@72.3)		Rb-6 : SNP11391 (6@188.3)		combination of the 3 QTLs (Rb-1, Rb-5, Rb-6)	
		Yolo Wonder	Perennial	Yolo Wonder	Perennial	Yolo Wonder	Perennial	Yolo Wonder	Perennial
Rb-3 : (3@43.1)	Yolo Wonder	0.37% (34)	0.30% (39)	0.17% (39)	0.47% (33)	0.18% (37)	0.36% (39)	0.18% (9)	0.36% (7)
	Perennial	42.0% (37)	15.6% (34)	41.8% (32)	18.0% (31)	36.2% (40)	22.6% (30)	69.7% (7)	5.3% (6)

This table presents the effect of the three significant epistatic interactions between loci and their combination for the resistance breakdown frequency of *pvr2³*. The names and the positions of the markers in interaction are indicated. The numbers correspond to the average of the *pvr2³* resistance breakdown (RB) frequency (in %) for each allelic combination and the number between brackets corresponds to the number of lines belonging to each allelic class.

DISCUSSION

Durable management of plant resistance to pathogen populations was mostly promoted through the spatio-temporal management of resistance genes, but can also be achieved through management of resistance genes at the genotype scale: the cultivar genotype construction. In this paper we firstly succeeded to identify plant QTLs for a trait directly related to the evolution of the viral population: the frequency of emergence of viral variants adapted to the resistance. The collocation of the plant genetic factors modulating this viral evolution with the QTLs decreasing the virus accumulation further indicated that the more efficient the more durable was the resistance, which was not obvious since higher resistance levels are also expected to increase the selection pressure. Consequently, the high durability of the resistance of the Perennial pepper landrace was shown to result from a complex genetic architecture, including alleles at a major gene and at QTLs from the genetic background with complementary but also contradictory effects, which address the question of the selection processes leading to this combination in this landrace.

Heritability of the breakdown frequency of a major resistance gene

In this study, we measured the resistance breakdown (RB) frequency of a major resistance gene (*pvr2³*) in a set of pepper DH lines carrying *pvr2³* but segregating for the genetic background. The observed RB frequency directly resulted from the frequency of appearance and from the

accumulation dynamics of RB variants in the resistant plants, two steps of the virus evolution toward RB that are considered as major components of the resistance durability (Fabre *et al.*, 2009). Previous studies of this pathosystem showed that the RB mutation most probably occur in the resistant pepper host which allowed for weak PVY multiplication and that the genetic background acts at several levels, including the mutational pathways of the virus to overcome resistance and the speed of selection of the RB variants (Montarry *et al.* 2011, Quenouille *et al.* 2013). The genetic analysis showed that the RB frequency of *pvr2*³ was highly variable between genotypes (from 0% to 93.2%) and highly heritable ($h^2=0.87$). Because of the non-normal distribution of data for RB frequency, the h^2 value is most probably overestimated. However, it fits the Spearman correlation value between the 2 repeats of the test (0.75), demonstrating the genetic control of this trait. This result is not trivial. Indeed, during the RB processes, several factors could have increased the heterogeneity of results between plants and experiments and consequently could have decreased the heritability of RB frequency. Among these factors are the stochastic nature of (i) the appearance of RB mutations in PVY genome, (ii) the genetic drift acting on virus during systemic invasion of plants (Garcia-Arenal *et al.*, 2001; French and Stenger, 2003) and (iii) the fact that at least 9 different VPg mutations, with contrasted effects on PVY competitiveness, can be involved in *pvr2*³ breakdown (Ayme *et al.*, 2006; Montarry *et al.*, 2011). Our study demonstrated that measuring the resistance breakdown frequency with a reasonable number of plants (60 per genotype) compensates these factors of heterogeneity and allows the genetic analyses. The genetic background dependency of *pvr2*³ durability could be attributed to 4 distinct genetic factors with quantitative effects: one major-QTL (RB-3) and three additional QTLs (RB-1, RB-5 and RB-6) acting additively and/or in interaction with RB-3. The discovery of such genetic factors shows that direct selection for alleles increasing the durability of a major resistance is feasible and opens new tools for durable resistance breeding. Nevertheless, the successful detection of such genetic factors affecting the RB frequency of a major resistance gene depends on the ability to generate RB strains at high frequencies through experimental evolution. For many virus diseases, the experimental conversion of an avirulent virus to an RB variant is rather easy and was commonly used to determine viral determinants of pathogenicity (Ayme *et al.*, 2006; Margaria *et al.*, 2007; Hajimorad *et al.*, 2010; Lacroix *et al.*, 2011). Moreover, the RB frequency of resistance genes is highly correlated with the durability of these genes in the field (Ayme *et al.*, 2006; Janzac *et al.*, 2009; Lacroix *et al.*, 2011). In tobacco, field observations revealing that *va*²-RB isolates were more frequent than *va*⁰-RB isolates (Verrier and Doroszewska, 2004) were confirmed by results from experimental evolution (Lacroix *et al.*, 2011). Moreover, the distributions of RB mutations in viral genomes are usually similar in field and laboratory observations (Moury *et al.*, 2004; Ayme *et al.*, 2006; Hajimorad *et al.*, 2010). However, for some resistance genes which proved very durable in the field, it may be difficult to generate RB variants by direct inoculation of avirulent strains. In such cases, alternative protocols imposing strong inoculum pressures like graft-inoculation or inoculation through agroinfiltration can generate sufficient inoculum pressure to increase RB frequency (Bruun-Rasmussen *et al.*, 2007; Janzac *et al.*, 2009). These experimental procedures make it possible to measure the breakdown frequency of major resistance genes in many plant-virus pathosystems, providing powerful tools for improving the durability of virus resistance in plants. Considering other pathogens, experimental evolution may not be so relevant, due to their longer generation time, smaller populations and smaller mutation rates. However, if confirmed as a generic result, the relationship between RB QTLs and QTLs controlling the level of quantitative resistance may provide alternative selection criteria.

Alleles increasing quantitative resistance are favourable to the durability of major resistance genes.

Using the same pepper DH progeny, we performed a QTL analysis of quantitative resistance. To get rid of the *pvr2*³ major effect and to reveal the quantitative resistance conferred by the genetic

background, we used a *pvr2*³-RB PVY variant (the “CI Chimera VPg-N”) which differs from the PVY variant used for the measurement of *pvr2*³ RB frequency by a single nucleotide substitution. This mutation restores the interaction between the PVY VPg and the pepper eukaryotic translation initiation factor 4E1 (eIF4E1) encoded by *pvr2*³, causing the breakdown of the *pvr2*³-mediated resistance (Charron *et al.*, 2008). Using this PVY variant, four QTLs affecting virus accumulation and/or symptom expression were detected. These four QTLs were also previously detected for the AUDPC trait with two other PVY strains, with identical allele directed effects: resistance-favourable Perennial alleles on chromosomes P1 and P6 (LG3 and LG10 in Caranta *et al.*, 1997) and favourable Yolo Wonder alleles on chromosomes P9 and P3 (LG9 and LG4 in Caranta *et al.*, 1997). In this experiment, distinct natural PVY isolates with distinct *pvr2*³ breaking down mutations in their VPg were used, indicating no differences between the QTLs detected using distinct VPg mutants. Our present results show that three of the four QTLs affecting the RB frequency of *pvr2*³ collocate with QTLs affecting quantitative resistance (viral accumulation and/or AUDPC). These collocations may result from genetic linkage between the different traits or from a pleiotropic effect of quantitative resistance factors on the RB frequency of *pvr2*³. Looking at the effect of parental alleles that collocate on chromosomes P1, P3 and P6, all the alleles that increased the quantitative resistance also decreased the breakdown frequency. Moreover, the QTL on chromosome P3 displayed the major effect on both virus accumulation and RB frequency. This concordance between parental allele effects for the different traits strongly suggests that collocations are due to pleiotropic effects and that alleles increasing the resistance level also increase the resistance durability. This is also consistent with the initial observations, in 3 pathosystems, that the durability of major genes is enhanced in partially resistant cultivars (Palloix *et al.*, 2009; Brun *et al.*, 2010; Fournet *et al.*, 2012). However, it may look contradictory since higher resistance levels are expected to increase the selection pressure on the pathogen and the risk of selection of adapted pathogen populations (Poland *et al.* 2009). In a recent study of the different mechanisms that could account for the higher durability of polygenic resistance, Quenouille *et al.* (2013) showed the selection of adapted RB mutants by the polygenic resistance of Perennial was slower than by the monogenic resistance (*pvr2*³ in a susceptible genetic background). In this same report, the additional quantitative resistance carried by the genetic background explained a large part of the variation in RB frequency. Dissecting the genetic background into QTLs indicate that breeding for quantitative resistance alleles will enhance the durability of major genes. Such a strategy is achievable for most pathosystems where breaking down variants or experimental evolution tests are not available.

Resistance durability as the result of a combination of mechanisms

At the landscape scale, it was proposed previously that diversifying the methods for pathogen control is an efficient strategy to limit adaptation of pathogen populations (Mundt *et al.*, 2002; Pink *et al.*, 2002). Diversifying resistance mechanisms at the genotype scale can also have a positive effect on durability. In accordance with Quenouille *et al.* (2013), our results raise the main impact of viral accumulation on the RB frequency of *pvr2*³, since all QTLs involved in viral accumulation are also involved in the RB frequency of *pvr2*³. Viral accumulation was measured in apical leaves and a decrease of viral accumulation can result from decreases of the efficiency of virus replication, cell-to-cell and/or systemic movement. Combining a major resistance gene with another gene involved in viral accumulation has already proved to enhance resistance durability in the PVY/tobacco system (Acosta-Leal and Xiong, 2008). In their study, Acosta-Leal and Xiong (2008) showed that the RB frequency of the *va* recessive gene which confers a resistance to PVY cell-to-cell movement in tobacco was reduced when *va* was combined with a second recessive gene (*va2*) conferring resistance to within-cell PVY accumulation. Two additional QTLs affected the RB frequency but had no detectable effect on viral accumulation (RB-1 and RB-5), suggesting distinct modes of action. QTL RB-1 collocates with QTL A-1 controlling symptom intensity and dynamics.

In plants, a decrease of symptom intensity can result from resistance and/or tolerance mechanism. Here we consider tolerance in the sense of Schafer (1971) in which tolerant plants endure high pathogen loads without severe losses in field. Thus, QTL A-1 can be considered as a tolerance QTL since it reduces the AUDPC but it does not significantly affect viral accumulation. To our knowledge, mechanisms involved in plant tolerance are still unknown but they could also have an effect on the *pvr2*³ RB frequency. Finally, QTL RB-5 did not colocalize with any quantitative resistance QTL, suggesting that a third mechanism, independent of quantitative resistance, affects the *pvr2*³ RB frequency. Quenouille *et al.* (2013) also showed that beside virus accumulation, the plant genetic background can control two additional modes of action: making the mutational pathways for viral resistance breakdown more complex and unlikely and modulating the selection pressure exerted on viral populations. QTLs RB-1 and RB-5 may be candidates for such modes of actions.

A genetic trade-off between broad-spectrum resistance and resistance durability that can be solved by breeding strategy

Among the 4 additive and/or epistatic QTLs affecting the RB frequency, the alleles from Perennial were shown to decrease the risk of *pvr2*³ breakdown at 3 of these QTLs (RB-1, RB-5 and RB-6), whereas the Perennial allele increased strongly this risk for the major QTL RB-3. This result looks surprising since the *pvr2*³ allele was previously shown to be highly durable in the Perennial landrace (Palloix *et al.*, 2009). It is noteworthy that the peak of QTL RB-3 was localized at the *pvr6* locus which encodes the eIF(iso)4E, an isoform of eIF4E1 encoded by *pvr2*. It was previously shown that the Perennial allele at *pvr6* included a deletion of 82 nucleotides followed by a premature stop codon, compared to the *pvr6*⁺ allele from Yolo Wonder. The eIF(iso)4E protein encoded by the Perennial *pvr6* allele was therefore non-functional (Ruffel *et al.*, 2006) and can be considered as a deleted allele of *pvr6*⁺. When combined with different *pvr2* alleles, the Perennial *pvr6* allele was shown to enlarge the spectrum of resistance to additional potyviruses, including *Chilli vein mottle virus* (ChiVMV) and *Pepper vein mottle virus* (PepVMV) (Caranta *et al.*, 1997; Moury *et al.*, 2005; Rubio *et al.*, 2009). Hence, the Perennial allele at the *pvr6*/RB-3 locus contributes to a gain of resistance against PepVMV and ChiVMV but also to a decrease of the durability of the *pvr2*³ resistance to PVY together with a decrease in quantitative resistance traits VA and AUDPC. The Perennial inbred line was selected from a landrace issued from farm selection in India, where several potyviruses including ChiVMV are prevalent in pepper crops and it is highly likely that *pvr2*³ and *pvr6* have been selected locally to enlarge the spectrum of resistance against these potyviruses. Whether *pvr6* is the functional gene for the RB-3 QTL remains to be determined through functional validation. However, the tight colocalization between *pvr6* and RB-3 already reveals a trade-off between a large resistance spectrum against potyviruses when combined with a *pvr2* resistance allele, and the durability of the *pvr2*³ PVY resistance in the Perennial genotype.

Such a trade-off looks unusual since resistance gene pyramiding and broad-spectrum resistance are often considered favourable to durability (McDonald and Linde, 2002; Kou and Wang, 2010). This statement was recently reinforced by Le Van *et al.* (2013) who showed that broad spectrum resistance QTLs in apple did not exert any differential selection pressures on a mixed *Venturia inaequalis* inoculum, whereas narrow spectrum QTLs did. In pepper, the combination of *pvr2* resistance alleles (particularly *pvr2*¹ and *pvr2*² alleles) with *pvr6* is already used in breeding programs to create resistant varieties against a large range of potyviruses (Rubio *et al.*, 2009) and such cultivars may lead to a premature breakdown of *pvr2*-mediated resistance by PVY. More generally, translation initiation factor-mediated virus resistance is widespread (Robaglia and Caranta, 2006) and combination between mutated or knock-out alleles at different genes coding for

translation initiation factors is expected to provide a way to breed for broad spectrum resistance. In tomato, the narrow range resistance conferred by a knock-out allele of eIF4E1 was enlarged when combined with a knock-out of eIF4E2 (Mazier *et al.*, 2011). Our study indicates combining mutant alleles from genes belonging to the same multigenic family should be used carefully, since the trade-off observed in pepper between large-spectrum and durability of resistance, may also occur in other plants species.

Our previous observations indicate introgressing resistance alleles from the plant germplasm into new elite cultivars with susceptible genetic backgrounds can endanger the long-term use of these genes as well as provide an evolutionary springboard to the pathogen for resistance breakdown (Palloix *et al.*, 2009). Investigating the background-dependence of the durability of *pvr2*³ led to an improved understanding of how selection acted on this gene/genetic background combination (Chandler *et al.*, 2013). In mass selection of plant populations by farmers, only the seeds from the healthiest and most productive individuals participate to the next cultivated generation. In North-West India where Perennial originated, plants combining *pvr2*³ and *pvr6* certainly gain a selective advantage in presence of ChiVMV. However, the rapid adaptation of PVY and/or other potyviruses would have counterselected this allelic combination in a few cultivation seasons. Over a few plant generations, and in heterogeneous plant populations, this multi-pathogen selection pressure promoted the recombinant individuals carrying favorable alleles at additional QTLs (RB-1, RB-5 and RB-6) which, in interaction with RB-3, decreased the risk of *pvr2*³ resistance breakdown. Such a polygenic combination of co-adapted alleles contributed to increase resistance in efficiency, spectrum and durability.

DATA ARCHIVING

Data available from the Dryad Digital Repository:

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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