QTL mapping to anthracnose leaf blight resistance in tropical maize

Cristiane Romanek¹, Rodrigo Rodrigues Matiello¹*, Caroline de Jesus Coelho¹, Lilian Schafascheck¹, Danilo Fernando Guimarães Silva¹ and José Raulindo Gardingo¹

Abstract: The objectives were to map genomic regions associated with quantitative trait loci (QTL) to anthracnose leaf blight (ALB) in tropical maize, and to determine the effects of the loci on resistance to the pathogenic fungus Colletotrichum graminicola (Ces.). QTL analysis to ALB was carried out in a population of $F_{2:3}$ progenies resulting from the cross between the contrasting lines $L_{04-2}$ and $L_{95-1}$. Seventeen QTL were located by mapping analysis by composite interval on eight chromosomes. Four QTL mapped on chromosomes 9 (1 QTL) and 10 (3 QTL) were the most stable, detected in at least two experiments, and represented the most part of the phenotypic variation (27.7 to 54.3%) of resistance. The majority of QTL mapped were associated to specific severity evaluations/development stage of maize growth, and may explain the quantitative inheritance of resistance of tropical maize to ALB.

Key words: Disease resistance, Colletotrichum graminicola, composite interval mapping, Zea mays, AUDPC.

INTRODUCTION

Anthracnose leaf blight (ALB), caused by fungus Colletotrichum graminicola (Ces.) G. W. Wils., is present in all regions of maize cultivation, especially in hot and humid environments (Bergstrom and Nicholson 1999). The pathogen can infect different parts of the plant at several growth stages. Stalk rot and ALB are the most important forms of the disease (Badu-Apraku et al. 1987). The presence of ALB or stalk rot limits the high yield potential of susceptible genotypes (Matiello et al. 2013, Rezende et al. 2004). Smith (1976) reported reduction in grain yield caused by ALB ranging from 19 to 28% in maize hybrids and lines, respectively. On the other hand, higher reductions are estimated with stalk rot, which can amount 40% yield losses (Matiello et al. 2013).

Genetic resistance for maize ALB can be considered a more efficient and less costly control method when compared to fungicide control (Coêlho et al. 2001). However, the development of cultivars with genetic resistance to disease is costly and laborious, since it involves numerous selection cycles and favorable environmental conditions for the pathogen's development, as well as for the expression of resistance in segregating populations. Thus, QTL mapping (Quantitative Trait Loci) to disease resistance using molecular markers may accelerate the development of resistant genotypes, maximizing the yield potential of crops by means of marker-assisted selection (Pozar et al. 2009).
literature indicates that a few genes with additive and dominant gene action control resistance to ALB (Badu-Apraku et al. 1987). Rezende et al. (2004) reported a major gene as well as genes with minor additive and dominant controlling the resistance.

Studies aiming to know the nature and magnitude of the effects of resistance genes are important to guide the breeding procedures in order to introduce new resistance alleles in susceptible germplasm, and maximize the use of genetic variability in populations under selection (Pozar et al. 2009). Thus, determining the number, the nature of gene action, and the position of QTL in the maize consensus map can significantly contribute to the clarification of the genetic basis of resistance to ALB in tropical maize germplasm. In this sense, the objective of this study was to map and estimate the effects of resistance QTL to ALB in tropical maize germplasm in different environments and evaluation periods during the disease life cycle.

MATERIAL AND METHODS

Plant material

Inbred lines of tropical maize, belonging to the C. graminicola resistance breeding program of State University of Ponta Grossa, originated from south of Brazil maize landraces, were initially screened for stalk rot and ALB (Prochno et al. 2016). The ALB contrasting inbred lines L₉₀₄-2 (resistant) and L₉₉₅-1 (susceptible) were crossed to obtain the segregating population of F₂ generation. Individuals were selfed to generate the F₂₃ progenies, which were evaluated for resistance to ALB in three field experiments.

Evaluation of maize resistance to anthracnose leaf blight

F₂₃ maize progenies were evaluated in three experiments sown on 01/10/2012, 11/19/2013 and 01/17/2014, at the Farm School “Capão da Onça” (lat 25° 05’40” S, long 50° 03’ 13” W and alt 1017 m asl), Ponta Grossa, PR, Brazil. Treatments consisted of 121 F₂₃ progenies plus the parents (L₀₄-2 and L₉₅-1) and F₁ generation, which were used as experimental control. Experiments were carried out in a randomized block design with three replications. The plot consisted of a 3 m row, spaced 0.90 m between rows, with approximately 15 plants per plot.

The isolated of C. graminicola was given by Dow AgroSciences Ltda. (Jardinópolis, São Paulo), as culture medium discs form with fungus colonies. The multiplication of pathogen was realized in oat-agar culture medium (10 g of oat flour, 2.5 g of agar and 250 mL of distilled water), being maintained at room temperature on the lab bench, until complete sporulation of colonies. For inoculum preparation, 20 mL sterile distilled water were added to Petri dishes containing the sporulated colonies, followed by surface scraping in order to release conidia. The suspension was adjusted to 6 x 10⁵ conidia mL⁻¹ concentration with a Neubauer chamber. A drop of spreader Tween 80° was added to each liter of the conidial suspension. Plants were inoculated twice between the stages V6 and V7, by spraying 7 mL of the spore suspension in the plant whorl, using a knapsack sprayer calibrated at a constant pressure of 35 lbs pol⁻², by compressed CO₂.

The evaluation of maize progenies regarding resistance to ALB was carried out from the appearance of the first disease symptoms, at the fifteenth day after the first inoculation. Three severity evaluations were carried out at

![Figure 1. Anthracnose leaf blight (Colletotrichum graminicola) symptoms. Recognition of score scale (1 to 6) used in field evaluation.](image-url)
VT stage (tasseling), R2 stage (blister), and R3 stage (milk) (Ritchie and Hanway 1989). The characterization of resistance of maize progenies to ALB was carried out by the rating scale (% severity) proposed by Silva et al. (1986), ranging from 1 (highly resistant) to 6 (highly susceptible) (Figure 1). From the means of the scale grades of the three evaluations, was calculated the area under the disease progress curve (AUDPC). Afterwards, was carried out the frequency distribution analysis of maize progenies for AUDPC. The number of phenotypic classes (k) was estimated by the formula \( k = \sqrt{n} \), in which \( n \) refers to the total progenies (\( n = 121 \)). The phenotypic range class (PRC) was determined by \( PRC = \frac{R}{k} \), in which \( R \) is the range of the AUDPC found among the progenies. Phenotypic means of progenies for the 1st, 2nd, 3rd evaluations and for the AUDPC in the respective experiments were used for QTL mapping. To verify the distribution normality was applied Shapiro-Wilk test (SAS Institute 2015).

**Molecular markers**

Genomic DNA was extracted from leaves of individual plants of the \( F_2 \) generation, of the inbred lines (L\(_R\)04-2 and L\(_S\)95-1), and of the \( F_1 \) generation (L\(_R\)04-2 x L\(_S\)95-1), following the protocol of Hoisington et al. (1994). Eighty-eight SSR loci, distributed in 10 maize chromosomes, were amplified in the parental inbred lines in order to select polymorphic loci. SSR loci were selected from the Maize Genetics and Genomics Data Base - MaizeGDB (http://www.maizegdb.org). Polymerase chain reaction (PCR) was prepared in 0.2 mL microcentrifuge tubes containing: 1X buffer Invitrogen\textsuperscript{TM}; 2.0 mM MgCl\(_2\); 0.1 mM dNTP solution (100 mM, Amresco\textsuperscript{®}); 0.2 µM of each forward and reverse primer; 0.75 U Taq DNA polymerase (5 U/µL, Invitrogen\textsuperscript{TM}); and 20 ng DNA template for a total volume of 20 µL. The amplified samples were stained with 4 µL GelRed 0.1X plus loading buffer (1: 1), and separated by electrophoresis on 2% agarose gel. Of the 88 SSR loci, 29 were used to genotype the mapping population. The 121 individuals of the mapping population (\( F_2 \)) were genotyped for each polymorphic locus with codes. (1) Amplified fragment corresponding to the SSR allele from the resistant line (L\(_R\)04-2), (-1) amplified fragment corresponding to the SSR allele from the susceptible line (L\(_S\)95-1) and (0) presence of the two fragments corresponding to the alleles of the parental lines L\(_R\)04-2 and L\(_S\)95-1.

The amplification methodology of the AFLP marker was performed according to Vos et al. (1995). Initially, it was tested 40 combinations of the primers EcoRI and MseI in order to select those with the best amplification pattern, higher polymorphism between the parental lines and between the DNA bulks, according to the methodology of bulked segregant analysis (BSA) proposed by Michelmore et al. (1991). Of the 40 primer combinations, 19 were selected for genotyping the mapping population: E32-M48, E32-M49, E32-M50, E32-M51, E32-M52, E32-M53, E32-M55, E32-M59, E32-M60, E35-M50, E35-M56, E40-M50, E40-M51, E40-M51, E40-M60, E42-M50, E42-M51, E42-M60, E44-M51, E44-M56 [such as the primers sequence in Keygene (2004)]. Only the polymorphic combinations of each EcoRI + MseI combination were genotyped in (1) presence and (0) absence.

**Linkage analysis and composite interval mapping (CIM)**

SSR and AFLP loci that presented expected Mendelian segregation (undistorted) by Chi-square at 5% were used for the construction of linkage map with the aid of the Mapmaker software version 3.0 (Lander et al. 1987). The linkage group construction was based on command “group” using a LOD of 3.0 (maximum likelihood method) and maximum recombination frequency of \( \theta = 0.20 \). The recombination frequencies were converted on map distances by Haldane (1919)’s mapping function. After the linkage groups were defined, the markers inside each group were ordered by “order” command with LOD higher than 3.0. To ensure the correct marker order, the “ripple” command was used.

Analyses of the QTL mapping were carried out using the means of the disease severity grades of the three evaluations and of the AUDPC of the 121 \( F_{2.3} \) maize progenies in the three experiments. The location of the QTL was carried out through the analysis by the composite interval mapping (CIM), using the QTL Cartographe\textsuperscript{®} software (Basten et al. 2003). Linkage map was used for the QTL positioning. The QTL position, genetic effect, and phenotypic variation (partial correlation, %) were established at the 3.0 LOD peak.

**RESULTS AND DISCUSSION**

**Resistance of progenies to anthracnose leaf blight**

The analysis of frequency distribution confirmed the phenotypic contrast for resistance to ALB between the lines
L₀ 04-2 and Lₛ 95-1, regardless of the experimental year (Prochno et al. 2016). The frequency distribution for AUDPC presented a normal distribution by Shapiro-Wilk test at 5% of probability. Also was observed an asymmetric pattern, evidencing a trend of F₂:₃ progenies to greater resistance to ALB in the three experiments, with higher frequency between the 2nd and 4th classes of AUDPC. In the three experiments, highly resistant maize progenies were found, similarly to the source of resistance L₀ 04-2 (Figure 2).

Linkage map

Of the 413 polymorphic fragments (SSR and AFLP), only 54% with Mendelian segregation (undistorted) were used in the construction of the linkage map, avoiding errors in the position of marks on map and generate false positive associations (Liu 1998). The 11 SSR loci allowed co-locating AFLP fragments in the maize consensus map by the linkage analysis. Results allowed assigning chromosomal reference to the ten maize linkage clusters by the presence of at least one SSR locus per chromosome (Figure 3). The range of the length of the linkage clusters (107.5 to 244.3 cM) and the total length of the linkage map (1710.1 cM) are in agreement with other genetic maps of tropical maize (Mangolin et al. 2004). The linkage map of Castiglioni et al. (1999) demonstrated greater coverage of the genome, with 2057 cM, and with markers distributed every 6.6 cM. This result can be attributed to the greater number of individuals genotyped (229 families), enabling larger gametes sampling, and consequently greater probability in the detection of recombination events. The mean distance between the markers obtained in the present study was 11.4 cM, which shows good coverage of the genome analyzed in the segregating population (L₀ 04-2 x Lₛ 95-1).

QTL mapping

The composite interval mapping (CIM) allowed the mapping of a large number of genomic regions (17 QTL) associated with resistance to ALB. In some intervals, coincident QTL were mapped for the different forms of disease quantification, as well as between the evaluation environments (Table 1). The absence of coincident QTL in the different evaluation environments characterizes the occurrence of QTL x environment interactions, since the mapped QTL at a given interval cannot be expressed in other evaluation environment. Higher accuracy of QTL effects by the CIM is expected for several evaluation environments, since the phenotypic values of progenies in these environments are simultaneously taken into consideration, enabling the quantification of the QTL x environment interactions (Jiang and Zeng 1995).

There are few published studies on QTL mapping to ALB in maize. On the other hand, there are several studies on stalk rot; however, they are not conclusive. The literature shows wide divergence in the number of genes that control resistance to stalk rot in maize. Carson and Hooker (1982) mapped genes or gene units on chromosomes 1, 4, 6 and 8 using the reciprocal translocation technique. However, Jung et al. (1994), by using the molecular marker RFLP, identified only one genomic region located on chromosome 4 with strong association to a QTL responsible for over 50% of the phenotypic variation in resistance to anthracnose.

In recent years, several studies have been carried out aimed at identifying genomic regions associated with pathogen
Table 1. LOD score, estimates of additive effect, and proportion of phenotypic variation explained by the QTL mapped by the mapping interval composite methodology

<table>
<thead>
<tr>
<th>QTL Number</th>
<th>Chr</th>
<th>Markers Interval/Position (cM)</th>
<th>1st Experiment</th>
<th>2nd Experiment</th>
<th>3rd Experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>E1</td>
<td>E2</td>
<td>E3</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>E32M48_308/177.9 E42M50_174/189.4</td>
<td>LOD R² (%)</td>
<td>31.3</td>
<td>90.6</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>E35M56_680/0.0 E35M56_112/14.1</td>
<td>LOD R² (%)</td>
<td>31.6</td>
<td>94.1</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>E42M51_162/0.0 E42M50_76/7.6</td>
<td>LOD R² (%)</td>
<td>10.8</td>
<td>70.3</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>E32M48_167/51.0 E32M59_104/61.4</td>
<td>LOD R² (%)</td>
<td>10.7</td>
<td>72.5</td>
</tr>
<tr>
<td>5</td>
<td>4</td>
<td>E35M60_87/0.0 E32M60_185/10.4</td>
<td>LOD R² (%)</td>
<td>15.0</td>
<td>72.6</td>
</tr>
<tr>
<td>6</td>
<td>4</td>
<td>E32M52_73/15.3 E44M51_84/34.8</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>7</td>
<td>4</td>
<td>Umc1511/88.1 E32M53_434/119.3</td>
<td>LOD R² (%)</td>
<td>15.0</td>
<td>72.6</td>
</tr>
<tr>
<td>8</td>
<td>4</td>
<td>E32M53_434/119.3 E44M51_135/137.7</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>9</td>
<td>5</td>
<td>E32M48_532/242.0 E32M50_139/244.3</td>
<td>LOD R² (%)</td>
<td>15.0</td>
<td>72.6</td>
</tr>
<tr>
<td>10</td>
<td>8</td>
<td>E35M60_80/0.0 E32M50_100/23.9</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>11</td>
<td>8</td>
<td>E32M60_94/57.3 E32M50_248/74.1</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>12</td>
<td>8</td>
<td>E35M60_86/85.5 Pho15/107.5</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>13</td>
<td>9</td>
<td>E32M48_562/126.4 E32M48_97/157.7</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>14</td>
<td>9</td>
<td>E32M51_314/179.1 E35M56_174/201.1</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>15</td>
<td>10</td>
<td>E32M49_698/28.3 E32M59_207/58.7</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>16</td>
<td>10</td>
<td>E32M50_118/85.4 E44M56_81/109.1</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
<tr>
<td>17</td>
<td>10</td>
<td>E32M59_76/161.6 Umc1084/191.7</td>
<td>LOD R² (%)</td>
<td>43.1</td>
<td>94.0</td>
</tr>
</tbody>
</table>

Chr: Chromosome; E1: 1st evaluation; E2: 2nd evaluation; E3: 3rd evaluation; AUDPC: area under disease progress curve; LOD: LOD score that sustain the QTL position; R²: proportion of phenotypic variation explained by QTL at position considering H3 (a ≠ 0, d ≠ 0) in relation to H0 (a = 0, d = 0); a: estimative of additive effect of QTL under H3; d: estimative of dominance deviations effect under H3.
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resistance alleles in agricultural species (Wisser et al. 2006). For McMullen and Simcox (1995), loci that determine disease resistance are not randomly distributed in the genome, but in the form of clusters. Often the same chromosomal segments have been associated with resistance to multiple pathogens in maize. For instance, on chromosome 3, in bins 3.04 and 3.05, QTL were mapped for resistance: to *Puccinia sorghi*, *Cercospora zeae-maydis*, *Cochliobolus heterostrophus*, and *Exserohilum turcicum* (Wisser et al. 2006). This fact suggests that phytochemical compounds which stimulate resistance to different diseases can be produced by the same QTL, although it is possible the existence of QTL specific for a given pathosystem in maize (McMullen and Simcox 1995).

Prior knowledge on different QTL that have already been mapped in the 10 maize chromosomes/bin (chromosome splitting) enables establishing possible relations between the QTL to ALB mapped in this work. Thus, the QTL 1 of Chr 1 can be analyzed in relation to the position of the interval of the marker Bnlg1007, located in the bin 1.02 (Table 1 and Figure 3). The distance between the interval to the marker Bnlg1007 is between 109 cM and 120.5 cM, which is approximately six bins, since one bin corresponds to approximately 20 cM (McMullen and Simcox 1995). Thus, the QTL would be located near the bin 1.08, which is a region where QTL to stalk rot caused by *Gibberella zeae* (Pé et al. 1993) and *Cochliobolus heterostrophus* (Balint-Kurti and Carson 2006) had already been mapped.

The position of the QTL mapped on chromosome 2 (QTL 2), can be related to the locus Bnlg125 (bin 2.02). The distance between the interval and the Bnlg125 is between 68.7 cM and 82.8 cM, corresponding to approximately four bins, and thus it is possible that this QTL is near the bin 2.06 (Table 1). In this region, QTL have been mapped for important foliar diseases, such as *Exserohilum turcicum* (Welz et al. 1999a) and *Cercospora zeae-maydis* (Lehmensiek et al. 2001). On Chr 3 two QTL (QTL 3 and 4) were mapped on 2nd evaluation at 2nd experiment explaining from 75 to 90.6% of phenotypic variation (Table 1).

Four QTL were mapped on Chr 4 (intervals E35M60_87 – E32M60_185, E32M52_73 – E44M51_84, Umc1511 – E32M53_434, and E32M53_434 – E44M51_13S), which demonstrated specificity to severity evaluation or experiments (Table 1). To infer the approximate position of the QTL 6, was taken into account the position of this interval in relation to the locus Umc1511 (bin 4.05). The markers of this interval are at 72.8 and 53.3 cM, representing approximately three four bins. According to these relations, this QTL may be located near the bin 4.01/4.02. In this chromosome, QTL to stalk rot caused by *Gibberella zeae* was mapped in bin 4.01 (Pé et al. 1993), and by *Colletotrichum graminicola* next to bin 4.08 (Jung et al. 1994, Broglie et al. 2011), as well as to foliar lesions caused by *Phaeosphaeria maydis* (Moreira et al. 2009) and *Cercospora zeae-maydis* (Lehmensiek et al. 2001). In addition, the QTL 7 is possibly next to bins 4.05 or 4.06, which are genomic regions known for QTL to Cercospora leaf spot (Bubeck et al. 1993) and northern leaf blight in maize (Welz et al. 1999b).

The CIM analysis mapped a QTL on Chr 5 on markers interval E32M48_532 – E32M50_139, which explain 61.8% of phenotypic variance of resistance to ALB only on 2nd evaluation at 2nd experiment (Table 1). On Chr 8 were mapped three QTL. From these, the QTL 11 was the most stable, being mapped at the three experiments, responsible for the major part of resistance (23.4 to 94.0%) (Table 1 and Figure 3). This QTL is located at a distance of 33.4 cM to 50.2 cM for the Phi015 microsatellite locus (8.03), close to the bins 8.05/8.06. The region of Chr 8 that comprises these bins is known to be associated with QTL and resistance genes to various pathogens in maize. QTL in this region were mapped in different populations for resistance to *Exserohilum turcicum* (Welz et al. 1999a, b, Zwöhniter et al. 2010), *Cochliobolus heterostrophus* (Balint-Kurti et al. 2008), *Cercospora zeae-maydis* (Bubeck et al. 1993), *Puccinia sorghi* (Brown et al. 2001, Kerns et al. 1999), *Ustilago maydis* (Lüebberstedt et al. 1998), and to aflatoxin accumulation in maize caused by *Aspergillus flavus* (Paul et al. 2003).

From two QTL mapped on Chr 9, the QTL 13 was the most important for resistance in all study environments, presenting significant additive effect to reduce the disease severity (-0.90 to -26.79). These QTL is located at a mean distance of 47.5 cM from the Phi065 marker (9.03) (Table 1 and Figure 3). Thus, this QTL would be located around the bins 9.05/9.06. In this bin, it was mapped QTL for *Exserohilum turcicum* (Brown et al. 2001), and in the regions 9.03/9.04, it was mapped QTL for *Cochliobolus heterostrophus* (Zwöhniter et al. 2010).

The QTL 15 on Chr 10 has the SSR Umc1084 (10.07) marker as reference (Table 1 and Figure 3). The mean distance of the interval for the SSR is 148.2 cM, which is approximately 6 bins. Therefore, the QTL is possibly located near the bin 10.01, which is the region where QTL were mapped to common rust (Kerns et al. 1999). The QTL 17 (10.07) was the
most stable, because was associated to resistance in 8 from 12 evaluations (considering the three experiments). Since there are no reports of other QTL in the region, possibly the resistance locus of bin 10.07 is specific to ALB in maize. Additionally, in this chromosome, it was detected other resistance loci, such as in bin 10.05/10.06 to common rust (Lübberstedt et al. 1998), to Cercospora leaf spot (Bubeck et al. 1993), and to stalk rot caused by Giberella zeae in 10.05 (Pé et al. 1993). Other QTL (16) was mapped on this chromosome, responsible from 29.6 to 52.1% of ALB resistance, highlighted with negative additive effect.

In order to use efficiently a scheme of selection assisted by molecular markers in a breeding program, it is necessary to have markers strongly associated with the QTL of interest, which is a pronounced effect on the phenotypic expression, and stable for the evaluation environments. In this study, it was mapped four stable QTL for the different evaluation environments.

**Figure 3.** LOD profiles and 1-LOD and 2-LOD support intervals for E2 (2nd evaluation), E3 (3rd evaluation) and AUDPC (area under disease progress curve) QTL on chromosome 1, 8, 9, and 10. Only QTL present in at least two experiments were represented. The dotted line at LOD 3.0 represents the LOD threshold.
environments for AUDPC on Chr 9 and 10 (Table 1 and Figure 3). These QTL may be strong candidates for assisted selection schemes for resistance to ALB in tropical maize germplasm. Tropical maize populations have broad genetic basis compared with those of temperate climate zones. Besides, tropical areas are more likely to undergo environmental stress than temperate areas (Ribaut et al. 1997). Thus, QTL mapping in tropical genotypes enabled identifying new genomic regions and resistance alleles, which are possibly not yet known to the scientific community. In Brazil, this is the first report of quantitative inheritance to ALB resistance from tropical maize germplasm. Possibly the wide genetic base of this germplasm would explain the large number of QTL for resistance to ALB (Prochno et al. 2016).

Interactions of QTL x severity evaluations can be explained by climatic differences of the 1st and 3rd experiment in relation to the 2nd field experiment. High relative humidity / temperature conditions and higher concentration of inoculum of C. graminicola in the experimental areas of the 1st and 3rd experiments enabled the CIM analysis to the most same QTL to ALB resistance in both experiments. Although the AUDPC use is more efficient in relation to specific quantifications of the disease, its application in breeding programs is too restrictive by difficulty of performing several disease evaluations (at least three) in the field (Lopes et al. 2011). Thus, the identification of the ideal time for severity evaluation, which enables to adequately represent the reactions of different genotypes during the development of the epidemic could be the better strategy for selecting genotypes with high genetic resistance to ALB in this pathosystem. Our results demonstrated that the ALB evaluation from R2 (Blister) stage on second growing season at South of Brazil, enabled the better genetic resistance discrimination between individuals of segregating population (L²₉ 04-2 x L₉5-1), being confirmed by similarity of results from QTL mapped for AUDPC in relation to observed for 2nd evaluation. Thus, to improve the efficiency on selection of ALB resistant genotypes by South of Brazil maize breeding programs we recommend at least one evaluation at R2 stage in segregating populations inoculated with C. graminicola.

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REFERENCES


Jung M, Weldekidan T, Schaff D, Paterson A, Tingeys S and Hawk J (1994) Generation means analysis and quantitative trait locus mapping
of anthracnose stalk rot genes in maize. *Theoretical and Applied Genetics* 89: 413-418.


