Combining Ability Analysis for Northern Leaf Blight Disease Resistance on Tanzania Adapted Inbred Maize Lines

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Abstract

Northern leaf blight (NLB) disease incited by Exserohilum turcicum has increased in incidence and severity. Inbred lines combining ability and its interactions to the environment are required for the development of NLB disease resistance. The specific objectives were to estimate the combining ability for NLB disease resistance, determine maternal effects, and determine the heterosis in the F1 hybrids by using a full 11 × 11 diallel cross. The resulting 110 F1 hybrids with the 11 parents were evaluated together with 9 commercial varieties at three Agricultural Research Institutes: Tumbi, Uyole and Selian which represent diverse environments. Breeding materials were planted in 13 × 10 alpha lattice design with two replications per site. Top ten experimental hybrids in each site had negative mid parent heterosis for NLB disease severity. Heterosis for NLB disease severity ranged 94-362%. The overall mid parent heterosis means for yield across sites was 152%. Maternal effects had non-significant (P>0.005) influence on the inheritance of the NLB disease severity. Mean sum of squares for GCA was highly significant (P<0.001) on disease severity indicating additive genetic action. Mean sum of squares for SCA were highly significant on disease severity and yield implying non-additive gene action. The mean squares for reciprocals effects were highly significant for yield and non-maternal effects sums of squares had significant effect (P<0.05) on yield. The GCA effects contribution was high for disease severity (91%) and lesion number (85%). With the exception of CML 395 and KS03-0B15-12 parents which were susceptible, all GCA effects were negative implying the contribution to disease resistance in their progenies. Due to preponderance of the additive gene action, recurrent selection could be used to improve the resistance of inbreds while the non-additive gene action could be exploited in breeding for disease resistant high yielding hybrids.

Keywords: Combining ability; Maize; Northern leaf blight; Resistance; Tanzania

Introduction

Northern leaf blight disease (NLB) also known as Turcicum leaf blight or Northern corn leaf blight incited by the fungus Exserohilum turcicum (syn. Helminthosporium turcicum), Teliospore Setopheria turcica, is an endemic foliar maize disease in the world. The past decades witnessed the disease concentrating in the highlands of the world. However, now it affects both highlands and mid altitude maize growing areas and causing a significant yield reduction [1,2]. Maize yield losses caused by NLB disease vary depending upon location, pathogen virulence, pathogenicity, plant growth stage, number and position of leaves affected, relative humidity, temperature and susceptibility or resistance of the host plant. Raymundo [3] narrates the grain yield loss of over 50% in USA, while in India, the yield reductions of about 60% have been reported [4]. In East Africa, a maize grain yield reduction with the range of 16-24% have been reported [5].

Breeding for host plant resistance remains the most reliable and steadfast method of controlling the disease. Hence, host plant resistant (HPR) is considered as the best option and alternative to deal with the NLB disease problem [6,7]. However, breeding for HPR largely depends on correct methods of selecting suitable parents to be candidates of breeding for disease resistance. To study the potential, suitability and applicability of breeding materials, breeders apply different mating designs to achieve this purpose. One of the common mating designs is the diallel cross which have been extensively used in evaluations of breeding materials potential in various crops [8-12]. The diallel cross design enables breeders to estimate general combining ability (GCA) and specific combining ability (SCA) which are frequently used in genetic studies. Literature survey shows that, GCA for NLB disease is normally significant to denote its importance in additive gene action contribution. Vivek et al. [13] further contend that, GCA in NLB disease is area specific and is affected by environmental effects. This means that resistance to NLB disease varies from one location to another unless is monogenic before breaking down [13,14].

The majority of resistance of maize diseases is quantitatively inherited. In maize, NLB disease shows vertical and horizontal resistance inheritance mechanisms [15]. NLB disease is reported to be controlled by six dominant Ht1, Ht2, Ht3, HtN, NN and HtM and one recessive ht4 genes [16-20]. All these provide qualitative inheritance in the form of dominance or partial dominance. According to Pataky et al. [21] the HtN gene confers partial resistance to NLB disease. Other researchers have reported on the durable resistance to NLB conferred by major genes [22]. Ogliari et al. [23] reports on dominant Hph genes inducing resistance to NLB pathogen and recessive rt genes inducing resistance to specific NLB pathogen races. Several types of gene action are involved in controlling the inheritance of NLB disease in maize. Additive, dominance, and epistatic gene action have been reported in controlling the disease expression. However, additive gene action was found as the most important [23,24]. Maternal effects are less important for the traits associated with the inheritance of NLB. For example, Sigulas et al. [25] found non-significant maternal effect on 16 maize genotypes. Other researchers have reported non-significant importance of cytoplasmic and maternal effects on the inheritance of NLB resistance in maize genotypes [26].

Cultivars grown by farmers in Tanzania are potentially vulnerable to the NLB disease. Breeding for additional resistant varieties is needed. But, genetic information like GCA and SCA on the available inbred...
lines which is adapted to Tanzania conditions to be used as sources of NLB disease resistance and hybrid development is not known. Although various studies have been conducted in the world to estimate gene action related to NLB disease resistance and revealed useful information, these findings are limited to specific crosses and in some cases area bounded [27]. There is still potential room to widen the resistance genetic base as reports of new occurrence, distribution and resurgence of NLB disease are escalating worldwide. Another challenge facing the NLB disease struggle is the presence of pathogen races. The Common known races include 0, 1, 2, 3, 4, 12, 23 and 23N. To make the matter more complex, there is emergence of new races. These create a new dimension to fight war against the pandemic. The recent studies in Kenya by Muiru et al. [28] revealed 0, 1, 2, 3, N, 12, 13, 13N, 3N, 123, 23 and 23N races to connote the constantly and non-stopping breeding for NLB disease in the area. Therefore, the study on NLB disease resistance was initiated in Tanzania. The overall objective was to study the gene action of NLB disease resistance in inbred maize lines adapted to Tanzania conditions. The specific objectives were to- 1) estimate the combining abilities for NLB resistance of 11 maize inbred lines adapted to Tanzania condition, 2) determine maternal effects which are involved in NLB disease resistance in maize germplasm and, 3) determine the heterosis in the F$_1$ hybrids.

Materials and Methods

Sources and characteristics of breeding materials

Germplasm used in this study were obtained from screening for NLB disease resistance of 70 breeding materials at ARI-Tumbi in the growing season 2008/2009. The screening study resulted to the selection of 11 parents with various reaction types to NLB disease which were selected for the study (Table 1). Cob sizes were assessed by the developed scale of 1-5 where 1=very small, 2=small, 3=average, 4=big, and 5=excellent while NLB disease reactions assessment was performed according to procedures developed by Reid [29].

F$_1$ hybrid development

Breeding materials used in this study were developed from an 11 × 11 full diallel cross mating design. Crosses were performed at Selian Agricultural Research Institute (SARI) in Arusha in Tanzania. These crosses resulted in 55 F$_1$ and 55 reciprocal families.

Field evaluation

The resulting 110 F$_1$ progeny and 9 commercial hybrids (Kilima-ST, Kito-ST, Lishe-H1, Lishe-K1, Selian-H208, Selian-H308, Stuka-1, TMV-1 and Vumilia-k1) were planted in 13 × 10 alpha lattice design in three locations with two replications per site. The breeding materials were evaluated in three sites namely ARI-Tumbi, Selian and Uyole in the 2010/2011 growing season. One F$_1$ hybrid CML 159 × VL 05616 was doubled per replication to make 130 maize genotypes for evaluation. An experiment involving 11 inbred materials was set adjacent to the hybrid trial to avoid inter-plot competition and was planted on the same day to avoid biasness [30]. Inbred lines were planted in RCBD with two replications per site. Maize was planted at a spacing of 0.75 m × 0.30 m, which gives a plant population density of 44,444 plants per hectare. Basal fertilizer was applied at a rate of 40 Kg P ha$^{-1}$ and Murate of Potash was applied at the rate of 40 Kg K$_2$O ha$^{-1}$. Top dressing was done by using UREA (46%) to make a recommended rate of 100 Kg N ha$^{-1}$. Disease field assessment was conducted at about three weeks after silking of each particular genotype.

Inoculation procedures

To ensure uniform disease infestation, artificial inoculation was conducted according to procedures described by Reid [29] as follows- A sample grinder machine [Laboratory mill model-4, Thomas-Wiley, Thomas scientific (TM), and USA] was used to grind infected leaves into powder form. Leaves were collected from Tabora, Arusha and Mbeya, where trials were conducted and then mixed. A bazooka (Sistrunk Inoculators, Starkville, MS 39759) was used to apply the powder in the whorl of the plants. One dose of powder from the bazooka application amounted to 0.1 g of leaves powder. Two applications, one at 6-8 leaf stage and the second at 11-12 leaf stage were conducted. Furthermore, in the second rows of a spreader local variety (Situka-1) which is highly susceptible to NLB disease was planted around the field and after every 10 rows of the test materials. The uses of spreaders have been used successfully in screening germplasm for other disease studies [19]. In addition to NLB disease resistance, maize genotypes were assessed for other agronomic characteristics. Other collected agronomic data included- Days to 50% anthesis, days to 50% silking, plant height, ear height, husk cover, kernel type, grain texture and yield. Grain yield was estimated using the following formula-

$$ \text{Grain yield (t/ha)} = \frac{\text{FW} \times (100 - \text{MC}) \times \text{SH}\% \times 10,000}{85 \times \text{PS}} $$

Where, FW=field weight of unshelled cobs (kg)
MC=grain moisture content (%)
SH%=Shelling percentage (expressed as a fraction)
$10,000=1$ ha=$10,000$ m$^2$
PS=Plot area (m$^2$)

| Name       | Source                  | Grain colour | Cob size (1-5) | R×n |
|------------|-------------------------|--------------|----------------|-----|
| KS03-GB15-126 | SARI-Tanzania            | White        | 4              | R   |
| EB04-0A01-304 | SARI-Tanzania (QPM line) | White        | 4              | R   |
| CML 159    | CIMMYT-Mexico           | White        | 5              | R   |
| KS03-OB15-2 | SARI-Tanzania            | White        | 4              | R   |
| KS03-OB15-45 | SARI-Tanzania            | White        | 4              | R   |
| VL 05616   | CIMMYT-Zimbabwe (Vivek)  | White        | 5              | R   |
| KS03-OB15-47 | SARI-Tanzania            | White        | 5              | R   |
| CML 395    | CIMMYT-Zimbabwe          | White        | 5              | S   |
| KS03-OB15-12 | SARI-Tanzania            | White        | 4              | S   |
| CML 442    | CIMMYT-Zimbabwe          | White        | 4              | R   |
| CKL 05007-B-B | CIMMYT-Kenya (KARI)     | White        | 4              | R   |

Where, T=No.treatment, R × n=No.reaction, Num=No.treatment number, cob size 1=very small, and 5=bigger.

Table 1: Characteristics of 11 parents used in a diallel mating.
85% = grain moisture content adjustment factor. Grain yield was adjusted to 15% (100-15). Data were recorded according to IBPGR [31] maize descriptors list. Yield and disease resistance data were validated and analyzed in Genstat statistical computer software [32].

Data analysis

Collected data were first analysed by analysis of variance [33] then by using Genstat computer software according to Payne et al. [34] to test the existence of significant differences on the measured maize traits. Gene action component effects were estimated by application of SAS 05 computer programme using Griffing's [9] Method 1, Model 1 (fixed model) as follows:

\[ Y_{ijkl} = \mu + g_i + r_j + s_k + e_{ijkl} + \frac{1}{b} \sum_{i=1}^{b} \sum_{j=1}^{a} e_{ijkl} \]

\[ Y_{ik} = \text{value of F}_i \text{ of a cross of the i}^\text{th} \text{ female and the j}^\text{th} \text{ male in the k}^\text{th} \text{ block and i plot/observation} \]

\[ \mu = \text{population mean; } i=j=1, 2, \ldots, n. \]

\[ g_i = \text{GCA effect of } i^\text{th} \text{ parent} \]

\[ r_j = \text{reciprocal effects} \]

\[ s_k = \text{SCA effect} \]

\[ e_{ijkl} = \text{error effect for } ijk \text{ observation} \]

\[ b = \text{number of replications} \]

\[ c = \text{number of plants/plot} \]

The ratio of GCA/SCA was also assessed. For the ratio greater than 1 indicated additive genetic effect while the ratio lower than one denoted dominance gene action for the particular traits. The ratio closer to one implies the possibility of prediction based on GCA component only. The use of Griffing's [9] Method 1, Model 1 provides similar results as other approaches [34,35].

Heterosis estimation

Mid parent heterosis was calculated according to Falconer and Mackay [36] and Saleh et al. [37] as follows:

\[ MPH = \frac{F_i - \text{MP}}{\text{MP}} \times 100 \]

Where, \( F_i = \) mean of the \( i \) hybrids; \( MPH = \) mid-parent heterosis and \( MP = \) mean of two parents.

Results

Estimation of general combining ability (GCA) and specific combining ability (SCA)

The model of analysis explained more than 60% of the variations (R²). R² was 85, 64 and 73% for disease severity, yield and lesion numbers respectively to indicate that, the model was adequate (Table 2). The environment mean square was highly significant (P<0.001) for severity, yield and lesion number to imply the significant impact of environment on genotypes. The same trend was recorded on genotypes and on replications with the nested environment. The interaction of Environment and hybrids were highly (P<0.001) significant on disease severity only. In all three sites, all GCA had highly significant (P<0.001) effects on disease severity. With the exception of CML 395 and KS03-0B15-12, all CGA effects were negative. Parent EB04-0A01-304 had significant positive GCA effects on yield while VL 05616 and CKL 05007-B-B had significant negative GCA effects on lesion number (P<0.001), while parent CML 442 had negative significant (P<0.05) effects on the same trait.

In all three sites, all GCA had highly significant (P<0.001) effects on disease severity. With the exception to CML 395 and KS03-0B15-12, all CGA effects were negative. The same parents had significant positive GCA (P<0.001) effects on yield (Table 5).

Three hybrids CML 159 × CML 395, VL 05616 × KS03-0B15-12 and KS03-0B15-2 × CML 395 had positive significant (P<0.05) SCA effects on disease severity while hybrid CML 395 × CKL 05007-B-B possessed

\[ \text{Severity} \times \text{Yield} \times \text{Lesion number} \]

Table 2: Mean squares partial analysis of variance on NLB disease severity (%), yield and lesion number of maize hybrids.

| Source       | Severity (%) | Yield | Lesion number |
|--------------|--------------|-------|---------------|
| Source       | DF | MS | DF | MS |
| Env.         | 2  | 8846.89*** | 92.37*** | 1241.12*** |
| Rep (Env.)   | 3  | 1675.42*** | 86.82*** | 765.43*** |
| Genotypes    | 120 | 770.3***   | 11.72*** | 74.28***  |
| Genotypes    | 240 | 110.28*** | 2.19      | 63.9      |
| Corrected total | 725 | 0.85 | 0.64 | 0.73 |

\[ R^2 = 0.85 \]

\[ * \quad ** \quad *** \text{indicates significance level at 0.05, 0.01 and 0.001 respectively.} \]

Table 3: Diallel analyses of hybrids for disease reaction and yield over three sites.

\[ \% \text{GCA contribution} \]

\[ \% \text{SCA contribution} \]

\[ * \quad ** \quad *** \text{indicates significance level at 0.05, 0.01 and 0.001 respectively.} \]
positive highly significant SCA effects on yields (Table 6). Parents KS03-0B15-126 and CML 395 were susceptible to NLB disease. On yield, hybrid VL 05616 × KS03-0B15-12 had positive significant SCA effects (P<0.05) while hybrid CML 395 × CKL 05007-B-B expressed positive highly significant SCA (P<0.001) effects. Positive significant SCA is desired in yield of maize trait. All hybrids had non-significant SCA effects on lesion numbers.

Reciprocal, maternal, nonmaternal effects on yield

Mean squares of SCA and maternal effects had highly significant positive (P<0.001) effects on yield while, GCA showed positive significant (P<0.005) effects on yield (Table 7). Reciprocal effects were also highly significant (P<0.001). The hybrids with positive highly significant effects were KS03-0B15-126 × KS03-0B15-12 and CML 159 × KS03-0B15-12 indicating the contribution of cytoplasmic gene and nuclear gene effects. Maternal effects were also observed. Five parents showed positive significant maternal effects at different dosage levels. For example parent KS03-0B15-126 showed positive highly significant (P<0.001) while KS03-0B15-47 showed positive significant (P<0.05) maternal effects. Non-maternal positive significant effects were observed in four hybrids. With the exception of KS03-0B15-126 × KS03-0B15-12 which showed positive highly significant effects (P<0.001), the remaining EB04-0A01-304 × KS03-0B15-45, EB04-0A01-304 × KS03-0B15-45 and CML 159 × KS03-0B15-12 showed positive significant (P<0.05) maternal effects.

Heterosis

The studied F1 hybrids showed variations in the enhanced expression of mid parent heterosis (MPH) for NLB disease severity. At Tumbi, the mean was 12.97% and it was ranged 93.46-361.99%, while at Uyole, the mean was 13.22% and ranged 92.80-178.81%. At Selian, the mean was 11.14% and the ranged 76.60-144.19%. Each site ranked different F1 hybrids to indicate that, The F1 hybrids performed differently across sites (Table 8). The maximum mid parent heterosis for yield was higher than 350% in all sites.

At Tumbi, the mean was 133.60 and ranged 27.13-367.37 while at Uyole, the mean was 141.62 and ranged 45.30-352.98. At Selian, the mean was 180.32 and the ranged 12.55-460.84. The F1 hybrids behaved differently in terms of yield heterosis among sites (Table 9).

Yield performance and NLB disease reactions of parent materials

There were no significant differences (P>0.05) in yield performance across test sites implying stability of inbred lines across environments (Table 10). On severity, testing environments differed significantly (P<0.001) with ARI Selian site recording the highest severity (22.9%) followed by ARI-Uyole 17.95%. Genotype performance across sites showed significant difference (P<0.001) on NLB disease lesion number with ARI-Uyole recording the highest (8.68) while ARI-Tumbi showed the lowest (1.55). On lesion length, there were also highly significant differences (P<0.001) across sites with ARI-Selian recording the highest (156.5 mm) followed by ARI-Uyole 26.1 mm.)
Estimation of general combining ability (GCA) and specific combining ability (SCA)

The analysis model of ($R^2$) explained 85, 64 and 73% effects for disease severity, yield and lesion numbers respectively to indicate that, the model was adequate. Level of significant differences among genotypes indicated by mean squares were also tested to justify the use of GCA and SCA and found to be highly significant ($P<0.001$) which fulfilled conditions suggested by Griffing [9]. Results revealed highly significant ($P<0.001$) environment mean squares for disease severity, yield and lesion number implying the significant impact of environment on genotypes. This could be caused by weather, edaphic factors and initial inoculum concentrations in the test sites which call for area specific cultivar recommendations as suggested by Macharia et al. [38].

Mean sum of squares of GCA was highly significant ($P<0.001$) on disease severity and lesion numbers and significant on yield ($P<0.05$) to denote the additive gene action expression. At the same time SCA mean squares were highly significant on disease severity and yield only to imply non-additive gene expression on those traits. These findings suggest that, both additive and non-additive gene effects are involved in the expression of NLB disease resistance in maize. Other researchers have reported on the additive and non-additive gene actions to express NLB disease resistance in maize [14,23,39,40]. The percentage of GCA contribution was 90.81% and 85.41% for disease severity and lesion numbers respectively. By considering levels of significant through mean squares and percentage of GCA contribution, it was obvious that additive gene action was more important than non-additive gene action for the expression of NLB disease resistance in maize germplasm which is in accordance to the findings of Sigulas et al. [25].

Parents combining ability on disease severity and yield

In disease resistance studies, negative GCA is desired while positive GCA effects are not desired. The GCA of nine parents had negative highly significant ($P<0.001$) effects on disease severity indicating additive gene effects for resistance. On yield, parent EB04-0A01-304 had significant positive GCA effects. Parents with positive GCA have the potential to impart high yielding characteristics to the next generation. Therefore, the combination of parents with negative GCA for disease severity and positive GCA for yield are likely to produce high yielding F$_1$ hybrid.

Individual sites analysis also showed similar results for NLB disease severity inheritance. All parents had highly negative significant effect for NLB disease severity with the exception to CML 395 and KS03-0B15-12. However, on yield the same CML 395 and KS03-0B15-12 parents had significant positive GCA ($P<0.001$) effects for yield. This means that these parents contributed to NLB disease susceptibility at the same time contributed to high yield potential. Based on individual site analysis, parents like CML 395 and KS03-0B15-12 need to be improved for NLB disease resistance so as to have parents with both high yielding and resistant potentials.
Specific combining ability (SCA) on NLB disease severity on $F_1$ hybrids

The positive significant ($P<0.005$) SCA effects on disease severity showed by $F_1$ hybrids indicates the non-additive gene action as expressed in CML 159 × CML 395 and VL 05616 × KS03-0815-12 hybrids which could be probably due to masked susceptibility or maternal cytoplasmic effects of VL 05616 and CML 395 parents. However, the clear reason was not established by this study.

Reciprocal, maternal and non-maternal effects on yield

This study revealed that reciprocal effect which is associated with maternal and non-maternal effects were involved in yield expression. These gene actions have the tendency of reducing breeding efficiency through masking genetic variance [41]. The significant contribution of reciprocal, maternal and non-maternal effects on yield indicates the contribution of cytoplasmic genes and nuclear gene effects to bring an impact on maize yield. Some environmental factors like drought can enhance maternal effects as recorded by Derera et al. [42]. Thus, appropriate mating designs such as diallel cross and North Carolina design II can be employed to improve maize breeding procedures which encompass estimation of reciprocal effects depending on the number of parents to be used in the study [43].

Heterosis

Based on mid parent heterosis (MPH), $F_1$ hybrids showed variations in the enhanced expression of disease severity and yield. The MPH was significantly higher for disease severity across sites to imply
dominance gene action effects on the studied germplasm. Negative heterosis is desired in the studies of NLB disease resistance. All top ten experimental hybrids in each site had negative heterosis. On yield, a positive heterosis is desired. High positive heterosis on yield indicates the superiority performance for NLB disease reactions of F1 hybrids. The overall mid parent heterosis mean across sites was 151.84%. These results agree with Saleh et al. [37] who recorded much higher heterosis in maize.

Parents, commercial cultivars and F1 hybrid breeding potentials

The high variations in terms of NLB disease reactions and yield associated traits showed by parents implied high potential for breeding progress and yield increase in maize production. The selected inbred lines clearly expressed their traits across three diverse agro-ecological zones to show genetic potential for NLB disease reactions. The mean severity was higher at ARI Selian (22.9%) and ARI-Tumbi had the lowest (10.45%). The higher disease prevalence on parents and hybrids at ARI-Uyole and ARI-Selian could be attributed to weather conditions, land use and increased pathogen pathogenicity. Uyole is located in the highlands with favorable rainfall and temperature for disease development while Selian records more rainfall than Tumbi site. Uyole and Selian sites are characterized by land shortage which results in intensive land use with limited crop rotation flexibility. These practices result in accumulation of inoculums in maize stovers and alternative hosts. On the other hand ARI-Tumbi has less rainfall compared to the other two sites. At the same time, Tumbi site is characterized by abundant land to allow crop rotation, natural fallow, improved fallow and improved woodlots [44]. Frequent bush fire and free range animal grazing system which are common practices at Tumbi area could be other factors leading to relatively low level of NLB disease infestation in that area.

All commercial hybrids showed susceptibility to NLB disease with the scale of more than five. According to Ngugi et al. [45] and Reid [29], the detected commercial cultivar susceptibility can be classified as medium to high. Thus, all those commercial cultivars succumbed to NLB disease indicating their susceptibility. This implies that, farmers in Tanzania are growing NLB disease susceptible cultivars which could justify the disease resurgence and outbreaks in the near future if appropriate measures are not put in place. However, there was differential performance of individual hybrids to signify the importance of interactions which calls for area specific breeding procedures.

Conclusions

The following conclusions were drawn from this study-

a. This study showed the predominance of additive gene action for controlling the NLB disease resistance in maize. In the study the mean sum of squares for GCA was highly significant (P<0.001) on disease severity indicating the predominance of additive gene action. This was further emphasized by the high GCA contribution for disease severity (91%) and lesion number (85%). The majority of parents had negative CGA to imply contribution to disease resistance on their progenies.

b. The mid parent heterosis for NLB disease severity ranged 93.46- 361.99%. Genotypes with negative heterosis to NLB disease is desired because they imply superiority performance of progenies towards resistance direction.

c. In this study reciprocal and maternal effects had non-significant (P>0.05) effects on the inheritance of the NLB disease severity.

d. Due to preponderance of the additive gene action, recurrent selection could be used to improve the resistance of inbreds while the non-additive gene action could be exploited for breeding resistant hybrids in maize breeding program.
|         | Tumbi       | Uyole       | Selian      |
|---------|-------------|-------------|-------------|
| **Top ten** |             |             |             |
| 7 × 11  | 4.65        | 6 × 10      | 5.30        |
| 11 × 4  | 5.85        | 1 × 7       | 6.70        |
| 6 × 5   | 3.75        | 8 × 5       | 7.32        |
| 7 × 5   | 6.30        | 11 × 6      | 4.76        |
| 9 × 11  | 4.35        | 7 × 10      | 4.16        |
| 4 × 5   | 5.70        | 2 × 10      | 8.84        |
| 8 × 2   | 5.40        | 3 × 2       | 7.20        |
| 3 × 2   | 4.80        | 5 × 4       | 3.90        |
| 5 × 9   | 5.10        | 4 × 5       | 8.16        |
| 3 × 10  | 6.45        | 11 × 10     | 7.02        |
| **Middle five** |         |             |             |
| 9 × 4   | 3.00        | 9 × 5       | 7.20        |
| 10 × 4  | 7.05        | 10 × 4      | 6.70        |
| 8 × 1   | 6.00        | 1 × 5       | 5.90        |
| 9 × 5   | 4.50        | 7 × 1       | 5.72        |
| 4 × 3   | 3.75        | 8 × 4       | 5.50        |
| **Bottom five** |       |             |             |
| 5 × 10  | 3.90        | 8 × 11      | 3.76        |
| 3 × 7   | 7.65        | 10 × 1      | 3.70        |
| 7 × 6   | 4.20        | 7 × 8       | 7.40        |
| 4 × 8   | 6.30        | 11 × 4      | 6.80        |
| 4 × 6   | 4.50        | 6 × 8       | 5.84        |

|         | Tumbi       | Uyole       | Selian      |
|---------|-------------|-------------|-------------|
| **Commercial checks** |             |             |             |
| Kilima-ST | 6.90        | 5.40        | 5.16        |
| Kito-ST   | 2.25        | 2.15        | 3.15        |
| Lishe-H1  | 3.75        | 6.80        | 5.25        |
| Lishe-K1  | 3.30        | 4.92        | 5.62        |
| Selian-H208 | 5.85      | 7.70        | 6.19        |
| Selian-H304 | 4.20      | 6.30        | 5.88        |
| Stuka-1   | 3.40        | 5.32        | 4.76        |
| TMV-1     | 4.20        | 5.32        | 5.81        |
| Vumilia-k1 | 5.70        | 5.90        | 5.48        |

|         | Mean        | Lsd         | Cv          |
|---------|-------------|-------------|-------------|
|         | 5.04        | 6.31        | 4.377       |
|         | 3.348       | 3.895ns     | 3.895ns     |
|         | 33.60       | 35          | 32.5        |
|         | 0.542ns     | 0.293ns     | 0.581ns     |

Where, 1=KS03-0B15-126, 2=EB04-0A01-304, 3=CML 159, 4=KS03-0B15-2, 5=KS03-0B15-45, 6=VL 05616, 7=KS03-0B15-47, 8=CML 395, 9=KS03-0B15-12, 10=CML 442 and 11=CKL 0507-B-B

|         | Tumbi       | Uyole       | Selian      |
|---------|-------------|-------------|-------------|
| **Middle four** |       |             |             |
| 4 × 9   | 2.03        | 3 × 1       | 3.74        |
| 2 × 7   | 3.57        | 3 × 6       | 3.74        |
| 10 × 6  | 1.71        | 10 × 6      | 3.77        |
| 3 × 5   | 2.16        | 2 × 11      | 3.81        |
| **Bottom four** |       |             |             |
| 10 × 1  | 1.98        | 8 × 9       | 6.98        |
| 5 × 3   | 1.87        | 8 × 7       | 7.07        |
| 10 × 7  | 2.21        | 6 × 9       | 7.1         |
| 2 × 1   | 2.55        | 8 × 3       | 7.24        |

|         | Kilima-ST   | 5.09        | 6.51        |
| Kito-ST  | 5.56        | 6.04        | 5.60        |
| Lishe-H1 | 5.14        | 6.12        | 6.74        |
| Lishe-K1 | 5.24        | 6.40        | 5.08        |
| Selian-H208 | 5.57      | 6.85        | 6.70        |
| Selian-H308 | 5.72      | 5.90        | 6.93        |
| Stuka-1  | 4.85        | 7.19        | 5.66        |
| TMV-1    | 6.18        | 6.17        | 5.53        |
| Vumilia-K1 | 5.43      | 6.20        | 5.29        |

|         | Grand mean  | 3.03        | 4.39        |
|         | Lsd         | 2.251       | 2.2048      |
|         | Cv          | 37.50       | 25.40       |
|         | Prob        | <0.001      | <0.001      |

Table 12: NLB disease severity of hybrids across three sites.

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