# IDENTIFICATION OF SOME FABA BEAN (Vicia faba L.) GENOTYPES USING MORPHOLOGICAL AND MOLECULAR CHARACTERS

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### **ABSTRACT**

Field and laboratory experiments were carried out at the Farm of El-Gemmeza Agricultural Research Station, Gharbia Governorate and Seed Technology Research Department, ARC, Egypt, during 2010/2011 and 2011/2012 seasons to identify and discriminate ten faba bean genotypes using morphological characters and molecular marker. The results revealed that some morphological characters such as pinnul shape, lines density of flag flower, pod color at maturity, testa shape and color were useful to identify some genotypes from each other, while they were not enough for identifying other genotypes. By using Inter-simple sequence repeat (ISSR-PCR) technique, it was possible to determine the genetic diversity and relationships of the ten faba bean genotypes included in this study. A total of 71 amplified bands were generated with five ISSR primers, of which 59 (83.1%) were polymorphic which represent a relatively high polymorphism level. These results are important in protecting of plant breeders rights and at releasing these genotypes as a new varieties.

## INTRODUCTION

The morphological, quantitative and biochemical characters study was designed to find out distinguished characters of faba bean genotypes. Morphological characterization is the first step in the classification and description of any crop germplasm. Nevertheless, the qualitative traits are often used for separating varieties when a limited range of quantitative traits are found in certain group. Furthermore, morphological description is a precondition for the protection and registration of varieties (UPOV 2002). Germplasm evaluation is considered the first step in plant breeding program and it is commonly based on a simultaneous examination of large number of populations for several characters of both agronomic and physiological interest (Pezzotti et. al. 1994). Cultivars within species are normally discriminated by morphological descriptors. Many tools are now available for studying genetic variability among accessions including total seed protein, isozymes and various types of molecular markers. Rehab, Tawdy (2007), identified ten vicia faba varieties based on morphological differences in seed, seedling and adult plant such as days from planting to flowering and maturity, anthocyanin coloration, color of testa, number of pods and 1000 seed weight), in addition to biochemical variability of genomic fingerprinting. Zubair et. al., (2007) evaluated 14 quantitative traits for forty diverse mungbean [Vigna radiata (L.) Wilczek] genotypes, they found that medium to high variance was observed for days to flower initiation, days to flowering, days to maturity, plant height and pods per plant. Meanwhile, small variance was observed for seeds per pod and 100 seed weight. Lalazar (2012) evaluate

the diversity of phenology and morphology traits such as days from germination to flowering and maturity, flower bed length, flowers number and length, flower pedanlel length and 1000 seed weight in 11 genotypes, he reported that the highest days from germination to flowering was in Potomak and Harvester genotypes and the lowest in Wadekh genotype (31 day). The highest number of pod was in Potomak and Harvester genotypes and the lowest in Saksa b/v 615 and Oltin genotypes. The highest 1000 seed weight was noticed from Potomak genotype and same genotype had the lowest days from germination to maturity (50 day). Mudzana et al. (1995) reported that the morphological characters such as plant height, number of days to 50 per cent flowering, flower length, pod length, number of seeds per pod could be used for variety identification of faba beans. Bonetti et al. (1995) reported that 17 bean cultivars were grouped based on pod length (very short, short, medium, long, very long), maturity (early, medium, late) and time of flowering (early, medium and late). Ashok et al. (2008) grouped seven french bean varieties based on hilum color and seed shape. On the other hand, the commonly used polymerasechain reaction (PCR)-based marker systems for genetic diversity and relationships in faba bean species are randomly amplified polymorphic DNA (RAPD) (Link et. al., 1995), amplified fragment length polymorphism (AFLP) ( Duc et al., 2010) and species specific repeats (SSR) (Zeid et. al., 2009). The main limitations of these methods are low reproducibility of RAPD, high cost of AFLP and the necessity to know the flanking sequences to develop species specific primers for SSR polymorphism (Belaj et. al., 2003; Jabbarzadeh, et. al., 2010). Inter-simple sequence repeat (ISSR-PCR) is aroute that overcomes most of these technical limitations (Chen et. al., 2008). ISSR markers have been widely applied to characterize plant germplasm and to demonstrate its effectiveness in assessments of plant genetic diversity (Galvain et. al., 2003, Pharmawati et. al., 2005 and Bhagyawant and Srivastava 2008).

The purpose of the present study was to identify ten faba bean genotypes by using some morphological traits and describe the genetic diversity of these by using Inter-Simple Sequence Repeats (ISSRs) and classify these genotypes.

## **MATERIALS AND METHODS**

Field and laboratory experiments were carried out at Gemmeiza Agriculture Research Station, Gharbia Governorate, and Seed Technology Research Department, ARC, Giza during 2010/2011 and 2011/2012 seasons. Seeds of the studied genotypes were received from Legume Research Dep., Field Crops Research Institute, Agricultural Research Center. The experimental design was a Randomized Complete Block design with three replications. Seeds were inoculated and hand planted. Sowing date was 15<sup>th</sup> November in the first season, while it was 20<sup>th</sup> November in the second season. All the agronomic practices were conducted as recommended and the studied traits were as follows:-

## Morphological characters

Qualitative traits were visually recorded using scales reported by IBPGR (1984). These characters included; anthocyanin coloration, leaf size spots on ears, pinnul shape, color of stem, color of flower ground, lines density on flag flower, wing color of flower, pod corner with stem, pod shape, reversal of pod surface, pod color at maturity, pods division on stem, color of testa, color of Hilum, seed shape and testa shape.

## **Quantitative characters**

These characters included; days from sowing to 50% flowering of plants with at lest one flower and days from sowing to maturity, stem thickness, time of beginning of flowering, time of maturity, highest of the first pod (cm), number of pinnule/ leaf, branching grassroots, number of flowers/ marble, number of pod at nods, number of seeds/ pod, plat height (cm) and seed index. Collected data for each season were statistically analyzed and the Least Significant Difference (L.S.D.) was used to compare among them (Gomez and Gomez, 1984).

#### Molecular markers

#### Plant material and DNA extraction:

DNA was extracted from the tissue of young, healthy leaf which was selected from each genotype, using the DNA extraction kit (Quigen Inc., Cat.no.69104, USA). DNA quality was tested using 1% agarose gel electrophoresis and its concentration was determined spectrophotometrically. **ISSR-PCR analyses** 

Five ISSR primers were selected for testing the genetic diversity between these genotypes. Names and sequences of these primers are shown in Table (1). The PCR reaction was carried out in a 25 µl volume of a mixture containing 25 ng of genomic DNA, 0.1 mM dNTPs, 2.5 mM MgCl2, 1 unit Taq polymerase, 10x Taq buffer and 0.6 µM primer. DNA amplification was carried out using the thermocycler model PTC 200 (MJ Research, Watertown, MA, USA). The amplification program included a denaturing step at 94 °C for 5 min, followed by 35 cycles with a denaturing step at 94 °C for 1 min, an annealing step at 53 °C for 1 min and an extension step at 72 °C for 2 min. After the last cycle, samples were kept at 72 °C for 5 min.

#### Gel electrophoresis

Gel electrophoresis was applied according to Sambrook *et al.* (1989). Agarose (1.2 %) was used for resolving the PCR products. Bands were detected on UV–transilluminator and photographed by Gel documentation 2000, Bio- Rad. Similarity and dendrogram tree was performed using the SPSS program version 10.

Table (1): Names and sequences of the five primers used for ISSR-PCR analyses.

Primer name	Sequence
SH6	5 <sup>'</sup> CGCGATAGATAGATA 3 <sup>'</sup>
SH7	5 <sup>'</sup> GACGATAGATAGATA 3 <sup>'</sup>
SH8	5 <sup>/</sup> AGACAGACAGACGC 3/
SH9	5' GATAGATAGATAGC 3/
SH10	5/ GACAGACAGACAAT 3/

#### RESULTS AND DISCUSSION

Some morphological characters of the different faba bean genotypes are presented in Table (2). The genotypes ( $G_{429}xG_{40}$ ) and ( $H_8xG_{461}$ ) identified with anthocyanin coloration mixture and mixture violet, respectively while in the other faba bean genotypes it was absent, the genotypes ( $G_{Blanka}xG_2$ ) and ( $H_{10}xG_{461}$ ) have weak violet.

Table (2): Some morphological characters of faba bean genotypes (combined dataof 2010/ 2011 and 2011/2012 growing seasons).

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Characters Genotypes	Anthocyanin coloration	Leaf size	Spots on ears	Pinnul shape	Color of stem	Color of flower ground
G <sub>461</sub> x G.Blanka	Absent	Large	Present	Semi Flat	Light brown	Light brown
T.W. x G.Blanka	Absent	Medium	Absent	Round	Brown	Light brown
G <sub>716</sub> x G <sub>402</sub>	Absent	Small	Present	Round	Brown	Light brown
G <sub>461</sub> x G <sub>402</sub>	Absent	Medium	Present	Round	Light brown	White
G.Blanka x G <sub>2</sub>	Weak Violet	Small	Absent	Narrow	Brown	White
G <sub>429</sub> x G <sub>2</sub>	Absent	Medium	Present	Narrow	Light brown	White
G <sub>429</sub> x G <sub>40</sub>	Mixture	small	Present	Narrow	Brown	White
H <sub>8</sub> x G <sub>461</sub>	Medium violet	Large	Absent	Narrow	Very light brown	White
H <sub>10</sub> x G <sub>461</sub>	Weak Violet	Medium	Present	Round	Light brown	White
T.W.G <sub>461</sub> xEgypt <sub>1</sub>	Absent	Large	Present	Round	Dark brown	White

Leaf size can divide the tested genotypes into three groups; small  $(G_{716}xG_{402},\ G_{Blanka}xG_2,\ and\ G_{429}xG_{40}),\ medium\ (T.W.x\ G_{Blanka},\ G_{461}xG_{402}\ and\ G_{429}xG_{40})$  $H_{10}xG_{461}$ ) and large  $(G_{461}xG_{Blanka}, H_8xG_{461}, \text{ and } (T.W.x G_{461})xEgypt1).$ Regarding Spots on ears, they were present in the genotypes (G<sub>461</sub>xG<sub>Blanka</sub>,  $G_{716}xG_{402},\ G_{461}xG_{402},\ G_{429}xG_{2},\ G_{429}xG_{40},\ H_{10}xG_{461}\ and\ (T.W.x\ G_{461})xEgypt_{1}$ and absent in the genotypes (T.W.x G<sub>Blanka</sub>, G<sub>Blanka</sub>xG<sub>2</sub>, and H<sub>8</sub>xG<sub>461</sub>). The genotype (G<sub>461</sub>xG<sub>Blanka</sub>) identified with pinnul shape (semi flat). The genotypes (H<sub>8</sub>xG<sub>461</sub> and (T.W.x G<sub>461</sub>)xEgypt<sub>1</sub>) were identified with color of stem (very light brown and dark brown), respectively. Color of flower ground characters divided faba bean genotypes under studied into two classes, the first class, concluded the genotypes ( $G_{461}xG_{402}$ ,  $G_{461}xG_{Blanka}$ , T.W.x  $G_{Blanka}$  and  $G_{716}xG_{402}$ ) light brown. Meanwhile, the second class (white) was concluded the genotypes ( $G_{461}xG_{402},\ G_{Blanka}xG_2,\ G_{429}xG_2,\ G_{429}xG_{40},\ H_8xG_{461},\ H_{10}xG_{461}$ and (T.W.xG<sub>461</sub>)xEgypt<sub>1</sub>). These results are in agreement with Rehab, Towdy (2007) she reported that, anthocyanin coloration was absent in Sahel 1 while in the other varieties was present, Intensity of anthocyanine coloration was medium in Mesr 1 but was slight for the rest of varieties and showed that, Mesr 1 and Nubaria 1 varieties were short in leaflet size while the other varieties were medium.

Data in Table (3), show some morphological characters of different faba bean genotypes included in this study. The genotype (G<sub>461</sub>xG<sub>Blanka</sub>) identify with lines density on flag flower (dense). Regarding wing color of flower, all faba bean genotypes were black spot. Pod corner with stem can divide the tested genotypes into two groups; the genotypes (G<sub>461</sub>xG<sub>Blanka</sub>,  $G_{429}xG_2$ ,  $G_{429}xG_{40}$ ,  $H_8xG_{461}$  and  $(T.W.x\ G_{461})xEgypt_1$ ) were mixture and the genotypes (T.W.x  $G_{Blanka}$ ,  $G_{716}xG_{402}$ ,  $G_{461}xG_{402}$ ,  $G_{Blanka}xG_2$  and  $H_{10}xG_{461}$ ) were existing. The genotypes under study divided into three class according to pod shape the genotypes (G<sub>461</sub>xG<sub>Blanka</sub>, T.W.x G<sub>Blanka</sub>, G<sub>716</sub>xG<sub>402</sub>, G<sub>429</sub>xG<sub>2</sub>, G<sub>429</sub>xG<sub>40</sub> and  $H_8xG_{461}$ ) have semi-cylindrical while, the genotypes ( $G_{461}xG_{402}$  and (T.W.x G<sub>461</sub>)x Egypt<sub>1</sub>) were narrow extrovert and the genotypes (G<sub>Blanka</sub>xG<sub>2</sub> and H<sub>10</sub>xG<sub>461</sub>) were extrovert non narrow under pod shape. Regarding reversal of pod surface, the genotypes ( $G_{461}xG_{Blanka}$ ,  $G_{429}xG_{2}$ ,  $G_{429}xG_{40}$  and (T.W.xG<sub>461</sub>)x Egypt<sub>1</sub>) have shiny surface and the genotypes (T.W.x G<sub>Blanka</sub>,  $G_{716}xG_{402}$ ,  $G_{461}xG_{402}$ ,  $G_{Blanka}xG_{2}$ ,  $H_8xG_{461}$  and  $H_{10}xG_{461}$ ) have matted pod surface. The genotypes  $(G_{461}xG_{Blanka},\ G_{429}xG_2,\ G_{429}xG_{40},\ H_8xG_{461}$  and  $(T.W.xG_{461})x$  Egypt<sub>1</sub>) can identified with pod color at maturity Dark black, Black, Mixed, White yellow and White brown respectively, while other genotypes were brown.

Table (3): Some morphological characters of faba bean genotypes (combined data of 2010/ 2011 and 2011/2012 growing seasons).

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Characters Lines densition on fla flower		Wing color of flower	Pod corner with stem	Pod shape	Reversal of pod surface	Pod color at maturity	
G <sub>461</sub> xG.Blanka	Dense	Black spot	Mixture	Semi- cylindrical	Shiny	Dark black	
T.W.xG.Blanka	Medium	Black spot	Existing	Semi- cylindrical	Matted	Brown	
G <sub>716</sub> xG <sub>402</sub>	Slight	Black spot	Existing	· · ·		Brown	
G <sub>461</sub> xG <sub>402</sub>	Slight	Black spot	3		Matted	Brown	
G.BlankaxG <sub>2</sub>	Slight	Black spot	Existing	Extrovert non narrow	Matted	Brown	
G <sub>429</sub> xG <sub>2</sub>	Medium	Black spot	Mixture	Semi- cylindrical	Shiny	Black	
G <sub>429</sub> xG <sub>40</sub>	Slight	Black spot	Mixture	Semi- cylindrical	Shiny	Mixed	
H <sub>8</sub> xG <sub>461</sub>	Medium	Black spot	Mixture	Semi- cylindrical	Matted	White yellow	
H <sub>10</sub> xG <sub>461</sub>	Medium	Black spot	Existing	Extrovert non narrow	Matted	Brown	
T.W.G <sub>461</sub> xEgypt <sub>1</sub>	Slight	Black spot	Mixture	Narrow extrovert	Shiny	White brown	

Also some morphological characters of the studied faba bean genotypes are given in Table (4). It is clear that, pods division on stem divided the genotypes under studied into two gropes, the genotypes  $(G_{461}xG_{Blanka},\ T.W.xG_{Blanka},\ G_{716}xG_{402}\ and\ G_{461}xG_{402})$  were both sides of meanwhile, the other genotypes were homogeneous. The genotypes  $(T.W.x\ G_{Blanka}\ and\ H_{10}xG_{461})$  identified with color of testa (White yellow and yellow, respectively). Color of hilum of all genotypes under study was black. Regarding seed shape, the genotypes  $(G_{461}xG_{Blanka},\ T.W.xG_{Blanka},\ G_{716}xG_{402},\ G_{Blanka}xG_2,\ H_8xG_{461},\ H_{10}xG_{461}\ and\ (T.W.xG_{461})xEgypt_1)\ have extrovert and the genotypes <math display="inline">(G_{461}xG_{402},\ G_{429}xG_2)\ and\ G_{429}xG_{40})\ have\ mixture\ shape.$  The genotype  $(G_{429}xG_{40})\ identity\ with\ testa\ shape\ (small\ patches)\ while,\ the\ other genotypes\ were\ fate.\ These\ results\ consent\ with\ Rehab\ tawdy\ (2007),\ she\ found\ that\ color\ of\ testa\ was\ beige\ in\ all\ the\ varieties\ of\ faba\ bean\ under\ studies\ except\ for\ G.717\ which\ was\ semi\ beige\ green.$ 

Table (4): Some morphological characters of faba bean genotypes (combined data of 2010/2011 and 2011/2012 growing seasons).

Pods division on stem	Color of testa	Color of Hilum	Seed shape	Testa shape
Both sides of	Dark green	Black	Extrovert	Fate
Both sides of	White yellow	Black	Extrovert	Fate
Both sides of	Wight green	Black	Extrovert	Fate
Both sides of	Mixed	Black	Mixture	Fate
Homogeneous	Wight brown	Black	Extrovert	Fate
Homogeneous	Wight brown	Black	Mixture	Fate
Homogeneous	Mixed	Black	Mixture	small
				patches
Homogeneous	Wight brown	Black	Extrovert	Fate
Homogeneous	Yellow	Black	Extrovert	Fate
Homogeneous	Dark green	Black	Extrovert	Fate
	Both sides of Homogeneous Homogeneous Homogeneous Homogeneous	Both sides of Dark green Both sides of White yellow Both sides of Wight green Both sides of Mixed Homogeneous Wight brown Homogeneous Mixed  Homogeneous Wight brown Homogeneous Wight brown Homogeneous Wight brown Homogeneous Wight brown Homogeneous Yellow	Both sides of Dark green Black Both sides of White yellow Black Both sides of Wight green Black Both sides of Mixed Black Homogeneous Wight brown Black Homogeneous Mixed Black Homogeneous Wight brown Black Homogeneous Wight brown Black Homogeneous Wight brown Black Homogeneous Wight brown Black Black Homogeneous Wight brown Black Homogeneous Wight brown Black	stemHilumBoth sides ofDark greenBlackExtrovertBoth sides ofWhite yellowBlackExtrovertBoth sides ofWight greenBlackExtrovertBoth sides ofMixedBlackMixtureHomogeneousWight brownBlackExtrovertHomogeneousWight brownBlackMixtureHomogeneousMixedBlackMixtureHomogeneousWight brownBlackExtrovertHomogeneousYellowBlackExtrovert

Combined data of the quantitative characters of the studied faba bean genotypes are presented in Tables (5 and 6). It is clear that, two characters namely height of the first nod and time of beginning of flowering were to some exent effective tools to differentiate between genotypes. Whereas as studied characters were almost similar. The genotype ( $H_8 \times G_{461}$ ) gave the highest stem thickness (0.9 cm) and seed index (89.5 gm). The genotype ( $G_{429} \times G_{40}$ ) recorded maximum of the first nod (31.1 cm) while, the genotype ( $H_8 \times G_{461}$ ) recorded the lowest height of the first nod (24.5 cm). The genotype ( $G_{429} \times G_{420} \times G_{42$ 

shortest pod length was noticed by (G<sub>461</sub>xG.Blanka). Concerning time of beginning of flowering and maturity the genotype (T.W.G<sub>461</sub>xEgypt<sub>1</sub>) was the latest among all the genotypes (55 and 156 days from begging planting) respectively, while the genotype (H<sub>10</sub> x G<sub>461</sub>) was the earliest for beginning of flowering and maturity 46 and 152 days from begging planting, respectively comparing with the other genotypes. The genotypes (G.Blanka x  $G_2$  and  $H_{10}$ x G<sub>461</sub>) recorded highest No. of pinnule/ leaf (5 pinnule) but, the genotype (G<sub>461</sub> x G<sub>402</sub>) have little No. of flowers/ marble (3 flowers) comparing with the other genotypes. No. of seeds/ pod was (4 seeds) at (G<sub>429</sub> x G<sub>40</sub> and T.W.G<sub>461</sub>xEgypt<sub>1</sub>) genotypes, while it was (3 seeds) at all the other genotypes. Similar results were in agreement with Ibrahim (2010) and Al Barri and Mungez (2013), they revealed that VF-14, VF-10 and VF-12 lines significantly longer days to flowering and differed from all the other genotypes, VF-8, VF-12 and VF-13 lines were significantly the tallest plants while, VF-6 and VF-7 were the shortest, VF-11 line showed the lowest average of pod height and VF-19 gave the highest value, VF-4 and VF-10 lines showed the highest pod length, while VF-2 genotype gave the lowest pod length, VF-10 line significantly had the highest average seed number per pod while VF-7 and VF-19 significantly showed the lowest average seed number per pod and VF-4 genotype gave significantly the highest 100-seed weight while, VF-17 and VF-2 gave the lowest 100-seed weight.

Table (5): Some quantitative characters of faba bean genotypes (combined data of 2010/ 2011 and 2011/2012).

Characters  Genotypes	Stem thickness (cm)	Height of the first nods (cm)	Time of beginning of flowering	Time of maturity	No. of pinnule/ leaf
G <sub>461</sub> xG.Blanka	0.8	26.5	48	154	4
T.W.xG.Blanka	0.8	29.0	47	154	4
G <sub>716</sub> x G <sub>402</sub>	0.7	25.5	49	155	4
G <sub>461</sub> x G <sub>402</sub>	0.8	26.6	48	153	4
G.Blanka x G <sub>2</sub>	0.8	29.4	50	154	5
G <sub>429</sub> x G <sub>2</sub>	0.8	26.9	47	153	4
G <sub>429</sub> x G <sub>40</sub>	0.8	31.1	47	153	4
H <sub>8</sub> x G <sub>461</sub>	0.9	24.5	50	153	4
H <sub>10</sub> x G <sub>461</sub>	0.8	26.6	46	152	5
T.W.G <sub>461</sub> xEgypt <sub>1</sub>	0.7	29.1	55	156	4
L.S.D. at 5%	0.1	2.7	0.95	1.45	1.0

Table (6): Some quantitative characters of faba bean genotypes (combined data of 2010/ 2011 and 2011/2012).

Characters Genotypes	No. of flowers/marble	No. of pods at nods	No. of seeds/ pod	Pod length (cm)	Plant height (cm)	Seed ় index (gm)
G <sub>461</sub> xG.Blanka	4	2	3	8.6	100.3	83.8
T.W.xG.Blanka	4	2	3	9.4	99.6	87.5
G <sub>716</sub> x G <sub>402</sub>	4	1	3	9.2	98.6	79.5
G <sub>461</sub> x G <sub>402</sub>	3	2	3	9.1	102.0	87.2
G.Blanka x G <sub>2</sub>	5	2	3	9.9	96.8	83.6
G <sub>429</sub> x G <sub>2</sub>	4	1	3	9.0	102.5	83.8
G <sub>429</sub> x G <sub>40</sub>	5	1	4	9.5	96.0	82.3
H <sub>8</sub> x G <sub>461</sub>	5	2	3	8.9	101.5	89.5
H <sub>10</sub> x G <sub>461</sub>	5	2	3	9.0	98.0	87.7
T.W.G <sub>461</sub> xEgypt <sub>1</sub>	4	1	4	8.9	96.3	85.2
L.S.D. at 5%	1.0	1.0	1.0	0.6	4.7	1.9

#### Molecular marker

In the present study, five selected primers of ISSR were used to differentiate between the ten faba bean genotypes (Figure 1). Inter-Simple Sequence Repeat (ISSR) technique yield more polymorphisms than other molecular techniques. The five primers amplified different numbers of bands and revealed various levels of polymorphism. A total of 71 ISSR loci were observed and 59 (83.1%) of them were polymorphic. these primers yielded 14, 11, 18, 14 and 14 bands, respectively (Table 7). The percentage of polymorphism was 92.9 %, 90.9 %, 72.2 %, 85.7 % and 78.6 %, respectively. The primer SH8 yielded the largest number of bands (18 band) while, primer SH7 had fewer number of bands. Among all the ISSR loci observed, 11 were unique. The highest number of unique band can observed in primer SH8 which produced four markers. While, the lowest number can observed in primer SH7 and SH10 which produced one marker. The ISSR primer method is reported to produce more complex marker (Parsons et. al., 1997 and Chowdhury et. al., 2002), which is advantageous when differentiating closely related cultivars.

Table (7): Levels of polymorphism and unique genotypes specific bands for ten faba bean genotypes by five ISSR primers.

Bands	Total	Polymorphic	Monomorn	Polymorphism	Unique k	pands	
Primer	bands	bands	hic bands	%	Genotypes	Ms (bp)	
					3	938	
SH6	14	13	1	92.9	5	341	
					9	228	
SH7	11	10	1	90.9	3	800	
					9	1493	
SH8	18	13	5	72.7	8	927	
					3	644	
					6	266	
SH9	14	12	2	85.7	5	681	
					4	556	
SH10	14	11	3	78.6	2	729	
Total	71	59	12	83.1			

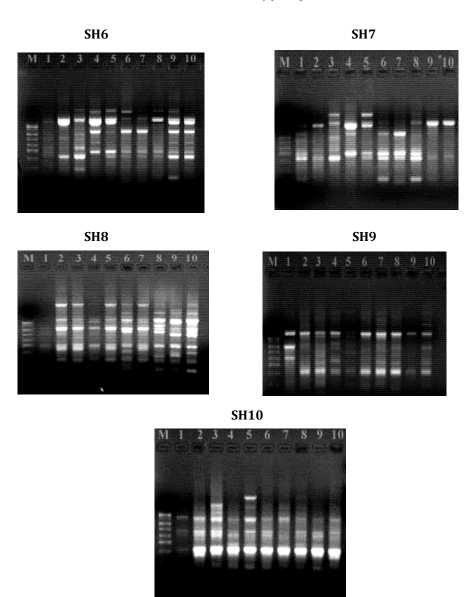


Figure (1): ISSR fingerprinting of ten faba bean genotypes, Lines from left to right: M= Marker,  $G_{461}xG_{Blanka}$ , T.W. $xG_{Blanka}$ ,  $G_{716}xG_{402}$ ,  $G_{461}xG_{402}$ ,  $G_{Blanka}xG_2$ ,  $G_{429}xG_2$ ,  $G_{429}xG_{40}$ ,  $H_8xG_{461}$ ,  $H_{10}xG_{461}$  and (T.W. $xG_{461}$ )x Egypt<sub>1</sub>.

## **Genetic diversity**

Standard genetic distances could be estimated between all the genotypes showed in Table (8). The genetic similarity ranged from 81% to 53% with an average of 67%. The closest distance was found between the genotypes ( $G_{429}xG_2$ ) and (T.W.x  $G_{461}$ )x Egypt<sub>1</sub>) and also between ( $H_{10}xG_{461}$ )

and  $(T.W.x \ G_{461})x \ Egypt_1$  (81%), while the longest was between the genotypes  $(G_{716}xG_{402})$  and  $(H_{10}xG_{461}$  (53%). The results of clustering genetic distance using the unweighted pair group method separated genotypes to two main clusters with many subclusters, where, the genotype1  $(G_{461}xG_{Blanka})$  was in a separate subcluster, while, the other genotypes were in the second subcluster (Figure 2). Salem *et. al.*, (2011) subjected Thirty-four faba bean (*Vicia faba* L.) to molecular diversity assessment using 12 inter-simple sequence repeat primers and found that there is a high genetic variability related to collection sites and it should be utilised in faba bean improvement. Also, Abdel-Razzak *et. al.*, (2012) clarified that ISSR markers and protein analysis were helpful to recognize genetic variation among faba bean cultivars.

Table (8): Similarity matrix among ten faba bean genotypes based on ISSR analysis.

on 1991 analysis.									
Genotypes	G <sub>461</sub> X G <sub>Blanka</sub>	T.W.x G <sub>Blanka</sub>	G <sub>716</sub> x G <sub>402</sub>	G <sub>461</sub> x G <sub>402</sub>	G <sub>Blanka</sub> x G <sub>2</sub>	G <sub>429</sub> x G <sub>2</sub>	G <sub>429</sub> x G <sub>40</sub>	H <sub>8</sub> x G <sub>461</sub>	H <sub>10</sub> x G <sub>461</sub>
G <sub>461</sub> xG <sub>Blanka</sub>									
T.W.x G <sub>Blanka</sub>	0.64								
G <sub>716</sub> xG <sub>402</sub>	0.64	0.74							
G <sub>461</sub> xG <sub>402</sub>	0.63	0.60	0.61						
$G_{Blanka}xG_2$	0.56	0.69	0.63	0.62					
G <sub>429</sub> xG <sub>2</sub>	0.70	0.64	0.61	0.71	0.72				
G <sub>429</sub> xG <sub>40</sub>	0.69	0.74	0.73	0.70	0.68	0.77			
H <sub>8</sub> xG <sub>461</sub>	0.65	0.68	0.72	0.69	0.69	0.78	0.79		
H <sub>10</sub> xG <sub>461</sub>	0.60	0.61	0.53	0.69	0.66	0.73	0.60	0.68	
(T.W.x G <sub>461</sub> )x Egypt <sub>1</sub>	0.66	0.74	0.61	0.70	0.68	0.81	0.76	0.72	0.81

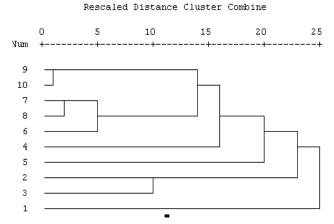


Figure (2): Dendrogram of the genetic distances among the ten faba bean genotypes based on ISSR analysis.

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التمييز المورفولوجى والجزيئى لبعض التراكيب الوراثية من الفول البلدى أحمد عبد اللطيف محمد الإمام\*، السيد محمد ربيع\*، عزيزة محمد حسنين\* ومجدى ابراهيم العبادى\*

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تهدف هذه الدراسة الى استخدام بعض الصفات المور فولوجية والجزيئية لتمييز عشرة تراكيب وراثية من الفول البلدى من خلال أجراء بعض التجارب الحقلية والمعملية بمحطة البحوث الزراعية بالجميزة ومعامل قسم بحوث تكنولوجيا البذور بالجيزة – مركز البحوث الزراعية ، خلال موسمي ٢٠١١/٢٠١، الوريقة قسم بحوث تكنولوجيا البذور بالجيزة – مركز البحوث الضفات المور فولوجية مثل شكل الوريقة الوسطى، كثافة الشرائط (الخطوط) على علم الزهرة، لون القرن عند النضج وشكل ولون قصرة البذرة كانت الوسطى، كثافة الشرائط (الخطوط) على علم الزهرة، لون القرن عند النضج وشكل ولون قصرة البذرة كانت أمكن تمييز الاختلافات الوراثية بين التراكيب الوراثية الني اشتملت عليها الدراسة على المستوى الجزيئي وذلك باستخدام خمسة من البادنات العشوانية وتكنيك ISSR-PCR. وقد بلغت نسبة التشابه من (٥٠-١٨%) بمتوسط قدرة (٦٧%). والنتائج المتحصل عليها من هذه الدراسة ذات أهمية كبيرة في حفظ حقوق مربو النباتات عند تسجيل التراكيب الوراثية كأصناف تجارية جديدة إلا أنة علي مربي النبات الانتخاب من قاعدة وراثية عريضة حتى يمكن الحصول علي صفات مور فولوجية مميزة للسلالات الجديدة عن الأصناف المنزرعة المسجلة وذلك عند تسجيلها كأصناف جديدة مما يسهل التحقق من نقاوة الصنف الجديد أثناء مراحل اكثارة المختلفة.