Heterosis and character association of mid altitude adapted quality protein maize (Zea mays L.) hybrids at Bako, Western Ethiopia

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Abstract

The production of hybrid Quality Protein Maize (QPM) is started very recently in Ethiopia that requires extensive research to exploit heterosis to increase the productivity of the crop. Exploitation of heterosis through the production of hybrids is proven breeding method to improve the yield of maize. Accordingly, the aims of this study were to estimate the magnitude of heterosis of QPM inbred lines and to determine the association among morph-agronomic traits. The crossing and experiment were conducted at Bako National Maize Research Center during the 2014 and 2015 main cropping season, respectively. Forty-five F₁ hybrids obtained by crossing 10 inbred lines in diallel fashion were planted in two separate trials on adjacent experimental blocks. The experimental design used was alpha lattice for the hybrid trial and randomized complete block design for inbred lines trial with three replications. Each entry was planted on a one-row plot of 5.1 m length with 0.75 m and 0.3 m spacing between rows and plants, respectively for both trials. Data on phenology, growth, grain yield and yield related traits, and disease reaction were collected at the appropriate plant growth stages. The mean squares due to hybrids and parents were highly significant for all traits except common leaf rust in the hybrids trial and diseases parameters, stem lodging, bad husk cover and ear rot indicating the existence of genetic variability for most traits. The highest positive mid and better parent heterosis was observed for grain yield and yield related traits indicating the possibility of increasing grain yield and yield related characters via hybridization. Negative heterosis was observed for phenology and growth traits and diseases parameters as well. Grain yield had highly significant positive association with plant height, ear plant⁻¹, ear length and number of kernel row⁻¹ and negatively correlated with anthesis silk interval, plant and ear aspect, common lea rust, phaeosphaeria leaf spot and ear rot at genotypic and phenotypic level. In addition, plant height, ear plant⁻¹, ear length, number of kernel row⁻¹ had positive direct effects and anthesis silk interval, plant aspect, phaeosphaeria leaf spot and ear rot had negative direct effects on grain yield at both genotypic and phenotypic level. While, common leaf rust had negative direct effects at genotypic and days to anthesis, ear height, root lodging and ear aspect had negative direct effects on grain yield at phenotypic level. This indicated that, by increasing the positive and decreasing the negative direct effects of the traits grain yield could increase invariably.

Introduction

Maize (Zea mays L.) is the basic staple cereal grain and currently produced on above two million hectares and the 2nd most widely grown crop in Ethiopia and stands first in terms of production (9.5 million ton) and productivity 4 ton per ha. It is produced by about nine million farmers indicating its popularity and importance to the livelihoods of more than 70% of the farming households in Ethiopia [1].

The conventional maize grown widely around the world supplies many macros and micronutrients necessary for human metabolic needs; however, it lacks B vitamins and the essential amino acids lysine and tryptophan [2–4]. Consequently,
Improving the quality of maize protein for human consumption by increasing its lysine and tryptophan through decreasing the zeins protein fraction content has therefore been a long-term goal of several maize breeding programs [7, 8]. The bio–fortification of conventional maize by doubling the amount of lysine and tryptophan through the identification of a mutant genotype, popularly known as opaque–2 (o2), with reduced zeins protein fraction and improved agronomic performance gave rise to newly commercialized maize types called Quality Protein Maize (QPM) [9].

In most maize hybrid breeding programs, the main objective is to develop improved inbred lines that can form best hybrids upon crossing [10]. Sustainable production of QPM in Ethiopia and elsewhere is dependent not only on its nutrition benefit but also on how much can be harvested per unit area of land. Unless the productivity of QPM is comparable or better than the conventional maize varieties currently in use, farmers may be reluctant to produce it. This requires a rigorous inbred line development and evaluation work to identify potential parental lines for hybrid variety development.

The concept of heterosis is practically exploited to develop hybrid varieties. Heterosis may be defined as the increase in size, vigor, fertility, and overall productivity of a hybrid plant, over the mid parent value (average performance of the two parents) and over the performance of best parent. It is occurred when two inbred lines of out bred species are crossed, as much as when crosses are made between pure lines [11].

Although yield is usually the primary trait of interest, maturity, stand–ability, grain quality, stem quality, and resistance to major diseases and insects are all corollary traits that the maize breeder must consider for eventual usefulness of genotypes evaluated for yield [12].

Correlation is the degree to which two or more variables are related and change together [13]. Usually more than one trait is measured on progenies evaluated either for a specific trait in cyclical selection programs or in applied breeding programs that require a combination of traits to satisfy growers. In genetics, there are two main causes of correlation between characters, genetic and environmental. The genetic cause of correlation is chiefly pleiotropy, though linkage is a cause of transient correlation, particularly in population derived from crosses between divergent strains [14]. It has established in classical genetics that many genes have manifold effects; i.e., some genes seem to affect traits that are unrelated. Genes that have manifold effects are pleiotropic, i.e., the same gene affects different traits in a complementary way. The existence of pleiotropic effects of genes in different classical genetic studies showed the presence of pleiotropy in different quantitatively inherited traits. Then it is possible that selection may be exerted on secondary traits that have greater heritability than the primary trait. Indirect selection will be effective if the heritability of the secondary trait is greater than that of the primary trait and the genetic correlation between them is substantial [12]. In maize, both genetic and environmental correlations have been extensively studied by various researchers and their importance with respect to a particular trait has been well documented [15,16].

Correlation coefficients do not give a complete picture of the causal basis of association and selection based on correlation coefficients without taking into consideration the interaction between the component traits could be misleading. Therefore, to design appropriate breeding strategies for improvement in yield through selection, it would be desirable to conduct both correlation and path coefficient analysis [17]. Path coefficient can be defined as a ratio of the standard deviation of the effect due to a given cause to the total standard deviation of the effect [18].

Therefore, this study is initiated to support the quest for better mid–altitude adapted QPM hybrid varieties in Ethiopia with the following specific objectives: to estimate the magnitude of heterosis in crosses derived from mid altitude QPM inbred and the association of traits with grain yield.

Materials and methods

Description of experimental site

The experiment was conducted at Bako National Maize Research Center (BNMRC), which is located in Western Ethiopia. Bako Maize Research Center lies between 9°6’ North latitude and 37°09’ east longitude at an altitude of 1650 meters above sea level (m.a.s.l.) in the sub–humid agroecology of Ethiopia.

Experimental materials

Ten white–grained Quality Protein Maize (QPM) inbred lines obtained from BNMRC were crossed using diallel mating design during the main cropping season of 2014 and forty–five single cross hybrids were generated. The parental inbred lines were selected based on their tryptophan and lysine content and per se performance history for grain yield and yield related traits. The inbred lines which contain good level of essential amino acids, lysine (4% in whole grain) and tryptophan (>0.8% in whole grain) were selected [19]. The parental inbred lines and the resulting hybrids (45) were organized into two separate sets of trials and tested in adjacent blocks (hybrids and inbred lines trials) at Bako trials evaluating site to avoid the unbalanced competition between the hybrids and inbred lines.

Experimental design

The 45 F₁ hybrids and the 10 inbred lines were planted following experimental design 9 x 5 alpha–lattice (0.1) for the hybrid trial Patterson and Williams, [19] and a randomized complete block design (RCBD) for the inbred line trial each with three replications. Each entry (the hybrids and parental inbred lines) was planted in a one–row plot of 5.1 m length and 0.75 m between rows and 0.3 m distance between plants in a row.
Data collected

Phenology and growth data: 1. Days to anthesis (AD): The number of days from planting to when 50% of the plants in a plot shed pollen. 2. Days to silking (SD): The number of days from planting to when 50% of the plants in a plot 2–3 cm long silk. 3. Anthesis Silking Interval (ASI): Recorded as the number of days between days to silking and days to anthesis (ASI= SD – AD). 4. Ear Height (EH): The height from the ground level to the base of the upper most ear-bearing node of five randomly taken plants from each experimental unit was measured in centimetres. The measurement was made two weeks after pollen shedding has ceased.5. Plant Height (PH): The height from the soil surface to the base of the first tassel branch of five randomly taken plants from each experimental unit was measured in centimetres. Like ear height, this was also measured two weeks after pollen shedding had ceased from the same plants that were used to measure ear height.

Grain yield and yield related traits data: 1. Grain yields (GY), at harvesting ears were removed from all plants in each plot leaving other crop residues (husk, leaf, stem and tassel) intact. The total field weight from all the ears of each experimental unit was measured. This was adjusted to 12.5% moisture (electronically determined using digital moisture tester) level and 80% shelling percentage to estimate grain yield in tons (t ha⁻¹) for each genotype. 2. Root Lodging (RL): Prior to harvest, the number of plants inclined more than 30° from the vertical axis were counted and recorded. The data was calculated as (number of root lodged plants)/ (total plants per plot) ×100. 3. Number of Ears Per Plant (EPP): The total number of ears harvested from a plot divided by the number of plants in that particular plot at harvest. 4. Ear Length (EL): Length of the ear from the base to tip. It has been measured as the average length of five randomly taken ears from each experimental unit in centimeter. 5. Ear Diameter (ED): This was measured at the mid-way along ear length, as the average diameter of five randomly taken ears from each experimental unit in centimeter. 6. Number of kernel rows per ears (NKR): was calculated as (number of root lodged plants)/ (total plants per plot) ×100. 7. Number of Kernels Per Row (NKPR): This was recorded as the average number of kernels per row from 10 randomly taken ears. 8 Plant Aspect (PA): Recorded using 1–5 scale; where, 1 is best genotype (considering the overall performance of the plant in the field) and 5 is the genotypes have undesirable overall plant appearance. 9. Ear Aspect (EA): Record using 1–5 scale; where, 1 refers to good ears (considering the overall appearance of the ear) and 5 refers to poor ear with undesirable ear characteristics. 10. Number of Rotten Ears (ER): counting and recording the number of ears rotten from the total number of harvested ears per plot. Reaction of the genotypes to Turcicum Leaf Blights (TLB) and Phaeosphaeria Leaf Spot (PLS) were evaluated. The disease severity scores were rated at 1–5 rating scale was used where, 1 is the best and 5 the worst in terms of reaction to the diseases. In addition, the intermediate ratings between two numerals (1.5, 2.0, 2.5, 3.0, 3.5, 4.0 and 4.5) have also been used for all the diseases under study.

Data analysis

Mid Parent Heterosis (MPH) and Better Parent Heterosis (BPH) in percent were calculated for those parameters that showed significant differences among F₁ hybrids and parental lines following the method suggested by [14]. The estimate of mid and better parent heterosis, was done only when both parents and crosses had showed significance difference for the respective traits.

\[
MPH (\%) = \frac{F_1 - MP}{SE(d)} \times 100
\]

\[
BPH (\%) = \frac{(F_1 - BP)}{BP} \times 100
\]

Where, \( F_1 \) = Mean value of the cross

\( MP \) = Mean value of the mid-parent

\( BP \) = Mean value of the better parent

\( SV \) = Mean value of standard variety

Significance of heterosis has been tested using the t-test against the critical difference (CD). The CD for testing the significance of mid parent (MP) and better parent (BP) was calculated as suggested by [20,21], Cochran and Cox [20] and Singh and Chaudhary [21] as follows:

Critical difference (CD) for heterosis over MP:

\[
CD\ for\ MP = \pm (\sqrt{\frac{3eMS}{2r}} \times t)
\]

\[
SE\ (d)\ for\ MP = \pm (\sqrt{\frac{2eMS}{r}})
\]

Critical difference for heterosis over better parent.

\[
CD\ for\ BP = \pm (\sqrt{\frac{2eMS}{r}} \times t)
\]

\[
SE\ (d)\ for\ BP = \pm (\sqrt{\frac{2eMS}{r}})
\]

Association of characters

Genotypic and phenotypic correlation coefficients were calculated according to Al–Jibouri, et al. [22], from the analysis of variance and covariance as follow:

\[
Genotypic = \frac{\sigma_{g1}^2}{\sqrt{(\sigma_{e1}^2)(\sigma_{g1}^2)}}
\]
Phenotypic correlation was calculated using the formula:

$$r_{xy} = \frac{\sigma_{p12}}{\sqrt{(\sigma_{p1}^2)(\sigma_{p2}^2)}}$$

where $\sigma_{p12}$ is the phenotypic covariance between the two traits, $\sigma_{p1}^2$ is the phenotypic variance of the first trait and $\sigma_{p2}^2$ is the phenotypic variance of the second trait.

The phenotypic correlation coefficients were tested for traits of significance with 't' table for sample correlation coefficients at n-2 degree of freedom, as suggested by [13]; while the genotypic correlation coefficients were tested for their significance using the formula:

$$t = \frac{r_{gxy}}{SE_{gxy}}$$

$$SE_{gxy} = \sqrt{\frac{(1-r^2)^2}{2h^2x h^2y}}$$

The 't' value, calculated using the above formula, was compared with 't' tabled at (g-2) degree of freedom at 1% and 5% levels of significance; where, $r_{gxy}$ is the genotypic correlation between x and y traits; g = number of genotypes, h2x and h2y are heritability for traits x and y, respectively.

**Path coefficient**

A path coefficient analysis was computed according to [17].

A path coefficient analysis is simply a standardized partial regression coefficient. The general formula used was:

$$r_{ij} = p_{ij} + \sum r_{ik} p_{kj}$$

Where, $r_{ij}$ = mutual association between the independent character (i) and dependent character (j) as measured by the correlation coefficients; $p_{ij}$ = components of direct effects of the independent character (j) as measured by the path-coefficients, and $\sum r_{ik} p_{kj}$ = summation of the components of indirect effects of a given independent character (i) via all other independent characters (k).

The residual effect was computed as:

$$R = \sqrt{1 - \sum p_{ij} r_{ij}}$$

Where, $R$ is residual, $p_{ij}$ is direct effect, and $r_{ij}$ is the correlation coefficients.

**Results and discussions**

**Analysis of variances**

Analysis of variance conducted for both hybrid and inbred lines trials showed the existence of significant differences among genotypes for all traits Table 1 below.

The significant genotypic mean squares observed for most traits in both sets of trials indicated the existence of appreciable level of differences in the performances of hybrids and inbred lines for those traits. This indicates the possibility of making selection for further improvement of both sets of genotypes.

| Trait  | Hybrids (Df = 44) | Replication (Df = 2) | Blocks (Df = 18) | Error (Df = 78) | Lines (Df = 9) | Error (Df = 18) |
|--------|-------------------|----------------------|------------------|----------------|----------------|----------------|
| GY     | 3.0**             | 4.41                 | 0.97             | 0.64           | 7.31**         | 0.09           |
| EL     | 3.49**            | 36.6                 | 1.24             | 0.92           | 15.0**         | 1.17           |
| ED     | 0.31**            | 32.7                 | 0.03             | 0.03           | 0.49**         | 0.02           |
| EA     | 0.35**            | 0.1                  | 0.23             | 0.09           | 0.96**         | 0.09           |
| NKPR   | 3.5**             | 8.63                 | 0.81             | 0.59           | 2.56**         | 0.63           |
| AD     | 10.3**            | 366.7                | 9.76             | 6.05           | 77.1**         | 6.11           |
| ASI    | 34.0**            | 1.23                 | 3.27             | 1.63           | 77.7**         | 1.49           |
| PH     | 8.7**             | 0.21                 | 0.43             | 1.16           | 22.8**         | 2.19           |
| NEPP   | 262.0**           | 604.6                | 245.1            | 141.7          | 2225.9**       | 63.04          |
| EH     | 697.1**           | 202.3                | 106.4            | 96.7           | 593.8**        | 82.9           |
| NEPP   | 0.19**            | 0.05                 | 0.05             | 0.03           | 0.40**         | 0.04           |
| PA     | 0.31**            | 0.03                 | 0.18             | 0.14           | 0.73**         | 0.12           |
| RL     | 260.5**           | 377.4                | 149.9            | 146.3          | 4264.2**       | 136.9          |
| ER     | 218.9**           | 53.7                 | 41.3             | 50             | 293.6          | 136.8          |
| TLB    | 0.23**            | 0.47                 | 0.13             | 0.13           | 0.70           | 0.11           |
| PLS    | 0.31**            | 0.05                 | 0.14             | 0.16           | 0.41**         | 0.11           |

** and * signiﬁcant at probability level of P < 0.01 and P <0.05, respectively; Df: Degrees of freedom; GY: Grain yield, EL: Ear Length, ED: Ear Diameter; EA: Ear Aspect; NKPR: Number of Kernels Per Rows; AD: days to anthesis; ASI: Anthesis Silking Interval; PH: Plant height; EH: Ear Height; NEPP: Number of Ears Per Plant; PA: Plant Aspect; RL: Root Lodging; ER: Ear Rot; TLB: Turcicum Leaf Blight; PLS: Pheasphorial Leaf Spot and Values with no asterisk are non-signiﬁcant.
The non-significant differences observed among the parental inbred lines for TLB, PLS and ER might attribute to the fact that the selected inbred lines were relatively resistant to these diseases as a result of prior selection as compared with the hybrids.

Similar to the current study, other authors have reported the existence of significant differences among hybrids and inbred lines Habtamu, et al. [23]. This is because the selection of inbred lines for higher performances and for different traits to exploit hybrid vigor/heterosis in the crosses made among them.

**Heterosis estimates**

For GY, all F1 hybrids showed positive and highly (P<0.01) significant mid and better parent heterosis except cross L5×L2, L8×L7 and L9×L5 crosses were showed significant (P<0.05) better parent heterosis, and L5×L3 that revealed non-significant (P >0.05) better parent heterosis. Mid and better parent heterosis for GY ranged from 115% and 25.67 to 95.19%, respectively. While a better parent and mid parent heterosis for this trait. In contrast to the current study, Habtamu, et al. [23], reported low mid and better parent heterosis values for late maturity group of Maize inbred lines. The results presented in Table 2. With respect to days to Anthesis (AD), values of mid and better parent heterosis for all traits are positive and significant mid parent and better parent heterosis ranging from positive to negative for EL. Among 80 F1 crosses evaluated, Gudeta [29], reported that more than 61% of the crosses had positive and significant heterosis over the better parent while more than 98% of the crosses showed positive and significant heterosis over the mid parent for EL. He also reported that most of the F1 crosses had positive and significant better parent and mid parent heterosis. The heterosis observed for EL and ED could be exploited in mid-altitude quality protein maize breeding program to develop desirable genotypes.

All the F1 crosses showed highly significant (P<0.01) and positive mid and better parent heterosis for number of kernels row⁻¹ (NKPR). Mid parent heterosis were ranged from 12.39% (L8×L7) to 66.4% (L9×L1), whereas better parent heterosis ranged from 1.08% (L7×L5) to 28.26% (L9×L1). The results of present study corroborate with the findings of Jehan, et al. [30], who observed high heterosis for NKPR in diallel crosses of maize inbred lines. Bayisa, et al. [31], also reported that 98% of the crosses they evaluated showed positive mid parent heterosis while, 65% of the same crosses had positive better parent heterosis for this trait. In contrast to the current study, Habtamu, et al. [23], reported low mid and better parent heterosis values for late maturity group of Maize inbred lines.

The disparity among the results of these studies might be attributed to the differences in the type and maturity group of the materials used.

Most F1 crosses evaluated in the current study had negative mid and better parent heterosis for number of ears per plant (EPP). Mid–parent heterosis ranged from −34.2 (L6×L5) to 55.9% (L8×L3), whereas better parent heterosis ranged from −49.9 (L6×L5) to 33.5% (L8×L3). Out of 45 F1 hybrids, only five hybrids showed significant and positive mid parent heterosis, which is desirable as it indicates the proliﬁcacy of the F1 progenies as compared to the parental lines up on hybridization. On the other hand, negative heterosis indicates that parents bear a greater number of EPP than their F1 progenies. This result agrees with the ﬁndings of Bello and Olayiwaju [27], who observed only six crosses with signiﬁcant and positive heterosis for number of EPP out of 28 crosses evaluated. The present result is contrasting with the ﬁndings of Dagne, et al. [26], who reported higher number of F1 hybrids with positive mid and better parent heterosis but few hybrids with negative mid and better parent heterosis among the hybrids of 15 QPM inbred lines.

Mid–parent heterosis value for ear length (EL) and ear diameter (ED) ranged from 13.12 (L8×L5) to 62.64% (L7×L2) and from 7.49 (L8×L3) to 46.08% (L4×L2), respectively, indicating that all F1 crosses showed highly significant and positive mid parent heterosis for these traits. Better parent heterosis for both EL and ED varied between 0.54 (L10×L8) to 62.24% (L7×L2) and 1.73 (L8 × L6) to 36.37% (L4×L2), respectively. About 80 percent of the F1 crosses revealed signiﬁcant and positive better parent heterosis for EL, while, about 88% of the F1 hybrids showed signiﬁcant and positive better parent heterosis for ED. In line with the present ﬁndings, Berhanu [28], reported positive and signiﬁcant mid and better parent heterosis for ED in most F1 hybrids studied. Habtamu, et al. [23], reported positive and significant mid parent and better parent heterosis for Ear Height (EH) and Plant Height (PH). Mid parent heterosis for EH and PH ranged from 36 to 115% and 25.67 to 95.19%, respectively. While a better parent heterosis varied between 22.79 to 97.47% and 13.48 to 74.89% for EH and PH in that order. The positive and significant heterosis observed for PH conﬁrms the increase in hybrid vigor up on hybridization as previously reported by Berhanu [28].
### Table 2: Mid and better parent heterosis for grain yield and yield related traits in QPM hybrids evaluated at Bako during the 2015 main season.

| Cross   | Mid parent heterosis | Better parent heterosis |
|---------|-----------------------|--------------------------|
| Code    | GY EPP EL ED NKR GY EL ED NKR |
| L2 x L1 | 218.3** -13.3 49.9** 30.0** 11.6** | 136.6** 44.3** -22.5 26.3** 7.42 |
| L3 x L1 | 183.2** 18.5* 56.4** 24.0** 0.0 | 144.1** 51.4** 10.9 16.1** 0.0 |
| L3 x L2 | 237.2** -9.26 53.6** 38.7** 2.78 | 128.1** 43.3** -23.4* 33.4** 1.83 |
| L4 x L1 | 381.3** 21.8* 51.3** 32.4** 4.19 | 272.8** 42.5** 13.8 20.5** 2.75 |
| L4 x L2 | 507.7** 13.6 56.0** 46.1** 3.46 | 473.1** 41.3** 8.24 36.4** 2.28 |
| L4 x L3 | 136.8** 14.5 26.7** 18.4** -0.82 | 65.5** 22.6** 0.67 14.8** -1.09 |
| L5 x L1 | 118.7** -16.5* 34.3** 23.4** 3.5 | 63.8** 20.1** -32.2** 20.7** 2.3 |
| L5 x L2 | 97.2** -28.8** 28.0** 20.2** 8.84* | 22.5* 10.6* -46.9** 17.8** 5.73 |
| L5 x L3 | 38.0** -18.9* 18.9** 17.6** 4.02 | 16.5 9.49 -30.4** 11.5** 2.65 |
| L5 x L4 | 125.2** -13.2 42.2** 20.2** 14.0** | 119.1** 37.0** -25.0* 3.43 8.91 |
| L5 x L5 | 175.0** -3.6 35.0** 23.8** 12.6** | 89.2** 26.0** -5.6 36.4** 2.28 |
| L6 x L1 | 146.0** -14.6 42.3** 23.2** 12.9** | 107.2** 41.5** -21.7 12.6** 8.23 |
| L6 x L2 | 94.6** -34.2** 25.8** 22.3** 0.0 | 31.7** 11.9* -49.9** 10.3** 2.67 |
| L6 x L3 | 52.4** -34.2** 25.8** 22.3** 0.0 | 31.7** 11.9* -49.9** 10.3** 2.67 |
| L6 x L4 | 175.0** -3.6 35.0** 23.8** 12.6** | 89.2** 26.0** -5.6 36.4** 2.28 |
| L7 x L1 | 146.0** -14.6 42.3** 23.2** 12.9** | 107.2** 41.5** -21.7 12.6** 8.23 |
| L7 x L2 | 507.7** 13.6 56.0** 46.1** 3.46 | 473.1** 41.3** 8.24 36.4** 2.28 |
| L7 x L3 | 136.8** 14.5 26.7** 18.4** -0.82 | 65.5** 22.6** 0.67 14.8** -1.09 |
| L7 x L4 | 125.2** -13.2 42.2** 20.2** 14.0** | 119.1** 37.0** -25.0* 3.43 8.91 |
| L7 x L5 | 175.0** -3.6 35.0** 23.8** 12.6** | 89.2** 26.0** -5.6 36.4** 2.28 |
| L8 x L1 | 146.0** -14.6 42.3** 23.2** 12.9** | 107.2** 41.5** -21.7 12.6** 8.23 |
| L8 x L2 | 94.6** -34.2** 25.8** 22.3** 0.0 | 31.7** 11.9* -49.9** 10.3** 2.67 |
| L8 x L3 | 52.4** -34.2** 25.8** 22.3** 0.0 | 31.7** 11.9* -49.9** 10.3** 2.67 |
| L8 x L4 | 175.0** -3.6 35.0** 23.8** 12.6** | 89.2** 26.0** -5.6 36.4** 2.28 |
| L9 x L1 | 200.5** -4.34 46.1** 27.5** 0.72 | 175.8** 35.3** -25.0* 14.7** 4.66 |
| L9 x L2 | 242.2** 14.0 46.1** 27.5** 0.72 | 175.8** 35.3** -25.0* 14.7** 4.66 |
| L9 x L3 | 144.8** -27.8** 42.9** 22.1** 4.87 | 181.8** 35.4** 18.5 7.65* 3.19 |
| L9 x L4 | 196.0** 34.4** 38.2** 23.1** 4.87 | 98.1** 11.9 36.9** 15.8** 3.19 |
| L9 x L5 | 63.3** -11.1 16.9** 21.9** -1.4 | 24.5** -19.2** 9.93 19.3** -1.85 |
| L9 x L6 | 113.1** -12 39.9** 23.3** 5.26 | 93.1** 28.9** 9.44** 2.61 |
| L9 x L7 | 150.2** 2.86 45.2** 18.9** 0.72 | 132.2** 30.1** 13.7** 0.48 |
| L9 x L8 | 94.9** 2.9 27.4** 13.8** 5.94 | 61.2** 11.9** 12.7** 3.88 |

** and * significant at probability level of P < 0.01 and P <0.05, respectively; Df = Degrees of freedom, CD= critical difference, GY=Grain yield, EPP= number of ears per plants, EL= ear length, ED=ear diameter, NKP= number of kernels row, and Values with no asterisk are non-significant.
Table 2: Continued.

| Cross     | AD    | EH    | PH    | NKPR  | AD    | EH    | PH    | NKPR  |
|-----------|-------|-------|-------|-------|-------|-------|-------|-------|
| L2 x L1   | -10.3** | 62.0** | 62.5** | 55.9** | -3.62* | 52.5** | 38.7** | 24.5** |
| L3 x L1   | -9.20** | 80.0** | 66.0** | 50.8** | -9.06** | 73.2** | 50.5** | 27.7** |
| L3 x L2   | -12.2** | 72.0** | 82.7** | 31.5** | -5.43** | 56.4** | 70.7** | 18.3** |
| L4 x L1   | -11.5** | 101.0** | 75.4** | 61.7** | -9.06** | 95.4** | 69.1** | 26.8** |
| L4 x L2   | -14.5** | 115.0** | 95.2** | 50.5** | -5.43** | 97.5** | 71.9** | 20.8** |
| L4 x L3   | -8.60** | 85.0** | 73.4** | 21.3** | -6.27** | 83.4** | 62.6** | 5.8** |
| L5 x L1   | -9.84** | 50.0** | 36.5** | 35.2** | -5.98** | 33.0** | 24.3** | 26.4** |
| L5 x L2   | -9.45** | 49.0** | 47.8** | 14.0** | -6.79** | 26.3** | 16.9** | 15.2** |
| L5 x L3   | -9.20** | 49.0** | 43.9** | 13.8** | -5.13** | 37.0** | 20.0** | 16.0** |
| L5 x L4   | -8.77** | 73.0** | 51.6** | 22.6** | -2.14 | 58.2** | 36.6** | 18.0** |
| L6 x L1   | -10.6** | 64.0** | 50.2** | 52.8** | -3.64* | 58.0** | 43.5** | 25.0** |
| L6 x L2   | -13.8** | 45.0** | 60.7** | 29.1** | -13.6** | 41.5** | 42.7** | 14.1** |
| L6 x L3   | -14.1** | 68.0** | 69.2** | 35.3** | -7.27** | 56.3** | 60.1** | 11.9** |
| L6 x L4   | -14.3** | 84.0** | 73.4** | 21.3** | -6.00** | 72.9** | 52.9** | 16.0** |
| L6 x L5   | -7.93** | 43.9** | 34.1** | 18.1** | -5.00** | 22.8** | 17.3** | 2.04** |
| L7 x L1   | -12.2** | 57.0** | 58.9** | 66.4** | -9.24** | 44.1** | 50.2** | 27.1** |
| L7 x L2   | -12.0** | 71.0** | 63.9** | 51.7** | -6.80** | 48.9** | 47.0** | 20.1** |
| L7 x L3   | -9.53** | 62.0** | 73.3** | 37.4** | -6.30** | 53.6** | 65.8** | 11.0** |
| L7 x L4   | -11.5** | 83.0** | 78.6** | 45.3** | -5.88** | 72.2** | 74.9** | 17.5** |
| L7 x L5   | -9.32** | 36.0** | 36.1** | 19.5** | -8.55** | 31.3** | 17.8** | 1.08** |
| L7 x L6   | -8.30** | 68.0** | 60.4** | 39.3** | -4.55** | 49.6** | 58.6** | 14.7** |
| L8 x L1   | -11.3** | 74.0** | 63.9** | 51.7** | -9.09** | 48.9** | 43.6** | 27.5** |
| L8 x L2   | -10.6** | 85.0** | 73.8** | 23.2** | -6.34** | 61.8** | 49.6** | 21.4** |
| L8 x L3   | -9.9** | 76.0** | 66.4** | 18.4** | -7.44** | 68.1** | 52.1** | 14.3** |
| L8 x L4   | -12.2** | 73.0** | 62.3** | 23.8** | -7.44** | 64.0** | 58.0** | 20.9** |
| L8 x L5   | -9.24** | 40.0** | 25.7** | 12.9** | -7.69** | 35.1** | 13.5** | 8.23** |
| L8 x L6   | -9.09** | 61.0** | 56.9** | 23.4** | -4.55** | 43.3** | 51.3** | 19.9** |
| L8 x L7   | -5.83** | 51.0** | 47.9** | 12.4** | -5.04** | 50.3** | 41.1** | 17.5** |
| L9 x L1   | -11.6** | 60.0** | 60.1** | 64.5** | -10.55 | 50.7** | 48.0** | 28.3** |
| L9 x L2   | -11.3** | 75.0** | 82.3** | 44.6** | -5.88** | 65.1** | 67.0** | 19.5** |
| L9 x L3   | -11.7** | 75.0** | 75.4** | 38.7** | -10.55 | 51.6** | 71.7** | 13.0** |
| L9 x L4   | -11.2** | 92.0** | 69.5** | 57.5** | -7.66** | 67.0** | 62.3** | 23.0** |
| L9 x L5   | -7.68** | 53.0** | 44.9** | 21.2** | -5.13** | 23.2** | 23.1** | 3.24** |
| L9 x L6   | -8.12** | 68.0** | 59.3** | 41.6** | -2.27 | 55.8** | 53.9** | 17.1** |
| L9 x L7   | -11.9** | 62.0** | 58.9** | 41.9** | -10.11** | 34.4** | 55.3** | 19.0** |
| L9 x L8   | -12.7** | 60.0** | 57.7** | 23.2** | -11.66** | 33.3** | 47.1** | 2.28** |

** and * significant at probability level of P < 0.01 and P < 0.05, respectively; CD= critical difference, AD=days to anthesis, PH= plant height, EH= ear height, NKPR= number of kernels per rows, and Values with no asterisk are non-significant.

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**Association of characters**

Association of yield and other attributes assumed special importance as basis for selection of desired strain. Genetic correlation between different characters can be often also because of either.

**Genotypic and phenotypic correlation**

The values of estimated genotypic and phenotypic correlation coefficient between pair of characters in all possible combination are presented in Table 3. It was found that the genotypic correlation coefficients were higher than the corresponding phenotypic correlation coefficient for all traits in similar direction. Similar with the present study, Assaduzzaman [34], reported the genotypic correlation coefficients were higher than their corresponding phenotypic correlation for all traits studied on fourteen Lablab genotypes. The superiority of genotypic correlation coefficients indicated the less influence of environmental factors on expression of the traits and phenotypic correlation indicated the influence of environmental factors. Therefore, the result showed that a fairly strong inherent association between the characters studied Munawar, et al. [35].

Genotypic correlation coefficient is the heritable association between two variables. However, phenotypic correlation includes both phenotypic and environmental effect. Hence, significant phenotypic correlation without significant genotypic correlation has no value Bekele and Rao [36]. This explaining why genotypic correlation showed more significant difference between the pairs of traits than phenotypic correlation.

In this study, Grain Yield (GY) had exhibited strong significant positive genotypic and phenotypic correlation coefficient with Ear Length (EL), number of kernels row-1 (NKPR), Ear Plant-1 (EPP). It had also highly (P <.01) significant positive association with plant height (PH) and Ear Height (EH) at phenotypic correlation level and with PH which had significant (P <.05) positive association at genotypic level. Ear height and days to Anthesis (AD) had positive significant correlation with GY only at phenotypic level. Such results could help the breeder to select high grain yielder varieties through selection for one or more of these characters. The selection for long ear, more NKPR, EPP, taller PH and high ear placement may be accompanied by increasing maize GY. Similar results were reported by, Akeel, et al. [37] and Nzuve, et al. [38]. Other research work agreed with this study was showed that the strong correlation EH and PH had with grain yield suggested that, tall plants with high ear placement gave better yields to the shorter plants with lower ear placement Nzuve, et al. [38]. This indicated that by increasing these attributes in growth parameter especially plant height would help photosynthetic apparatus to synthesize more assimilates and hence production of higher yield.

It is only at phenotypic level, GY had significant (P <.05) positive association with the AD. In contrast to the present findings, Shashidhara [39], proven that, GY had negatively associated with AD at both genotypic and phenotypic level, whereas, PH, EL and NKPR had significant positive correlation again at both genotypic and phenotypic level. In addition, Number of Kernels Row (NKR) had positive correlation at phenotypic level with GY. This is according to various research work reported by Amini, et al. [40], Bekele and Rao [36] and Adesoji, et al. [41], GY had significant and positive association with EH, PH, EL, NKPR, NKR, 100 seed weight and total dry weight by the studied in different locations and year.

**Table 3: Phenotypic (above diagonal) and genotypic (below diagonal) correlation coefficient of yield and yield related traits of 45 single crosses.**

| Traits  | GY  | AD  | ASI  | EH  | PH  | RL  | EPP  | EL  | ED  | NKPR | PA  | EA  | TLB  | CLR  | PLS  | ER  |
|---------|-----|-----|------|-----|-----|-----|------|-----|-----|------|-----|-----|------|------|------|-----|
| GY      | 0.19*| -0.28**| 0.25**| 0.27**| -0.19*| 0.39**| 0.40**| -0.08| 0.32**| -0.37**| -0.39**| -0.16| -0.15| -0.19*| -0.25**|
| AD      | 0.26| -0.47**| 0.45**| 0.41**| 0.19*| 0.58**| 0.19*| -0.29**| 0.11| 0.05| -0.03| -0.16| -0.15| -0.26**| -0.09|
| ASI     | -0.34*| -0.51**| -0.27**| -0.09| 0| -0.45**| -0.05| 0.11| -0.07| 0.08| -0.07| 0.02| -0.06| 0.14| 0.06|
| EH      | 0.28| 0.51**| -0.37*| 0.59**| 0.27**| 0.43**| 0.28**| -0.25**| 0.16| 0.03| 0.05| 0.03| 0.02| -0.04| -0.01|
| PH      | 0.37*| 0.55**| -0.17| 0.68**| 0.22| 0.34**| 0.35**| -0.27**| 0.24**| 0| -0.22**| -0.05| -0.06| 0.01| 0.1|
| RL      | -0.22| 0.25| 0.05| 0.33*| 0.26| 0.1| 0.12| -0.14| 0.07| 0.23**| 0.15| 0.04| 0.08| -0.18*| -0.1|
| EPP     | 0.43**| 0.71**| -0.55**| 0.56**| 0.47**| 0.24| 0.11| -0.20*| 0.07| -0.03| -0.05| -0.04| -0.09| -0.16| -0.17**|
| EL      | 0.40**| 0.28| -0.03| 0.32*| 0.43**| 0.08| 0.18| -0.42**| 0.63**| -0.13| -0.21*| -0.13| 0.04| -0.07| 0.05|
| ED      | 0.15| -0.59**| 0.2| -0.47**| -0.34*| -0.42**| -0.5*| -0.4**| -0.5**| -0.11| 0.02| -0.13| -0.04| 0.03| 0.0|
| NKPR    | 0.48**| 0.21| -0.11| 0.18| 0.27| -0.01| 0.16| 0.74**| -0.21| 0.20*| -0.19*| 0.01| -0.02| -0.05| -0.03|
| PA      | -0.57**| 0.12| 0.08| -0.04| -0.1| 0.29| -0.07| -0.18| -0.37*| -0.33*| 0.42**| 0.03| 0| 0.01| 0.22**|
| EA      | -0.45**| -0.03| -0.12| 0.09| 0.33*| 0.17| -0.06| -0.26| 0.21| -0.26| 0.58**| 0.03| 0.16| 0.11| 0.4**|
| TLB     | -0.37*| -0.21| 0.12| -0.04| 0.09| 0.02| -0.12| -0.45**| 0.11| -0.44**| 0.26| 0.21| 0.07| 0.04| -0.11|
| CLR     | -0.32*| -0.23| 0.01| 0.03| 0.08| 0.12| -0.08| -0.11| -0.11| -0.12| 0.16| 0.28| 0.23| 0.42**| 0.03|
| PLS     | -0.31*| -0.35*| 0.19| 0| -0.02| 0.14| -0.24| -0.15| 0.05| -0.11| 0.02| 0.2| 0.11| 0.58**| 0.16|
| ER      | -0.37*| -0.09| 0.02| 0.17| -0.08| 0.26| -0.25| 0.04| 0.37*| 0.44**| -0.04| -0.07| 0.14|

** and * significant at probability level of P < 0.01 and P < 0.05, respectively. GY: Grain yield, EL: Ear Length, ED: Ear Diameter; EA: Ear Aspect; NKPR: Number of Kernels Row; NKR: Number of Kernels Per Rows; AD: days to anthesis; ASI: Anthesis Silking Interval; PH: Plant height; EH: Ear Height; EPP: Number of Ears Per Plant; PA: Plant Aspect; RL: Root Lodging; ER: Ear Rot; TLB: Turcicum Leaf Blight; PLS: Phaeosporial Leaf Spot and Values with no asterisk are non-significant.

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In disagreement of the present findings, Aminu and Izge [42], reported AD, EH and PH had exhibited negative correlation with GY and suggested that these traits were not closely associated and therefore, they may not be jointly selected. The difference results were found due to the genotypes was evaluated under drought condition. On the other hand, Berhanu [28], Bello, et al. [43] and Kinfe, et al. [44], reported that, GY had significant and positive phenotypic correlation with EH, PH, EL, EPP, NKPR and AD in agreement with this study and proven the existence of direct association between the traits.

Other traits i.e., plant Aspect (PA), Ear Aspect (EA) and Anthesis Silking Interval (ASI) had strongly significant negative association with grain yield at genotypic and phenotypic level, except ASI had moderately significant at phenotypic level. Where, ASI, Ear Rot (ER) and Phaeosphaeria Leaf Spot (PLS) had significant negative association at genotypic and phenotypic level, and Turcicum Leaf Blight (TLB) had at genotypic level. This revealed that, by decreasing these attributes, could consistently increase grain yield. The selection made to improve yield of maize genotype may be useful through decreasing these traits. In line with the current study, Hadji [45], observed ASI, Ear Husk Cover (HC) and EPP had showed significant negative association with GY at genotypic level. This showed that, these genotypes had short days to ASI, husked cobs and a smaller number of EPP had potential to give high grain yield.

However, non-significant correlations were observed between GY and other traits due to masking effects of environment. This is indicating that selection for increase the level of these traits may not bring significant change in GY.  

Path coefficient

Phenotypic and genotypic path coefficient analysis is a proved effective means of separating direct and indirect effect of associated traits on yield. The analysis using grain yield as a dependent variable was conducted for the traits that exhibited significant genotypic and phenotypic association with yield. The phenotypic and genotypic direct (bold) and indirect effects of twelve and eleven traits on grain yield were presented in Tables 4,5 below, respectively.

Days to Anthesis (AD) had a negative direct effect on Grain Yield (GY) at phenotypic level. The correlation coefficient between the two traits was positive and significant (P < 0.05). Moreover, the negative indirect contribution of AD to GY was through number of Ears Per Plant (EPP) at phenotypic level. Since correlation is positive, but the direct effect is negative, the indirect effects seem to be cause for correlation. This result agrees with some earlier findings, Saleem [46], reported AD had negative direct on grain yield by the study on ten S1 families evaluated under irrigated and drought condition. Therefore, in such situations, the indirect causal factors are to be considered simultaneously for selection.

At both genotypic and phenotypic level, Anthesis Silking Interval (ASI) had showed highly significant negative correlation and negative direct effect on grain yield. Ear Height (EH) had highly significant positive correlation and direct effect on GY at phenotypic level; while, the genotypic correlation is positive and statistically non–significant. In contrast to this finding, Muhammad et al. [15] reported that, EH had negative direct effect on GY by the study on eight local hybrids Maize. The different result was obtained may be due to the use of different source materials for the study. In agreement with the current finding, Hadji [45], observed, EH had exerted positive direct effects on GY at phenotypic level.

Other works agreed with this finding reported by Munawar, et al. [35], showed that the number of kernels row–1 (NKPR) had positive direct effect followed by EH for seven exotic hybrids maize sourced from different seed companies in Pakistan. Similarly, Sreckov, et al. [47], they reported EH had positive direct effect on GY, however, it was non–significant.

Table 4: Direct and indirect effects of yield contributing traits, phonological and growth and disease parameters on yield at phenotypic level.

| Traits | AD  | ASI | EH  | PH  | RL  | EPP | EL  | NKPR | PA  | EA  | PLS1 | ER  | Rp   |
|--------|-----|-----|-----|-----|-----|-----|-----|------|-----|-----|------|-----|------|
| AD     | -0.190 | 0.072 | 0.05 | 0.03 | -0.047 | 0.171 | 0.055 | 0.005 | -0.008 | 0.004 | 0.039 | 0.010 | 0.26 |
| ASI    | 0.09 | -0.153 | -0.031 | -0.006 | 0.00 | -0.132 | -0.014 | -0.003 | -0.011 | 0.011 | -0.021 | -0.007 | -0.34 |
| EH     | -0.09 | 0.042 | 0.112 | 0.043 | -0.068 | 0.127 | 0.080 | 0.007 | -0.004 | -0.007 | 0.007 | 0.002 | 0.28 |
| PH     | -0.08 | 0.014 | 0.067 | 0.072 | -0.055 | 0.099 | 0.100 | 0.011 | 0.000 | 0.033 | -0.001 | 0.012 | 0.37 |
| RL     | -0.04 | 0.00 | 0.03 | 0.016 | -0.252 | 0.029 | 0.035 | 0.003 | -0.034 | -0.022 | 0.028 | 0.011 | -0.26 |
| EPP    | -0.11 | 0.069 | 0.049 | 0.024 | -0.025 | 0.294 | 0.031 | 0.003 | 0.004 | 0.008 | 0.024 | 0.020 | 0.43 |
| EL     | -0.04 | 0.007 | 0.032 | 0.025 | -0.031 | 0.032 | 0.286 | 0.029 | 0.020 | 0.031 | 0.011 | -0.006 | 0.40 |
| NKPR   | -0.02 | 0.011 | 0.018 | 0.017 | -0.018 | 0.02 | 0.18 | 0.045 | 0.029 | 0.028 | 0.008 | 0.003 | 0.48 |
| PA     | -0.01 | -0.012 | 0.003 | 0.00 | -0.057 | -0.008 | -0.038 | -0.009 | -0.149 | -0.062 | -0.001 | -0.27 | -0.57 |
| EA     | 0.01 | 0.011 | 0.005 | -0.016 | -0.036 | -0.015 | -0.06 | -0.008 | -0.062 | -0.15 | -0.016 | -0.048 | -0.45 |
| PLS1   | 0.05 | -0.021 | -0.005 | 0.00 | 0.046 | -0.046 | -0.021 | -0.002 | -0.001 | -0.016 | -0.151 | -0.019 | -0.31 |
| ER     | 0.02 | -0.009 | -0.002 | -0.007 | 0.024 | -0.005 | 0.014 | -0.001 | -0.033 | -0.060 | -0.024 | -0.120 | -0.37 |

Residual effects = 69.5%

AD: days to anthesis; ASI: Anthesis Silking Interval; PH: Plant height; EH: Ear Height; EPP: Number of Ears Per Plant; PA: Plant Aspect; RL: Root Lodging; ER: Ear Rot; TLB: Turcicum Leaf Blight, PLS: Pheosphaeria Leaf Spot; EA: Ear Rot and Rp: Phenotypic correlation.

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The magnitude of direct effect of Plant Height (PH) on grain yield was very small at phenotypic level, where both genotypic and phenotypic correlations were positive and statistically significant. Therefore, the direct effect and correlation coefficient explain the true association between the two characters and selection for having tallest plant feature will improve GY. Similar with the current study, Adesoji, et al. [41], reported that, growth parameters such as plant height had positive direct effect on GY by the study on two maize varieties at Nigeria. However, there is a contrasting finding with the present findings was reported by Munawar, et al. [35] and Zarei, et al. [48]; Adesoji, et al. [41] and Kinfe, et al. [44], also reported similar result with the current findings for EL and NKPR. These traits are therefore, very important components of GY and should be given high emphasis in any selection process aimed at improving grain yield in maize.

In agreement of the present study, Muhammad, et al. [15], reported similar results and they concluded that, the effective selection for superior genotypes is possible considering, ear length and number of kernels row\(^{-1}\). Researchers like, Rafiq, et al. [49], Zarei, et al. [48]; Adesoji, et al. [41] and Kinfe, et al. [44], also reported similar result with the current findings for EL and NKPR. It is therefore, concluded that these GY related agronomic parameters could be considered as important selection criteria in improving hybrid and open pollinate maize varieties Bello, et al. [43].

Turchicum Leaf Blight (TLB) had a positive lesser direct effect on GY at genotypic level. The genotypic correlation coefficient between the traits and GY was statistically significant and negative. The correlation explains, low disease severity level or tolerant genotypes had a potential to yield more. While the direct effect was negligible, the indirect effect seemed to be cause of correlation. In such conditions, the other indirect causal factors are to be considered simultaneously for selection.

Common Leaf Rust (CLR) had negative direct effect on GY at genotypic level. The genotypic correlation coefficient was negative as well. On the other hand, phenotypic correlation showed non-significant association with GY. The negative indirect effect CLR exerted on GY was enhanced by negative indirect effect through Phaeosphaeria Leaf Spot (PLS). The correlation results explain that, improving the resistance level of genotypes was boosting GY. Therefore, selecting of

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**Table 5: Direct and indirect effects of yield contributing traits, phonological and growth and disease parameters on grain yield at genotypic level enhance.**

| Traits | ASI | PH | EPP | EL | NKPR | PA | EA | TLB | CLR | PLS1 | ER | Rg |
|--------|-----|----|-----|----|------|----|-----|-----|-----|------|----|----|
| ASI    | -0.190 | -0.018 | -0.066 | -0.003 | -0.019 | -0.026 | 0.00 | -0.008 | 0.020 | -0.012 | -0.018 | -0.34 |
| PH     | 0.03 | 0.106 | 0.056 | 0.042 | 0.047 | 0.034 | -0.001 | 0.006 | 0.017 | 0.001 | 0.034 | 0.37 |
| EPP    | 0.10 | 0.049 | 0.121 | 0.017 | 0.027 | 0.024 | 0.00 | 0.008 | 0.015 | 0.015 | 0.052 | 0.43 |
| EL     | 0.01 | 0.045 | 0.021 | 0.10 | 0.129 | 0.060 | -0.001 | 0.031 | 0.022 | 0.009 | -0.025 | 0.40 |
| NKPR   | 0.02 | 0.029 | 0.019 | 0.073 | 0.175 | 0.113 | -0.001 | 0.030 | 0.024 | 0.007 | -0.008 | 0.48 |
| PA     | -0.02 | -0.011 | -0.009 | -0.018 | -0.058 | -0.338 | 0.002 | -0.018 | -0.032 | 0.001 | -0.075 | -0.57 |
| EA     | 0.02 | -0.034 | -0.007 | -0.026 | -0.045 | -0.194 | 0.003 | -0.015 | -0.057 | -0.012 | -0.089 | -0.45 |
| TLB    | 0.02 | -0.009 | -0.009 | -0.011 | -0.021 | -0.053 | 0.001 | -0.016 | -0.203 | -0.036 | 0.014 | -0.32 |
| CLR    | -0.04 | -0.002 | -0.028 | -0.015 | -0.019 | 0.007 | 0.001 | -0.007 | -0.117 | -0.062 | -0.028 | -0.31 |
| PLS1   | -0.02 | -0.018 | -0.031 | 0.012 | 0.007 | -0.126 | 0.001 | 0.003 | 0.014 | -0.008 | -0.202 | -0.37 |

Residual effects = 58.8%

ASI: Anthesis Silking Interval; PH: Plant Height; EPP: Number of ears plant\(^{-1}\); EL: Ear length; NKPR: Number of kernels row\(^{-1}\); PA: Plant Aspect; EA: Ear Aspect; TLB: Turchicum Leaf Blight; CLR: Common Leaf Rust; PLS: Phaeosphaeria Leaf Spot; ER: Ear Rot and Rg=Genotypic correlation.

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genotypes for resistant to CLR could be effective by considering indirect causal factors at the same time.

The other maize foliar disease was PLS which had significant negative genotypic and phenotypic correlation coefficient with GY. In addition, the magnitude of the direct effect was negative at both genotypic and phenotypic level. The correlation coefficient at both genotypic and phenotypic level showed that, the genotypes with minimum disease reaction score could have a potential to boost the yield. Again, the selection for disease resistant high yielding hybrids should consider the indirect causal factors which could enhance yield and reduce disease development.

The other bad character encountered GY was Ear Rot disease (ER). It had negative and highly significant (P <0.01) correlation at phenotypic level and significant (P <0.05) correlation at genotypic level. At both genotypic and phenotypic level, it had negative direct effect on GY. Like other disease parameter, the selection of genotypes with a smaller number of ears susceptible to ER could help to get the varieties giving high yield and vice-versa. Therefore, the selection for ER free hybrids will be valuable if it considered other indirect causal factors in addition to the disease.

The residual effect estimation of 69.51% indicated that the causal variables explained only about 30.49% of the variability in grain yield and the remaining 69.51% of variability stays unexplored at phenotypic level. On the other hand, the residual effect of 58.84% exhibited that the fundamental variables elucidated only about 41.16% of the variability in grain yield and about 58.84% of the variability remain uninvestigated at genotypic level of path coefficient analysis. In contrast with the current study, Hadji [45], reported small residual effect 44% at phenotypic and 11% very small genotypic level for QPM inbred lines evaluated at the same location with the current study. Similary, Adesoji, et al. [41] and Kinfe, et al. [44], found small residual effects as compared with the current finding.

The reason seems to be very low variability in the present study were due to non-significant correlations coefficient of the remaining traits with the causal factor, GY at both genotypic and phenotypic level. Besides, some other factors which have not been considered here need to be included in this analysis to account fully for the remaining variation in grain yield. It means that, the high value obtained in residual effects indicated that other factors and variables not considered in this study were of high effect on grain yield. Similar with this work, Oad, et al. [50] found the maximum (79%) residual effect for twenty-two selected F4Chickpea genotypes obtained from ICARDA. However, it was not reported for maize, Abebe [52,53], found maximum residual effects (72.48 percent) at phenotypic level for Ethiopian Mustard.

Summary and conclusion

In this study, almost all crosses were showed positive and highly significant mid and better parent heterosis for all traits assessed. In both cases, the same cross, L4×L2 and L5×L3 consistently manifested the highest and lowest percentage of heterosis respectively.

The estimation of phenotypic and genotypic correlation coefficient between pair of characters in all possible combination revealed the presence of a true association among traits. At phenotypic correlation level, GY had significant positive association with, EL, NKPR, AD, PH, EH, EPP and TGB. However, it had significant and negative correlation with EA, ASI, PA, RL, ER and PLS. The positive phenotypic correlation showed as by increasing these traits there is a possibility to increase GY while the negative association indicated as, increasing effects of these traits may resulted in decreasing GY considerably.

Similarly, at genotypic level, GY had significant positive correlation with, EL, NKPR, PH and EPP which indicated that, breeding program engaged to increase these traits will increase GY production. On the other hand, it had significant and negative correlation with EA, PA, ER, TLB and PLS. This showed that, by decreasing the effects of bad traits, invariably increase yield and since little score was given for highly attractive cobs and for plants with good stature.

Traits like, EPP and EL had the maximum positive direct contribution to GY at both genotypic and phenotypic level. Whereas, PA, ER and ASI had negative direct effect on grain yield at both genotypic and phenotypic. In addition, PH and EPP had positive direct contribution at both genotypic and phenotypic correlation, however, EA and EH had positive direct effect on GY at genotypic and phenotypic correlation level respectively. Where, AD, RL and EA had negative direct influences on GY at phenotypic level and TLB and CLR had negative direct effect on GY at genotypic level. So, the positive and negative direct effects were nullified and enhanced by their respective negative and positive indirect effect to make counter balanced the effect on GY via other characters indirectly.

In general, it can be concluded that the research results suggested the importance of continuous and extensive research on quality protein maize best fit to mid altitudes of the country to generate information that can be used to design breeding strategy.

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