Quantitative Trait Loci for Tomato Yellow Leaf Curl Virus and Tomato Mottle Virus Resistance in Tomato

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ABSTRACT. The genetic basis of resistance to tomato yellow leaf curl virus (TYLCV) and tomato mottle virus (ToMoV) was studied in three different mapping populations of tomato (Lycopersicon esculentum Mill.). Bulked segregant analysis (BSA) was used to identify random amplification of polymorphic DNA (RAPD) markers linked to TYLCV and ToMoV resistance. Segregated RAPD markers associated with resistance were linked to morphological markers self-pruning (sp) and potato leaf (p) on chromosome 6. RAPD genetic linkage maps of chromosome 6 were constructed for each of the three populations. Common mapped markers revealed straightforward homologies between the chromosome 6 linkage group of the three populations. Multiple-QTL mapping (MQM) was used to identify quantitative trait loci (QTL) for resistance linked to chromosome 6. These revealed that the resistance against TYLCV and ToMoV was mainly explained by two QTL in two populations and one QTL in another. For all of the resistance QTL detected, the favorable allele was provided by the resistant parents. The presence of three different sources of TYLCV and ToMoV resistance, and the markers in tight linkage with them, provide a means of systemically combining multiple resistance genes. Successful cloning of the R gene from tomatoes would lead to deeper understanding of the molecular basis of resistance to TYLCV and ToMoV, and might also shed light on the evolution of resistance genes in plants in general.

Received for publication 4 May 2005. Accepted for publication 16 Aug. 2005.
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J. Amer. Soc. Hortic. Sci. 131(2):267–272. 2006.

ABSTRACT. The genetic basis of resistance to tomato yellow leaf curl virus (TYLCV) and tomato mottle virus (ToMoV) was studied in three different mapping populations of tomato (Lycopersicon esculentum Mill.). Bulked segregant analysis (BSA) was used to identify random amplification of polymorphic DNA (RAPD) markers linked to TYLCV and ToMoV resistance. Segregated RAPD markers associated with resistance were linked to morphological markers self-pruning (sp) and potato leaf (p) on chromosome 6. RAPD genetic linkage maps of chromosome 6 were constructed for each of the three populations. Common mapped markers revealed straightforward homologies between the chromosome 6 linkage group of the three populations. Multiple-QTL mapping (MQM) was used to identify quantitative trait loci (QTL) for resistance linked to chromosome 6. These revealed that the resistance against TYLCV and ToMoV was mainly explained by two QTL in two populations and one QTL in another. For all of the resistance QTL detected, the favorable allele was provided by the resistant parents. The presence of three different sources of TYLCV and ToMoV resistance, and the markers in tight linkage with them, provide a means of systemically combining multiple resistance genes. Successful cloning of the R gene from tomatoes would lead to deeper understanding of the molecular basis of resistance to TYLCV and ToMoV, and might also shed light on the evolution of resistance genes in plants in general.
categories, which revealed that ToMoV resistance was controlled by more than one gene (Scott et al., 1995). ToMoV resistance was found to be correlated to indeterminate (sp+) plants, indicating possible linkage of resistance to the sp locus on chromosome 6. Later, this was confirmed by Griffiths and Scott (2001), who found that two resistance loci on chromosome 6 with additive effects conferred resistance from L. chilense accession LA 1932. ToMoV-resistant inbreds generally have been resistant to TYLCV, as found by Griffiths and Scott (2001).

The identification of molecular markers linked to resistance genes is particularly useful for both marker-assisted selection (MAS) and map-based gene cloning ( Tanksley et al., 1995). So far, a number of restriction fragment length polymorphism (RFLPs) and RAPD markers have been found to be linked to TYLCV and ToMoV resistance genes. The partially dominant major gene Ty-1 has been mapped to the 13-cM interval between RFLP markers TG97 and TG25 on the short arm of chromosome 6 (Zamir et al., 1994). Mapping a wild tomato L. hirsutum introgression associated with TYLCV resistance in a cultivated tomato (L. esculentum) line showed that chromosome 11 spanned RFLP markers TG36 to TG393, a distance of 14.6 cM (Hanson et al., 2000). Bulked segregant analysis (BSA) was used to identify four RAPD markers linked to TYLCV resistance in a 17.3-cM region on chromosome 6 (Chagué et al., 1997). Also, BSA was applied in a recent study (Griffiths and Scott, 2001) to identify RAPD markers linked to ToMoV resistance derived from L. chilense accession LA1932. Twelve markers segregated into two linked regions flanking either side of the morphological markers self-pruning (sp) and potato leaf (c) on chromosome 6. Griffiths (1998) also found that a third region encompassing the Ty-1 gene was associated with resistance to ToMoV in accessions LA 2779 and LA 1938. The objective of this present study was to search for evidence of novel QTL affecting TYLCV and ToMoV resistance from different tomato resources by applying MQM. This study was also performed to develop fine-scale genetic maps of chromosome 6 in each of three F2 populations.

Materials and Methods

Three tomato F1 mapping populations were developed for this study, using three different genetic resistance sources. Resistant inbreds derived from L. chilense genotypes LA1932, LA2779, and LA1938/Tykings were each crossed with susceptible Florida heat-tolerant breeding line Fla 7324. The developed populations were designated as 503, 506, and 509, respectively. Subsequently, F1 seed of each was bulked to obtain the three F2 progenies. Parents and F2 plants were grown in two sets of Todd planter flats (Speedling, Sun City, Fla.). TYLCV and ToMoV were field screened in Fall 2000 and Spring 2001 in the three F2 populations. One set was inoculated with whiteflies viruliferous for TYLCV, and the other set with whiteflies viruliferous for ToMoV, using the method described by Griffiths and Scott (2001). In this method, plants 20 d past the cotyledon stage are challenged with viruliferous whiteflies for 14 d. After inoculation, the whiteflies were killed and plants were planted in separate field plots for each disease. The plants were set on 84-cm-wide × 20-cm-high beds of EauGallie fine sand. The beds were spaced on 150-cm centers with seepage irrigation ditches spaced six beds apart. The blocks were arranged parallel to the beds and the irrigation ditches. The plants were spaced at 46 cm, with 91 cm between plots within the bed. Standard growing practices were followed (Hochmuth et al., 1999). Plots were grown randomly with 10 plants per plot for the parents, and 45 per plot for the F2 plants. Plants were rated for disease severity 45 d after the inoculation began. The rating scale was from 0 to 4, where 0 = no symptoms and 4 = severe symptoms and stunting.

DNA was extracted from the young leaves of 90 F2 plants of each population as described by Griffiths and Scott (2001). The strategy for identifying RAPD markers linked to resistance genes was described previously (Agrama and Moussa, 1996; Michelson et al., 1991). The polymorphic markers revealed by the previous work of Griffiths (1998) and Griffiths and Scott (2001) were used in this study to construct genetic linkage maps for the three populations and to integrate these maps.

Data were compiled from each population, and two-point linkages between all RAPD markers and the phenotypic sp marker were estimated. JoinMap 3.0 (van Ooijen and Voorrips, 2001) was used to construct maps of chromosome 6 for each of the populations and to construct an integrated map. To construct an integrated map of the three individual maps, the pairwise recombination frequencies for each cross were combined and one linkage group for the merged set was formed at logarithmic odds (LOD) ≥ 3.0. Threshold for declaring linkage was a LOD score of 4.0. Map units (cM) were derived using the Kosambi mapping function (Kosambi, 1944).

A computer software package, MapQTL version 3.0 (van Ooijen and Maliepaard, 1996), was used for interval mapping (Lander and Botstein, 1989). In the region of the putative QTLs (LOD > 2.5), the markers with the highest LOD values were taken as co-factors for running a MQM program method (Jansen, 1993; Jansen and Stam, 1994). A LOD value of 2.5 was chosen as the significant threshold value for declaring a QTL. A QTL was retained when statistics exceeded the threshold defined above for both the Kruskal–Wallis test and the QM. We arbitrarily considered that two QTLs had the same map position when LOD-score peaks were less distant than 20 cM (Lespinasse et al., 2000). QTLs were named by a five-letter abbreviation for the virus name followed by the number of QTL found.

Results

The three resistant parents were highly resistant to both TYLCV and ToMoV, as expected (Table 1). Since disease severity for some of the susceptible parents was less than 3, there were

| Genotype | Resistance source | Disease severity (mean ± se) |
|----------|-------------------|-------------------------------|
|          | ToMoV             | TYLCV                         |
| 503P1    | LA1932            | 0.0 ± 0.00                   |
| 503P2    | 3.5 ± 0.27        | 1.8 ± 0.65                   |
| 506P1    | LA2779            | 0.0 ± 0.0                    |
| 506P2    | 2.5 ± 0.23        | 2.6 ± 0.48                   |
| 509P1    | LA1938/Tykings    | 1.0 ± 0.05                   |
| 509P2    | 3.5 ± 0.00        | 3.3 ± 0.34                   |
| 503 F    | LA1932            | 1.4 ± 0.22                   |
| 506 F    | LA2779            | 1.6 ± 0.24                   |
| 509 F    | LA1938/Tykings    | 2.3 ± 0.17                   |

Rating scale was from 0 to 4, where 0 = no symptoms and 4 = severe symptoms and stunting.
some plants that escaped infection; it can be assumed that the F2 plants also had some susceptible escapes. The 506 F2 population segregated in a 3:1 ratio for indeterminate (sp); determinate (sp) plant habit with a χ² value of 0.010 (P = 0.76). Frequency distributions of TYLCV and ToMoV severity ratings 80 d after inoculation for the three F2 populations and their parents are presented in Fig. 1. Disease severity to TYLCV and ToMoV in the three F2 populations approximated normal distributions (Fig. 1).

BSA identified 83 repeatable polymorphisms among the three sources. Present study of these 83 loci yielded a total of 55 polymorphic markers among the three populations. JoinMap was used to generate three individual maps of chromosome 6. Of the 55 polymorphic markers, 28, 25, and 27 markers associated with TYLCV and/or ToMoV resistance on chromosome 6 in population 503, 506 and 509, respectively (Fig. 2). Population 506 was evaluated with the 25 linked RAPD markers together with the sp locus that is assigned to chromosome 6. When the three maps were compared with each other, the 506 linkage map was shorter than the other two.

RAPD bands that migrated to identical positions on the gel and segregated in both pairs of parents were considered to be homologous markers. Six mapped markers were homologous in all three populations. Twelve, nine, and 14 RAPDs were homologous in two of three populations: either 506 and 509, 503 and 506, or 503 and 509, respectively. The positions of homologous markers in the three F2 population linkage maps showed close conservation of marker order even though the populations were derived from different backgrounds. The marker order between maps was highly similar, with some minor rearrangements of markers at small intervals of less than 4.0 cM.

The traits were coded in the mapping analysis according to the observed disease severity. As a wide and continuous range of resistance scores in F2 plants was observed, the conclusion was drawn that the resistance was a multigenic trait. Scores for the disease screening were analyzed with Kruskal–Wallis rank test performance, which analyses only evaluated effects at the marker loci using a single QTL model. In the three genotypes, 503 (LA1932), 506 (LA2779), and 509 (LA1938/Tyking) differed in disease score. Significance ranged from P < 0.05 for TYLCV in LA2779 to P < 0.0001 for TYLCV2 in the 503 and 509 populations (Table 2). The unlinked markers did not show any association with the resistance in the three populations. To dissect and map the underlying genes, we performed QTL mapping on each data set for each disease separately. We applied MQM methods to map QTLs for resistance on chromosome 6 in the three populations. These revealed that the resistance against TYLCV and ToMoV was mainly explained by two QTLs in the 503 and 509 populations and one QTL in 506 (Table 2, Fig. 2).

QTLs for TYLCV and ToMoV in the population 503 (LA1932) were mapped in two different regions ≈45 cM apart. One major QTL (TYLCV 2), peak at marker 389-855, was detected at a LOD score higher than 7.0. It contributed to at least 46% of the phenotypic variance. The other QTL, TYLCV1, was considered major with a LOD score of 3.9, and contributed 34% of the phenotypic variance. Two major QTLs were detected for ToMoV (1 and 2) in the same positions as TYLCV with an approximate LOD score of 6.0, and contributed 30.7% and 39.3% of the phenotypic variance, respectively (Table 2). Only one QTL was detected for resistance to both diseases in population 506 (LA2779) on the far end of chromosome 6, with LOD scores of 3.4 and 4.1 for ToMoV and TYLCV, respectively (Table 2, Fig. 2). The contributions of this QTL were 29.8% and 47.5% of the TYLCV and ToMoV phenotypes, respectively. Two major QTLs were detected for disease resistance in population 509 (LA1938/Tyking). QTLs for TYLCV1 and ToMoV1 were mapped 15 cM apart in the center region of the chromosome, with LOD scores of 5.7 and 5.3, respectively. The second QTL contribution to the phenotypic variance was 35% and 29.7% for TYLCV2 and ToMoV2 with LOD scores of 4.7 and 3.8, respectively.

In summary, three different regions were found to be associated with resistance to TYLCV and ToMoV on chromosome 6 (Fig. 2). Flanking RAPD markers 697-1225 and 053-760 in background genotype LA2779 are good candidates for resistance screening to both viruses.

![Fig 1. Frequency distribution of phenotype rates for the tomato yellow leaf curl virus (TYLCV) and tomato mottle virus (ToMoV) virus resistance in the three F2 populations derived from L. chilense genotypes LA1932 (genotype 503), LA2779 (genotype 506) and LA1938/Tyk (genotype 509). Values of the resistant (P2) and susceptible (P1) parental lines are shown by arrows. Plants were rated for disease severity on 45 d after the inoculation. The rating scale was from 0 to 4, where 0 = no symptoms and 4 = severe symptoms and stunting.](image-url)
Likewise 389-855 and 169-1100 are good candidates for TYLCV resistance derived from LA1932, while 371-690 and 112-850 are good candidates in LA1938/Tyking genetic backgrounds. Two peak markers (035-775, 061-1050) were found to be associated with ToMoV resistance in the LA1932 population. However, markers 462-875 and 389-855 are good candidates in screening for resistance to ToMoV in LA1938/Tyking backgrounds. As expected, for all QTLs the favorable allele (i.e., the allele providing resistance) was inherited from the resistant parents.

Discussion

Bulk segregant analysis was applied previously to map RAPD markers linked to ToMoV resistance (Griffiths and Scott, 2001). The resistance was found to be associated with markers mapped to chromosome 6. Conventional genetic analysis and molecular genetic mapping in three F2 populations have found three regions for resistance to TYLCV and ToMoV. To our knowledge, this is the first report of different sources for resistance to geminiviruses in L. chilense. Saturation mapping of RAPD markers on
chromosome 6 surrounding the resistance genes in *L. chilense* identified markers more tightly linked to resistance than those previously found in one population (Griffiths and Scott, 2001). Also, it enabled the construction of individual genetic maps for each of three populations derived from different sources. Common mapped markers revealed straightforward homologies between the chromosome 6 linkage group of the three populations.

The two-gene nature of the new source of resistance makes it particularly useful in breeding for virus resistance in light of the common accessions to tomato breeding. The major gene nature also means that a genetic MAS can easily be identified. MAS would thus become an available tool for breeding TYLCV and ToMoV resistance wherever the disease is found in the world. We applied interval mapping and MQM methods to map QTL for resistance on chromosome 6 in the three populations. A major improvement in the accuracy of QTL mapping was achieved by using MQM where the “peak” markers were taken as co-factors. Therefore, QTLs identified using MQM methods were considered to be the most reliable (Jansen, 1993; Jansen and Stam, 1994). The information from the current linkage groups will be used in a further QTL analysis study with emphasis on geminivirus resistance associated traits. An integrated genetic linkage map of chromosome 6 will be generated based on common markers. It can also be used as a starting point for a positional cloning strategy, as has been reported for *Arabidopsis thaliana* (L.) Heynh. (Cao et al., 1997; Leyser et al., 1993), tomato (Brommonschenkel and Tanksley, 1997; Martin et al., 1993), *Oryza sativa* L. (Song et al., 1995; Yoshimura et al., 1998), grapevines (Dalbó et al., 2000), and *Zea mays* L. (Agrama et al., 2002).

Assuming the estimated physical size of the tomato genome as $8.33 \times 10^8$ bp (Peterson et al., 1998) and the genome size as 1300 cM (Grandillo and Tanksley, 1996; Tanksley et al., 1992) leads to an estimate of 640 kb/cM. If this estimate applies to the genomic region surrounding the resistance genes (R genes), there would be an opportunity to clone R genes by chromosome-landing strategy (Tanksley et al., 1995). Also, an alternative strategy is being pursued to clone (R) genes, such as cloning resistance gene-homologous by PCR (Michelmore, 1996; Staskawick et al., 1995). Nevertheless, high-resolution mapping of R genes will facilitate this or other strategies, especially if resistance genes are clustered in linked arrays on chromosomes (Witsenboer et al., 1995).

The presence of three different sources of TYLCV and ToMoV resistance, and the markers in tight linkage with them, provide a means of systematically combining multiple resistance genes. Successful cloning of R genes from tomatoes would lead to deeper understanding of the molecular basis of resistance to TYLCV and ToMoV, and might also shed light on the evolution of resistance genes in plants in general. The current research identified diagnostic tools that could be developed to screen tomato populations for presence of R genes without expensive and time-consuming progeny testing. Furthermore, R genes might be introduced into genomes of other susceptible lines where no, or only weak, TYLCV and ToMoV genetic resistance has been identified.

### Table 2. Detected QTL for tomato yellow leaf curl virus (TYLCV) and tomato mottle virus (ToMoV) resistance in the three *L. chilense* populations with proportion of the explained phenotypic variance (Exp%) and additive effect (Add) of the resistance allele. QTL were identified if the Kruskal–Wallis test or restricted multiple-QTL mapping (MQM) mapping was exceeded. QTL are presented per virus type from the disease test with the highest logarithmic odds (LOD) score.

| Population | QTL name | QTL interval | Kruskal–Wallis significance test | Restricted MQM mapping |
|------------|----------|--------------|----------------------------------|------------------------|
|            |          |              | Peak LOD score | Exp% | Add |
| 503        | TYLCV1   | 50.8–54.2    | **            | 3.9  | 34.1| 1.2 |
|            | TYLCV2   | 95.1–99.8    | ****          | 7.5  | 46.4| 1.7 |
|            | ToMoV1   | 51.2–64.1    | ****          | 6.2  | 30.7| 1.1 |
|            | ToMoV2   | 103.4–107.6  | ***           | 5.8  | 39.3| 1.5 |
| 506        | TYLCV    | 0.0–3.7      | *             | 3.4  | 29.8| 0.8 |
|            | ToMoV    | 0.0–4.8      | ***           | 4.1  | 47.5| 1.9 |
| 509        | TYLCV1   | 52.0–60.9    | ***           | 5.7  | 30.3| 0.9 |
|            | TYLCV2   | 104.1–113.0  | ****          | 8.1  | 35.0| 1.2 |
|            | ToMoV1   | 67.5–74.1    | ***           | 5.3  | 24.6| 1.0 |
|            | ToMoV2   | 94.0–101.2   | **            | 3.8  | 29.7| 0.8 |

*, **, ***Significant at $p < 0.05$, 0.001, 0.0005, or 0.0001, respectively, by Kruskal–Wallis test.
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Lycopersicon chilense