Combining Ability and Gene Action for some Traits and level of Aflatoxin Contamination in Peanut

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ABSTRACT

The aim of this investigation was to study combining ability, gene action and heterobeltiosis for some traits and determination of aflatoxin contamination in peanut. A diallel cross, without reciprocals, among five parents was done in 2013. Data revealed that the mean squares of genotypes, parents and crosses were significant for all studied traits in both of F_1 and F_2 generations. The analysis of variance for combining ability showed that mean squares due to general (GCA) and specific (SCA) combining ability were generally significant for all studied traits reflecting the importance of both additive and non- additive gene effects in the inheritance of these traits. The lines A1and 623 were good combiners for 100-pod weight, shelling percentage and pod yield feddan⁻¹ in the two seasons (one ardab=75kg and one feddan=4200m²). Genotypes 10A and 2A were good combiners for number of pods plant⁻¹, pod weight plant⁻¹, number of seeds plant⁻¹ and seed weight plant⁻¹ in the second season. Regression line intersects the Wr axis below the origin in shelling percentage in F_1 and F_2 generations and pod yield feddan⁻¹ in F_2 generation, reflecting over- dominance. On the other hand, pod yield feddan⁻¹ was controlled by partial dominance. Among these gene action partial dominance could easily be exploited through conventional breeding. Positive or negative heterosis over the better parent, i.e. heterobeltiosis was detected for all studied traits. Determination of aflatoxin contamination under normal storage conditions showed that the two crosses (P_3X, P_4) and P_3X, P_5 had total aflatoxins of 10.6, 20.1ppb, respectively. Meanwhile, total aflatoxins were not detected in parents and other F_2 crosses.

Keywords: Peanut, Combining ability, Gene action, Vr–Wr graph, heterobeltiosis, Aflatoxins.

INTRODACTION

Groundnut or peanut (Arachis hypogaea L.), is an annual legume. It is one of the world's most important oilseed crops, (Dwivedi et al., 2003). Peanut ranks the 13th among the most important food crops and the 4th among the most important oilseed crops in the world (Surendranatha et al., 2011). Seeds contain 45-60% oil, 25-30% protein and 20% carbohydrate (Singh and Singh, 1991). Aflatoxin contamination is one of the most obstacles facing peanut producers for exportation to the world market (Xue et al., 2003). Combining ability analysis is considered the quickest method of understanding the genetic nature of quantitatively inherited traits, and gives essential information about the selection of parents which in turn throw better segregants. The knowledge of the type of gene action involved in the expression of yield and yield components is essential to choose an appropriate breeding strategy to isolate desirable segregants in the later generations, John and Reddy (2015).

Several investigators studied combining ability and gene action in peanut. Shabana et al. (992) in Egypt, studied yield and its contributing traits. They applied the graphical approach suggested by Hayman (1954). In Pakistan Naazar et al. (1995) and Naazar et al. (2001) reported that estimates of general combining ability were significant for 100-pod weight, pod length and shelling percentage in F₁. Meanwhile, estimates for specific combining ability were significant for 100-seed weight in F₂ generation. Sanun et al. (2005) showed that estimates of both general and specific combining ability were significant for number of pods, pods kg⁻¹ and 100-seed weight, whereas estimates of GCA were greater than SCA estimates. In Egypt, Abd El-Aal (2008) and Abd El-Aal et al. (2013) found that pod and seed traits were largely controlled by additive gene action, while pod number plant ¹ and pod weight plant⁻¹ were controlled by non-additive genetic effect. Both genetic effects were equally important for shelling percentage. Alam *et al.* (2013) reported that the analysis of combining ability suggested that both additive and non-additive gene actions were involved in genetic system. The number of pods plant⁻¹, plant height, 100-pod weight and pod yield plot⁻¹ were preponderant by additive gene action. Meanwhile, primary branches plant⁻¹ and 100-seed weight were preponderant by non- additive gene action. Vaithiyalingan (2016) observed that additive gene action was predominant for all studied traits, except harvest index and single plant yield.

Information on variation, heritability and nature of gene action controlling the various agronomic and physiological traits in crop plants is of crucial importance to breeders in elaborating a suitable breeding program for crop improvement.

The present study was undertaken to detected the magnitude of both general and specific combining ability (GCA and SCA), heritability, gene action and heterosis for pod yield and some traits in F_1 and F_2 progenies of a five parent diallel cross (excluding reciprocals) of peanut genotypes. Aflatoxin contamination rate under storage conditions was also determined.

MATERIALS AND METHODS

The present study was carried out at Ismailia Research Station, ARC, Egypt during 2013, 2014 and 2015. Five peanut genotypes out of around 600 germblasm accessions were used in this study viz; line $329(P_1)$, line $10A\ (P_2)$, line $2A\ (P_3)$, line $1A\ (P_4)$ and line $623(P_5)$. These parents were randomly chosen, representing a wide range of variability in most traits (Table 1).

Table 1. Parents used and their origin

Table 1.17	Table 1:1 arens used and then origin										
Parent	Name	Origin	Seed color								
1	Line 329	China	Purple								
2	Line 10A	Egypt	white								
3	Line 2A	Egypt	Red								
4	Line 1A	Egypt	pink								
5	Line 623	U.S.A	pink								

A diallel - mating excluding reciprocals was carried out among the five peanut genotypes in 2013season. In 2014, the parental genotypes were planted again then re-hybridized to secure more F₁ hybrid seeds and the F_2 seeds were obtained from the F_1 plants. In 2015, an experiment was conducted in open field that included five parents, 10 F₁'s and 10 F₂'s. A randomized complete block design with three replications was used. Each entry was represented by one row in parents and F₁'s and four rows in F₂'s. Seeds were planted in rows 3 m long 60 cm apart in single seeded hills spaced 20 cm apart. Cultural practices were applied as recommended. At harvest ten guarded plants were taken at random from each experimental plot in parents and F₁'s and 30plants in F₂'s. The data recorded were plant height (cm), number of branches plant⁻¹, number of pods plant⁻¹, pod weight plant⁻¹ (g), number of seeds plant⁻¹, seed weight plant⁻¹ (g), 100-pod weight (g), 100-seed weight (g), shelling percentage (%) and pod yield ardab feddan⁻¹ (one ardab of pods= 75kg and one feddan=4200m²).

Data were analyzed according to Griffing (1956), model 1, method 2. In this approach, the combining ability variances and effects were estimated. Partitioning of genetic variance was calculated according to the procedure outlined by Hayman (1954). Heterobeltiosis percentage was determined for individual cross deviation from better parents according to Bhatt (1971).

Aflatoxins were determined according to Roos *et al.* (1997) and A.O.A.C (2006) using monoclonal antibody columns for total aflatoxins (VCAM Science Technology, Water Town, MA, USA). Aflatoxin identification was preformed by a modified HPLC. AFLATEST procedure Agillent 1200 series USA. HPLC equipment with two pumps, column (18, Lichiospher 100RP-18, 5umX25cm) was used. The mobile phase consisted of water, methanol a cetonitrile (54:29:17, V/V/V), at flow rate 1ml/min. The excitation

and emission lengths for all aflatoxins were 362 and 460nm (Fluorescence detector), respectively.

RESULTS AND DISCUSSION

Analysis of variance

The analysis of variance for plant height, number of branches $p\Gamma^1$, number of pods $p\Gamma^1$, pod weight, number of seeds $p\Gamma^1$, seed weight $p\Gamma^1$, 100- pod weight, 100-seed weight, shelling percentage and pod yield feddan⁻¹ are presented in Table (2). The results reflected significant differences among genotypes mean squares for all the above mentioned traits in F₁ and F₂ generations. Moreover, mean squares due to parents as well as differences among crosses were significant for studied traits. These data suggested that the parental genotypes were mostly different in their mean performance. The analysis of combining ability revealed that variance associated with general and specific combining ability reached the level of significance for all studied traits in both F₁ and F₂ (Table 2). The significant variances due to both general and specific combining abilities reflect the importance of additive and non-additive types of gene actions. However, general combining ability effects which were extremely of high magnitude for number of branches plant⁻¹, number of pods plant⁻¹, pod weight plant⁻¹ and number of seeds plant-1 in F2 generations suggested the predominant role of additive gene action. This result supported by the over unity of GCA and SCA values, indicating that additively play a considerable role in the inheritance of these characters. Therefore, selection in the early generation could be successfully practiced to improve these traits. The importance of additive and non-additive gene action for such traits are also reported by Shabana et al. (1992), Ruraswamy et al. (2001), El-Sawy (2006) and Abd-El-Aal et al. (2013).

Table 2. Mean squares of five peanut parents and their crosses for 10 traits.

S.O.V		Plant height (cm)		No. of branches pl ⁻¹		No. of pods pl ⁻¹		Pod weight pl ⁻¹ (g)		No. of seeds pl ⁻¹	
5.0.	d.f	$\overline{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2
Rep.	2	5.45	0.42	0.016	0.08	0.3	1.8	6.3	8.1	4.7	1.5
Genotypes	14	2.87**	85.36**	2.2**	24.42*	114.8**	329.7**	995.6**	1697,5**	11.1**	1275.7**
Parents	4	10.1**	66.0**	2.4**	54.43*	10.9**	111.0**	855.7**	303.0**	22.2**	243.5**
Crosses	9	0.3**	90.1**	2.2**	3.62*	796.9**	303.9**	1168.2**	1809.6**	1.6	1327.2**
P vs crosses	1	19.1**	119.0**	1.4**	91.53*	75.2**	1436.8**	0.8	6266.6**	65.6**	4940.2**
Error	28	1.8	0.3	0.056	0.16	0.4	0.8	6.3	14.6	2.7	1.1
GCA	4	20.7**	16.3**	0.95	38.68	5.41**	123.08**	12.4**	583.5**	36.2	450.7**
SCA	10	54.3**	33.2**	0.64**	26.87**	52.4**	104.64**	459.6**	558.7**	275.8**	415.1**
GCA/SCA		0.38	0.49	1.48	1.43	0.10	1.17	0.02	1.04	0.13	1.08
S.O.V	Seed weight pl ⁻¹ (g)		100-pod w	100-pod weight (g)		100-seed weight (g)		Shelling percentage (%)		rdab feddan ⁻¹	
5.U.V	d.f	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_2	F ₁	\mathbf{F}_2
		* I	- 2	* I	- 4	* I	1. 5	r 1	- 2	1 1	- 4
Rep.	2	5.5	2.4	56.26	13.3	7.2	1.5	4.53	3.12	0.21	0.7
Rep. Genotypes	2 14	-			13.3			-			
	_	5.5	2.4	56.26	13.3	7.2	1.5	4.53	3.12	0.21	0.7
Genotypes	14	5.5 739.0**	2.4 1266.1**	56.26 1270.53**	13.3 5424.0** 1842.0**	7.2 78.6	1.5 1446.7**	4.53 186.03**	3.12 198.87**	0.21 67.57**	0.7 45.6**
Genotypes Parents	14 4	5.5 739.0** 623.0**	2.4 1266.1** 128.9**	56.26 1270.53** 385.25**	13.3 5424.0** 1842.0**	7.2 78.6 22.20	1.5 1446.7** 530.8	4.53 186.03** 162.72**	3.12 198.87** 154.05**	0.21 67.57** 56.98**	0.7 45.6** 16.8**
Genotypes Parents Crosses	14 4	5.5 739.0** 623.0** 816.4**	2.4 1266.1** 128.9** 1199.6**	56.26 1270.53** 385.25** 1642.51**	13.3 5424.0** 1842.0** 7569.3**	7.2 78.6 22.20 112.3**	1.5 1446.7** 530.8 1873.4**	4.53 186.03** 162.72** 118.82**	3.12 198.87** 154.05** 117.13**	0.21 67.57** 56.98** 53.80**	0.7 45.6** 16.8** 35.3**
Genotypes Parents Crosses P vs crosses	14 4 9 1	5.5 739.0** 623.0** 816.4** 506.0**	2.4 1266.1** 128.9** 1199.6** 6412.7	56.26 1270.53** 385.25** 1642.51** 1464.1**	13.3 5424.0** 1842.0** 7569.3** 444.9**	7.2 78.6 22.20 112.3** 1.08 11.3**	1.5 1446.7** 530.8 1873.4** 1270.9**	4.53 186.03** 162.72** 118.82** 884.23**	3.12 198.87** 154.05** 117.13** 1113.73**	0.21 67.57** 56.98** 53.80** 233.9**	0.7 45.6** 16.8** 35.3** 253.8**
Genotypes Parents Crosses P vs crosses Error	14 4 9 1 28	5.5 739.0** 623.0** 816.4** 506.0** 4.9	2.4 1266.1** 128.9** 1199.6** 6412.7 16.8	56.26 1270.53** 385.25** 1642.51** 1464.1** 19.37	13.3 5424.0** 1842.0** 7569.3** 444.9** 31.0	7.2 78.6 22.20 112.3** 1.08 11.3** 15.68**	1.5 1446.7** 530.8 1873.4** 1270.9** 5.9	4.53 186.03** 162.72** 118.82** 884.23** 5.94	3.12 198.87** 154.05** 117.13** 1113.73** 8.81	0.21 67.57** 56.98** 53.80** 233.9** 3.4	0.7 45.6** 16.8** 35.3** 253.8** 0.3

^{*,**} significant at 0.05 and 0.01 levels of probability, respectively.

Mean performance

The results of means for pod yield clearly indicated the differences among parents, F_1 's and F_2 's (Table 3). Significant differences between parents and F_1 's and parent and F_2 's were found for all traits, except for number of pods plant⁻¹ among parents, F_1 's and F_2 's, revealed the existence of genetic variability in the

materials and the possibility of estimating combining ability effects. Results indicated that parents P_1 , P_2 and P_5 and crosses ($P_2 \times P_4$), ($P_3 \times P_4$) and ($P_4 \times P_5$) showed higher mean performance in most traits in both of F_1 and F_2 generations. The crosses showed higher means in most cases compared to its parent.

Table 3. Mean performance of five peanut parents and their crosses.

Construe	Plant he	ight (cm)	No. of branches pl ⁻¹		No. of	No. of pods pl ⁻¹		Pod weight pl ⁻¹ (g)		No. of seeds pl ⁻¹	
Genotype	$\mathbf{F_1}$	\mathbf{F}_{2}	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_2	
P1	25.03	22.93	4.0	4.27	31.23	26.07	85.83	55.6	50.27	50.20	
P2	25.27	28.60	4.0	3.47	23.23	33.20	73.33	60.4	38.80	41.27	
P3	16.20	18.57	5.8	5.20	20.43	26.13	56.33	66.7	32.30	49.67	
P4	22.30	26.43	5.6	6.17	27.40	20.40	67.37	42.2	44.93	33.20	
P5	16.30	17.97	5.5	4.60	34.73	17.40	100.03	46.3	68.10	30.80	
P1XP2	32.3	30.27	4.8	5.37	22.73	47.53	62.93	96.6	42.77	83.60	
P1XP3	19.2	20.93	4.4	3.70	38.40	48.00	109.83	109.0	77.57	87.20	
P1XP4	23.1	23.63	6.0	6.07	25.67	21.00	68.80	40.1	48.97	29.60	
P1XP5	29.4	25.80	6.3	6.80	17.87	23.20	48.63	57.6	28.53	40.73	
P2XP3	13.7	28.60	5.0	4.87	26.87	46.80	65.83	100.2	42.73	85.13	
P2XP4	24.9	17.23	3.9	3.27	39.13	30.67	110.60	65.9	75.27	59.27	
P2XP5	26.5	23.53	6.6	5.73	26.07	32.80	68.07	113.7	50.47	57.73	
P3XP4	35.9	24.60	5.2	4.47	32.47	45.60	76.63	78.0	57.33	86.07	
P3XP5	29.1	18.57	5.9	5.93	31.77	34.47	79.33	59.5	60.80	47.27	
P4XP5	34.5	33.17	5.3	4.77	26.07	36.20	72.20	72.3	50.40	55.93	
L.S.D at 0.05	0.92	0.99	0.67	0.64	-	-	2.64	6.4	4.10	1.78	
Construe	Seed wei	ght pl-1 (g)	100-pod	weight (g)	100-seed	weight (g)	Shellingpe	rcentage (%)	Pod yield a	rdab feddan ⁻¹	
Genotype	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_{2}	$\mathbf{F_1}$	F ₂	$\mathbf{F_1}$	$\overline{\mathbf{F}_2}$	

Conotypo	Seed weight	ght pl-1 (g)	100-pod	weight (g)	100-seed	weight (g)	Shellingpo	ercentage (%	b) Pod yield a	rdab feddan ⁻¹
Genotype	$\overline{F_1}$	\mathbf{F}_{2}	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$
P1	48.83	39.20	269.13	213.60	97.20	77.40	56.83	70.50	14.83	12.93
P2	38.40	32.73	298.37	185.23	101.53	75.83	52.37	54.03	12.90	15.17
P3	30.83	45.50	275.80	252.03	95.30	94.97	54.63	68.13	17.53	16.67
P4	44.63	28.70	273.53	200.60	99.41	86.37	66.20	67.97	19.40	14.11
P5	69.10	33.27	277.93	215.90	101.47	107.93	69.03	71.90	24.17	19.03
P1XP2	38.17	79.90	263.27	203.33	89.27	97.23	60.73	82.67	26.73	22.20
P1XP3	80.63	88.83	293.30	233.83	103.90	101.83	73.37	81.43	18.03	18.83
P1XP4	50.77	31.13	265.10	199.77	103.70	107.10	73.73	77.47	23.53	20.83
P1XP5	27.23	42.43	301.37	245.13	95.60	109.03	56.00	73.60	15.17	15.87
P2XP3	44.77	69.43	238.57	201.87	104.77	72.93	67.97	68.80	18.67	18.53
P2XP4	78.43	55.33	276.43	205.50	104.17	93.37	70.93	84.03	26.60	24.23
P2XP5	46.60	89.20	265.97	346.47	91.40	154.43	68.47	78.43	23.97	19.57
P3XP4	58.17	50.60	233.43	170.93	101.50	58.73	75.83	64.87	20.60	16.20
P3XP5	57.07	46.17	243.10	172.53	93.87	97.63	71.90	77.67	27.27	25.87
P4XP5	52.90	59.00	288.00	222.07	104.93	105.43	73.23	81.63	25.47	24.07
L.S.D at 0.0	5 2.34	6.85	4.80	9.31	3.51	4.08	4.96	4.96	0.67	0.86

General combining ability effects

The combining ability analysis gives useful information regarding the nature and magnitude of gene action involved in the expression of quantitative traits (Dhillon, 1975) which helps in selecting appropriate breeding method for crop improvement. The estimates of GCA for five parents are presented in Table (4). High positive and significant values were recorded for p₄ and P₅ for 100-pod weight (g), shelling percentage and pod yield feddan⁻¹ in both seasons, revealing the importance of these parents as donors for favorable alleles for these agronomic traits. Also P2 and P3 had positive and significant GCA for number of pods plant⁻¹, pod weight plant⁻¹, number of pods plant⁻¹ and seed weight plant⁻¹ in second season. It could be observed that the pervious conclusion was in harmony with the mean performance of parental genotypes indicating the efficiency of phenotypic performance for detecting the potentiality of parents for inclusion in cross breeding programs. Similar results were observed by Sanun et al. (2005), El-Baz *et al.* (2006), Yadav *et al.* (2006) Vishnuvardhan *et al.* (2011) and Abd-El-Aal *et al.* (2013).

Specific combining ability effects

Specific combining ability effects can be defined as the magnitude of deviation exhibited by the parental line in the cross from its expected performance on the basis of its general combining ability (GCA) effects. A significant deviation from zero in cross would indicate specially high or low specific combining ability (SCA) according to the sign whether positive or negative. Results given in Table (5) showed the estimates of SCA for the studied characters in ten crosses in both F₁ and F₂ generations. These results indicated that the crosses (P₁xP₂, P₁xP₄ and P₂xP₅) showed significant specific combining ability effects for number of branches plant⁻¹. The crosses $(P_4xP_5, P_2xP_4, P_2xP_5, P_3xP_5 \text{ and } P_4xP_5)$ exhibited highly significant SCA positive effects for shelling percentage and pod yield feddan⁻¹. Also, both crosses (P1xP3 and P3xP4) showed the best SCA for number of pods plant⁻¹ and number of seeds plant⁻¹.

Moreover, the cross P₁xP₃ exhibited positive and highly significant SCA effects for 100-pod weight and 100-seed weight. These crosses could account for the highest

average performance of the respective traits. In such hybrids, desirable transgressive segregates would be expected in the subsequent genotypes.

Table 4. Estimates of general combining ability (gi) effects of five peanut parents for the studied traits.

Conotypo	Plant height (cm)		No. of br	No. of branches pl ⁻¹		pods pl ⁻¹	Pod weig	ght pl ⁻¹ (g)	No. of se	eds pl ⁻¹
Genotype	2014	2015	2014	2015	2014	2015	2014	2015	2014	2015
P1	0.65**	-0.67**	-0.27**	-0.88**	-0.36	-0.56**	0.51	-1.59*	-1.33**	0.92**
P2	-0.21	0.23	-0.42**	-1.49**	-1.19**	4.06**	-0.60	10.24**	-2.69**	4.74**
P3	-2.75**	-1.83**	0.11**	-1.04**	0.11	4.48**	-2.00**	7.78**	-0.67	9.99**
P4	1.93**	2.34**	0.04	-0.82**	1.22**	-3.07**	0.67*	-12.13**	2.02	-5.40**
P5	0.37**	-0.08	0.53**	-0.60**	0.23	-4.90**	1.43**	-4.30**	2.67**	-10.26**
S.E (gi)	0.57	0.61	0.11	0.18	1.26	0.90	1.12	3.95	1.10	0.48
	Seed weig	ght pl ⁻¹ (g)	100-pod	weight (g)	100-seed	weight (g)	Shellingpe	rcentage (%)	Pod yield are	dab feddan ⁻¹
Genotypes	2014	2015	2014	2015	2014	2015	2014	2015	2014	2015
P1	-1.74**	0.59	5.14**	0.25	-1.19**	-0.87	-2.71**	2.13**	-1.83**	-1.43**
P2	-3.12**	6.11**	2.24**	2.87**	-0.36	-0.92	-3.38**	-2.75**	-0.60**	0.17
P3	-0.62*	4.21**	-9.33**	-3.47**	-0.08	-7.86**	0.26	-1.75**	-0.90**	-0.13
P4	3.27**	-9.02**	-2.18**	-15.44**	2.56**	-5.53**	4.23**	0.38	1.29**	-0.01
P5	2.20**	-1.89**	4.14**	15.78**	-0.92*	15.19**	1.60**	1.98**	2.04**	1.40**
SE(gi)	0.99	1.83	1 97	2.49	2 17	2.52	1 57	3.06	0.20	0.53

Table 5. Estimates of specific combining ability for ten peanut crosses.

Table 3.1	Table 5.12 timates of specific combining ability for ten peanut crosses.												
Conotypo	Plant he	ight (cm)	No. of branches pl ⁻¹		No. of p	ods pl ⁻¹	Pod weigh	nt pl ⁻¹ (g)	No. of se	eds pl ⁻¹			
Genotype	$\overline{\mathbf{F_1}}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	\mathbf{F}_2			
P1XP2	6.98	5.50**	0.29*	1.64**	-3.99**	11.40**	-13.36**	17.01**	-4.49**	22.09**			
P1XP3	-3.65	-1.77**	-0.64**	-0.49*	10.38**	11.45**	34.94**	31.86**	28.28**	20.44**			
P1XP4	-4.43	-3.24**	0.97**	1.66**	-3.46**	-8.00**	-8.76**	-17.09**	-3.00**	-21.77**			
P1XP5	3.46	1.35**	0.81**	2.18**	-10.28**	-3.97**	-29.69**	-7.45**	-24.09**	-5.78**			
P2XP3	-8.25	-6.37**	0.07	1.29**	-0.32	5.63**	-7.95**	11.27**	-5.19**	14.56**			
P2XP4	-1.73	-4.24**	-0.92**	-0.52*	10.84**	-2.95**	34.15**	-3.12	24.66**	4.08**			
P2XP5	1.43	-0.76**	1.29**	1.72**	-1.24**	1.01**	-9.14**	36.79**	-0.79	7.40**			
P3XP4	11.81	7.45**	-0.18	0.22	2.87**	11.57**	1.58	11.37**	4.70**	25.63**			
P3XP5	6.53	6.64**	0.06	1.47**	3.16**	2.26**	3.52**	-14.96**	7.51**	-8.31**			
P4XP5	7.29	6.93**	-0.50**	0.08	-3.65**	11.55**	-6.28**	17.75**	-5.57**	15.75**			
S.E.(si-j)	0.85	0.92	0.37	0.61	1.89	1.36	3.89	5.93	3.80	1.65			
Conotypo	Seed weight pl ⁻¹ (g)		100-pod weight (g)		100-seed v	veight (g) S	Shellingpero	entage (%)	Pod yield ard	lab feddan ⁻¹			
Genotype	$\mathbf{F_1}$	\mathbf{F}_{2}	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$			
P1XP2	-8.08**	20.44**	-15.00**	-17.71**	-8.38**	3.01**	0.74	9.74**	8.17**	4.52**			
P1XP3	31.88**	31.27**	26.60**	19.13**	5.97**	14.55**	9.73**	7.51**	-0.23	1.45**			
P1XP4	-1.87	-13.20**	-8.74**	-2.97	3.13**	17.49**	6.13**	1.41	3.08**	3.34**			
P1XP5	-24.33**	-9.03**	21.20**	11.18**	-1.49	-1.30	-8.97**	-4.06**	-6.03**	-3.04**			
P2XP3	-2.60*	6.35**	-25.22**	-15.46**	6.01**	-14.30**	5.00**	-0.25	-0.83**	-0.46*			
P2XP4	27.18**	5.48**	5.50**	0.14	2.77**	3.81**	4.00**	12.86**	4.91**	5.13**			
P2XP5	-3.58**	32.22**	-11.30**	109.89**	-6.51**	44.15**	4.17**	5.66**	1.54**	-0.95**			
P3XP4	4.41**	2.64	-25.94**	-28.08**	-0.18	-23.89**	5.25**	-7.31**	-0.78**	-2.60**			
P3XP5	4.38**	-8.91**	-22.60**	-57.70**	-4.33**	-5.71**	3.96**	3.89**	5.14**	5.65**			
P4XP5	-3.67**	17.14**	15.16**	3.80	4.10**	-0.24	1.32*	5.73**	1.15**	3.74**			

3.25

3.77

2.36

Estimation of genetic component and heritability

1.83

6.13

8.62

0.99

S.E.(si-j)

The calculated values for the degree of dominance are listed in Table (6). This value reveals whether the different traits show an additive or nonadditive gene action. In descending order, the following characteristics showed degree of dominance for pod yield and its components in peanut The component of variation due to additive gene effects (D) was significant or highly significant in F1 and F2 for number of branches plant-1, shelling percentage and pod yield feddan⁻¹, indicating that the additive gene action was more important than the non-additive in controlling the inheritance of these traits. In contrast, Shabana et al. (1992) found that additive effects (D) was not significant for the number of branches plant⁻¹. This may be due to the differences in the parents used in the two researches. Genetic components due to dominant effects (H₁ and H₂) were highly significant for most studied

traits in both F₁ and F₂ generations. The magnitude of H₁ was greater than H₂ in all traits which indicated that the positive and negative alleles were not equal in proportion in the parents at any locus. It was also obvious that the magnitude of dominance (H₁) genetic component was higher than the magnitude of additive one (D) for all studied characters indicating the important role of dominance genetic variance. The h² values, over all dominance effect of heterozygous loci was positive and highly significant for number of branches plant⁻¹, number of pods plant⁻¹, pod weight plant⁻¹, number of seeds plant-1 and seed weight plant⁻¹ in F₂ generation and for shelling percentage and pod yield feddan⁻¹ in both F₁ and F₂, indicating that most of the dominant genes had positive effects. The ratio $(H_1/D)^{0.5}$ which measures the average degree of dominance was more than unity for all studied traits, indicating that over - dominance is controlling these

4.60

1.00

0.80

traits. To improve these traits, pedigree selection could be applied. Proportion of genes with asymmetry positive and negative effects as $(H_2/4H_1)$ was lower than 0.25 for all studied characters. The ratio of total number of dominance to recessive genes in all parents (KD/KR) was greater than unity for all studied characters in both F_1 and F_2 generations, indicating that dominant alleles were found in all parents for these characters. Heritability estimates in broad sense (H_b) were high for all studied traits and ranged from 50.16% for shelling percentage to 98.75% for plant height. Narrow sense heritability (h_n) were low in most characters to

moderate for pod weight plant⁻¹, seed weight plant⁻¹, shelling percentage and pod yield feddan⁻¹. The low value of narrow sense heritability are mainly due to dominance components accounted for a great portion of the genetics of these characters. Different estimates of heritability in narrow sense and in the broad sense were recorded by some researchers Shabana *et al.* (1992), Ayub-Khan *et al.* (2000), Yogendra *et al.* (2002), El-Baz *et al.* (2006), Abd-El-Aal (2008), Abd-El-Aal *et al.* (2013), Alam *et al.* (2013), John and Reddy (2015) and Vaithiyalingan (2016).

Table 6. Estimates of genetic components and their	derived parameters 1	for some peanut traits.
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Genetic	Plant he	ight (cm)	No. of bra	nches pl ⁻¹	No. of p	ods pl ⁻¹	Pod weigh	ht pl ⁻¹ (g)	No. of se	eds pl ⁻¹
parameter	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$	$\mathbf{F_1}$	$\mathbf{F_2}$
D±S.E	20.22	20.21	0.797**	0.76**	33.04	33.34	284.38	280.37	183.87	185.47
F±S.E	34.84	34.80	0.792	0.65	87.02	88.19	693.35	677.31	406.38	412.78
$H_1\pm S.E$	212.94**	214.46	2.718**	3.28**	234.36**	238.97	2019.22**	2068.85	1181.59**	1193.07
$H_2\pm S.E$	180.59**	180.59**	2.119**	2.12**	171.72**	171.72**	1592.77**	1592.77**	934.22**	934.22**
h^2	87.04	120.38	0.345	93.07**	3.95	1468.01**	-0.34	6354.53**	110.36	5053.97**
E±S.E	0.10	0.12	0.018	0.05	0.55	0.26	0.88	4.89	1.98	0.38
(H1/D)0.5	3.25	1.63	1.847	1.04	2.66	1.34	2.66	1.36	2.53	1.27
H2/4H1	0.21	0.21	0.195	0.16	0.18	0.18	0.20	0.19	0.20	0.20
KD/KR	1.72	3.24	1.736	2.41	2.96	166.61	2.69	17.07	2.55	15.32
Hn	52.83	48.54	35.53	52.64	70.6	67.66	58.9	55.8	67.9	60.5
Hb	98.35	98.75	97.90	80.29	80.7	82.9	80.5	88.60	70.8	67.7
Genetic	Seed wei	ght pl ⁻¹ (g)	100-pod v	veight (g)	100-seed v	veight (g)	Shelling perce	entage (%)	Pod yield ard	lab feddan ⁻¹
parameter	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2	$\mathbf{F_1}$	\mathbf{F}_2
D±S.E	206.97	202.07	125.39	118.09	5.86	5.42	53.43**	51.31*	18.94*	18.90**
F±S.E	476.39	456.79	313.99	284.82	18.46	16.69	47.79	39.29	20.03	19.90
$H_1\pm S.E$	1406.28**	1461.82	2093.25**	2202.84	131.11**	155.94**	204.60**	235.56**	89.36**	90.42**
$H_2\pm S.E$	1089.38**	1089.38**	1626.90**	1626.90**	100.55*	100.55*	166.04**	166.04**	78.21**	78.21**
h^2	129.09	6495.06**	372.87	323.42	-0.71	1276.05	225.84**	1102.88**	59.85**	258.77**
E±S.E	0.69	5.59	3.03	10.32	1.54	1.98	0.81	2.94	0.06	0.09
(H1/D)0.5	1.46	1.34	4.09	2.16	4.73	2.68	1.96	1.07	2.17	1.09
H2/4H1	0.14	0.19	0.19	0.18	0.19	0.16	0.20	0.18	0.22	0.22
KD/KR	2.31	11.54	1.88	3.53	2.00	3.69	1.59	2.11	1.64	2.86
KD/KR Hn	2.31 52.84	11.54 53.64	1.88 58.5	3.53 60.5	2.00 48.02	3.69 44.67	1.59 34.84	2.11 48.84	1.64 50.8	2.86 56.7

86.76

84.6

Graphical (wr/vr) analysis.

58.59

Hb

Graphical presentation (Vr,Wr) of different traits in both generations are given in Figures 1 and 2. The regression coefficient significantly differed from zero but not from unity for F_1 and in F_2 , indicating that the genetic system could be deduced to be additive without the complication of non-allelic interaction. For the other cases, regression slope differed from unity, indicating that a complementary type of epistasis was involved.

56.29

95.5

90.8

The regression line intersected the Wr below the point of origin in shelling percentage in both generations and pod yield faddan⁻¹ in the F_2 , revealed the presence of over - dominance. Meanwhile, it intersects the Wr axis above the origin in pods yield in ardab faddan⁻¹ in the F_1 reflecting partial dominance. However, the regression line intersected the Wr below the point of origin in the remaining cases, indicating an over - dominance in the inheritance of these cases.

This contradiction between the two types of analysis might be an expected result of the presence of complementary type of non-allelic interaction which inflated the ratios of H_1 to D and distorted the Vr,Wr (Hayman, 1954 and Mather and Jinks, 1982). However, the regression line intersected the Wr below the point of origin in the remaining cases, indicating an overdominance in the inheritance of these cases. The array

points scattered along the regression line for these traits in both generations indicating genetic diversity among the parents. The low magnitude of correlation coefficient between parental mean (Yr) and the (Wr+Vr) might be due to a presence of non-allelic interaction in some parental line.

50.16

80.5

85.9

86.48

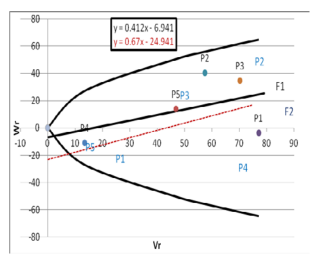


Fig 1. Wr/Vr graph for shelling percentage -1 in F_1 and F_2 generations.

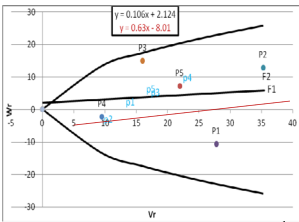


Fig 2. Wr/Vr graph for pods yield in ardab fad⁻¹ in F_1 and F_2 generations.

The parental lines P_1 and P_2 for shelling percentage trait in the F_1 and the F_2 included the largest number of recessive genes. On the other hand, P_2 for pod yield faddan-¹ in the F_1 had the highest number of recessive genes. The P_4 and P_5 were high for shelling percentage in the F_1 and the F_2 generations and P_2 , P_4 in the F_1 , F_2 for pod yield faddan-¹ i.e, they contained greater number of dominant allels for those cases.

Heterobeltiosis

Physical manifestation of the beneficial effects of hybridization between diverse parents is usually termed as heterosis and is referred as heterobeltiosis and relative heterosis based on F₁ superiority over better parent and/or mid - parental value, respectively. In plant breeding programmes, useful heterosis is referred to

denote the expression of increased vigor of a hybrid over its better parent. Heterosis is a complex biological phenomenon often manifested in the superiority of a hybrid over parental forms according to the rate of development of one or more complex characters (Konarev, 1974). Estimates of heterotic effects for the F₁ crosses are shown in Table (7). Significantly positive heterobeltiosis effects relative to better parent values may be considered favorable for most traits under investigation. Highly significant negative (desirable) heterotic effects relative to the best parent were noticed for plant height in crosses (P₁xP₃, P₁xP₄, P₂xP₃ and P₂xP₄). Significant or highly significant positive heterotic effects were found for number of branches plant⁻¹ in the four crosses $(P_1xP_2, P_1xP_4, P_1xP_5)$ and P_2xP_5 and number of pods plant⁻¹ and number of seeds plant⁻¹ in four crosses $(P_1xP_3, P_2xP_3, P_2xP_4)$ and P_3xP_4 , pod weight plant⁻¹ in two crosses (P₁xP₃ and P₂xP₄). Highly significant positive heterobeltiosis was recorded for 100-pod weight in two crosses $(P_1 x P_3 \text{ and } P_1 x P_5)$. Highly significantly positive heterotic effects were found for seed weight plant⁻¹ in the $(P_1 x P_3, P_1 x P_4,$ P₂xP₃, P₂xP₄ and P₃xP₅) crosses, 100-seed weight in the $(P_1xP_3$ and $P_1xP_4)$. All crosses except (P_1xP_5) and P₂xP₅) revealed significant and highly significant positive heterobeltiosis for shelling percentage and pod yield feddan⁻¹. These results for most cases are in harmony with that reached by El-Sawy (2006), El-Baz et al. (2006), Abd-El -Aal (2008), John et al. (2012) and Abd- El-Aal et al. (2013).

Table 7. Heterobeltiosis % of the studied traits of peanut F_1 crosses.

Character crosses	Plant height (cm)	No. of branches Pl ⁻¹	No. of pods pl ⁻¹	Pod weight pl ⁻¹ (gm)	No. of seeds pl ⁻¹	Seed weight pl ⁻¹ (gm)	•	100- seed weight (gm)	Shelling %	Pod yield Ardab Feddan ⁻¹
P1XP2	28.0**	19.8**	-27.2**	-26.6**	-14.9**	-21.8**	-11.7**	-12.0**	6.86**	80.22**
P1XP3	-23.4**	-23.6**	22.95**	27.9**	54.31**	65.1**	6.35*	6.89**	29.09**	2.85**
P1XP4	-7.9**	5.9**	-17.8**	-19.8**	-2.59	4.0**	-3.08	4.32*	11.37**	21.31**
P1XP5	17.4**	15.2**	-48.5**	-51.3**	-58.1**	-60.6**	8.43**	-5.78**	-18.88**	-37.24**
P2XP3	-45.8**	-13.8**	15.64**	-10.2**	10.14**	16.6**	-20.0**	3.18	24.40**	6.46**
P2XP4	-1.5**	-30.2**	42.82**	50.8**	67.51**	75.7**	-7.35**	2.59	7.15*	37.11**
P2XP5	4.9**	21.3**	-24.9**	-31.9**	-25.8**	-32.6**	-10.8**	-9.98**	-0.82	-0.83*
P3XP4	61.0**	-10.3**	18.49**	13.76**	27.60**	30.3**	-15.3**	2.10	14.55**	6.19**
P3XP5	78.3**	2.3	-8.54**	-20.6**	-10.7**	-17.4**	-12.5**	-7.49**	4.15**	12.83**
P4XP5	54.7**	-5.9**	-24.9**	-27.8**	-25.9**	-23.4**	3.62	3.42	6.08**	5.38**
L.S.D at 0.05	1.22	5.5	2.71	3.51	5.53	3.11	6.37	4.55	3.38	0.89

Determination of aflatoxins

Results in Table (8) showed that the two crosses (P_3X P4 and P_3X P) had a total aflatoxins 10.6, 20.1ppb, respectively. Meanwhile, total aflatoxins were not detected in all other parents and F_2 crosses. These results are in harmony with those found by Mahmoud *et al.* (2006) who found no cultivar completely resistant to aflatoxin contamination production and invasion with aflatoxigenic fungi while, there was a significant difference in genotype ability to allow invasion and aflatoxin production. The variable amount of aflatoxin in contaminate peanut genotypes and may be due to the environmental factors, nature of the fungal strains (Anderson *et al.*, 1995). Furthermore, the resistance of

peanut seeds to A. flavus and/or A. parasiticus invasion might be due to genetic and/or biochemical composition of the seed or appears to be associated with certain structural and biochemical characters of the pod and seed and there is a possibility that genotypes may have differential effects up on the population of aflatoxigenic fungi in geocar posphere (Holbrook et al., 2000). Also, Liang et al., (2009) concluded that the resistance has been associated with testa wax and presence of cutin layer, active oxygen and membrane lipid peroxidation, phytaolexin accumulations and antifungal proteins in the peanut seeds. Sharaf et al., (2011) concluded that B-1-3 glucanases enzyme has a role in the defense of peanut against the infection by A. flavus and the

resistant peanut mutants for *A. flavus* were identified by analyzing B-1-3 glucanases activities using polyacrylamide gel electrophoresis (PAGE). They found that these mutants have the ability to reduce the aflatoxins accumulation and RAPD-PCR showed pattern can be used as marker assisted selection (MAS) for the resistance of the fungus.

Table 8. Aflatoxin contamination of some peanut genotypes under field conditions.

	genery pe			/X	
Genotype	A	flatoxin	contami	nation p	pb
Genotype	\mathbf{B}_{1}	\mathbf{B}_{2}	G_1	G_2	Total
$\overline{P_1}$	ND	ND	ND	ND	ND
\mathbf{P}_2	ND	ND	ND	ND	ND
P_3	ND	ND	ND	ND	ND
\mathbf{P}_4	ND	ND	ND	ND	ND
P_5	ND	ND	ND	ND	ND
$P_1 \times P_2$	ND	ND	ND	ND	ND
$P_1 \times P_3$	ND	ND	ND	ND	ND
$P_1 \times P_4$	ND	ND	ND	ND	ND
$P_1 \times P_5$	ND	ND	ND	ND	ND
$P_2 \times P_3$	ND	ND	ND	ND	ND
$P_2 \times P_4$	ND	ND	ND	ND	ND
$P_2 \times P_5$	ND	ND	ND	ND	ND
$P_3 \times P_4$	5.8	1.3	2.6	0.9	10.6
$P_3 \times P_5$	11.5	2.8	4.2	1.6	20.1
$P_4 \times P_5$	ND	ND	ND	ND	ND

ND = Not detected

CONCLUSION

In light of the present findings it is evident that both additive and non-additive gene effects were important. Parental lines A1 and 623 were good combiners for 100-pod weight, shelling percentage and pod yield feddan⁻¹ in both seasons revealing the importance of these parents as donors for favorable alleles for these traits. Five crosses (P₄xP₅, P₂xP₄, P₂xP₅, P₃xP₅ and P₄xP₅) showed significant and desirable SCA effects and heterobeltiosis for shelling percentage and pod yield feddan⁻¹. Meanwhile, total aflatoxins were not detected in all other parents and F₂ crosses. These results seem to be useful for peanut breeding programs in making a proper decision when initiating a crossing plan.

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القدرة علي التألف والفعل الجيني لبعض الصفات ومدي التلوث بالأفلاتوكسين في الفول السوداني رحاب حمدان عبد الكريم عبد الرحمن'، خالد مصطفي المليجي' و وفاء وهبه محمد شافعي'' فسم المحاصيل الزيتية _ معهد المحاصيل الحقلية _ مركز البحوث الزراعية _ مصر ' المركز الاقيليمي للاغذية والاعلاف '' المركز الاقيليمي للتصميم والاحصاء ـ مركز البحوث الزراعية _ مصر

يهدف هذا البحث الي در اسة القدرة علي الائتلاف و تحديد الفعل الجيني لبعض الصفات وقوة الهجين للأب الأفضل و تقدير مستوي الأفلاتوكسين في بذور الفول السوداني المخزنه تحت الظروف العادية. وقد تم التهجين بين خمسه أباء هي سلالة ٢٢٩، سلالة أ١٠ سلالة أ١٠ سلالة أ١ مسلالة أ١ وسلالة ٢٢٣ متباينة في صفاتها باستخدام نظام الهجن الدائرية ما عدا الهجن العكسية، وقد تمت الزراعة خلال ثلاثة مواسم صيفية هي ٢٠١٠ و ٢٠١٥ متباينة في صفاتها باستخدام نظام الهجن الدائرية ما عدا النهجن العكسية، وقد تمت الزراعة خلال ثلاثة مواسم صيفية هي ٢٠١٥ و ٢٠١٥ بمحطة البحوث الزراعية بالاسماعيلية. وقد أظهرت النتائج تباينا معنويا لكل الصفات تحت الدراسة في كل من الجيل الأول والثاني، كما كان تحليل التباين القرة العامة والقرة الخاصة علي التألف معنويا لكل الصفات المدروسة مشيرة الي أهميه كلا من الفعل الجيني المضيف وغير المضيف في وراثة الصفات. كما كان التركيبان الوراثيان 623 مماء خامة علي التألف لصفات عدد القرون/النبات ووزن القرون بالاردب الفدان في كلا الموسمين، و كان التركيبان أ 7وأ ١٠ ذات قدرة عامة علي الصفات المدروسة قيم عاليه لكفاءة التوريث بمعناها العام في كلا الجيلين، واظهرت صفات ال ١٠٠٠ جزرة ونسبة التصافي قيما منخفضة لكفاءة التوريث بمعناها الضيق كما أظهرت باقي الصفات قيما متوسطة. أظهرت قوة الهجين قيما معنوية سلبا وايجابا عن الأب الأب الأعلي مدي التلوث بالأفلاتوكسين فقط أظهرا قابلية التلوث بالأفلاتوكسين مدي التلوث بالأفلاتوكسين ليذور الجيل الثاني والأباء تحت ظروف التخزين العادي أن هجينين فقط أظهرا قابلية للتلوث بالأفلاتوكسين.