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Estimates of genetic parameters using six populations in two bread wheat crosses

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Abstract

Six populations of two wheat hybrids, namely Sids $14 \times \text{Giza 171}$ and Sids $14 \times \text{Gemmeiza 11}$ were established during three successive seasons of 2017/2018, 2018/2019 and 2019/2020. The results showed that there are significant differences between the mean of six populations in the cross- crossing of wheat crosses for most the studied traits, it is possible to obtain sufficient amounts of genetic variation for these traits in the studied material. The additive gene effects were significant for all traits studied except for DH in cross 1 and NSP in cross 2. The highest significant effects of the dominant genes were found to be highly significant for most traits under study, except for the number of grains/spike in the hybrid 2. The additive \times additive type of gene effects was positive and highly significant for plant height and weight of 100 grains in the two crosses, and 50% flowering to heading and grains yield/plant in cross 1. High expected genetic advance was recorded in the second generation (F₂) for grain yield per plant in the two hybrids, The results showed that there is an improvement in the genotypes possible through early selection of the higher crop.

Keywords: wheat, six population analysis, genes, heterosis, heritability, genetic advance.



1. Introduction

Wheat (Triticum aestivum L.) is first food crops in Egypt. About 730.3 million tons of wheat were produced from 217 million hectares in 2010 and 2017/2018 at a productivity level of 3 tons/ha⁻¹ (Brown et al., 2010; FAO, 2012; 2019). It is a food staple for millions of people because it provides 50% of the calorie and protein requirements of a large number of the world's population. The grain yield can be improved through indirect selection on the basis of yield components. The breeder is focusing on improving wheat yield potential by developing now divergent genotypes with a trait that may have a positive and negative effect on traits of other components (Chandara et al., 2004). Besides, factors controlling the reported heritability estimates largely indicate that some morphological traits affecting grain vield in wheat are more heritable than the crop itself (Fatehi and Mohamed, 2010). Generation mean analysis techniques were used to obtain significant information about the types of action of genes controlling yield and yield components as well as traits. The main objective of this research is to study the relative importance of genetic action, genetic progression, inbreeding depression and the degree of control of sexual traits in two bread wheat crosses using their six groups, i.e., P₁, P₂, F_1 , BC_1 , BC_2 and F_2 .

2. Materials and methods

The search has been performed in 2017/2018, 2018/2019, and 2019/2020

growing seasons, at the Experimental Farm of Faculty of Agriculture, Al-Azhar University, Assiut, Egypt. Three bread wheat cultivars were used. The name, pedigree origin and of these broad wheat genotypes are presented in Table (1). In 2017/2018 season the parents were crossed produce F1 hybrid grains and designated as follows: In 2018/2019 season some, F1 plants were back crossed to their parents to produce $BC_1(F_1 \times P_1)$ and BC₂ ($F_1 \times P_2$) generations. Also, crosses were made to produce more F₁ grains. At the same time, some F_1 hybrids were selfed to produce F₂ generation. In 2019/2020 season the six populations, i.e., P_1 , P_2 , F_1 , F_2 , Bc_1 and Bc_2 of the two hybrids were sown in a randomized complete blocks design with three replicates. Each replicate consisted of 44 rows (16 rows for F_2 , 8 rows for each BC_1 and BC₂ 4 rows each for P_1 , P_2 and F_1). Each row was 5.0 m long, 60 cm apart and 20 cm between plants. The agriculture practices recommended for wheat production were adopted in all the Characters growing seasons. were assesses using individual plants from the six populations (30 plants for each P_1 , P_2 , and F₁, 60 plants for each BC₁ and BC₂ and 120 plants for each F₂ population) the following characters were studied characters, *i.e.*, days to 50% heading (DH), plant height (PH) (cm), number of (NSP), spike/plant number of kernels/spike (NKS), 1 weight of 100 grains (GW) (g) and (GYP) grain yield/plant (g).

Table (1): The name, pedigree and origin of genotypes used in the two bread wheat crosses.

Cross		Parent	Pedigree				
Cross 1	P1	Sids- 14	BOW"S"/Vee"S"//Bow"S"/TSI/3/Beni Sewef 1 SD293-1SD-2SD-4SD –Osd				
	P2	Giza-171	Sakha93 / Gemmeiza 9				
Cross 2	P1	Sids- 14	BOW"S"/Vee"S"//Bow"S"/TSI/3/Beni Sewef 1 SD293-1SD-2SD-4SD -Osd	Egypt			
	P2	Gemmeiza 1 1	BOW"S"/KVZ"S"//7C/SERI-82/3/GIZA 168/SAKHA 61 GM7892-2GM-1GM-2GM-1GM-0GM	Egypt			

2.1 Statistical and genetic analysis

All the genetic analysis were done using generation means, the scaling tests (A, B and C) were applied according to Mather and Jinks (1982) to test the presence of non-allelic interaction as following:

$$\begin{split} A &= \ 2 \ \overline{B_1} - \overline{P_1} - \overline{F_1} \\ B &= \ 2 \ \overline{B_2} - \overline{P_2} - \overline{F_1} \\ C &= \ 4 \ \overline{F_2} - 2 \ \overline{F_1} - \overline{P_1} - \overline{P_2} \end{split}$$

Those parameters genetic model (m, a, h, aa, ad and dd) were according to Jinks and Jones (1958) and Hayman (1958):

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\begin{split} \mathbf{m} &= \text{ mean} \\ \mathbf{a} &= \text{ additive effect } = \overline{\mathbf{P}_1} - \overline{\mathbf{B}_2} \\ \mathbf{h} &= \text{ dominance effect } = \overline{\mathbf{F}_1} - 4 \ \overline{\mathbf{F}_2} - \frac{1}{2} \ \overline{\mathbf{P}_1} - \frac{1}{2} \ \overline{\mathbf{P}_2} + 2 \ \overline{\mathbf{B}\mathbf{C}_1} + 2 \ \overline{\mathbf{B}\mathbf{C}_2} \\ \mathbf{aa} &= \text{ additive } \times \text{ additive gene interaction } = 2 \ \overline{\mathbf{B}\mathbf{C}_1} + 2 \ \overline{\mathbf{B}\mathbf{C}_2} - 4 \ \overline{\mathbf{F}_2} \\ \mathbf{ad} &= \text{ additive } \times \text{ dominance } = \ \overline{\mathbf{B}\mathbf{C}_1} - \frac{1}{2} \ \overline{\mathbf{P}_1} - \overline{\mathbf{B}\mathbf{C}_2} + \frac{1}{2} \ \overline{\mathbf{P}_2} \\ \mathbf{dd} &= \text{ dominance } \times \text{ dominance } = \ \overline{P_1} + \overline{P_2} + 2 \ \overline{F_1} + 4 \ \overline{F_2} - 4 \ \overline{\mathbf{B}\mathbf{C}_1} - 4 \ \overline{\mathbf{B}\mathbf{C}_2} \end{split}
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The genetic components of variance were calculated by the formulas of F2 variance were obtained according to Mather and Jinks (1982) as:

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E (environnemental variance) = ½ (VP1 + VP2 + VF1)
D (additive variance) = 4 VF2 - 2 (VBC1 + VBC2)
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H (dominance variance) = 4 (VF2 – $\frac{1}{2}$ VD – VE)

The significant of the genetic components were tested using the t test, where t = effect / (variance effect)^{1/2}.

2.2 Heterosis

Estimate of heterosis (%) were calculated as the percent deviation of F_1 mean performance from the mid-parent or better parent as follows:

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Heterosis from the mid – parent % (M. P) = (F_1 - \overline{MP}) / \overline{MP}) \times 100
Heterosis from the better – parent % (BP) = (\overline{F_1} - \overline{BP}) / \overline{BP}) \times 100
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2.3 Inbreeding Depression (I. D %)

Its values measured from the following equation:

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I. D % = (\overline{F}_1 - \overline{F}_2 / \overline{F}_1) \times 100
Variances of I. D deviation = \overline{VF}_1 + \overline{VF}_2
T: I. D = \overline{F}_1 + \overline{F}_2 / (V. I. D)^{0.5}
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Phenotypic and genotypic coefficients of variability were calculated as outlined by Burton (1952):

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PCV = (\sqrt{VP} / \overline{X}) \times 100GCV = (\sqrt{VG} / \overline{X}) \times 100
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2.4 The average degree of dominance (\bar{a})

The average degree of dominance (\bar{a}) : was calculated by the formula presented by Mather and Jinks (1982):

 $\bar{a} = (H/D)^{1/2}$

Complete dominance is considered when $\bar{a} = \pm 1.0$, partial dominance is indicated when a lise between ± 1.0 , while overdominance is considered if lies the ratio exceeded ± 1.0 . if degree of dominance value is equal to zero, it indicates the absence of dominance. The positive and negative signs indicate the direction of dominance.

2.5 Heritability

2.5.1 Heritability in broad-sense (h^2_b)

Heritability in broad-sense (h^2_b) was estimated according to the following formula presented by Mather and Jinks (1982):

 $h^{2} {}_{b} \% = (VG / VP) \times 100$

2.5.2 Heritability in narrow-sense (h^2_n)

It was estimated according to the following formula presented by Mather and Jinks (1982):

 $h_{n}^{2} \% = (\frac{1}{2} D / VP) \times 100$

2.6 Expected gain from selection (G.S)

The expected gain from selecting (G.S)

was calculated according to Allard (1960):

G.S % = [(K × σ ph × h²_n) / F₂] × 100

3. Results and Discussion

The mean values and standard error for the sixth generation were calculated with the two hybrids for six traits and presented in Table (2). The results showed that there are significant differences between the mean of six populations in the crosscrossing of wheat crosses for all studied characters, which indicates the presence of a sufficient amount of genetic variation for these traits in the studied material. Results showed that the mean values of F_1 were higher than for all parents, BC₁ and BC₂ for 50% flowering of heading, NSP, NKS and GYP in the two hybrids, excluding PH and weigh of 100 grains in both hybrids. The backcrossed mean values were larger than both parents in the two crosses for days to 50% heading and GYP. The backcrossed mean values were higher than both parents in the two hybrids for NSP and weigh of 100 grains in second cross. The mean values of the first and second back - generations were higher than in P₂ for number of grains per spike for both hybrids. Regarding the mean value for the second generation were lower than the median values F_1 50% flowering of heading, NSP, NKS and GYP in two hybrids, except, PH and weigh of 100 grains in the two hybrids, Genetic variance is high in all traits, with PH and 100-grain weight excluded in two hybrids, while the F-value was lower than in both parents. Similar results were obtained with Awad (1996), Amin (2013), Al-Masry and Al-Nahhas (2018).

Table (2): Mean performance of parents, F_1 , F_2 , and back crosses generations in two bread wheat crosses for all studied traits.

Characters generation	Days to 50 heading	Plant height	No. spike/plant	No. kernels/spike	Weight 100 - grains	Grain yield				
Cross 1 (Sids 14 × Goemiza 11)										
P1	94.43±0.509	111.3±0.536	7,03±0.040	62,04±1.07	4,79±0.008	18.46±0551				
P2	97.53±0.408	110.83±0.561	6.33±0.021	58.20±0.965	5.00±0.011	16.60±0.609				
F1	106.46±0.433	91.13±0.782	7.37±0.034	73.34±1.82	4.58±0.130	25.66±0.967				
F ₂	99.85±0.660	93.17±0.586	7.26±0.80	73.15±1.27	4.16±0.07	18.71±0.603				
BC_1	106.44±0.738	104.95±0.757	6.84±0.034	66.07±1.81	5.15±0.009	19.23±0.647				
BC ₂	98.68±0.679	100.16±0.851	6.33±0.029	71.55±1.72	4.54±0.005	22.36±0.802				
L. S.D. _{0.05}	1.70	1.60	0.860	2.36	0.555	1.62				
		Cros	s 2 (Sids 14 × Giza 17	1)						
P1	94.43±0.509	111.3±0.536	7,03±0.040	62,04±1.07	4,79±0.008	18.46±0551				
P2	93.79±0.469	111.51±0.874	6.57±0.436	73.31±1.33	4.78±0.10	24.33±0.559				
F ₁	99.76±0.769	108.49±0.954	10.07±0.445	76.96±2.00	4.70±0.180	29.66±1.128				
F ₂	96,73±0.669	109.19±0.678	7.26 ±0.168	74.83 ±1.11	4,86 ±0.062	25.45 ±0.632				
BC1	97.76±0.890	108.75 ± 0.907	9.21±0.345	76.55±1.50	5.49±0.082	25.91±0.653				
BC ₂	96.11±0.764	116.35±0.720	9.98±0.322	66.11±1.46	5.20±0.086	26.29±0.820				
L. S.D. _{0.05}	1.71	1.72	1.05	2.21	0.522	1.66				

Results of scaling test (A, B and C) together with the six parameters model and type of epistasis are calculated and given in Table (3). It is worthy to mention that the least one of the A, B and C tests were significant for all studied traits except, 50% flowering to heading in cross 1. It can be said from the obtained results that the six parameters model is valid to explain the nature of the gene. Furthermore, the A, B, or C measures were not significant, indicating that the interactive model failed to explain the type of gene action in this case. These results are in general agreement with those of Shafey *et al.* (1993), Tammam (2005), Kattab *et al.* (2010), El-Aref *et al.* (2011), Zaazaa *et al.* (2012), and Amin (2013) for NSP and NKS and GYP and Moussa (2010) for DH, PH, and 100-grain weight by Lal *et al.* (2013). On the other hand, Abdel-Radi (2018) explained that the scaling test indicated a non-allelic interaction for all studied traits except for NSP in hybrid 2 and NGS and GYP in hybrid 1 under normal conditions.

Table (3): The scaling test and estimates of the additive (D), dominance (H) and interaction parameters in two bread wheat hybrids for all studied traits.

C1t	Cross	Genetic parameter								
Character		А	В	С	М	А	Н	Aa	Ad	dd
D	1	12.03**±1.61	6.36**±1.48	-5.5±2.86	99.85**±0.66	7.78**±1.00	21.38**±3.07	10.9**±3.33	9.33**±1.05	-16.30**±4.94
Days to 50 % neading	2	1.43±2.00	-1.33±1.77	3.28±3.16	96.73*8±0.669	1.65±1.17	6.52*±3.65	0.82±3.56	1.38±1.22	92±5.65
Dlant hainht	1	7.46**±1.78	-1.63±1.95	-31.7**±2.97	93.17**±0.586	4.78**±1.13	17.60**±3.12	37.53**±3.26	4.54**±1.20	-43.36**±5.41
r lant neight	2	-2.29±2.11	12.7**±1.93	-3.55±3.47	109.19**±0.678	-7.60**±1.15	10.52*±3.72	13.44**±3.56	-7.49**±1.26	-23.85**±5.79
Number of miles/alast	1	-0.730±0.461	-1.03**±0.418	0.96±0.92	7.26**±0.168	0.50*±0.27	-2.52**±0.889	-2.73**±0.860	0.15±0.31	3.56*±1.45
Number of spikes/plant	2	1.32±0.862	3.32**±0.89	-2.18±1.43	7.46**±0.251	-0.77±0.477	11.81**±1.47	8.54**±1.39	-1.00±0.536	13.18**±2.37
	1	6.18**±3.07	20.97**±2.99	44.53**±5.68	73.15**±1.27	-5.47**±1.88	-13.58*±5.95	-17.37**±6.32	-7.39**±2.01	-9.78±9.42
Number of kernels/spike	2	14.1**±3.75	-18.05**±3.79	-10.87±6.22	74.83**±1.11	10.44**±2.100	-4.71±6.49	-14.00**±6.12	16.07**±2.26	17.95**±2.26
Wighet of 100 - grains	1	0.914.**±0.211	-0.517*±0.213	-2.32** ±0.394	4.16**±0.07	0.611**±0.119	2.40**±0.368	2.721**±0.374	0.716**±0.139	-3.11**±0.621
	2	1.48**±0.217	0.92**±0.22	-1.12**±0.35	4.86**±0.062	0.29*±0.119	1.85**±0.367	1.94**±0.344	0.28*±0.137	-4.34**±0.595
Grain yield/plant	1	-5.65**±1.70	2.45±1.97	-11.53*±3.29	18.71**±0.603	-3.12**±1.03	16.47**±3.23	8.34**±3.17	-4.05**±1.110	-5.15±5.22
	2	3.71**±1.81	-1.41±2.06	7.17**±3.84	25.45**±0.632	-0.371**±1.04	10.88**±3.49	2.61±3.28	-4.34**±0.559	-4.92±5.45

3.1 Gene effects

The six parameters of gene effect, i.e., means (m), additive (d), dominance (h), additive \times additive (aa), additive \times dominance (ad) and third type of epistatic effect dominance × dominance (dd) are presented in Table (3). Average is high and significant in all traits in the two hybrids. The additive gene effects (d) all traits studied were significant (*) and either positive or negative except for 50% flowering to heading in hybrid 1 and number of spikes/plant in hybrid 2. These result agreement with those obtained by Khattab et al. (2010), Abd El-Rahman (2013), Amin (2013) and Elmassrya and El- Nahas (2018). Also, Zaazaa et al. (2012) found that the additive gene effects were not significant except NGS and GYP in hybrid III and NGS in hybrid I. The value dominance gene effects (h) were found to be significant (*) and highly significant (**) for most studied traits except for NKS in hybrid 2. While it was negative and highly significant for NSP and NKS in cross1. The additive gene effects (d) less than dominance effects (h). These results are in harmony with those obtained by Khattab et al. (2010) and Zaazaa et al. (2012) for PH, NSP, NKS, 100 grains weight and GYP. Dominance effects were generally greater than additive, except for NKS in cross 2. These results are in accordance with those by Khaled (2013). The values additive \times additive (aa) type of gene effects was positive and highly significant for PH and 100-grain weight in two hybrids, and DH and GYP in cross 1. While it was negative and highly significant were by obtained by NKS in two crosses and not significant for days to DH and GYP in hybrid 2. So, early generations selection for these characters might be effective for weight breeding program where it was found that the inter action between the additional alleles is of very importance, especially in autologous crops such as wheat, unlike the hybrid crops in which the interaction between the dominant alleles is of great importance because it depends on the strength of the hybrid. This is a very crop. These accordance with those by Amin (2013). On the other hand, Akhtar and Chowedry (2006) revealed that the negative $(d \times d)$ type of gene effects, were recorder for PH and GYP. The parameters additive x dominance (ad) was Significant and positive highly significant for DH, PH in hybrid 1and NKS in cross 2 and 100grain weight in two crosses, while it was negative and highly significant were by obtained by GYP in two crosses and PH and NKS in cross 2. The additive \times dominance gene effects was not significant for DH in cross 2 and NSP in two crosses. Similar results have been reported by Abd El- Rahman and Hammad (2009) for number of kerlines per spike and kernel weight. The dominance × dominance (dd) gene significant (*) or interactions was significant (**) and positive in cross 1. for NSP and number of NKS in two crosses. Similarly, Fethi and Mohamed (2010) found that the parameters dominance effects and third types of epistatic 335

dominance × dominance (dd) epistasis were more important than additive effects and other epistasis components for NKS. A significant or highly significant and negative $d \times d$ (dd) gene interactions were obtained in two for plant height and 100grain weight and days to 50% heading in hybrid 1. Also, found that the dominance × dominance (dd) gene effects were not significant for DH and NKS in hybrid 1 and GYP in the two hybrids. These results agree with those reported by El-Aref et al. (2011) and Amin (2013). The type of epistasis determined was as complementary when dominance (d) and third type of epistatic effect (dd) gene effects have same sign and duplicate epistasis when the sign was different. Thus, selection in the early generations is effective when the additive effects is greater than the non-additive effect, also, the non-additive portion are greater than additive, the improvement of the characters needs intensive selection through later generations. These results are in harmony with those reported by Kattab et al. (2010), Amin (2013) and Abd El- Rady (2018).

3.2 Heterosis, inbreeding depression (%) and phenotypic (PCV) and genotypic (GCV) coefficient of variation

Heterosis, inbreeding depression (%) and phenotypic (PCV) and genotypic (GCV) coefficient of variation in two bread wheat hybrids for all traits studied are presented in Table (4). Heterosis estimates relative to MP and BP were found to be highly significantly positive for days to 50% heading, NSP and GYP in both crosses and NKS in cross 1. Heterosis above mid parent recorded for number of kernels per spike in cross 1. While it was negative it was significantly negative for 100 kernels weight in both crosses and plant height in cross 1. While it was non significantly for plant height in cross 2. These results are in harmony with obtained by Kattab et al. (2010), Zaazaa et al. (2012), Abd Alla and Hassan (2012) and Elmassry and El-Nahas (2018). Kumar *et al.* (2018) reported that significant and positive mid parent (M.P) and better parent (B.P) heterosis were observed in four hybrids for grain yield per plant. Concerning inbreeding depression value, positive and highly significant most traits in two hybrids, however. it was highly significant and negative for plant height in the two hybrids. However, these results the expected as the expression of heterosis in F_1 will be followed by considerable reduction in F2 due to homozygosis. The results are in similar with Zaazaa et al. (2012) and El massry and El-Nahas (2018) for NSP, NKS and GYP and Moussa (2010) for days to heading Kumar et al. (2018) found that significant inbreeding depression (I.D) was recorded frequently for yield and vield contributing traits. The phenotypic coefficient (PCV) of variability values were higher than (GCV) for all traits studied in the two crosses (Table 4). Results indicated that the PCV and GCV values were much close, these revealed the major proportion of the observed variation was contributed by the genetic

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factor in additive genetic variance in most values for phenotypic coefficient and genotypic coefficient of variability. Therefore, these traits were highly affected by environmental factors. The results agreed with those obtained by Zaazaa *et al.* (2012).

3.3 Genetic variance and of three parameters model

The assessment of genetic variance, additive (D)and dominance (H) gene

effects in Table (5). The results revealed that the (a) genetic variance was less than (d) for all traits in two hybrids, while the (a) genetic variance was higher than (d) for DH and GYP (g) in hybrid 1, indicated that the additive gene effects play the main role in the inheritance of these traits and using selection in early segregating generations could be effective to isolate lines characterized by high grain yield under this study. Similar results were reported by El-Aref *et al.* (2011), Amine (2013) and Abd El- Rady (2018).

Table (4): Heterosis, inbreeding depression %, phenotypic (PCV) and genotypic (GCV) coefficient of variation in two bread wheat hybrids for all characters studied traits.

Chamatan	Commun	Heterosis %		International demonstration 9/	DCV9/	CCV %	
Characters	Crosses	MP	BP	inbreeding depression %	F.C. V 70	G.C. V. 70	
Dave to 50 % handing	1	10.92 **	9.15 **	6.21**	7.31	6.90	
Days to 50 % neading	2	6.05 **	5.75**	3.03 **	7.57	6.80	
Blant hai abt	1	-17.94 **	-17.77**	-2.24 **	6.89	5.62	
riant neight	2	-2.61	-2.70	-0.64**	6.80	5.46	
Northan of miles /alout	1	3.24 **	8.94 **	1.40 **	30.49	27.35	
Number of spikes/plant	2	48.08 **	43.24 **	25,91 **	36.88	23.23	
Number of large algorithm	1	6.30**	9.80**	6.33 **	16.83	14.19	
Number of kernels/spike	2	13.71**	4.97	2.76**	16.30	11.95	
Wighet of 100 - grains	1	-6.36**	-8.32**	9.25 **	18.92	12.87	
	2	-1.87 **	-2.08 **	9.15 **	13.98	8.19	
Conin stield/alast	1	46.38**	54.54 **	27.07 **	35.34	26.52	
Grain yield/plant	2	38.63 **	21.08 **	14.19 **	27.23	21.20	

* and ** Significant and high Significant at 0.05 and 0.01 level of probabilities.

The average degree of dominance $(H/D)^{0.5}$ given in Table (5) revealed that partial dominance gene effects was presented for all traits in hybrid 1 and weight of 100 grains in cross 2. While it was, revealed that over dominance towards the BP for all traits in hybrid 2 and plant height in hybrid 1 except, number of kerlines per spike in cross 2. Khattab *et al.* (2010), Amin (2013) and El-Gammaal and Yahya (2018) found the same results. On the other hand, Abd-Allah and Mostafa (2011) found that the complete dominance was found for NSP and 100-kernel weight

in the first cross whereas, over dominance (p>1) was detected for grain yield and its attributes in the 2^{nd} cross.

3.4 Heritability in broad (Hb) and narrow (Hn) senses and genetic advance

Heritability estimates indicate that the progress from selection for plant characters is relatively easy or difficult to make in breeding program. Heritability estimates in broad and narrow-sense and genetic advance (G.S %) are presented in Table (5). Heritability values in broad sense were moderate to high for all studied traits in the two hybrids ranging from 34.34% for 100 kernels weight in cross1 to 89.04 for plant height in cross 1. Narrow sense heritability values were moderate to high in most traits except PH in cross 1, NSP in two crosses, NKS in cross 2 and 100 kernels weight in cross 1, indicating that these traits were greatly affected by additive and non-additive effects and there is appreciable amount of heritable variation. Meanwhile, the remaining traits which had low narrow sense heritability estimates, show that selection for these traits will be difficult and high environmental influence well be a problem. These results are in accordance with the findings of El -Aref *et al*. (2011), Amin (2013), Abd El -Rady (2018), El Massry and El-Nahas (2018), and El-Gammaal and Yahya (2018). El-Said, Rania (2018) reported that the heritability in broad-sense were high values detected for all the studied traits, indicated that these traits are more genetic, while narrow sense heritability were low for NSP and GYP, so the role of additive part is low.

Table (5): Genetic variance, broad (H_b) and narrow (h_n) sense heritability and expected genetic advance (G.S%) for traits in two crosses.

Chamaatana	Crosses	Genetic variance			(II/D)1/2	Heritability		G.S %	
Characters		D	Н	Е	(H/D)	Hb	Hn	Hb	Hn
Dava to heading	1	92.47	5.23	5.84	0.237	89.04	86.60	7.55	6.89
Days to heading	2	49.78	179.66	10.49	1.89	80.48	46.29	8.84	5.08
Plant height	1	9.11	91.77	13.71	3.17	72.22	11.07	15.41	2.36
F failt fielght	2	59.88	171.30	19.63	1.69	64.43	54.21	6.35	5.35
Number of nilses/alout	1	2.01	11.59	0.96	0.425	80.45	21.14	35.57	9.46
Number of pikes/plant	2	3.48	23.97	4.56	2.62	39.67	22.98	21.12	12.29
Number of kernels/enike	1	157.23	649.82	38.64	0.830	71.11	41.21	17.35	10.06
Number of kernels/spike	2	66.04	493.49	68.60	2.73	53.77	22.18	12.71	5.24
Wight of 100 grains	1	0.138	1.47	0.30	3.26	34.34	15.00	17.07	12.68
wighet of 100 – grains	2	0.470	2.08	0.378	0.475	62.14	46.17	6.96	3.04
Grain viold/plant	1	47.43	3.84	19.11	0.284	67.54	54.11	34.34	27.67
Gram yield/pialit	2	60.24	143.16	18.99	1.54	62.68	60.63	24.75	23.94

Predicted genetic advance as percent of F_2 ranged from (2.36) for PH in hybrid 1 to (27,67) for GYP in hybrid 1 and ranged from 5.08 for DH to 23.94 for GYP in cross This led to a sufficient 2. improvement in its variable characteristics and these results indicated the possibility of practicing selection in early generations and obtain high yielding genotypes. Moreover, the remaining traits, which found the low values expected genetic suggesting the advance. role of environmental factors and dominance gene action in inheritance system of these traits. These results are in accordance with the findings of Khattab *et al.* (2010), El- Aref *et al.* (2011), Amin (2013), Abd El- Rady (2018) and Elmassry *et al.* (2018), Hassan (1993) and Shafey *et al.* (1993) found that the highest genetic advance was obtained for plant height, weight of 1000 grains and GYP. On the other hand, Kuobisy (2011) reported that the genetic advance was generally low for all studied characters in the two hybrids, while, PH, and NKS in two crosses and GYP in cross 1 were

moderate of genetic advance values. These results are of great interest for wheat breeders to improve yield potential and evacuation new wheat genotypes and enhancement of Egyptian wheat germplasm.

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