USING MULTIVARIATE STATISTICAL TECHNIQUES AND ESTIMATION OF SOME GENETIC PARAMETERS IN THIRTEEN FABA BEAN GENOTYPES

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ABSTRACT

Two field experiments were conducted at the Agricultural Research Station of gemmeiza, ARC, El-Gharbia Governorate during 2011/2012 and 2012/2013 seasons in order to evaluate the performance of thirteen Faba bean genotypes for seed yield per plant and its related characters. Four statistical procedures, *i.e.* correlation and multiple regression analysis, factor and cluster analysis were applied to determine the main contributing characters in seed weight per plant variation. Genotype 2 possessed the earliest flowering plants and Genotype 9 (mutant 1) possessed the earliest matured plants. Genotype 8 followed by Genotype, 6, 4 and 1 showed the tallest plants, highest number of pods, seeds, seeds yield per plant and 100 seed weight. Highly significant and positive associations were obtained between seed weight/plant and each of number of seeds/plant and number of pods/plant. The full model regression gave R-square of 0.995, which means that 99.5% of the variation in Faba bean yield is explained by the independent variables in the model. Step-wise regression analysis for seed yield / plant as dependent variable and the other traits as independent, revealed that number of seeds, 100seed weight and number of branches are the most important seed yield / plant components. These traits accounted for 99.26% of total variation exist in seed yield / plant. Factorial analysis classified the eight studied characters, as yield per plant components, into three main factors explained 79.49% of the total variability in the dependent structure. Factor 1 was responsible for 35.71% of the total variation and included number of pods/plant, number of seeds/plant and seeds yield /plant. Factor 2 included100seed weight, days to maturity and plant height contributed by 24.49% of the total variation. Number of branches/plant and days to flowering accounted for 19.29% of the total variation. Graphically, showing eigen values in response to number of components for the estimated variables of Faba bean. The hierarchical cluster analysis permitted the classification of the genotypes according to similarity groups. Analysis of variance for traits studied showed significant differences among the genotypes. Phenotypic coefficient of variation for most characters was closer to the corresponding genotypic coefficient of variation values, showing little environmental effect on the expression of these characters. The highest values of phenotypic and genotypic coefficients of variation were obtained with number of branches/plant, seeds yield/ plant, number of seed and number of pods/plant indicating a wide range of variation which provided a good opportunity for yield improvement. Also, large proportions of heritability coupled with high values of genetic advance (% of mean) were recorded for the number of branches/plant, seeds yield/ plant, number of seeds and the number of pods/plant, explaining that these traits have more chance for Faba bean yield development among the tested genotypes.

Key words: Faba bean, genetic parameters, multivariate statistical analysis.

1. INTRODUCTION

Faba bean, broad bean or field bean (*Vicia* faba L.) is a major legume that is used as food owing to the high nutrient components in seeds (Duc, 1997). It is largely consumed in the Middle East. Genetic variability of Faba bean is quite large. *Vicia faba* is a considerable source

of energy: 344 K cal/100 g and can efficiently replace animal protein in poor countries (Bond *et al.*, 1985 and Duke, 1981). It is a significant source of protein rich food in developing countries. It is also used both as a human food and a feed, horses, poultry and pigeons in industrialized countries (Duke, 1981). Its characteristics are suitable for sustainable agriculture (Nadal *et al.*, 2003). Faba bean is also grown for green manure and can significantly enhance yields of cereals or other crops (Wani *et al.*, 1994). Several studies were concentrated about the impact of each yield component on the yield.

The main goal when growing crops everywhere is to maximize net profit mainly through increasing seed yield. Different types of multivariate analysis such as factor and cluster analysis can be used to identify groups of genotypes that have beneficial traits for breeding and instructing the patterns of variation in genotype accession, to recognize relationships among accessions and possible gaps (Cowen and Frey, 1987). Though simple correlation coefficient is very useful in quantifying the size The objectives of this work were to study the genetic parameters and identify the interrelationships among seed yield and its related characters using correlation and regression analysis and cluster and factor analysis. The results may be helpful to plan appropriate selection strategies for improving seed yield in Faba bean.

2. MATERIALS AND METHODS

2.1. Location of the study and plant materials

The present study was carried out at the Agricultural Research Station of Gemmeiza (ARC) to evaluate the performance of 13 Faba bean genotypes (crosses resulted from F7) for yield and its related characters. The details of pedigree for the tested genotypes are presented in Table (1).

Genotype	Genotype	Pedigree
number	name	
1	Cross 1	Giza 461 × Gizablanka
2	Cross 2	T.W. × Gizablanka
3	Cross 3	Giza 716 × Giza 402
4	Cross 4	Giza 461 × Giza 402
5	Cross 5	(Gizablanka × Giza 2) × misr 1
6	Cross 6	(Giza 429 × Giza 2) × misr 1
7	Cross 7	(Gizablanka × Giza 40) × misr 1
8	Cross 8	Giza 461 × Giza 3
9	Mutant1	Giza 461 M.
10	Cross 9	(TripleWhite × Giza429) × Misr 1
11	Giza 834	561/2076/85× 461/845/83
12	Misr 1	Derived from G3× 123A/45/76
13	Giza 3	Cross (Giza 1× Dutch)

Table (1): Name and pedigree of the studied faba bean genotypes.

and direction of association, it may not always be effective because it provides only one dimensional coefficient neglecting important and complex interrelationships among other traits (Kang, 1994).

The factor analysis is also a data reduction method to reduce the number of variables into factors. Cluster analysis is a multivariate method for grouping the individuals or objects according to their traits, and classifying the individuals with similar types into the same group or cluster (Hair *et al.*, 1995). Chaieb *et al.* (2011) used the hierarchical cluster analysis for classification of 14 Faba bean genotypes according to some morphological traits. Sarparast *et al.* (2011) used cluster analysis to investigate genotypic structure of nine Faba bean genotypes.

2.2. Experimental design and plot arrangement

The experiments were designed in a Randomized Complete Blocks Design (RCBD) with three replications. Each replicate consisted of thirteen plots, each devoted to one genotype. Plots were 11.2 m^2 including four ridges, 2.8 m long spaced 0.7 m apart, in hills spaced 0.2 m apart. The two season experiments were sown on 15^{th} and 20^{th} of November for 2011/2012 and 2012/2013 seasons respectively. All the recommended cultural practices were followed up to harvest and all agronomic practices were carried out uniformly for all treatments.

2.2.1. Data collection

Data on different agronomic traits were collected on both plant and plot basis.

At maturity ten plants were taken at random from the inner two ridges in each plot to measure the following traits: days to 50% flowering, days to maturity, plant height (cm), number of branches per plant, number of pods per plant, number of seeds per plant, 100-seed weight (g) and seed yield per plant(g).

2.3. Statistical analysis and estimation of genetic parameters

Individual analysis of variance was performed for all the studied traits on each season according to the procedure described by Snedecor and Cochran (1989) for a randomized complete blocks design with three replications. The combined data across the two seasons were subjected to statistical analysis using MSTAT- C computer software. Data for the two years were tested for homogeneity error variance using Ftest and it was found to be homogeneous so the Least data were combined for analysis. significant differences LSD test at 5% level of probability was used to compare means.

Four statistical procedures, differing in mathematical background, goals and final output, were used to explore the relationships between seed yield/plant and other traits of Faba bean. The used models are summarized as follows:

Simple correlation coefficients were computed among seed yield/plant and its components (Gomez and Gomez, 1984).

Multiple regression analysis according to Draper and Smith, (1966) was done to develop equations to predict yield.

Factor analysis: This approach was applied according to Cattell (1965) to reduce a large number of correlated variables to a much smaller number of independent clusters of variables called factors. After the loading of the first factor was calculated, the process was repeated on the residual matrix to find further factors. When the contribution of a factor to the total percentage of the trace was less than 10%, the process was stopped. After extraction, the matrix of factor loadings was submitted to a Varimax Orthogonal Rotation, as applied by Kaiser (1958). The purpose of rotation was to rebuild the larger loadings in each factor and to suppress the minor loading coefficient so as to improve the opportunity of achieving meaningful biological interpretation of each factor.

Cluster Analysis (CA) was used to arrange a set of variables into clusters. The aim was to establish a set of clusters so that cases within a cluster are more similar to each other than between clusters. This description may be abstracted through the use of the particular to the general class or type (Polignano *et al.*, 1989). Twin span (two way indicator species analysis) was used to classify variables producing an order two way table of their occurrence. The cluster analysis was performed using a measure of similarity levels and Euclidean distance (Everitt, 1993 and Eisen *et al.*, 1998).

The genotypic and phenotypic variances (σ_g^2 and σ_{ph}^2) were calculated from the results of analysis of variance as follows:

1- Genotypic variance (σ_g^2) was calculated from the formula outlined by

Comstok and Moll (1963):

 $\sigma_g^2 = (Mg - Msg)/rs$

Where σ_{g}^{2} = Genotypic variance

Mg = Mean square of genotypes

Msg = Mean square of the seasons x genotypes interaction.

r = Number of replications.

s = Number of environments (seasons).

2- Phenotypic variance (σ_{ph}^2) was computed according to the following equation as applied by Mathur *et al.* (1971) and Verma and Singh (1971):

$$\sigma_{ph}^2 = \sigma_g^2 + (\sigma_{sg}^2/r) + (\sigma_e^2/rs)$$

Where $\sigma_{sg}^2 = (Msg-Me)/r$.

 $\sigma_e^2 = Me$ (error mean square).

Genotypic and phenotypic coefficients of variation were calculated according to Burton (1951) as follows:

G. C. V.= σ_g / \overline{x} .100 and P. C. V. = $\sigma_{ph} / \overline{x}$.100. Broad sense heritability (H) was calculated as described by Hanson (1963) using the following formula:

 $h_b\% = (\sigma_g^2 / \sigma_{ph}^2) . 100$

The expected genetic advance under selection (Gs) was calculated from the following formula as suggested by Fehr (1987):

 $Gs = K. \sigma_{ph}. h_b$

Where σ_{ph} = The phenotypic standard deviation and

K = The selection differential in standard deviation units. In this investigation, the value used for K is 2.06, which corresponds to selecting the best 5% of the population.

3. RESULTS AND DISCUSSION

3.1. Combined analysis of variance

The results of combined analysis of variance after homogeneity test for error variances are summarized in Table (2). F-test of different sources of variation revealed that there were no significant differences of the year effects for all E. M. Rabie and H. E.A. Ibrahim.....

traits except for plant height, while the effect of genotypes was significant for all traits except for plant height. cross 8 possessed the highest values for the number of pods and seeds per plant and recorded (27.50) and (79.57 g), respectively. On the other hand, cross 6 possessed the least values for the

	Wean squares										
SOV	df	Days to	Days	Plant	Number	Number	Number of	Seed yield	100-seed		
1		50%	to	height	of	of	seeds/plant	/plant (g)	weight		
		Flowering	Maturity	(cm)	branches/	pods/plant			(g)		
					plant						
Rep	2	5.35	8.17	3.01	0.29	3.95	29.79	24.94	0.289		
Genotypes	12	21.72**	14.60**	53.45	7.78**	31.27**	386.65**	254.46**	84.89 **		
Error	24	1.26	1.35	28.22	0.37	3.62	25.79	12.14	2.61		

Table (2): Combined analysis of variance of the evaluated genotypes o	ver two seasons.
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3.2. Mean performance

The mean performances of thirteen Faba bean genotypes for different traits are given in Table (3). Based on two year data; results revealed that the cross 2 possessed the earliest flowering plants (47.33 days) and matured after157.17 days. On the other hand, the cultivar Giza 3 took 53.33 and 156.33 days to flower and maturity, respectively.

With respect to maturity date, M1 possessed the earliest matured plants (153.33 days). On the other hand, cross 5 possessed the latest ones (158.00 days). Regarding to plant height, Cross 6 possessed the tallest plants (103.17 cm) whereas, Cross 9 exhibited the shortest plants (94.17 cm). For the number of branches per plant, cross 2 showed the profuse plants (5.63) whereas the cultivar Giza 3 possessed the lowest branched plants (2.23). Results showed that the number of pods (19.05) and the number of seeds per plant (50.23). For seed yield per plant and 100-seed weight, cross 1 possessed the heaviest seed weight (60.57 and 84.75 g) among the thirteen genotypes. On the other hand, cross 5 possessed the lowest value of 100 seed weight (72.80 g), and cross 6 possessed the lowest value of seed yield per plant (41.1g)

According to the previous results, it could be concluded that the tallest plants, highest number of pods, seed yield per plant and 100 seed weight were obtained by cross 8 followed by cross 6, cross 4 and cross 1. These results reflect that the selection prospects within this genotype to improve the performance through breeding program.

These results are in harmony with those reported by Bastawisy *et al.* (2006), Salem (2011) and Ihsanullh (2010) who found

Genotype	Days to	Days to	Plant	Number of	Number	Number of	Seed	100-
	50%	Maturity	height	branches/plant	of	seeds/plant	yield	seed
	Flowering	r	(cm)		pods/plant		/plant (g)	weight
								(g)
Cross 1	52.00	157.17	99.67	5.50	24.47	71.60	60.57	84.75
Cross 2	47.33	157.17	97.33	5.63	23.33	62.78	48.75	77.85
Cross 3	51.00	155.83	96.6 7	5.50	24.28	68.75	52.50	76.48
Cross 4	50.50	157.33	102.50	4.20	21.65	61.78	51.15	82.92
Cross 5	50.50	158.00	98.00	4.98	24.60	67.07	48.78	72.80
Cross 6	51.00	157.83	103.17	4.63	19.05	50.23	41.10	80.37
Cross 7	48.33	155.17	102.50	3.98	19.32	52.73	39.20	74.40
Cross 8	52.83	156.00	102.17	3.85	27.50	79.57	59.97	75.47
M1	47.83	153.33	97.17	4.65	23.67	57.13	46.15	80.73
Cross 9	52.33	154.17	94.17	5.25	21.33	56.83	41.93	73.85
Giza 843	52.00	154.33	97.33	2.97	20.57	59.22	44.60	75.42
Misr 1	50.00	154.17	96.17	2.53	23.40	65.73	48.45	73.73
Giza 3	53.33	156.33	101.83	2.23	21.53	59.70	46.60	78.13
L. S. D 0.05	2.69	2.89	NS	1.52	2.93	6.69	6.19	4.18

Table (3): Mean	performance of	f 13 faba bean g	genotypes for	the studied traits ((Combined	data over two	years).
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significant differences among tested genotypes for seed yield and its related characters.

3.3. Correlation analysis

The simple correlation coefficients among seed yield/plant and the other related traits are presented in Table (4). Results showed that all traits are associated highly significantly with seed yield/plant. The most important positive relationships to Faba bean breeder are those between seed yield/plant and each of the number of seeds/plant ($r = 0.924^{**}$) and the number of pods/plant ($r = 0.801^{**}$).

On the other hand, the traits of yield components exhibited considerable trends of associations among them. Positive and highly significant correlations were detected between seed yield/plant and each of the number of pods/plant ($r = 0.801^{**}$) and the number of seeds/plant ($r = 0.924^{**}$). Also, significant positive correlation ($r = 0.267^{*}$) was observed between the number of pods/plant and the number of branches. The number of days to 50% flowering was found to be significantly and negatively associated with the number of

3.4. Multiple linear regression analysis

The obtained results showed that the prediction equation for seed yield per plant (\hat{Y}) is formulated using the Faba bean variables as follows:

Seed yield per plant (\hat{Y}) = - 49.2 + 0.0119 Plant height + 0.100 Number of branches + 0.0311 Flowering date - 0.0141 Maturity date, days -0.0266 Number of pods + 0.765 Number of seeds^{**} + 0.637 weight of 100-seed^{**}

R-Sq = 99.3% R-Sq (adj) = 99.2%

The seed yield per plant formula explains 99.3% of the total variation within the yield components, while the remaining 0.7% may be due to residual effects. The t-test showed that number of seeds and weight of 100-seed has contributed significantly towards seed yield per plant. While the other six variables did not. The overall results reflect the importance of the mentioned commensurate two variables in Faba bean selection for breeding programs. These findings are in accordance with the results obtained by Chaieb *et al.* (2011).

Table (4): Simple correlation coefficients among the eight studied traits (combined over two seasons).

Traits	Ph	NB	DF	DM	NP	NS	100 SW	SY/P
Ph	1							
NB	0.084	1						
DF	-0.089	-0.278*	1					
DM	0.155	0.191	0.106	1				
NP	-0.046	0.267*	0.036	-0.06	1			
NS	-0.127	0.156	0.212	0.059	0.872**	1		
100SW	0.180	0.095	-0.084	0.270	-0.153	-0.164	1	
SY/P	-0.045	0.204	0.181	0.162	0.801	0.924**	0.215	1

*, ** and ns indicates significant at the 0.05 and 0.01 level of probability and insignificant, respectively. Ph: Plant height (cm), NB: No. of branches/plant, DF: Days to 50% flowering, DM: Days to maturity, NP: No. of pods/plant, NS: No. of seeds/plant,100 SW:100 - seed weight (g) and SY/P: Seed yield/plant (g)

branches (r = -0.278^*). Similar results were observed by Hakan *et al.* (2003), Nawar *et al.* (2010), Talal and Shataya (2013) and Tafere *et al.* (2013).

3.5. Stepwise multiple linear regression analysis

Data presented in Table (5) show stepwise multiple regression analysis of the estimated

Table (5): Stepwise regression	between seed yield per	plant and some studied characters
in Faha bean.		

Step	1	2	3
Constant	3.854	- 48.905	- 48.866
No. of seeds/plant	0.7130	0.7606	0.7586**
100 - seed weight (g)		0.643	0.639**
R-Sq	85.41	99.24	99.26
R-Sq(adj)	85.22	99.22	99.23

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variables in predicting seed yield per plant. The obtained results showed that 99.26% of the total variation in yield resulted from the number of seeds and 100 seed weight, as being the most suitable inputs to the model.

The obtained results showed that the best prediction equation for seed yield/plant (\hat{Y}) is formulated as follows:

Seed yield / plant = $-48.866 + 0.7586^{**}$ number of seeds $+ 0.639^{**}$ weight of 100-seed.

Hence, it could be concluded that selection based on the number of seeds and 100 seed weight is more appropriate. These findings are in accordance with the results obtained by Golparvar (2012) and Mojgan *et al.* (2013),

3.6. Factor analysis

The factor analysis technique divided the eight studied Faba bean yield components into three independent groups or factors which explained 79.49 % of the total variability in the dependence structure. The factors were constructed by applying the principal component approach to establish the dependent relationship between seed yield attributes in Faba bean. For interpretation only factor loadings greater than 0.5 were considered important (Seiler and Stafford, 1985). Summary of the composition of variables of the three extracted factors with loadings are given in Table (6) and Fig. (1).

of the relationship between the factor and its related traits. It accounted for 24.49 % of the total variability in the dependence structure.

Factor 3 was responsible for 19.29 % of the total variability in the dependence structure. It included the two characters the number of branches/plant and days to 50% flowering. It contains the variable usually regarded as a growth and earliness factor. These results are in agreement with Mehasen and Mohamed (2004).

EL-Badawy (2006) found that using of factor analysis by plant breeders has the potential of increasing the comprehension of the casual relationship of variables and can help to determine the nature and sequence of traits to be selected in a breeding program.

3.7. Cluster analysis

Cluster analysis seemed to be an efficient procedure for extracting the structured relationships among accessions and provides a hierarchical classification of them (Polignano *et al.*, 1989). Hierarchical cluster analysis for clustering the investigated 13 genotypes (the three categories) obtained with the average linkage procedure; UPGMA (un-weighted pair group method using arithmetic average) developed by Sokal and Michener (1958) is illustrated in Table (7) and Fig. (2).

Variable	Loading	% Communality	Eigen values	% of variance
Factor 1:				
No. of pods/plant	0.924	0.884		
No. of seeds/plant Seed	0.935	0.955	2.856	35.71
yield/plant (g)	0.950	0.970		
Factor 2:				
100 - seed weight (g)	0.716	0.568	1.959	
Days to maturity	0.809	0.689		24.49
Plant height(cm)	0.808	0.828		
Factor 3:				
No. of branches/plant	0.862	0.840	1.544	19.29
Days to 50% flowering	0.698	0.625		
Cumulative variance				79.49

Table (6): Summary of factor loadings for eight characters of Faba bean.

Factor 1 included three variables which accounted for 35.71 % of the total variability. The three variables were the number of pods/plant, the number of seeds/plant and seed yield per plant. The three variables had high communality with factor 1. Therefore, this factor may be called seed yield factor.

Factor 2 was made up of the traits100-seed weight, days to maturity and plant height. The sign of the loading values indicates the direction

The clustering pattern revealed that the genetic divergences in 13 genotypes on ten characters were grouped in 3 main clusters (Table 7). The maximum number of genotypes (7) was grouped in cluster 2 followed by cluster 3 (4) and cluster 1 (2). Thus, in view of considerable genetic diversity in Faba bean found in the present study, their appearance had sufficient scope for genotypic improvement through hybridization between the divergent





Fig. (1): Scree plot showing eigen values in response to the number of components for the estimated variables of Faba bean.

Table (7): Level of different clusters for faba bean studied genotypes(combined data over two vears)

1

Similarity	Cluster	No. of genotypes	genotypes included
62.56	1	2	Cross1, Cross 8
55.89	2	7	Cross 2, M1, Cross 4, Giza 3, Cross 3, Cross5, Misr 1
44.73	3	4	Cross 6, Cross7, Cross9,Giza 483



Fig. (2): Distribution of thirteen Faba bean genotypes in three clusters 1=Cross1, 2 = Cross2, 3= Cross3, 4= Cross4, 5= Cross5, 6 = Cross 6, 7= Cross7, 8 = Cross8, 9 = M110 = Cross 9, 11= Giza 843, 12 = misr1, 13= Giza 3

genotypes (Eisen *et al.*, 1998), Chaieb *et al.*, 2011, Sarparast *et al.* (2011), Golparvar, 2012 and Peyman and Aminpana, (2014). The present investigation will be effective to assess the extent of available variability, which will be useful for selecting superior genotypes on the basis of their phenotypic expression so as to use them in breeding programs to improve the important characters.

3.8. Variability and genetic parameters

Genotypic and phenotypic coefficients of variation (GCV and PCV), broad sense of heritability (h^2), and genetic advance expressed as percentage of the mean for the studied traits are shown in Table (8) and Fig. (3).

Because the variance as a measure of variability (phenotypic or genotypic) is affected by the measure unit of traits, it is seldom to be used in comparisons. Accordingly, the comparison among yield traits concerning the degree of variability was made using the magnitude of coefficient of variation which is an absolute measure of variability.

The estimate of (GCV) was only high for the number of branches/plant (36.26) while

moderate values of (GCV) were recorded with seed yield/plant (19.09), No. of seeds/plant (16.35), number of pods/plant (14.01), 100-seed weight (g) (6.60) and days to 50% flowering (5.10). The remaining two traits showed low estimates of (GCV).

On the other hand, the values of (PCV) were slightly higher than the values of (GCV) for all traits which reflect somewhat the influence of environment on the expression of traits. In accordance, the selection would be effective to improve these traits among the studied genotypes. Similar findings were reported by Mehasen and Mohamed (2004) and Tafere *et al.* (2013).

It is important to emphasize that, without considering genetic advance, the heritability values (h^2) would not be practically important in selection based on phenotypic appearance. Johnson *et al.*, (1955) confirmed that heritability estimates along with genetic advance would give a more reliable index of selection value.

In the present investigation, high values of heritability coupled with high values of genetic advance (as % of mean) were recorded for seed

Character	Mean	GCV %	PCV %	h ² b	Genetic advance (GA)	Expected GA(% of mean)
Plant height (cm)	99.13	3.73	4.19	79.27	6.78	6.84
No. of branches/plant	4.31	36.26	36.75	97.34	3.18	73.69
Days to 50% flowering	50.69	5.10	5.15	98.17	5.28	10.14
Days to maturity	155.91	1.30	1.32	96.66	4.10	2.63
No. of pods/plant	22.73	14.01	14.36	95.14	6.40	28.14
No. of seeds/plant	69.17	16.35	16.55	97.57	23.01	33.27
100 - seed weight (g)	77.45	6.60	6.63	99.10	10.48	13.53
Seed yield/plant (g)	48.44	19.09	19.26	98.18	18.87	38.96

Table (8): Means, genotypic (GCV) and phenotypic (PCV) coefficients of variability, heritability (h²bs) and genetic advance (GA).



Fig. (3). Effect of GCV, PCV, GA, H% on seed yield/plant. ph: plant height (cm), nb: number of branches/plant, df: days to maturity, dm: days to maturity, np: number of pods/plant, ns: number of seeds/plant, 100sw 100 - seed weight (g) and swp: seed yield/plant (g).

yield/plant, the number of branches/plant, seed weight/plant, the number of seeds/plant and the number of pods/plant indicating the importance of the additive gene effects, so, selection would be effective in early generations for these traits. High values of heritability accompanied with low genetic advance for plant height, 100-seed weight, days to flowering and days to maturity are indicative of non-additive gene effects. Therefore, limited scope for improvement of these traits is expected under the tested genotypes. The current conclusions are supported by Mehasen and Mohamed (2004), who confirmed that plant breeders can safely make their selection when they take in consideration high values of heritability and genetic advance.

The estimates of GCV and PCV values give the extent of variability existing for various traits, but do not give any information about the heritable portion of it. Therefore, estimates of heritability and genetic advance would give better idea about the possible gains of selection (Chavan *et al.*, 1994).

Conclusion

Based on the results obtained in this study the following conclusions could be drawn:- Results from the present study indicate that cross 1 and cross 8 gave the highest seed yield per plant. Highly significant and positive associations were obtained between seed weight/plant and each of the number of seeds/plant and the number of pods/plant. The full model regression and stepwise regression analysis gave, R-square of 0.995 and 99.26% of the total variation exist in seed yield / plant trait. factor analysis classified the eight studied characters The hierarchical cluster analysis permitted the classification of the genotypes according to similarity groups. Large proportions of heritability coupled with high values of genetic advance (% of mean) were recorded for the number of branches/plant, seeds yield/ plant, number of seeds and the number of pods/plant. The information from this study would be a valuable to Faba bean breeder for developing high yielding genotypes.

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إستخدام أساليب التحليل متعدد المتغيرات وتقدير بعض المعالم الوراثية في الفول البلدي

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ملخص

أجريت تجربتين في محطة بحوث الجميزة خلال موسمي 2011 /2012 و2013/2012 لتقييم أداء ثلاثة عشر تركيب وراثي من الفول البلدي لمحصول بذور النبات والصفات المرتبطة به .تم استخدام أربعة طرق إحصانية هي تحليل الارتباط، والانحدار المتعدد، وتحليل العامل والتحليل العنقودي لتحديد مدى مساهمة الصفات في تباين وزن بذور النبات. تم تقدير درجة التوريث بمعناها الواسع ومعامل الإختلاف المظهري والوراثي ومعدل التحسين الوراثي المتوقع. أوضحت النتانج أن الهجين 2 كان أكثر تبكيرا في التزهيرو المهجين 9 (MI) كان أكثر تبكيرا في النضج. أظهرت النتانج أن أطول النباتات، وأكبر عدد من القرون والبذور ووزن محصول النبات ووزن 100 بذرة تم الحصول عليها من الهجن 8 و 6 و 4 و1 على التوالي. مما يشير الى تفوقها و كونها من التراكيب الوراثية المبشرة التي يوصبي بالتوسع في استخدامها في برامج تربية الفول البلدي. أظهرت نتانج تحليل الارتباط وجود إرتباط عالي موجب بين محصول النبات وكل من عدد البذور / نبات وعدد القرون / نبات. أشارت نتائج تحليل الانحدار بالطريقة العيارية (النموذج الكامل) أن معامل التحديد R² سجل 0.995 مما يعني أن 99.5٪ من التباين في محصول البذور للنبات يفسره المتغيرات المستقلة في النموذج. أوضحت نتائج تحليل الانحدار المتعدد المرحلي أن عدد البذور ووزن 100 بذرة وعدد الأفرع على النبات من أهم مكونات محصوَّل النبات وهذه الصفات تفسر 99.26٪ من التباين الكلي. اوضحت نتائج التحليل العاملي ان أكثر الصفات أهمية هي عدد القرون وعدد البذورو محصول النبات حيث تفع في العامل الأول. اشارت النتائج إلى ان التحليل العنقودي الهرمي قسم التراكيب الورائية إلى ثلاثة عناقيد وفقا لدرجة التشابه بينهم. أظهر معامل الإختلاف المظّهري والوراثي تباين لكل الصفات مما يشير إلى إمكانية تحسين صفات المحصول. اشارت النتانج إلى أن أعلى قيم من درجة التوريث في المعنى العام ومصحوبة بأعلى قيم للتحسين الوراثي المتوقع (% متوسط) كان لعدد الأفرع ومحصول النبات وعدد البذور وعدد القرون على النبات وأن هذه الصفات تساهم في تحسين محصول الفول للتر اكيب الور اثية التي تم إختبار ها.

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