Discovery of Genomic Regions Associated With Resistance to Late Wilt Disease Caused by *Harpophora Maydis (Samra, Sabet and Hing)* in Maize (*Zea mays* L.)

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Abstract

Late wilt disease (LWD) caused by *Harpophora maydis* (Samra, Sabet and Hing) is emerging as major production constraint in maize across the world. As a prelude to develop maize hybrids resistance to LWD, genetic basis of resistance was investigated. Two $F_{2:3}$ mapping populations (derived from CV156670 × 414-33 (P-1) and CV143587 × CV143587 (P-2)) were challenged with LWD at two locations (Kallinayakanahalli and Muppadighatta) during 2017 post-rainy season. Wider range of LWD scores were observed at both locations in both the populations. LWD response was influenced by significant Genotype × location interaction. Six and 56 $F_{2:2}$ progeny families showed resistance level better than resistant parent. 150 and 199 polymorphic SNP markers were used to genotype P-1 and P-2, respectively. Inclusive composite interval mapping was performed to detect significant QTL, QTL × QTL × Location interaction effects. Three major and four minor QTL controlling LWD resistance were detected on chromosome-1. Position and effect of the QTL varied with the location. Significant di-QTL interactions involving QTL with significant and/or non-significant effects) located within and between all the ten chromosomes were detected. Five of the seven detected QTL in our study showed significant QTL × location interaction. Though two major QTL ($q\text{-}lw\text{-}1.5$ and $q\text{-}lw\text{-}1.6$) with lower Q×L interaction effects could be considered as stable, their phenotypic variance is not large enough to deploy them in MAS. Based on these results, strategies to breed maize for resistance to LWD are discussed.

Introduction

Maize productivity in India is constrained by several prevalent and emerging insect pests and diseases. Of these, post-flowering stalk rot (PFSR) is a complex disease caused by three pathogens, namely, *Fusarium moniliforme* J. (Sheld), *Macrophomina phaseolina* (Tassi) Goid and *Harpophora maydis* (Samra, Sabet and Hing) (Khokhar et al. 2014). Stalk rot caused exclusively by seed-borne and soil-borne pathogen *H. maydis* causes premature wilting symptoms at post-flowering stage, a condition known as Late Wilt Disease (LWD). Typical symptoms of LWD include pre-mature wilting of leaves, discoloration of the stalk followed by stalk tissue disintegration and fibrousness in advanced stages (Samra et al. 1963). LWD has been reported in about 10 countries, with significant economic losses in Egypt (Sabet et al. 1961), India (Payak et al. 1970), Spain and Portugal (Molinero-Ruiz et al. 2010) and Israel (Drori et al. 2013). Based on survey, economic losses of 40% have been reported in Egypt (Labib et al. 1975; Galal et al. 1979), up to 100% in Israel (Degani et al. 2019) and up to 51% in India (Johal et al. 2004) due to LWD. However, 3.5 to 38.4% loss of grain yield attributable to soil inoculation by *H. maydis* has been reported based on an empirical study using a limited number of genotypes in Egypt (El-Naggar et al. 2015). The Egyptian, Indian, and Hungarian isolates have been reported to differ in their morphology, pathogenicity, and route of infection (Warren 1983). Very recently, we have reported 5.8 to 44.2% grain yield losses attributable to LWD by creating artificial epiphytotic condition through injecting *H. maydis* inoculum into the stem internode (Sunita et al. 2020) in India.

Genetic intervention is regarded as eco-friendly and cost-effective option to mitigate losses caused by LWD (El-Shafey et al. 1988; Zeller et al. 2000). As a prelude to genetic intervention, stable sources of resistance to *H. maydis* (Shekhar et al. 2010; Rakesh et al. 2016b; Aruna 2017; Mahesh 2019) have been identified. The inheritance of resistance to LWD has been reported as complex with significant genotype × environment interaction (Shekhar et al. 2010; Rakesh 2016; Aruna 2017; Mahesh 2019). Hence, direct selection for LWD resistance is likely to be less effective. Nevertheless, DNA markers could be used as effective surrogates of such traits in maize for which *apriori* identification and validation of closely linked markers is essential. As a step towards this, Quantitative Trait Locus/Loci (QTL) conferring LWD tolerance (Rakesh, 2016) on chromosomes 1,3,5,7 and 10 have been detected. The objectives of our study were to (i) validate the QTL detected by Rakesh (2016) (ii) to detect and estimate size effects of new LWD resistance governing QTL, if any in two different genetic backgrounds and (iii) detect and estimate size effects of their interaction with two spatial environments.

Material And Methods

Basic genetic material and development of mapping populations

The basic genetic material consisting of one LWD resistant inbred line, CV156670 and two LWD susceptible inbred lines, 414 – 33 and CV143587, developed and being maintained at Bayer Crop Science (erstwhile Monsanto India Ltd.) were used as parents to develop two connected mapping populations. For intellectual property protection reasons, the pedigree of these inbreds is not included. The two $F_1$’s (available with Bayer Crop Science), CV156670 × 414 – 33 and CV156670 × CV143587 were selfed during 2017 summer season to derive $F_2$ populations. Two $F_2$ populations derived from connected crosses CV156670 × 414 – 33 and CV156670 × CV143587 were selfed during 2017 rainy season. These $F_{2:3}$ populations constituted inter-connected mapping populations to detect QTL controlling LWD resistance.

Phenotyping mapping populations for responses to LWD

The mapping populations were evaluated at two locations, Kallinayakanahalli (KNH) 13.464013°N, 77.519178°E and Muppadighatta (MPG) 13.241161°N, 77.483637°E. 300 $F_{2:3}$ progeny families each from two $F_2$ populations, parental inbred lines and checks (susceptible and resistant check) were sown in single rows of 4 m length with 0.6 m spacing between rows and 0.2 m spacing between plants in two replications following randomized complete block design (RCBD) during 2017 post-rainy season. Recommended package of practices were followed to establish good crop stand till artificial inoculation. The $F_{2:3}$ progeny families were screened for responses to LWD using artificial inoculation protocol (Rakesh et al. 2016a).

Artificial inoculation of *H. maydis* inoculum

Naturally infested LWD stalk specimens collected from commercial maize production fields were used for isolation of pathogens and inoculum preparation. The inoculum concentration was assessed using haemocytometer and was adjusted to $4 \times 10^6$ spore ml$^{-1}$. A hole was made at the second internode on the stem of plants using a proprietary specialized stem jabbers and 2 ml of the *H. maydis* was injected to the holes using a syringe twice at 45 and 55 days after
sowing (DAS). From each progeny family, 20 plants were inoculated. All the recommended production practices were followed except for the application of potassium fertilisers and fungicides to maintain the plants after inoculation.

**Sampling and data recording**

The stalks of the artificially inoculated plants were split open at 110 DAS to examine the typical symptoms of LWD. Disease symptoms were manifested by discoloration, disintegration and fibrousness at the inoculated internode. Based on the coverage of disease symptom within an internode and its spread across internodes, the data on LWD severity was recorded on all the inoculated plants using modified 1–9 scale (Table 1) (Rakesh et al. 2016a).

**Table 1**

ANOVA of mean LWD scores in two F$_{2:3}$ populations across two locations

| Source of variation | CV156670 × CV143587 (P-1) | CV156670 × CV1414-33 (P-2) |
|---------------------|-----------------------------|-----------------------------|
|                     | Degrees of freedom | KNH | MPG | Degrees of freedom | KNH | MPG |
| Mean sum of squares | 'F' probability | Mean sum of squares | 'F' probability | Mean sum of squares | 'F' probability |
| Between F$_{2:3}$ families | 235 | 0.71** | <0.001 | 0.47 | 0.08 | 186 | 1.07 | <0.05 | 0.71 | <0.001 |
| Within F$_{2:3}$ families | 13 | 0.15 | 0.17 | 0.06 | 0.32 | 0.07 |

**Genotyping F$_{2:3}$ families**

Five F$_{2:3}$ seeds from each plant of two F$_2$ populations were pooled and genotyped using Monsanto proprietary-SNP markers through TaqMan assay (Semagn et al. 2015). A total of 150 and 199 SNP markers polymorphic between the parents of the two mapping populations were selected from proprietary database (unpublished) for genotyping the two mapping populations derived from CV156670 × 414−33 and CV156670 × CV143587, respectively.

**Statistical analysis**

**LWD responses**

LWD responses from 20 plants of each F$_{2:3}$ progeny family were recorded. Data were cured based on missing data points, outliers and correlation between replicates (Mailman et al. 2007; Rakesh 2016). Mean LWD scores of F$_{2:3}$ families were used for statistical analysis after data curation. Analysis of Variance (ANOVA) of F$_{2:3}$ progeny families was performed to partition the total variation in LWD response scores into those attributable to F$_{2:3}$ progeny families and within families. Additionally, pooled ANOVA was performed to test for the consistency of the F$_{2:3}$ progeny families for LWD response across two locations and to quantify the contribution of location and genotype × location interaction. Genotypic data of markers with >15% missing data and markers showing significant segregation distortion were not considered for statistical analysis.

**Detection of QTL controlling resistance to LWD**

Markers showing segregation distortion were removed (Supplementary table 1 and 2). The genotypic and phenotypic data of F$_{2:3}$ progeny families evaluated in two locations were integrated to detect QTL controlling LWD resistance, initially using single marker analysis. Subsequently, Inclusive Composite Interval Mapping (ICIM) was performed to detect and estimate size effects of QTL and QTL × QTL interactions (Li et al. 2015) controlling LWD resistance at individual locations and across locations (pooling data from both locations). The accuracy of QTL position and significance of size effect of QTL and QTL × Location interaction conferring LWD resistance was determined using data-driven estimates of threshold LOD scores obtained by 1000 permutations (Churchill and Doerge 1994). Similarly, significant QTL × QTL interactions controlling LWD resistance were detected at threshold LOD value of 5.0 and their size effects were estimated. All these statistical analyses were implemented using QTL ICIMapping software version 4.0 (Wang et al. 2016). The detected di-QTL interactions associated with resistance to LWD, dominance × dominance di-QTL interactions were interpreted based on theoretical investigations of Kearsey and Pooni (1996).

**Results**

**LWD responses of F$_{2:3}$ mapping populations**

The LWD response scores of the F$_{2:3}$ progeny families derived from CV156670 × 414−33 (P-1) and CV156670 × CV143587 (P-2) recorded at both KNH and MPG locations were normally distributed (Fig. 1). While LWD response scores of F$_{2:3}$ progeny families ranged from 3.17–9.00 and from 3.13–8.11 at KNH and MPG, respectively in P-1, the responses ranged from 2.38–8.38 and from 3.12–8.64 at KNH and MPG, respectively in P-2. Wider range of LWD scores at both locations in both the populations indicated sufficient LWD expression. Further, there were considerable numbers of plants towards both resistant and susceptible parents (Fig. 2). Six progeny families including two at KNH and four at MPG in P-1; 56 progeny families including 37 at KNH and 19 at MPG in P-2 showed resistance level better than resistant parent. Similarly, six progeny families including four at KNH and two at MPG in P-1; five progeny families including two at KNH and three at MPG in P-2 showed higher susceptibility than susceptible parent. The F$_{2:3}$ families derived from both the mapping populations differed significantly across both the locations except P-1 at MPG (Table 1). Further, LWD responses of F$_{2:3}$ families of both populations interacted significantly with location as evident from significance of genotypex location (G×L) interaction mean squares in pooled ANOVA (Table 2). While
mean squares attributable to location was non-significant, those attributable to genotype × location was significant in P-2. This trend was reversed in P-1 (Table 2).

| Source of variation | CV156670 × CV143587 (P-1) | CV156670 × CV414-33 (P-2) |
|---------------------|---------------------------|---------------------------|
|                     | Degrees of freedom | Mean sum of squares | F’ probability | Per cent contribution | Degrees of freedom | Mean sum of squares | F’ probability | Per cent contribution |
| Genotypes           | 235             | 1.52                | < 0.001        | 46.55             | 186             | 3.17              | < 0.001        | 77.89             |
| Locations           | 01              | 22.41               | < 0.001        | 2.92              | 01              | 0.01              | 0.971          | 0.00              |
| Genotype × Location (G × L) | 235 | 0.61                | 0.08            | 18.68             | 186             | 0.44              | < 0.001        | 10.81             |
| Error               | 470             | 0.52                | -              | 31.85             | 372             | 0.23              | -              | 11.30             |

### Detection of QTL controlling LWD resistance

#### QTL main effects

Seven QTL i.e., two (q-lw-1.1 and q-lw-1.2) in P-1 (Fig. 3) and five (q-lw-1.3, q-lw-1.4, q-lw-1.5, q-lw-1.6 and q-lw-1.7) in P-2 detected on chromosome-1 (Fig. 4) were found associated with LWD resistance with phenotypic variation explained (PVE) ranging from 4.34 to 13.06 % (Table 3). Among these, three major QTL (q-lw-1.4 and q-lw-1.6 at MPG; q-lw-1.5 at KNH) with > 10% PVE were detected in P-2. Among the remaining four minor QTL, two (q-lw-1.1 and q-lw-1.2) were from P-1 and other two (q-lw-1.3 and q-lw-1.7) were from P-2. Further, while five (q-lw-1.1, q-lw-1.3, q-lw-1.5, q-lw-1.6 and q-lw-1.7) of these seven detected QTL showed dominance effects, one (q-lw-1.4) showed additive effects in desirable direction (decreasing effects) for LWD resistance (Table 3).

### QTL × Location interaction

Pooled analysis across locations indicated significant QTL × location (Q×L) interaction. Of the seven detected QTL, five (q-lw-1.2, q-lw-1.4, q-lw-1.5, q-lw-1.6 and q-lw-1.7) interacted significantly with location with their size effects ranging from 0.04 to 6.80 % PVE (Table 3). Of these five interacting QTL, only q-lw-1.7 was not detected when individual location-wise analysis was performed. This means that, remaining four (q-lw-1.2, q-lw-1.4, q-lw-1.5 and q-lw-1.6) of these five QTL detected using pooled analysis were detected only in either of the locations but not in both the locations. These four QTL were also stable for their position but unstable for their effects (decreased size effects) as evident from pooled analysis.

Of the five QTL which showed significant Q × L interaction, one QTL, q-lw-1.4 (6.80% PVE Q × L) interacted with locations to a greater extent than the other four QTL (Fig. 5). None of the QTL detected through location-wise analysis were common in both locations. Two QTL (q-lw-1.3 and q-lw-1.5) flanked by same pair of markers, namely, Marker-151 and Marker-152 (with inter-marker distance of 23.8 cM) were detected in P-2. While the QTL, q-lw-1.3 was mapped at 11.40 cM
in KNH (in location-wise analysis), q-lw-7.5 was mapped at 9.40 cM in MPG (in location-wise as well as pooled analysis). Further, their position and effects differed (Table 3).

Di-QTL epistasis

A total of 48 significant di-QTL interactions with size effects (in % PVE) ranging from 4.00 to 19.03% were detected across both locations in P-1. Of these, ten were between QTL located within the same chromosome (Fig. 6a). Only nine of these 48 interactions involved QTL with significant effects. Similarly, 52 significant di-QTL epistatic interactions including ten within same chromosome were detected in P-2 with their size effects (in % PVE) ranging from 3.18 to 13.83 % (Fig. 6b). Only four of these 52 significant di-QTL interactions involved QTL with significant effects. Summation of direction and effects of dominance (h) and dominance × dominance (l) effects associated with all di-QTL interaction effects indicated duplicate di-QTL epistasis in both the populations. Duplicate di-QTL epistasis was in desirable direction (dominant decreasing effects) at KNH and MPG in P-1 and P-2, respectively (Table 4).

Validation of reported QTL conferring resistance to LWD

Rakesh (2016) reported QTL on chromosome-1 at 103.20 cM (IMD- 10.6 cM), flanked by MONIND14101828 and MONIND14262087. This QTL explained only 7.26 % of LWD resistance in mapping population derived by CV138811 × CV143587. In our study, one of these markers namely, Marker-161 (MONIND14101828) flanking the QTL, q-lw-1.6 could be validated in P-2 (CV156670 × CV143587). However, this QTL flanked by Marker-160 and Marker-161 was detected at position (83.40 cM) and phenotypic variation (13.06 % PVE at MPG and 4.35 % PVE across locations) different from that reported by Rakesh (2016) (Table 3).

Discussion

LWD is emerging as a major and serious biotic production constraint with a potential to reduce yield and quality of both grain and fodder (Sunitha et al. 2020; El-Naggarr et al. 2015; Drori et al. 2013; Johal et al. 2004). Developing LWD resistant cultivars is considered as an eco-friendly and sustainable approach to mitigate losses caused by LWD. Precise information on the number and mode of action of genes/QTL controlling resistance to LWD help devise suitable strategies to develop resistant cultivars. The results of our study indicate possible involvement of a large number of genes controlling resistance to LWD as evident from the normal distribution of LWD responses in F2:3 populations. Several previous researchers such as Shehata (1976), Nawar and Salem (1985), Abdel-Snbour and Bekhit (1993) and El-Hosary and El-Fiki (2015) have also reported the involvement of a large number of genes controlling LWD resistance in maize. Further, recovery of individuals that surpass the levels of resistant and susceptible parents indicate (i) presence of alleles controlling resistance even in susceptible parent and those controlling susceptibility even in resistant parent and (ii) complementation of parental alleles controlling resistance. Thus, F2:3 populations serve as potential source for recovering inbred lines with levels of resistance to LWD better than the resistant parent.

Most of the reports (Wu et al. 2020; Zhang et al. 2012; Yang et al. 2010; Suneetha et al. 2016) related to mapping QTL controlling resistance to stalk rot are predominantly on stalk rot caused by Fusarium sp. followed by Macrophomina phaseolina but not on the stalk rot caused exclusively by Harpophora maydis (Late wilt disease). This could be because of co-existence of Harpophora with Fusarium sp. and/or Macrophomina sp. which form a pathogen-complex causing FPSR (Degani et al. 2020; Khokhar et al. 2014). Additionally, soil-borne Fusarium sp. and/or Macrophomina sp. are known to invade the H. maydis infected stalks (Drori et al. 2013). This causes difficulty in phenotyping the symptoms caused exclusively by H. maydis. To address the issue of possible confounding effects of pathogens other than H. maydis, we used Indian pure isolates of H. maydis to inoculate the spores directly into the second internode of the stem (Rakesh et al. 2016) using stem jabbers (specialised injectors) twice at 45 DAS and 55DAS (Sunitha et al. 2020).

LWD resistance conferring QTL: Difficulty arising due to complexity of the disease and phenotyping could be the potential causes for sporadic reports on genetic basis of LWD resistance. The only report (Rakesh 2016) on mapping genomic regions controlling LWD resistance in two F2:3 populations (derived using CV138811 and CV143587 as donor and susceptible parents, respectively) resulted in the identification of three major QTL on chromosomes 3, 5 and 10; and six minor QTL on chromosomes 1, 2, 3, 5, 6 and 7. QTL analysis is not only intended to implement MAS for QTL but also for understanding the genetics of the quantitative trait. Hence, all the identified QTL, whether their effects are large or small and with or without environmental sensitivity are informative (Asins 2002). In our study, in addition to three major QTL detected either location-wise or across locations (pooled), four minor QTL were detected. In most of the studies, QTL with small effects go undetected due to bias caused by less-stringent threshold levels. To minimize such bias, we used LOD threshold derived from 1000 permutations (Churchill and Doerge 1994; Beavis 1994) in our study.

| Main QTL effect (h) | Dom × Dom effect (l) | CV156670 × CV143587 (P-1) | CV156670 × CV414-33 (P-2) | Interpretation on type of epistasis |
|---------------------|----------------------|--------------------------|--------------------------|----------------------------------|
| +                   | +                    | +                        | +                        | Complementary epistasis between dominant increasing effect QTLs |
| -                   | -                    | -                        | -                        | Complementary epistasis between dominant decreasing effect QTLs |
| +                   | -                    | +                        | +                        | Duplicate epistasis between dominant increasing effect |
| -                   | +                    | -                        | -                        | Duplicate epistasis between dominant decreasing effect |

Table 4
Direction of dominance effects and all possible dominance × dominance di-QTL interactions controlling resistance to LWD without considering their significance or otherwise
**QTL×Location interaction:** Position and effect of the QTL associated with LWD resistance varied significantly with the spatial environments represented by two locations in our study. If a quantitative trait exhibits significant genotype by environment interaction, it then follows that underlying QTL should also display significant interaction with either temporal or spatial environments (Bernardo 2020). In our study, the F2 progenies of both the crosses displayed significant interaction with the environment represented by two locations for responses to LWD (Table 2). Hence, detected significant interaction of LWD resistance governing QTL with location environments in our study did not surprise us. Several researchers have also reported significant QTL × location interaction associated with resistance response of maize diseases such as Fusarium ear rot (Wu et al. 2020, Robertson-Hoyt et al. 2006), northern leaf blight (Xia et al. 2020; Chen et al. 2016) and rough dwarf disease (Wang et al. 2019). The major QTL (>10% PVE) detected through location-wise mapping behaved as minor QTL (<10% PVE) when detected through pooled analysis. We could also observe shift in the position of QTL (q-lw-1.3 and q-lw-1.5) within the same flanking marker interval in different locations. However, such QTL despite showing significant G × E interaction, help select genotypes adapted to specific locations (Asins 2002). As expected based on previous reports by Shekhar et al. (2010), we could also observe significant influence of location and genotype × location interaction on the expression of LWD. Significance of G × E interaction for a trait reflects that underlying genomic regions exhibit significant QTL × Location interaction (Bernardo 2020). Among the three major QTL identified by us, although two QTL (q-lw-1.5 and q-lw-1.6) showed significant Q × L interaction, the magnitude of interaction effects was relatively low. These QTL with lower Q×L interaction effects could be considered as stable and deployed to implement MAS for LWD resistance after validation. This is because, the efficiency of MAS may get reduced if detected QTL controlling target trait exhibits significant interaction with either the environment or background genotype used (Asins 2002).

**QTL×QTL interaction:** The chances of recovering recombinant inbred lines (RILs) with dominant di-QTL interactions are fewer with any population other than F2. However, even a large F2 mapping population may contain fewer two-QTL RILs limiting the statistical power of detecting di-QTL interactions (Bernardo, 2020). Hence, there are chances that epistatic interactions go undetected. We addressed this limitation by increasing the LOD significance of > 5.0 and considering phenotypic data from replicated trials from F2:3. Some of the di-QTL interactions which involve QTL with their individual effects being non-significant are also informative, provided their di-QTL interaction effects are of higher magnitude. Under such instance, deployment of QTL alone in MAS without considering its significant di-QTL interaction would be ineffective. Hence, such QTL showing significant epistatic effects of large magnitude need to be introgressed together into recipient genetic background. However, there are chances that the magnitude of the significant di-QTL interaction may get reduced or get enhanced in recipient parent genetic background. The effects of such interacting QTL in recipient genetic background will be known after introgression. The direction of cumulative dominance effects associated with duplicate di-QTL epistasis varied with location. The duplicate di-QTL epistasis with decreasing dominance effects on LWD response is desirable. However, these di-QTL interaction effects need to be confirmed through extensive multi-location evaluation.

**QTL genetic background interaction**

Genetic background has considerable influence on the position and effect of QTL conferring disease resistance (Asins 2002; Awata et al. 2020). The same was evident in our study and none of the QTL was common between both populations, P-1 and P-2 (which differed for susceptible parent genome). However, we could validate only one of the reported markers, MONIND14101828 (Marker-161) flanking the reported QTL (Rakesh 2016) in P-2. While the QTL region having this marker, MONIND14101828 (Marker-161) in common is flanked by inter-marker distance of 11.8 cM in our study, it is flanked by inter-marker distance of 10.6 cM in previous report (Rakesh, 2016). Thus, this region needs fine mapping to identify QTL with narrow inter-marker distance for efficient deployment in MAS.

In summary, three major QTL, q-lw-1.4, q-lw-1.5 and q-lw-1.6 were identified from our study. However, PVE% explained by these three QTL is not large enough for use in MAS. Further, only one of these QTL, q-lw-1.6 could be validated. The three major QTL exhibited significant interaction with locations. Considering that the validated QTL reported by us and those reported by previous researcher exhibit significant interactions with locations, further research investigation is essential to identify stable QTL with large PVE for implementing MAS.

**Declarations**

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**Conflict of interest:** The authors declare that they have no conflict of interest.

**Availability of Data and material:** Not applicable

**Code availability:** Not applicable

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**Authors’ Contributions:** EG, VRPG and SR conceptualized the work. SNC conducted experiments and analyses. DS and SNC inoculation and screening for late wilt disease. SB, SR and SNC interpreted results. SNC and SR revised the manuscript. EG, SR, VRPG and HBH reviewed and edited the final version.

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**Figures**
Figure 1

Frequency distribution for responses to LWD of two F2:3 populations at Kallinayakanahalli and Muppadighatta locations
Figure 2

Patterns of expression of responses to LWD by two F2:3 populations (a. P-1 and b. P-2) across two locations
Figure 3

Genome-wide detection of QTL and their effects controlling LWD response mapped in F2:3 of CV156670 (R) × 414-33(S) (P-1) at (a) Kallinayakanahalli (b) Mppardighatta
Figure 4

Genome-wide detection of QTL controlling LWD response and corresponding additive and dominance effects mapped in F2:3 of CV156670(R) × CV143587(S) (P-2) at (a) Kallinayakanahalli (b) Muppadighatta
Figure 5

Genome-wide detection of QTL controlling LWD response and QTL × Location interactions in F2:3 of (a) CV156670(R) × 414-33(S)/ P-1 (b) CV156670(R) × CV143587(S)/ P-2
Figure 6

Genome-wide detection of significant di-QTL interactions controlling LWD response mapped in F2:3 of (a) CV156670(R) × 414-33(S) /P-1 (b) CV156670(R) × CV143587(S) /P-2.

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