Characterization of Hypersensitive Resistance to Bacterial Spot Race T3 (Xanthomonas perforans) from Tomato Accession PI 128216

Matthew D. Robbins, Audrey Darrigues, Sung-Chur Sim, Mohammed Abu Taher Masud, and David M. Francis

ABSTRACT

Several sources of resistance to spot have been identified but, in most cases, the resistance is race specific. The S. lycopersicum breeding line Hawaii 7998 (Ha7998) has demonstrated both a reduction in symptoms in the field (field resistance) (20) and a hypersensitive reaction (HR) in the greenhouse (11) to spot race T1 (X. euvesicatoria). Three quantitative trait loci (QTL), Rx-1, Rx-2, and Rx-3, were associated with the HR in a cross between Ha7998 and the S. pennellii accession LA716 (31,37). The Rx-3 locus was subsequently confirmed to provide HR as well as resistance in the field in a cross between Ha7998 and an elite breeding line, OH88119, and markers linked to Rx-3 were identified (36). A second breeding line, Hawaii 7981 (Ha7981), provides HR to race T3 by the interaction between avrXv3 and a single gene, Xv3, which is yet to be mapped (15,24). Field resistance to T3 from Ha7981 is quantitatively inherited and is based on Xv3 and modifier loci (21). Hypersensitivity (HR) to T4 race was identified from LA716 as Xv4 and mapped to chromosome 3 (1). The genetic characterization of this source of resistance requires clarification because the map position was not supported in subsequent studies (unpublished results). The S. lycopersicum var. cerasiforme accession, PI 114490, has shown resistance in the field to multiple races of spot (19,23). Markertrait analysis was used to identify a QTL on chromosome 11 that had a large effect (13 to 57% of the variation) on resistance for races T1, T2, T3, and T4 (35). Another major QTL on chromosome 1 contributed 3 to 11% of resistance against races T2, T3, and T4. Additional minor QTL were identified that collectively contributed to resistance to all four races.

Genetics and Resistance
An additional source of resistance to spot race T3 is the *S. pimpinellifolium* accession PI 128216. In a survey of several tomato accessions, PI 128216 and selections from the PI expressed HR when challenged by spot race T3 (22). Although some of the tomato accessions most resistant in the field also showed HR, the correlation between HR and field resistance was not complete (22). The clear HR response from PI 128216 to only race T3 has been used as a diagnostic to classify *Xanthomonas* strains into races (17). However, the genetic basis of resistance from PI 128216 to race T3 has not been described.

Given the complexity of the bacterial spot disease complex and the existence of multiple sources of resistance, it would be desirable to better characterize each source in order to facilitate pyramiding of genes effective against multiple races and against other bacterial diseases. The mapping of *Rx-3*, for example, provided an opportunity to combine this locus with *Pto*, which confers resistance to race 0 of bacterial speck (*Pseudomonas syringae* pv. *tomato*), using marker-assisted selection to identify coupling phase recombinants (34). Therefore, our goal was to describe the genetic basis of the HR reaction to spot race T3 in PI 128216, determine if HR provides resistance in the field, and pyramid this resistance with other bacterial resistance genes. We identified a locus on chromosome 11, *Rx-4*, that provides HR in greenhouse inoculations and resistance in the field. We combined this locus with *Pto* and *Rx-3* in order to develop breeding lines with resistance to multiple bacterial diseases.

**MATERIALS AND METHODS**

**Plant material.** We created an inbred backcross (IBC) population for simultaneous introduction and characterization of spot resistance (32). The IBC population was developed from a cross between OH88119 (*S. lycopersicum*) and PI 128216. OH88119 is an elite processing tomato parent chosen because of its early and concentrated fruit set as well as susceptibility to bacterial disease (2). PI 128216 is an accession of the United States Department of Agriculture (USDA) National Plant Germplasm System (NPGS) collection. Passport data in the Germplasm information network (GRIN) describes PI 128216 as an accession of *S. lycopersicum* collected in Bolivia in 1938. However, the accession morphologically appears to be *S. pimpinellifolium*, with characteristic foliage, flowers, current-sized fruit, and an indeterminate growth habit (22).

The two parents were crossed and the hybrid was backcrossed for two generations to OH88119 as the recurrent parent to create the BC$_2$ population, which consisted of 94 individuals. Two random progeny from each BC$_2$ individual were self-pollinated and the seed was saved separately to generate 188 BC$_3$S$_1$ lines. Individual families were then advanced by self-pollination and single-seed descent for four more generations. The BC$_3$S$_4$ population consisted of 178 lines because some were lost in the process of generating the succeeding filial generations (4). The theoretical expectation is that a low proportion (12.5%) of the donor *S. pimpinellifolium* parent is represented in each line and that each line is homozygous at 93.75% of its loci (32).

Additional segregating populations were screened for HR in the greenhouse to confirm marker-trait associations identified in the IBC population. Three individuals from the IBC population, 06.8040, 06.8068, and 06.8145, were chosen based on the presence of HR when inoculated with T3 strains and because they contained candidate loci associated with HR in the IBC population (Table 1). These three lines were individually backcrossed to OH88119 to develop three independent populations.

Progeny from four crosses were evaluated in the field for resistance to spot race T3. The four crosses used IBC lines as female parents and were 06.8145 × OH8245, 06.8145 × OH7530, 06.8145 × OH88119, and 06.8068 × OH88119. Cv. OH8245 was chosen based on its status as an elite parent for processing tomato hybrids (2,3) and observations that it possesses a nonblighting tolerance to spot when symptoms are present. Crosses were made to OH7530 in order to pyramid HR to race T3 with the resistance genes *Rx-3*, which provides an HR response to spot race T1, and *Pto*, which provides an HR response to bacterial speck race 0 (34). Hybrids from these crosses were self-pollinated to create F$_2$ populations for marker-assisted selection and disease evaluation.

**Experimental design.** Greenhouse evaluations of HR in segregating populations were based on comparisons to Ha7981 and PI 128216 as resistant controls and OH88119 as the susceptible control. The growth and care of greenhouse plants followed established methods (36). For the initial screening of the IBC population, 178 lines were evaluated in a randomized complete block (RCB) design with two replications. For confirmation of marker-trait associations, 48 individuals from the 06.8040 × OH88119 F$_2$ population, 48 individuals from the 06.8068 × OH88119 F$_2$ population, and 205 individuals from the 06.8145 × OH88119 F$_2$ population were evaluated in a completely randomized design with a single replication.

Disease evaluation was conducted in the field to test the hypothesis that the locus associated with the HR response in the greenhouse also provides resistance in the field. Plants were first sown in the greenhouse in 288-cell flats with plant care and growth conditions as previously described (36). Individual plants were genotyped with two markers, SL106151 and SL107371, which were associated with the HR response in the IBC population and confirming generations. Plants were then organized into genotypic classes (homozygous for the PI 128216 allele conditioning HR, heterozygous, and homozygous for the susceptible allele). Seedlings were spray inoculated (described below) and transplanted into the field in Wooster, OH 1 week following inoculation. To test the effect of genetic background, four segregating populations were tested, two in the OH88119 background and one each in the OH8245 and OH7530 background (as described above). The trial was established as a replicated RCB design with two replications consisting of 10 plants per plot.

**TABLE 1. P values of tests of marker association with hypersensitive reaction in the greenhouse at 24, 48, or 72 h after inoculation with bacterial spot (*Xanthomonas perforans*) race T3 in four segregating tomato populations.**

<table>
<thead>
<tr>
<th>Marker</th>
<th>Chromosome</th>
<th>24 h</th>
<th>48 h</th>
<th>72 h</th>
<th>06.8040</th>
<th>06.8068</th>
<th>06.8145</th>
</tr>
</thead>
<tbody>
<tr>
<td>LEOH200</td>
<td>6</td>
<td>0.0021</td>
<td>0.0039</td>
<td>0.0031</td>
<td>0.8234</td>
<td>NA</td>
<td>0.2507</td>
</tr>
<tr>
<td>LEOH112</td>
<td>6</td>
<td>0.0378</td>
<td>0.0067</td>
<td>0.0072</td>
<td>0.6772</td>
<td>NA</td>
<td>0.2932</td>
</tr>
<tr>
<td>SL107371</td>
<td>11</td>
<td>0.0434</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>NA</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>SL106151</td>
<td>11</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>NA</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>CoOH87</td>
<td>11</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>NA</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>SL20181</td>
<td>11</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>NA</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
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</table>

*a* Segregating populations are an initial BC$_2$S$_1$ population from OH88119 × PI 128216 (IBC population) or BC$_2$F$_2$ populations from backcrossing specific inbred backcross (IBC) lines 06.8040, 06.8068, and 06.8145 with OH88119. NA indicates that the marker was not segregating in the cross and could not be tested.

*b* Chromosome to which the marker maps based on a consensus linkage map derived from two F$_2$ mapping populations (Sun1642 × LA1589 and Yellow Stuffer × LA1589) (unpublished data).
Inoculum preparation and inoculation. The *X. perforans* race T3 strain used in the study was Xcv761 (16). Inoculum was produced by growing the bacteria on yeast, dextrose, and calcium carbonate (YDC) agar medium (14) at 28°C for 48 to 72 h. Bacterial cells were washed from the agar plates with sterile, double-distilled water and the suspensions were standardized to an absorbance at 600 nm of 0.15, a concentration of \( \approx 3 \times 10^8 \) CFU/ml. For evaluations of HR response in the greenhouse, inoculations were performed by leaf infiltration on two leaflets per plant. The bacterial suspension was injected on the underside of each young, fully expanded leaflet using a 3-ml syringe without a needle until the area of infiltration reached 1.5 cm in diameter. Control plants were also mock inoculated with sterile H₂O to provide a comparison. Plants were kept at 20 to 25°C in the greenhouse with supplemental lighting. For the field trial, plants were inoculated in the greenhouse 1 week prior to transplanting by covering the plants with a fine mist using a Preval Spray Gun (Preval Sprayer Division, Yonkers, NY) as previously described (34).

Disease scoring. Plants were scored for the HR response in the greenhouse using a qualitative scale whereas field symptoms were scored using a quantitative scale. In the greenhouse, plants were visually inspected for the HR response at 24, 48, and 72 h after inoculation. Each of the two inoculated leaves per plant was scored as HR or a susceptible response. The scores from both leaves were combined into a single score for each plant for data analysis. If one of the leaves showed no response and the other showed a clear response, the plant was scored according to the clear response. Single plants with ambiguous responses were excluded from the statistical analyses. In the field, plants were rated for disease severity on a per plot basis using the scale of Horsfall and Barratt (6), where 1 = 0, 2 = 0 to 3, 3 = 3 to 6, 4 = 6 to 12, 5 = 12 to 25, 6 = 25 to 50, 7 = 50 to 75, 8 = 75 to 87, 9 = 87 to 94, 10 = 94 to 97, 11 = 97 to 100, and 12 = 100% diseased tissue. Plots were rated twice, with the early rating timed to correspond to when 80% of plots had reached the mature green stage of fruit ripeness. The late rating was timed to correspond to when 80% of the fruit were ripe, the approximate time of harvest for maximum marketable yield and grade quality.

Molecular marker analysis and mapping. Procedures for DNA extraction, polymerase chain reaction (PCR), and electrophoresis used in the analysis of these populations were described previously (12,33,36). In total, 352 primer pairs, consisting of simple sequence repeat (SSR), insertion/deletion (indel), and single nucleotide polymorphism (SNP) markers, were screened for polymorphisms between the two parental lines of the BC₂S₅ population. We identified 70 markers polymorphic between OH88119 and PI 128216 and selected 53 markers scattered throughout the genome to genotype the IBC population. We identified 70 markers polymorphic between OH88119 and PI 128216 and selected 53 markers scattered throughout the genome to genotype the IBC population for maximum marketable yield and grade quality. Single plants with ambiguous responses were excluded from the statistical analyses. In the field, plants were rated for disease severity on a per plot basis using the scale of Horsfall and Barratt (6), where 1 = 0, 2 = 0 to 3, 3 = 3 to 6, 4 = 6 to 12, 5 = 12 to 25, 6 = 25 to 50, 7 = 50 to 75, 8 = 75 to 87, 9 = 87 to 94, 10 = 94 to 97, 11 = 97 to 100, and 12 = 100% diseased tissue. Plots were rated twice, with the early rating timed to correspond to when 80% of plots had reached the mature green stage of fruit ripeness. The late rating was timed to correspond to when 80% of the fruit were ripe, the approximate time of harvest

![Genetic linkage map location of Rx-4](image-url)
England BioLabs Incorporated, Ipswich, MA) in a cleaved amplified polymorphic sequence (CAPs) approach. Polymorphisms and sequence context for markers with the “SL” prefix are included in the supplementary material of Van Deynze et al. (28).

**Statistical analysis.** The extent of linkage and gametic phase disequilibrium was tested in the IBC population by calculating $R^2$ values and a $\chi^2$ goodness-of-fit test for independent assortment for all possible pairwise marker combinations. Pairwise $R^2$ values were calculated using GGT 2.0 software (27). Calculations for the $\chi^2$ test were based on an expected frequency of 0.125 for individuals homozygous for the PI 128216 allele at any marker locus in the IBC population because only 12.5% of the genome is expected to be from the donor parent in the BC$_2$S$_2$ generation. To visualize genomic locations in disequilibrium, the $P$ value for each pairwise test was coded by significance level and plotted in a heat map (Fig. 2).

In each segregating population, deviation from expected segregation ratios were tested using a $\chi^2$ goodness-of-fit test. In the F$_2$ populations screened for HR in the greenhouse, segregation of HR was tested against several ratios supporting both one-locus and two-loci hypotheses. These ratios were 3:1 (single locus, dominant gene), 9:7 (two loci, duplicate recessive), 15:1 (two loci, duplicate dominant), and 13:3 (two loci, dominant–recessive epistatic interaction) ratios.

**Fig. 2.** Heat map based on pairwise $\chi^2$ goodness-of-fit tests for independent assortment of markers in the PI 128216 inbred backcross population (BC$_2$S$_2$). Each column or row represents one marker and markers are ordered by genomic location on the reference map. Each cell indicates the $P$ value of the $\chi^2$ test between two markers as $P > 0.05$ (white), $0.05 \geq P > 0.01$ (light gray), $0.01 \geq P > 0.001$ (dark gray), or $P \leq 0.001$ (black). Several markers on chromosomes 6 and 11 failed to assort independently in the inbred backcross population.
Marker-trait associations were tested for significance using the nonparametric Kruskal-Wallis test as implemented in the SAS (version 9.1; SAS Software, Cary, NC) procedure NPAR1WAY with the Wilcoxon option specified. For the IBC population, each marker was tested against the 24-, 48-, and 72-h ratings for HR. To adjust for multiple comparisons, a marker had to have a P value < 0.01 in at least two time points to be declared significantly associated with the HR response. For the confirming generations in the greenhouse, the same analysis was performed except that each marker was tested against the 72-h ratings only.

Marker-trait associations were tested in the field evaluation using analysis of variance (ANOVA) implemented through the general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while genotypes were considered fixed effects, with the general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). 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Replications were considered random effects while general linear model (GLM) procedure in SAS (version 9.1; SAS Software). Replications were considere...
Three populations were developed from specific IBC lines to test the effect of the chromosome 6 and chromosome 11 locations on HR in the greenhouse. Of 205 plants in the 06.8145 × OH88119 population, 197 gave clear results. In both the 06.8040 × OH88119 and 06.8068 × OH88119 populations, 47 of the 48 plants tested were scored unambiguously. The HR response segregated in all three populations and did not significantly deviate from a 3:1 ratio in the 06.8040 and 06.8145 populations. A deviation from 3:1 ratio was suggested for the 06.8068 population \( (P = 0.005) \), with more susceptible progeny than expected. For this population, chromosome 11 marker distortion from the expected 3:1 ratio was also observed in the direction of the susceptible allele \( (P = 0.04) \). In the two small populations, we could not rule out alternative 13:3 ratios (06.8040 population, \( P = 0.118 \)) or 9:7 ratios (06.8068 population, \( P = 0.869 \)).

Nonparametric marker-trait analysis of the two populations segregating for PI 128216 alleles on chromosome 6 failed to confirm an association between the markers and the HR response (Table 1). All segregating markers on chromosome 11 were highly associated with HR \( (P < 0.0002) \) in all three confirming populations. Taken together, these results do not support a genetic effect from chromosome 6 but strongly implicate a locus on chromosome 11. The locus on chromosome 11, which we refer to as Rx-4, was sufficient to elicit an HR when challenged with the race T3 strain Xv/cs761.

In order to better characterize the location of Rx-4, we created a linkage map with additional markers on chromosome 11 (Fig. 1). The locus was mapped 20 centimorgans (cM) from SL10737i and 12 cM from SL10615i, the two markers initially associated with HR in the IBC population. Four markers (SSR637, SL10915, TOM144, and TOM196) mapped further away from Rx-4. CosOH57 mapped 7 cM away from the locus while 3 cM separated SL20181 and Rx-4. CosOH57 and SL20181 were tested for their association with HR in the IBC population and the three BC1F2 populations as well. As expected, these two markers were highly associated with HR in all populations where they segregated \( (P < 0.0001) \) (Table 1). The location of the markers linked to Rx-4 places the locus near the middle of the short arm of chromosome 11 (Fig. 1).

To determine whether Rx-4 also provides resistance in the field, four BC1F2 populations were grouped by genotype and evaluated for resistance to race T3. The severity of disease increased between early and late rating periods (Table 2). The susceptible control, OH88119, was the most susceptible while the resistant controls, PI 128216 and Ha7981, were among the most resistant in both ratings. The main effect of marker genotype was highly significant in both the early \( (P = 0.003) \) and late \( (P = 0.0001) \) ratings. The most resistant plots were either homozygous or heterozygous for the PI 128216 allele on chromosome 11, while the most susceptible were homozygous for the allele derived from OH88119, OH8245, or OH7530. In the early rating, plots homozygous for the PI 128216 allele had a mean rating of 1.9 and were more resistant than heterozygotes with a mean rating of 2.5 or homoygotes for the susceptible allele with a mean rating of 4.1.

In the late rating, however, PI 128216 homozygotes and heterozygotes were not significantly different, with mean ratings of 3.6 and 4.3, respectively. Both were significantly more resistant than homozygotes with the OH allele, with a mean rating of 7.9. The effect of Rx-3 was not significant in the 06.8145 × OH7530 F2 population in either rating \( (P > 0.78) \), nor was the interaction between Rx-3 and the PI 128216 locus on chromosome 11 \( (P > 0.70) \). These results confirm the race-specificity of these two loci.

The main effect of genetic background was also statistically significant in the early rating \( (P = 0.046) \). Regardless of marker class on chromosome 11, plots derived from crosses to OH8245 had a mean rating of 2.3 and were more resistant than the plots derived from crosses to OH88119, with a mean rating of 3.2. Plots derived from crosses to OH7530 had a mean rating of 2.5 and were not significantly different from the others. This effect was marginally nonsignificant \( (P = 0.108) \) in the late rating, with the mean ratings of the OH8245, OH7530, and OH88119 backgrounds as 4.7, 4.7, and 5.8, respectively (Table 2).

**DISCUSSION**

We identified molecular markers from PI 128216 that were associated with the HR response to bacterial spot race T3. Nonparametric single-marker analysis in IBC populations implicated two genomic locations. Subsequent testing of independent populations, however, confirmed the association of chromosome 11 with HR and did not support an effect from chromosome 6. Gametic phase disequilibrium between chromosome 6 and 11 in the IBC population provides an explanation for the spurious detection of association. These results illustrate the importance of confirming generations when mapping traits of interest.

In the two smallest confirming populations, 06.8068 × OH88119 and 06.8040 × OH88119, segregation ratios other than 3:1 could not be ruled out and 06.8068 × OH88119 did not fit a 3:1 ratio. Population size limited our ability to interpret gene numbers in these populations. A shift of only three individuals from the susceptible to the HR class would create a nonsignificant \( \chi^2 \) value relative to a 3:1 ratio. The 48 individuals screened from these populations were not sufficient to distinguish between a 3:1 and alternative segregation ratios. In addition, markers on chromosome 11 detected segregation distortion in the 06.8068 population in the direction of susceptible alleles. In contrast, when 200 individuals were screened from the 06.8145 × OH88119 population, the only ratio from which the population did not greatly deviate was 3:1, providing strong evidence for a single dominant gene. The locus on chromosome 11 alone was sufficient to confer the HR response, and no supporting data for a locus on chromosome 6 was evident. The importance of the locus on chromosome 11 was confirmed by the field results where the PI 128216 allele increased resistance to spot race T3. Taken together, these results indicate that a single, dominant gene on chromosome 11 from PI 128216 is sufficient to confer HR to bacterial spot race T3.

The nomenclature of *Xanthomonas* resistance loci in tomato is inconsistent in the literature. Current resistance loci include Rx-1, Rx-2, and Rx-3 which denote QTL from Ha7998 conferring HR resistance to race T1 (37). This designation was maintained in subsequent studies with regards to Rx-3, which was shown to confer resistance to race T1 in the field (36). A source of unmapped resistance to race T3 from Ha7981 has been named Xv3, where “Xv” refers to the species (*X. vesicatoria*) and the “3” refers to the race (15,24). This designation illustrates the problem with using taxonomic designations in the name because *Xanthomonas* taxonomy has been revised several times and T3 strains tend to be *X. perforans*, not *X. vesicatoria*. A source of HR to race T4 identified from LA716 has been named Xv4 (1). This locus name is also problematic for the reasons discussed above. In consultation with the tomato-breeding community, we propose that the locus identified in this study be denoted as Rx-4, which indicates resistance to *Xanthomonas*, locus number 4. This naming convention is not ideal because it does not provide information regarding the race or the species of bacteria. However, given that the taxonomy is in flux, we believe this name offers the best compromise.

Because Rx-4 conditions HR in the greenhouse, we designed an experiment to specifically test the effect of this locus under open-field conditions. This was accomplished by grouping individuals based on their genotype at markers linked to the locus. Our results indicate that Rx-4 also contributes to resistance in the field. In previous reports, HR was not highly correlated with resistance in the field (22). There are likely several factors, genetic and environmental, involved in field resistance. Our results from the early rating suggest, for example, that genetic background may
affect the level of resistance. Furthermore, data from the disease evaluation suggest that Rx-4 does not exhibit complete dominance for resistance in the field. It would be beneficial, therefore, to combine Rx-4 with other sources of resistance to Xanthomonas spp. as well as resistance to other bacterial pathogens.

The combination of Rx-4 with other sources of Xanthomonas resistance will require additional research in some cases. In this study, both PI 128216 and Ha7981 provide some resistance to spot race T3 in the field. Resistance to race T3 from Ha7981 has not yet been genetically characterized (7). Further characterization of the Ha7981 source is needed to determine how resistance can best be combined. Yang et al. (35) identified a QTL with large effect from PI 114490 that contributes to resistance to spot races T1, T2, T3, and T4 that was associated with the markers TOM144, TOM196, and SSR637 on chromosome 11. Although these markers are in the same general chromosomal region as Rx-4, the relationship between these loci is unknown. These two sources are most likely not allelic because resistance from PI 128216 is race-specific (T3) whereas resistance from PI 114490 is not race-specific and does not confer HR. Further analysis is required to determine the extent of genetic similarity between the two loci.

The markers identified in this study will be useful for marker-assisted selection for resistance to spot race T3. We demonstrated that families with PI 128216 alleles at SL10737i and SL10615i were more resistant in the field than families with susceptible alleles. This result suggests that gain can be made for resistance by indirect selection of marker loci. The use of CosOH57 or SL20181 is expected to be more efficient because they are more closely linked to Rx-4 than SL10737i and SL10615i.

Use of an IBC population to screen for traits for unimproved germplasm is beneficial because, once identified, the trait is in a genetic background that is more easily used in traditional breeding programs. Lines from our BCS5 population are expected to possess only 12.5% of their genome from the wild donor. The structure of the population also allows for replication, thus allowing more confidence in assessment of traits. We crossed resistant IBC plants with elite material for confirmation of marker-trait associations and to introduce the resistance into more diverse genetic backgrounds. This approach is sometimes referred to as simultaneous discovery and introgression. We have combined Rx-4 with Rx-3 and Pto. The tomato lines and markers reported here should be useful to develop resistant cultivars and to further investigate the basis of resistance.

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