ESTIMATION OF GENETIC COMPONENTS, HERTABILITY AND CLUSTER ANALYSIS FOR SOME AGRONOMIC CHARACTERS IN RICE.

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# ABSTRACT

Nine parents of rice were crossed to obtain 36  $F_1$  hybrids, through half diallel cross. These parents included one exotic variety, four commercial varieties and four promising lines. Hayman's 1954 method was employed for this purpose. The nine parents and their  $F_1$ s were grown in randomized complete block design with three replications at the Rice Research and Training Center Farm, Sakha, Kafr El-Sheikh, Egypt. Results revealed highly significant mean squares for all characters, suggesting the presence of genetic differences among genotypes for all studied traits. Significant or highly significant and positive of additive component (D) for all characters except for panicle weight and number of panicles plant<sup>-1</sup> were found. The dominance components  $H_1$  and  $H_2$  were highly significant for all characters except spikelet fertility% and grain yield plant<sup>-1</sup>. Heritability estimates in broad sense were high for all studied characters. The nine parents were distinctly classified into five clusters (below 25 Euclidean distance) according to the similarity.

## INTRODUCTION

Rice is one of most important grains in the world. It is not only a stable food, but also contributes major economic activity and a key source of income and employment for the rural population. Also, rice is considered the main food for all human ages because its grains are rich in protein, minerals, vitamins and fiber (Alam *et al*, 1998). The total rice production in Egypt reached 6.6 million tons with a national average of 10.00 ton/ha. This average ranked the first among the rice producing countries in the world. Rice production should be increased by about 60% by the year 2025 to feed the additional rice consumers (Duwayri *et al*, 1999)

### MATERIALS AND METHODS

These studies were conducted at Rice research and training center (RRTC), Sakha, Kafr El-Sheikh, Egypt during the Tow successive rice seasons 2010 and 2011 to estimate the genetic components, heritability and cluster analysis for agronomic characters in rice. The experimental materials were one exotic variety (HR5824-B-3-2-3), four commercial varieties (Giza 177, Sakha 101, Sakha 104 and Giza 171) and four promising lines (GZ 6910-28-1-3-1, GZ 7685-8-1-3-2, GZ 6522-15-1-1-3 and GZ 6903-1-2-2-1). A half diallel cross was conducted among the nine parents in 2010 season to produce 36 crosses in 2011 season. The parental varietes and the resulting 36 crosses were evaluated in a Randomized Complete Block Design (RCBD)

experiment with three replications. Each replication contained 25 individual plants with spaced at 20 cm among rows and 20 cm among plants. All agronomic practices such as land preparation, fertilizer application, weeding and pest control were done as recommended with rice crop during both seasons of the study. The nine parents were evaluated for following agromonic traits; days to maturing, plant height (cm), chlorophyll content, number of panicles plant<sup>-1</sup>, panicle length (cm), panicle weight (g), 1000-grain weight (g), spikelet fertility (%) and grain yield plant<sup>-1</sup>.

### Statistical analysis:

The genetic components of variation were calculated, by using Hayman's procedure (1954) and presented by Singh and Chaudhary (1977): D = Components of variation due to additive gene effects.

**F** = Mean of covariance of additive and dominance effects over the arrays

 $H_1$  = Component of variation due to dominant gene effects.

 $H_2$  = Dominance indicated asymmetry of positive and negative effects;

Where :

 $H_2 = H_1 \{1 - (U - V)^2\}$ 

**U** = Proportion of positive genes in the parents.

V = Proportion of negative genes in the parent { or of the genes with positive (negative) effects} and where **U+V** = 1.

 $h^2$  = Dominance effects (as the algebraic sum over all loci in heterozygous phase in all crosses).

**E** = Component of variation due to environmental effects.

The estimate of three components of genetic variation were determined using the following equations :

D = Volo - E

 $F = 2Volo - 4W_{olol} - 2(n-2) E/n,$ 

 $H_1 = Volo - 4W_{olol} + 4V_{ili} - (3n-2) E/n,$ 

$$H_2 = 4 V_{iii} - 4V_{oll} - 2E,$$

 $h^2 = 4(M_{Ii}-M_{Io})^2 - 4(n-1) E/n^2$ .

Where:

**E** = Environmental effect = 
$$\left[\frac{Error SS + \text{Re } p SS}{Error df + \text{Re } p df}\right] / No.of replications$$

**V**<sub>oLo</sub> = Variance among parents.

**Vr** = The variance of one array (r<sup>th</sup> array).

**V**<sub>iLi</sub> = Mean variance of the arrays.

Wr = The covariance between the parents and their off-spring in one array ( $r^{th}$  array).

**W**<sub>oLol</sub> = Mean covariance between the parents and arrays.

**V**<sub>oLI</sub> = Variance of the mean of arrays.

 $(\mathbf{M}_{Li}-\mathbf{M}_{Lo})$  = difference between mean of the parents and mean of their n<sup>2</sup> progeny.

Test of significance among parameter estimates were made using the standard error estimation technique suggested by Hayman (1954). The above estimates were then used in computing other parameters following Hayman (1958):

 $(H_1/D)^{1/2}$  = The mean degree of dominance at each locus.  $H_2/4H_1$ = The ratio of genes with positive and negative effects in the parents.  $K_D/K_R$  = {(4DH<sub>1</sub>)<sup>1/2</sup> + F}/{(4DH<sub>1</sub>)<sup>1/2</sup> - F}

= The ratio of dominance and recessive genes in the parents. **The estimates of heritability:** 

The heritability estimates were computed following the formula of Mathers and Jinks (1982).

### 1- Broad sense heritability (Hbs) :

$$\mathbf{H}_{bs} = \frac{1/2D + 1/2H1 - 1/4H2 - 1/2F}{1/2D + 1/2H1 - 1/4H2 - 1/2F + E}$$

2- Narrow sense heritability (Hns) :

$$\mathbf{H}_{ns} = \frac{1/2D + 1/2H1 - 1/2H2 - 1/2F}{1/2D + 1/2H1 - 1/4H2 - 1/2F + E}$$

## **RESULTS AND DISCCUSION**

### Mean performance:

Mean performance of the nine parents and their  $F_1$  of the diallel cross for the nine studied characters are presented in Table (1).

As revealed in the Table (1), the mean performance for the a forenamed characters, varied from cross to cross combination. For No. of days to maturity, the F<sub>1</sub> mean values of most of crosses (27 crosses) were intermediate between the two parents involving indicating partial or no dominance effect. It was noticed that four of the  $F_1$  mean values were in direction of earlier parents. This in turn suggested that earliness was dominant over lateness for these crosses. On the other hand, five crosses were towards the lateness. Complete to over dominance was observed in most of the crosses towards taller parents for plant height (21 crosses), put only two crosses exhibited dominance effect towards the shorter parents. However, plant height character of the rest 13 crosses was show intermediate between the two parents involved. Regarding the chlorophyll content character, different results were obtained. The F1 mean values of 20 crosses were towards higher parents, while three only were tended towards lower parents. However, 11 crosses were intermediate between the two parents involved.

Regarding the panicle length results in Table (10), clearly showed that out of 36 crosses, 23 have their mean values intermediate to the their respective parents. However, the  $F_1$  means of nine different crosses tended towards higher parents, while four crosses tended to the lower parents. For the spikelet fertility % (Table 10), results indicated that the three parents, Gz6522-15-1-1-3 (96%), Giza 177 (94%) and Gz6910-28-1-3-1 (92%) and the crosses, Giza 177 x Gz7685-8-1-3-2 (96%), HR5824-B-3-2-3 x Gz6522-15-1-1-3 (96%) and Gz7685-8-1-3-2 x Gz6903-1-2-2-1 (96%) showed the highest mean values in this respect.

								1000-	
Genotypes	DM	PH	CC	No.	PL	PW	SF	GW (g)	GYP
51		(cm)		pp	(cm)	(g)	(%)	(0)	(g)
\ Giza 177	170	1	55.77	19	۲۰	۳.٧٠	94	۲۸.۰۰	24.12
Y Sakha 101	151	٩٢	٤٨٩٧	20	۲٤	٤١٣	89	۲۷ ۳۰	71.17
Sakha 104     Sakha 10	100	۱.۸	٤٦,٦٣	۲٤	۲۲	۳.9٣	91	11.11	77.07
٤ Giza 171	101	137	50.45	۲۸	٣٣	٤١٣	90	۲٥	09.1.
• HR 5824-B-3-2-3	١٠٩	۸۳	TV 2T	۲۲	۲.	511	90	Y0 0Y	75.55
1 GZ 6910-28-1-3-	177	1.0	٤.٦.	۲٦	20	۳٩.		19 TV	٦
1							92		
Y GZ 7685-8-1-3-2	11.	99	٤. ٨.	۲.	۲.	r. 2 V	86	T V	01.95
^ GZ 6522-15-1-1-	170	۱.٤	50 1.	۲۱	22	٣.٧٠		77,77	75.9
3							96		
9 GZ 6903-1-2-2-1	171	٩٩	٤٢.٥٧	۲۸	۲٤	٤	86	۲٦_٦٧	75.07
F₁ hybrid									
1 X 2	139	117	٤٢.٤٧	۲۸	۲٤	٥.٩٠	93	۲٩ ٦٧	VA_VV
1 X 3	172	111	٤٢.٦٣	۲٤	۲۲	٤٢٠	93	14.14	٧٥.٤٧
1 X 4	147	185	٤١.٢٣	۳۷	20	٤٢٠	91	۲٦.٤٠	٦٦.٠٣
1 X 5	12.	1.7	źź VV	۲٩	۲۱	۳.۰۰	94	Y0. AV	07.17
1 X 6	150	۱.۷	٤٦ ٩٣	37	۲۳	0.77	94	29.01	09
1 X 7	11.	111	٤٢.0.	۳۸	۲۱	٤٣٧	96	۲۹ ۷۰	٦٥
1 X 8	122	1.0	٤٣.٥٣	źź	21	٤٤٠	93	19.1V	٧٠٩٠
1 X 9	177	90	٤٤٩٠	۳۷	۲ ٤	٦.٠٠	94	۳۰.٤٣	۷۱_٤٧
2 X 3	15.	۱.۷	٤٦.٢٣	۲۸	۲۲	0	88	۲۸ ۲۷	11.95
2 X 4	157	177	50.77	۳.	20	0.17	89	٢٤.٤٧	٨٤ ٧٠
2 X 5	15.	111	٤٢.٤٣	22	۲۳	٤٤٣	87	۲۹_۹۷	٦٣.٤٠
2 X 6	15.	1.1	٤٦.٦٣	٣٣	۲ ٤	٤٧٠	90	19.00	٥٤.٠٣
2 X 7	12.	1.4	٤٢.٨٠	۳۸	20	0.77	93	14.14	V9.1V
2 X 8	121	111	٤٨ ٨٠	39	20	٤.0.	93	11.07	11.01
2 X 9	182	111	٤٣.٤٧	٣٤	22	0.77	92	۲۸.۳۰	۷۱.۳۰
3 X 4	151	15.	٤0.1٣	٤.	22	٦.٠٠	94	11.11	۷۲.٤٣
3 X 5	120	1.5	٤٤.٢٠	۳۷	21	۳.٦٣	90	19 AV	٦٢_٦٣
3 X 6	121	1.0	٤٠.١٣	۳۸	22	۳٩٠	89	۳۱.۳۰	۳۵.۸۲
3 X 7	122	1.1	٤٣.٢٣	۳۷	۲٤	0.1.	94	14.11	V0.VV
3 X 8	172	1.9	٤٨ ٧٧	źź	۲۳	٤٧٠	92	۳۰ <u>۲</u> ۰	۸۱.٤۰
3 X 9	141	111	٤٣.٥٠	39	۲۳	٤.٢٣	87	11 <u>.</u> VV	۸۳.۸۷
4 X 5	121	11.	۳۹.۱۳	51	77	۳ <u>.</u> ۱۷	84	۲٤.٨٠	11.9.
4 X 6	157	129	٤٣.٧٧	٣ź	۲ź	0.27	92	۲٦.٤٠	V. 17
4 X 7	172	177	٤٤.٧٠	٤٢	17	٤.٩٠	95	15.15	15.01
4 X 8	140	189	20.77	٤.	10	٤٩٠	93	17	11.11
4 X 9	12.	1171	22	77	10	0.0.	92	11.1.	14.55
5 X 6	17.	47	2.47	77	11	r.rv	90	۲۷.۰۰	22.55
5 X 7	17.		21.57	۲.	11	۲.۱۰	90	14.54	0.9.
5 X 8	114	1.1	25.4.	11	11	2.5.	96	17.1	27 27
5 X 9	112	1.0	21 11	14	11	2.17	94	14.1.	2/11
6X/	110	1.0	21	۱۰ ۳۸	11	2. 11	91	1.2.	v1_z•
6 X 8	110	1.1	24.17	14	10	2.0.	90	11.11	v v . • •
6 X 9	114	1.0	27.15	1 V	10	2 71	90	11.1.	A • . Y •
/ X 8 7 X 8	11.	1.7	20,11		12	2.1.	8/	11.01	V 1 21
( 7 8	111	1 • 1	20,71	1 1	12	2,71	96	17.2.	17.1.
878	111	111	z V 1 •	17	١z	z.01	94	11.11	11.17

Table 1 : Mean performance of parental varieties and their  $F_1$  crosses for all characters under study.

Abbreviations: DM days to maturing; PH (cm), plant height; CC, chlorophyll content; No. PP, No-of panicles per plant; PL(cm), Panicle length; PW(g), Panicle weight; SF%, spikelet fertility%; 1000-GW(g), 1000-grain weight and GYP(g) grain yield per plant.

Regarding panicle weight results showed that, the three parents Sakha 101, Giza 171 and Gz6903-1-2-2-1 gave the highest mean values

(more than 4 g). The crosses, Giza 177 x Gz6903-1-2-2-1, Sakha 104 x Giza 171, Giza 177 x Sakha 101, Giza 177 x Gz6903-1-2-2-1, and Giza 171 x Gz6903-1-2-2-1 gave highest mean values of more than (5.50 g).

Concerning 1000-grain weight, the parents Gz7685-8-1-3-2, Gz6910-28-1-3-1 and Giza 177 showed the results clearly indicated that highest mean values of 30.07, 29.27 and 28.00 g, respectively, while the crosses, Sakha 104 x Gz6910-28-1-3-1, Giza 177 x Gz6903-1-2-2-1, Gz6910-28-1-3-1 x Gz7685-8-1-3-2 and Sakha 104 x Gz6522-15-1-1-3 gave heaviest weight of 1000-grain and ranged from 30.20 to 31.30 g and were superior in this respect. The highest grain yield per plant was obtained from the crosses, Sakha 101 x Giza 171 (84.70 g), Sakha 104 x Gz6903-1-2-2-1 (83.87 g), Sakha 101 x Sakha 104 (82.93g), Sakha 101 x Gz6522-15-1-1-3 (82.57 g), Sakha 104 x Gz6522-15-1-1-3 (81.40 g) and Gz6910-28-1-3-1 x Gz6903-1-2-2-1 (80.70 g). However, the lowest were estimated for the crosses, HR5824-B-3-2-3 x Gz6910-28-1-3-1, HR5824-B-3-2-3 x Gz6903-1-2-2-1 and HR5824-B-3-2-3 x Gz6522-15-1-1-3 with values of 44.33, 48.13 and 49.47 g per plant. The parental lines Sakha 101 and Sakha 104 manifested highest mean performance of 72.16 and 66.57 g, respectively, for the grain yield. Analysis of variances:

The analysis of variance for the studied characters calculated and the results presented in Table (2) showed that the genotypes mean square were found to be larger than their corresponding mean square for the error. Moreover, the results of F-test values. These findings indicated the presence of large variations among them. Similarly, the results indicated that the mean square of parents and crosses showed highly significant for all characters. These findings indicated overall wide differences among these populations. Parents Vs crosses mean squares indicated that the average heterosis could be highly significant in all crosses for all agronomic characters except days to maturing and panicle length.

s.o.v	d.f	Days to maturing (days)	Plant height	Chlorophyll content (SPAD)	No. of panicles /plant	Panicle length	
Replication	2	1.49	14.29	4.51	11.65	0.19	
Genotypes	44	241.77**	504.07**	24.43**	138.85**	14.41**	
parents	8	473.73**	626.04**	37.73**	35.01*	48.35**	
crosses	35	195.65**	441.19**	17.19**	87.02**	6.97**	
P vs F <sub>1</sub>	1	0.03	1729.35**	171.25**	2783.47**	3.42	
Error	88	1.40	12.58	1.73	14.68	1.00	

 Table 2 : The estimates of mean square of ordinary analysis for all characters under study.

\*, \*\* significant at 0.05 and 0.01 level of probability, respectively.

\$.0.V	d.f	Panicle weight	Spikelet fertility (%)	1000-grian weight	Grain yield/plant
Replication	2	0.02	821.34	0.52	120.25
Genotypes	44	1.77**	3612.00**	9.42**	400.04**
parents	8	0.31**	3961.68**	7.93**	386.83**
crosses	35	1.69**	2794.21**	9.58**	352.27**
P vs F <sub>1</sub>	1	16.19**	29437.35**	15.64**	2177.39**
Error	88	0.02	268.54	0.18	51.21

Table 2 : Continued.

\*, \*\* significant at 0.05 and 0.01 level of probability, respectively.

### Estimates of genetic components and heritability:

Table (3) and (4) present the genetic components and hereitability of nine agronomic characters under investigation. Significant or highly significant and positive values of additive component (D) for all characters except panicle weight and number of panicles plant<sup>-1</sup>. Moreover, the dominance component H<sub>1</sub> and H<sub>2</sub> were highly significant for all characters studied and ranged between  $(1.70\pm0.38)$  for panicle weight and  $(2053.43\pm311.85)$  for spikelet fertility % of H<sub>1</sub> and ranged between  $(1.47\pm0.33)$  and  $(1858.53\pm268.08)$  of H<sub>2</sub> for the former characters, respectively. These results indicated that both additive and dominance components were observed to contribute in the expression of all studied characters except for number of panicles per plant and panicle weight. The magnitude of H<sub>1</sub> was more than H<sub>2</sub> in all characters indiacting that at most loci the positive and negative alleles were in equal proportion. However, these results were in general agreement with those reported by El-Mowafi (1994), Hammoud (1996), El-Refaee (2002) and Hammoud (2004).

Dominance variance over all heterozygous loci ( $h_2$ ) was highly significant for all characters except days to maturing and panicle length indicating that dominance was unidirectional, while was not unidirectional for days to maturing and panicle length El-Refaee (2002). The estimates of (F) were insignificant and negative for plant height, panicle weight and grain yield plant<sup>-1</sup> and not significant and positive for number of panicles plant<sup>-1</sup> spikelet fertility % and 1000-grain weight suggested that dominant and recessive genes were in equal proportion. On the contrary, (F) value was significant and highly significant for days to maturing, chlorophyll content and panicle length, revealing that there was asymmetric gene distribution with an excess of dominant alleles as compared with recessive ones in the parents. The environmental component (E) was insignificant for all characters except spikelet fertility% and grain yield plant<sup>-1</sup> and differed in its magnitude indicating that these characters are affected by environmental components with different degrees.

Estimates of the mean degree of dominance  $(H_1/D)^{1/2}$  were over one for all characters studied except days to maturing, plant height and panicle length. This finding indicating that over dominance and epistatic gene effects played a major role in the inheritance of this traits. The estimates of  $(H_2/4H_1)$ ratio did not reach the ratio 0.25 in case of all the characters and ranged between 0.14 to 0.23 for panicle length and grain yield per plant, respectively, indicating that genes having positive and negative effects were equally distributed in the parents.

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The ratio of the total number of dominant to recessive alleles in the parents ( $K_D/K_R$ ) was less than unity for plant height (0.868), panicle weight (0.571) and grain yield plant<sup>-1</sup> (0.658) indicating an excess of dominant alleles as compared with recessive ones in the parents, while it was more than one for all other characters indicating that proportion of dominant alleles was greater.

Heritability estimates (Table 4) in broad sense were high for all studied characters while, heritability estimates in narrow sense was relatively high for days to maturing (85.69 %), plant height ((76.74 %) and grain yield plant<sup>1</sup> (60.22%). This further suggested that a major part of the total phenotypic variance for these characters was due to additive genetic variance. Accordingly, it is expected that an effective phenotypic selection for these characters can be achieved with satisfactory degree of accuracy. In case of chlorophyll content and number of panicles plant<sup>-1</sup>, the estimates of such parameter were low and gave the values of 31.70 and 20.54, respectively. This further suggested that a major part of the total phenotypic variance for these characters was due to dominance genetic variance and environmental effects. These finding led to conclusions the selection for such characters must be done in the late generations. These results in general agreement with those reported by El-Mowafi (1988), El-Mowafi (1994), Kumar et al. (1994), El-Refaee (2002), Hammoud (2004), Awad Allah (2006) and Abd Allah (2008).

Table	4 : The estimates of heritability in both	ו broad	and narrow	sense
	for all agronomic characters studi	ed.		

Agronomic characters	Broad sense heritability (Hbs)	Narrow sense heritability (Hns)
Days to maturing (days)	99.42	85.69
Plant height (cm)	97.62	76.74
Chlorophyll content (SPAD)	92.59	31.70
No. of panicles per plant	89.34	20.54
Panicle length (cm)	92.23	54.63
Panicle weight (g)	99.03	41.93
Spikelet fertility(%)	92.45	54.98
1000-grian weight (g)	98.20	52.82
Grain yield per plant (g)	88.00	60.22

#### Similarity matrix :

The similarity matrix based on the nine agronomic characters was used to establish the level of relatedness between the nine varieties. Pairwise estimates of similarity ranged from (1.10) to (9.25) table (5). Two varieties (Sakha 101) and (Giza 171) were the closest genotypes with the highest similarity index of (9.25). this was followed by (9.10) similarity between a pair of varieties (HR5824-B-3-2-3) and (GZ6903-1-2-2-1). The lowest level of similarity (1.10) was obtained between Giza 177 and Giza 171.

Entries	١	۲	٣	٤	٥	٦	۷	٨	٩
Giza 177	0.00								
Sakha 101	3.92	0.00							
Sakha 104	3.70	3.44	• • •						
Giza 171	1.10	9.25	٨٩٣	•.••					
HR5824-B-3-2-3	5.55	8.52	٨.٢٨	۱.٦٠	•.••				
GZ6910-28-1-3-1	3.66	3.21	۱.۸٦	1.10	٨.٣٩	•.••			
GZ7685-8-1-3-2	1.80	3.80	r.01	1.11	०.१२	۳.۷۰	•.••		
GZ6522-15-1-1-3	3.92	3.66	۲.۹۲	٨,٦١	٨. ٤٢	1.97	۳٩٠	•.••	
GZ6903-1-2-2-1	4.44	3.01	۲٫٤٧	Y. 10	۹.۱۰	1.99	٤٢٠	۲.۸۳	•.••

 Table 5 : Similarity matrix for nine varieties based on nine agronomic characters.

#### Cluster analysis:

The genetic divergence can provide a visual idea about variabilities presented in nine genotypes, in addition to assure the continued genetic improvement. The genetic divergences based on Euclidean distance using morphological characters (quantitative traits) among nine parents, were graphically illustrated as a dendrogram in (Figure 1).

The standardized Euclidean distance ranged from 16 to 78.12 in the nine parents, where the minimum distance was observed between Giza 177 and GZ 7685-8-1-3-2, while the maximum Euclidean distance was observed between Giza 171 and Sakha 101. The nine parents were distinctly classified into five clusters (below 25 Euclidean distance).



Figure (1): Cluster diagram for nine varieties classified by nine agronomic characters studied.

The first group included all varieties except Giza 171, which was long duration. The first group was divided into two sub groups based on maturing, plant height, spikelet fertility percentage and grain yield per plant. The first sub group, which was higher in these four characters, included seven varieties (Giza 177, GZ7685-8-1-3-2, Sakha 101, Sakha 104, GZ6910-28-1-

3-1, GZ6903-1-2-2-1 and GZ6522-15-1-1-3) except for HR5824-B-3-2-3, which was lower in these characters. The two varieties Giza 177 and GZ7685-8-1-3-2 of branch one were similar in all characters except grain yield per plant. Sakha 101 variety was separated alone into on branch because it has long maturing and highest grain yield per plant when compared with the rest five varieties. GZ6522-15-1-1-3 was separated from four varieties in one branch based on maturing, plant height and number of panicles per plant. GZ6903-1-2-2-1 genotype was separated from these three varieties based on maturing, plant height and spikelet fertility percentage. On the other hand, the remaining two varieties Sakha 104 and GZ6910-28-1-3-1 were similar in all characters except in grain yield per plant.

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تقدير المكونات الوراثيه ودرجه التوريث وتحليل الشجرة الوراثيه لبعض الصفات المحصوليه في الأرز عمرو فاروق عبد الخالق ،وائل حمدي محرم الكلاوى ، اشرف صلاح مصطفى عبداللطيف و جلال بكر أنيس مركز بحوث الأرز – معهد المحاصيل الحقلية – مركز البحوث الزراعية

أقيمت تجرب حقلية بهدف إجراء التحليل الدوري لتسعه أباء والحصول على ٣٦ هجين وكان إحدى هذه الآباء صنف مستورد وباقي الأصناف تجاريه وسلالات مبشرة وذلك من خلال استخدام طريقه ها يمان في عام (١٩٥٤). وتم زراعه الآباء والجيل الأول في تجربه قطاعات كاملة العشوائية في ثلاث مكررات عام ٢٠١١ بمزرعة مركز البحوث والتدريب في الأرز بسخا – كفر الشيخ. أخذت البيانات على ٩ صفات محصوليه. وأوضحت النتائج أن مجموع مربع متوسطات قيم الصفات المدروسة كانت عالية المعنوية مشيرة إلى وجود اختلافات وراثية كبيرة بين التراكيب الوراثيه في جميع الصفات المدروسة كانت عالية المعنوية الوراثي المضيف معنوي وعالي المعنوية لجميع الصفات المدروسة ما عدا صفات المدروسة. وكان التباين السنبلة. وكان التباين الوراثي السيادي عالي المعنوية لجميع الصفات المدروسة وزن السنبلة. وكان التباين الوراثي السيادي عالي المعنوية لجميع الصفات كما كان التباين البيئي غير معنوي لمعظم الصفات ما عدا صفه النسبة المنوية للحبوب الخصبة في السنبلة ومحصول النبات الفردي. وتبين من النتائج أن درجه التوريث بمعناها الواسع كانت عالية لجميع الصفات محل الدراسة وان تحلي من النتائج أن درجه التوريث بمعناها الواسع كانت عالية لجميع الصفات محل الدراسة وان تحليل النبات وراثيه قسم الأباء إلى حمس مجاميع رئيسيه وذلك طبقا لدرجه القرابة ما ين تلك الأباء.

وتوصى هذه النتائج إلى إمكانية استخدام هذه الهجن في برامج التربية والحصول منها على تراكيب وراثية متباينة تمكن من إجراء عملية الانتخاب الفردي في الأجيال التالي

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Table 3 : The estimates of genetic components for all characters studied.

	Estimates ± S.E.									
Genetic Parameters	Days to maturing (days)	Plant height (cm)	Chlorophyll content (SPAD)	No. of panicles /plant	Panicle length (cm)	Panicle weight (gm)	Spikelet fertility (%)	1000-grian weight (gm)	Grian yield/plant (gm)	
	**	**	**	ns	**	ns	**	**	**	
D	157.44±6.90 **	204.47±17.54 **	11.98±2.17 **	6.80±9.83	15.79±0.89 **	0.01±0.17	1226.95±141.29 **	2.58±0.55 **	111.36±21.09 **	
H1	66.37±15.23	186.72±38.71 **	28.70±4.78 **	141.04±21.70 **	10.99±1.96 **	1.70±0.38	2053.43±311.85 **	8.16±1.22	171.00±46.54 **	
H2	44.57±13.10 ns	147.56±33.27 **	19.69±4.11 **	125.80±18.66 **	6.31±1.69 ns	1.47±0.33	1858.53±268.08 **	6.36±1.05 **	162.85±40.01 **	
h2	-0.18±8.77	251.38±22.29 ns	24.82±2.75 **	405.35±12.50 ns	0.37±1.13	2.37±0.22 ns	4270.27±179.59 ns	2.26±0.70 ns	311.65±26.80 ns	
F	40.20±16.10	-27.57±40.91	15.87±5.06	3.27±22.94	15.90±2.07	-0.22±0.41	58.15±329.60	0.68±1.29	-56.96±49.19	
	ns	ns	ns	ns	ns	ns	*	ns	**	
E	0.47±2.18	4.20±5.55	0.60±0.69	4.87±3.11	0.33±0.28	0.01±0.06	93.61±44.68	0.06±0.18	17.58±6.67	
(H <sub>1</sub> /D) <sup>1/2</sup>	0.649	. 907	1.051	4.555	0.834	4.196	1.294	1.778	1.239	
(H <sub>2</sub> /4 H <sub>1</sub> )		. 191	. 171	0.223	0.143	0.217	0.226	0.195	0.238	
(K <sub>D</sub> /K <sub>R</sub> )	1.589		4.547	1.111	4.040	0.571	1.037	1.160	0.658	
(r)	. 911		. 209	0.449	0.913	0.769	0.847		0.351	

\*, \*\* significant at 0.05 and 0.01 level of probability, respectively. and ns : Not Significant.

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