# ACCEPTED ARTICLE Genetic Divergence for Different Yield Attributing Traits in Okra [Abelmoschus Esculentus (L.) Moench] Genotypes Grown in Himalayan Foothills Region Sudheer Kumar Yadav<sup>1</sup>, Udit Kumar<sup>1</sup>, K. Prasad<sup>2\*</sup>, Shubham Maurya<sup>1</sup>, Neetu Saroj<sup>1</sup> Department of Horticulture, Post-Graduate College of Agriculture, Dr. Rajendra Prasad Central Agricultural University, Pusa, Bihar-143121, India.

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13 ABSTRACT

The Himalayan foothills region of India is rich in the genetic diversity of okra which is yet to 14 be explored for its genetic divergence. To envisage the genetic diversity of this un-explored 15 varietal collection, the genetic divergence among 25 genotypes of okra was estimated using 16 Mahalanobis  $D^2$  statistic. The indigenous and exotic lines were grouped into 6 clusters using 17 Tocher's methods. Results revealed that a higher number of genotypes were recorded under 18 19 Cluster I (19) and cluster IV (2) while cluster II, III, V and VI were mono-genotypic. A higher intra- cluster distance was observed between clusters I (13.42) & IV (7.47), whereas, a higher 20 21 inter-cluster distance was found between clusters III and VI (111.03). The traits viz, YVMV incidence (44.67 %) was contributing the highest towards the total genetic divergence. The 22 present study revealed the detailed genetic divergence for different yield-attributing traits in 23 okra. This study presents a strong basis for the selection and evolving of better recombinants 24 for hybridization and quality improvement programme. This research bear utility in the form 25 of germplasm conservation and crop improvement for selected indigenous/exotic genotypes 26 grown in Himalayan foothills. 27

Key words: Okra, Himalayan foothill genotypes, divergence, cluster and recombinants.

# 1. Introduction

Okra [*Abelmoschus esculentus* (L.) Moench] is the most significant nutritive vegetable crop in India (Temam *et al.*, 2021). During the rainy and spring-summer seasons, okra is predominantly grown in the tropics and subtropics of the world for immature fragile green fruits as well as processed products (Temam, N., 2020). The okra crop, which belongs to the Malvaceae family and the genus *Abelmoschus*, is said to have originated in Ethiopia (Mohammed *et al.*, 2022). Although according to Sonwani *et al.*, 2022 it originated in the

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Hindustan region. Okra n=12 has a regular series of polyploides with somatic chromosomal
numbers 2n=72,109,120,132, and 144 (Raghuwanshi *et al.*, 2019).

The root and stems of the okra plant are commonly used to clarify sugarcane juice for the 39 manufacturing of brown sugar and gur (Makur et al., 2019). It is also utilised in the paper 40 industry due to the high mucilage and crude fibre content of the mature stems and fruits 41 (Haruna et al., 2016). Okra has a great medicinal value and is particularly useful in the 42 treatment of genito-urinary disorders such as haemorrhoids, ulcers, chronic dysentery, and 43 spermatorrhoea (Olaniyan and Omoleyomi, 2013). Okra fruits are rich in high iodine content 44 45 which helps to prevent goitre (Fayaz et al., 2022). Fresh okra vegetable plays a very crucial role in human nutrition, especially as sources of dietary fiber & vitamins (Prasad et al., 2018). 46 47 The tender green fruit is the potent source of vitamins A, B<sub>1</sub>, B<sub>3</sub>, B<sub>6</sub>, C, folic acid and essential minerals viz., Ca, K, Mg and P have an important role in human diet (Mkhabela et al., 2022). 48

49 The development of cultivars against target traits such as fruit yield and pest/disease resistance and their promotion is often a more rewarding and appropriate option for the 50 51 sustainability of smallholder farmers (Sandeep et al., 2022). This is especially relevant in developing and underdeveloped countries, where farmers often do not have the capability to 52 diagnose suitable preventive measures (Kumar et al., 2010). Yellow vein mosaic disease is the 53 most common biotic stress in okra under natural epiphytic conditions. The wild species are 54 widely acknowledged to be a major reservoir of resistance genes, particularly for yellow vein 55 mosaic disease (Gangopadhyay et al., 2017). 56

The concept of linking phenotypes with genotypes is somewhat basic and ubiquitous in plant 57 breeding, as well as in the history of plant domestication, which was accomplished through the 58 selection of superior plants for agricultural use (Kozak et al., 2011). Improving the varieties 59 with desired traits in terms of fruit size, shape and colour and resistance to biotic stress are 60 also very purposeful in the okras' export market. This characterization and genetic diversity 61 analysis was required against this backdrop. Traditionally, a combination of morphological 62 data and phenotypic traits significantly contributed towards the pattern of total genetic 63 64 diversity present in the population (Oppong-Sekyere et al., 2011). The genetic distances between the lines, determined by using these traits, can be treated as phenotypic similarities 65 between the lines (Kozak et al., 2011). 66

67 Genetic diversity refers to the heritable variation within and between groups of populations 68 (Alam *et al.*, 2014). The presence of genetic diversity between the genotypes from different 69 geographic regions is attributed to the selection and exchange of okra germplasm between farmers from different geographic locations and between ethnic groups. The duplications in the germplasm due the migrant farmers often carry seeds from their original growing location to their new growing locations (Oppong-Sekyere *et al.*, 2011). Genetic diversity exploits essential work in vegetable breeding because hybrids from diverse sources of germplasm exhibited a more opportunity for  $F_1$  hybrids development than closely related species (Alam *et al.*, 2014).

Genetic divergence has been an important topic in plant breeding in recent decades, and still 76 is. It should be defined as the divergence of the gene pool of a population from the gene pools 77 78 of other populations, which can occur as a result of mutation, genetic drift, and selection. The basis of morphological and genotypic divergences or similarities could be viewed as a 79 80 complicated genetic system (Kozak et al., 2011). Differences across the varieties of the same or different species may be linked to allelic variation and variable gene expression associated 81 82 with morphological and non-physiological features, in addition to the genotype-byenvironment interaction of each locus (Kozak et al., 2011 and Kyriakopoulou et al., 2017). 83

The D<sup>2</sup> statistics provides a magnitude estimate of diversity between two genotypes (Sruthi *et al.*, 2020). In okra, F<sub>1</sub> hybrids are more widespread, and new genotypes for heterosis are constantly being selected (Sandeep *et al.*, 2022). As a result, genetic divergence among existing genotypes must be investigated, and germplasm must be collected for identification (Sanwal *et al.*, 2012). When identifying relevant parental lines for heterosis breeding, grouping genotypes based on D<sup>2</sup> analysis will be advantageous (Mohammed *et al.*, 2022).

The clustering pattern indicated the existence of diverse forms in collections made from the close geographic location, indicating more opportunity to select the most desirable genotypes (Oppong-Sekyere *et al.*, 2011, Gangopadhyay *et al.*, 2017). A cluster diagram obtained from the phynotypic descriptors based on the simple matching coefficient (Oppong-Sekyere *et al.*, 2011).

The Pusa region of Bihar, India, represents Himalayan foothill Plain, and is well known for its distinctive soil, slope, altitudes, climate, and ecological diversity of okra germplasm which are yet un-explored for diversity trait in this region (Koku *et al.*, 2020). The genetic diversity of okra in this region is very high (Sonwani *et al.*, 2022). There is a necessity of developing high-yielding, better nutrition, post-harvest quality and biotic/abiotic resistant okra germplasm for this region.

101 The objective of the present study is to put a strong basis for the selection and evolving of 102 better recombinants for hybridization and quality improvement programme in the near future (Prasad *et al.*, 2018; Mkhabela *et al.*, 2022). This research also aims for germplasm
conservation and crop improvement for selected indigenous/exotic genotypes grown in
Himalayan foothills.

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# 2. MATERIALS AND METHODS

The research was carried out during academic session 2019-2020, at Dholi Farm, Tirhut 108 109 College Agriculture, a campus of Dr. Rajendra Prasad Central Agricultural University, Pusa, Samastipur- Bihar. Twenty-five genotypes of okra (Table 1), including both indigenous and 110 111 exotic lines were selected and sown on 25 Jun 2019 in randomized block design with each line consisting of three replications and last observation was taken on plant height at final harvest 112 113 on 20 September 2019. All the genotypes were procured from Indian Institute of Vegetable Research (IIVR), Varanasi, Uttar Pradesh-India, through materials transfer agreement (Table 114 1). A nationally adapted okra variety "Kashi Kranti" was also included as a check variety. A 115 standardized crop descriptor for okra genotypes was used to measure the various traits under 116 studies (IBPGR, 1991). The data was recorded from five randomly selected plants for different 117 yield attributing traits. The weather data were collected from Department of Agro 118 Meteorology, Tirhut College Agriculture, a campus of Dr. Rajendra Prasad Central 119 Agricultural University, Pusa, Samastipur-Bihar. The weather data was tabulated in (Table 2). 120 The Enlist of the studied phenotypic traits under present study were mention in (Table 3). The 121 description of studied phenotypic traits were summarized below on the following heads. 122

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## (A). Plant height (cm)

The measurement of plant height of all five selected and tagged plants in each treatment was done using a meter scale, from ground level to the apex of the plant at the final harvest. The measured value was presented in centimetre.

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## (B). Number of primary branches per plant

Primary branches per plant in each genotype of each plot were counted at the time offinal harvesting and the average branches per plant were determined.

# (C). Days to first flowering

The number of days from seed sowing to the anthesis of the first flower was counted.

138	(D). Days to first picking
139	The cumulative number of days from seed sowing to harvesting of the first fruit was
140	recorded.
141	
142	(E). Number of seeds per fruit
143	Five dried fruits were randomly collected from tagged plants in each plot and seeds were
144	extracted. The number of seeds were counted for the average.
145	
146	(F). Number of ridges per fruit
14/	Ridges per fruit were counted during the third pickings of each genotype in each plot and the
148	average number of ridges per fruit were calculated.
149	(G). Fruit length (cm)
150	The marketable fruits of each genotype were selected in each plot and the length of these
151	fruits, excluding fruit stalk, was measured with meter scale and the mean length per pod was
152	calculated.
153 154	(H). Fruit diameter (cm)
155	Five fruits of each genotype were randomly selected in each plot and diameter was measured
156	at the base of fruits with the help of venire caliper.
157	
158	(I). YVMV Incidence (%)
159	Under natural disease pressure conditions, all the plants in each plot were recorded for the
160	appearance of YVMV incidence percentage. Numbers of plants affected by YVMV were
161	counted at 30, 45, 60 and 75 days after sowing of crop in each genotype. The percent
162	incidence was determined based on the total number of plants per plot.
163	YVMV infestation of plants (%) = <u>Number of YVMV infected plants <math>\times</math> 100</u>
164	Total number of plants
165	(J). Average fruit weight (g)
166	Five edible fruits were selected randomly from tagged plants in each plot and their weight
167	(g) was taken using weighing balance.
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1/0	The number of fruits picked from all tagged plants in each picking were added and it
171	measured the total number of fruits per plant.
	5

172	(L). Fruit yield per plant (g)
173	The fruit yield per plant of the tagged plants was determined by addition of the total fruit
174	weight of all fruit pickings and is shown in grams per plant.
175	2.1. Genetic divergence
176	Genetic diversity was determined by Mahalanobis D <sup>2</sup> statistics based on their phenotypic
177	data (Sruthi et al., 2020; Kumar et al., 2021; Vaggar et al., 2022 and Meena et al., 2021).
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179	2.2. Cluster Analysis
180	The genotypes are grouped into various clusters based on their phenotypic similarity (Table-
181	3). $D^2$ values and Tocher's method were used for grouping of genotypes into different clusters
182	(Kumawat et al., 2020, Kumar et al., 2021; Vaggar et al., 2022).
183 184	2.3 Average intra and inter-cluster distance was used to figure out the summation of
185	distances between different genotypes
186	
187	2.4. Cluster diagram using "Dendrogram" is used to exhibit inter and intra cluster
188	distances between different clusters-based values of $D^2$ statistics helped in drawing a diagram.
189	(Yadav, 2020).
190	
191	2.5. Contribution of individual traits towards total divergence
192	Among all, the combination of genotypes <i>i.e.</i> n $(n-1)/2$ , each trait is ranked based on mean
193	difference, where n is the total number of genotypes. Using these ranks, the following table is
194	prepared to work out the percent contribution of each trait to the total divergence (table.5).
195	Percent Contribution by $Xp = Number of times appearing first in ranking by Xp \times 100$
196	n (n-1)/2
197	Where, Xp= individual traits
198	n= number of genotypes
199 200	2.6. Statistical analysis
201	The statistical analysis was carried out using Statistical Analysis System [SAS (9.4)] North
202	Carolina State University- United States America, OPSTAT (Operational Statistics-

tisticsdeveloped by Department Of Mathematics Statistics, Chaudhary Charan Singh Haryana 203 Agricultural University, Hisar, India) and GRAPES software (General R-shiny based 204

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#### 1. RESULTS AND DISCUSSION

209 The mean performance of morphological traits of different genotypes along with check variety (Kashi Kranti) is tabulated in (Table 4). The genotype SKY/DR/RS-13 (136.33 cm & 210 211 3.10) and IC- 43742 (133.07cm & 3.08) was reported for significantly higher plant height and number of primary branches per plant than the check value (116.67 cm & 2.30). The 212 213 genotypes VRO-214 (39.03 days & 45.33 days) were observed significantly at par near the check variety (40.57 days & 47.33 days) with respect to days to first flowering and days to 214 215 first picking. The genotypes namely IC-218872 (68.67) was observed for maximum number of seeds per fruit and VRO-177 (6.87) was recorded significantly superior for maximum number 216 of ridges per plant over the check variety (48.33 & 5.00). The significantly higher fruit length 217 were reported in EC- 015537 (14.53cm), EC-199367 (14.27cm), and VRO-107 (13.83cm), 218 fruit diameter was recorded higher in IC-43742 (1.81cm) than check variety (11.77 cm & 1.60 219 cm). The minimum YVMV incidence (%) was registered in genotypes EC- 199367 (9.33 %) 220 as compare to check variety (17.33 %). The significantly superior genotypes over check 221 variety, Kashi Kranti (11.22 g, 23.67 and 265.67 g) was recorded in VRO-956 (15.63 g) 222 followed by VRO-214 (13.18 g) and EC-199367 (12.91 g) with respect to average fruit 223 224 weight, IC-43742 (27.50) and EC-199367 (26.70) with respect to maximum number of fruits per plant, IC-43742 (312.33 g) followed by VRO-214 (320.33 g) and EC- 199367 (319.33 g) 225 for higher fruit yield per plant. In the consideration of per se performance of genotype existed 226 vast range of variation for different morphological traits among genotypes of okra under 227 228 present investigation indicated the evaluation of these genotypes in above respective morphological traits can be more useful in future breeding programme. The similar type 229 results were also reported by some previous researchers (Vaggar et al., 2022). 230

Analysis Platform Empowered by Statistics developed by Department of Agricultural

Statistics, College of Agriculture, Vellayani, Kerala Agricultural University- India).

In general, all the okra genotypes displayed relatively wide ranges of variation for all morphological traits evaluated (Oppong-Sekyere *et al.*, 2011). Higher diverse genotypes within the group of population exhibit better heterotic response (Suri *et al.*, 2022). The  $D^2$ analysis method is used to group the genotype based on quantitative traits. The genotypes were grouped in different clusters based on the genetic distance between them.

The 'cluster patterns', 'clusters mean', 'intra/inter cluster' and 'contribution percentage' were presented in Tables 5, 6, 7 and 8 respectively. The selected (25) genotypes were assessed for genetic divergence followed by division in 6 groups as per the 'Toucher' method (Table

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239 2). Cluster I had the maximum genotypes (19) followed by cluster IV (2), whereas, Clusters II,
240 III, V, and VI were monogenotypic, *i.e.* consisting of a single genotype. Our result got the
241 support from similar studies attempted earlier by researchers who worked on okra genetic
242 divergence (Prasad *et al.*, 2016; Alemu and Mohammed 2022; Carvalho *et al.*, 2022, and
243 Mohammed *et al.*, 2022).

Cluster means for twelve characters of twenty-five genotypes are presented in (Table 3). It 244 was recorded that the maximum cluster mean value for plant height (133.07cm), number of 245 primary branches per plant (3.10), fruit diameter (1.81 cm) and number of fruits per plant 246 247 (26.00) was recorded in cluster III. It indicates that if a breeding aim is to obtain long plant height, more number of primary branches per plant, fruit diameter and number of fruits per 248 249 plant then genotypes from these clusters would be selected. The minimum cluster mean value for days to first flowering (40.33 days), days to first picking (47.33 days), number of seeds per 250 251 fruit (28.33), fruit diameter (1.46 cm) and YVMV incidence (9.33%) was reported in cluster III, cluster II, cluster VI, cluster VI and cluster II respectively. It indicates that if a breeding 252 253 program aims to obtain desired days to first flowering, days to first picking, numbers of seeds per fruit, fruit diameter and YVMV incidence then genotypes from these clusters would be 254 255 selected. The maximum cluster mean value for number of seeds per fruit (63.00) and number of ridges per fruit (6.67) was observed in cluster IV and fruit length (14.27cm) and fruit yield 256 per plant (319.33g) was reported in cluster II and average fruit weight (15.63g) was found in 257 cluster V respectively. It reveals that if a breeding program is aimed to obtain desired number 258 of seeds per fruit, number of ridges per fruit, fruit length, fruit yield per plant and average fruit 259 weight, then genotypes from these clusters would be selected. During the investigations, the 260 identified clusters revealed crucial values for various factors under study. Among the clusters 261 (I, II, III, IV, V & VI), the diverse genotypes were compared for higher plant height, early 262 flowering, early picking & fruit yield per plant. Our results find support from the recent 263 reports on genetic divergence of okra crop (Kumar et al., 2021; Vaggar et al., 2022; Alemu 264 and Mohammed 2022 and Mohammed et al., 2022). 265

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Average distance in intra-cluster was varied from 7.47 to 13.42 (Table 4). However, cluster number (II, III, V and VI) had zero intra-cluster distance (0.00) indicated as these clusters were monogenotypic *viz.*, having a single genotype while, cluster number I (13.42) and cluster number IV (7.47) indicated maximum intra-cluster distance. The average inter - cluster distance ranged from 14.07 to 111.03. The maximum inter cluster distance reported between cluster number III and cluster number VI (111.03) followed by cluster II and cluster number

VI (90.76), cluster number IV and cluster number VI (90.10), cluster number III and cluster 272 number V (59.44), cluster number I and cluster number VI (48.80), cluster number II and 273 cluster number IV (48.75), cluster number IV and cluster number V (45.24), cluster number II 274 and cluster number V (40.40), cluster number III and cluster number IV (37.75), cluster 275 number V and cluster number VI (33.34). While, minimum inter- cluster distance reported 276 between cluster number II and cluster number III (14.02) followed by cluster number I and 277 cluster number II (23.07), cluster number I and cluster number V (23.63), cluster number I and 278 cluster number III (28.07) and cluster number I and cluster number IV (29.70). Similar 279 findings were reported by Carvalho et al. (2022), Alemu and Mohammed (2022) and 280 Mohammed et al. (2022) while working on cluster distance for genetic divergence of okra 281 282 crop.

The contribution of each character to 'overall genetic divergence' is presented in (Table 5). YVMV incidence (44.67 %) exhibited the highest contribution to the genetic divergence among all the studied traits. As yellow vein mosaic virus is a threat to okra cultivation all over India, the higher percentage of its contribution to the 'overall genetic divergence' could be an opportunity to identify the genotype immunity/resistance to it (Carvalho *et al.*, 2022).

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#### 2. CONCLUSIONS

Present study revealed the detailed genetic divergence for different yield attributing 290 traits in okra genotypes grown in Himalayan foothills. This study presents a strong basis for 291 the selection and evolving of better recombinants for hybridization and quality improvement 292 293 programmes. Based on the present investigation it can be concluded that for the character multiple branches per plant, genotype can be selected from cluster III, for the development of 294 295 hybrids. Similarly, to improve high yield per plant with lowest incidence of yellow vein mosaic virus, genotypes from cluster II are ideal for crossing or their derivatives for further 296 297 selection. This research bears utility in the form of germplasm conservation and crop improvement for selected indigenous/exotic genotypes grown in Himalayan foothills. 298

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<i>S. No.</i>	Genotypes	Source	<i>S. No.</i>	Genotypes	Source
1.	IC-43735	IIVR-Varanasi, India	14.	VRO-196	IIVR-Varanasi, India
2.	IC-43742	IIVR-Varanasi, India	15.	VRO-177	IIVR-Varanasi, India
3.	IC-218872	IIVR-Varanasi, India	16.	VRO-956	IIVR-Varanasi, India
4.	IC-105667	IIVR-Varanasi, India	17.	VRO-433	IIVR-Varanasi, India
5.	EC-199367	IIVR-Varanasi, India	18.	VRO-214	IIVR-Varanasi, India
6.	EC-015537	IIVR-Varanasi, India	19.	VRO-320	IIVR-Varanasi, India
7.	SKY/DR/RS-13	IIVR-Varanasi, India	20.	VRO-107	IIVR-Varanasi, India
8.	SKY/DR/RS-107	IIVR-Varanasi, India	21.	VRO-173	IIVR-Varanasi, India
9.	VRO-454-10-1	IIVR-Varanasi, India	22.	VRO-404	IIVR-Varanasi, India
10.	VRO-232-10-1	IIVR-Varanasi, India	23.	VRO-40	IIVR-Varanasi, India
11.	VRO-132-10-1,2	IIVR-Varanasi, India	24.	VRO-37	IIVR-Varanasi, India
12.	VRO-304-10-1	IIVR-Varanasi, India	25.	Kashi Kranti	IIVR-Varanasi, India
13.	VRO-109-1	IIVR-Varanasi, India			

**Table 1.** List of genotypes with their source of procurement for the study.

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\*IIVR: Indian Institute of Vegetable Research, Varanasi, Uttar Pradesh- India. https://iivr.icar.gov.in/.

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Table 2. Average monthly weather data on the field location during the okra growing season.

Month	Tempera	Temperature ( <sup>0</sup> C)		Humidity (%)		Rainfall (mm)	Evaporation
	Max.	Min.	Morning	Evening	<b>h</b> <sup>-1</sup> )		( <b>mm</b> )
Jun-2019	36.7	26.3	81	57	6.3	0.0	6.3
July-2019	30.4	25.2	92	76	4.2	138.7	4.2
August-2019	33.6	26.3	90	76	5.2	100.7	5.2
September-2019	32.4	25.9	89	80	2.8	74.2	2.8

414 Source: Dept. of Agro Meteorology, Tirhut College Agriculture, a campus of Dr. Rajendra Prasad Central Agricultural University, Pusa, Samastipur, Bihar- India.

S. No.	Traits	Sampling date				
1.	Plant height					
2.	Number of primary branches per plants					
3.	Days to first flowering					
3.	Days to first picking					
5.	Number of seeds per fruits	25 Jun 2019 to 20 September 2019				
6.	Number of ridges per fruits					
7.	Fruit length (cm)					
8.	Fruit diameter (cm)					
9.	YVMV incidence (%)					
10.	Average fruit weight (g)					
11.	Number of fruits per plants					
12.	Fruit yield per plant (g)					

**Table 3.** Enlist of the studied phenotypic traits under present study.

Genotypes	Plant	Number	Days to	Days to	Number	Number	Fruit	Fruit	YVMV	Average	Number	Fruit
Genergpes	height	of primary	first	first	of seeds	of ridges	length	diameter	incidence	fruit	of fruits	vield
	(cm)	branches	flowering	picking	per fruit	per plant	(cm)	(cm)	(%)	weight	per plant	per
	(•••••)	per plant	nowering	Prening	perman	per prane	(•••••)	(•••••)		(g)	Per prane	plant
		F F								(8)		(g)
IC- 43735	107.40	1.97	44.00	50.33	52.67	5.00	10.23	1.62	36.28	11.51	20.33	254.00
IC- 43742	133.07	3.08	40.33	48.00	56.67	5.00	12.00	1.81	12.33	12.07	27.50	312.33
IC-218872	108.33	2.20	42.33	53.00	68.67	6.47	10.13	1.68	27.67	11.78	21.97	265.33
IC- 105667	114.87	2.25	41.73	48.30	54.67	5.00	10.97	1.52	21.67	10.34	24.33	261.67
EC- 199367	117.20	2.57	41.33	47.33	39.33	5.00	14.27	1.63	9.33	12.91	26.70	319.33
EC- 015537	130.33	3.07	39.63	46.33	43.33	5.07	14.53	1.50	19.13	11.17	24.67	275.67
SKY/DR/RS-13	136.33	3.10	43.17	48.67	52.00	5.13	11.33	1.58	27.12	11.19	25.67	286.88
SKY/DR/RS- 107	110.00	2.27	41.27	47.67	42.33	5.00	10.63	1.71	33.33	11.55	22.00	256.67
VRO-454-10-1	100.00	2.27	41.33	48.67	48.00	5.00	10.03	1.53	38.12	11.23	21.50	250.00
VRO-232-10-1	95.67	1.90	43.33	51.00	51.33	5.00	9.23	1.62	24.83	10.52	21.00	255.00
VRO-132-10-1,2	116.33	2.25	40.83	48.67	35.33	5.13	13.67	1.64	21.00	10.96	25.67	296.67
VRO-304-10-1	91.33	1.87	42.50	52.33	52.67	5.00	9.62	1.80	32.45	11.73	20.00	247.67
VRO-109-1	101.00	1.90	44.33	49.67	37.67	5.00	9.60	1.58	34.78	11.05	19.00	236.67
VRO-196	119.33	2.40	41.33	47.33	42.33	5.00	13.00	1.68	15.18	11.98	24.00	285.33
VRO-177	96.67	1.77	47.33	55.33	57.33	6.87	10.53	1.72	27.76	10.33	20.67	230.67
VRO-956	107.67	2.08	42.40	49.67	44.00	5.13	13.17	1.69	37.39	15.63	20.67	287.67
VRO-433	118.33	2.73	42.93	50.67	28.33	5.00	10.90	1.46	51.67	11.35	22.33	266.00
VRO-214	133.00	3.00	39.03	45.33	46.00	5.13	12.50	1.79	14.67	13.18	24.00	320.33
VRO-320	120.07	2.80	41.37	48.00	47.00	5.07	10.13	1.72	24.33	11.77	24.00	283.33
VRO-107	119.00	2.53	40.47	47.33	30.67	5.00	13.83	1.49	25.00	11.80	25.33	302.67
VRO-173	122.67	2.37	46.00	53.00	48.33	5.07	10.17	1.64	30.97	11.43	24.33	282.67
VRO-404	111.67	2.25	41.67	47.00	54.00	5.00	10.70	1.59	41.00	11.73	20.60	256.67
VRO-40	121.00	2.43	44.33	50.33	40.33	5.07	10.83	1.49	29.71	10.67	22.67	238.67
VR0- 37	102.00	2.00	41.33	49.33	50.00	5.00	9.43	1.74	32.11	11.80	19.67	253.33
Kashi Kranti	116.67	2.30	40.57	47.33	48.33	5.00	11.77	1.60	17.33	11.22	23.67	265.67
CD (5%)	15.04	0.39	3.83	4.69	6.76	0.60	2.00	0.21	5.04	1.67	2.58	40.12

**Table 4.** Mean performance of morphological traits of twenty five okra genotypes.

	Table 5. Clustering pattern of okra genotypes selected for the study.						
Cluster	Number of genotypes	Genotypes in cluster					
I	19	VRO-304-10-1, VRO-37, IC-43735, VRO-232-10-1, SKY/DR/RS-107, VRO-454-10-1, VRO-173, VRO-404, VRO-40, VRO-109-1, VRO-320, IC-105667, Kashi Kranti, SKY/DR/RS-13, VRO-132-10-1,2, VRO-196, VRO-107, EC-015537, VRO-214					
II	1	EC-199367					
III	1	IC-43742					
IV	2	IC-218872, VRO-177					
V	1	VRO-956					
VI	1	VRO-433					



Table 6. Cluster mean for studied attributes of okra genotypes based on phenotypic similarity.

40	Characters	Mean of cluster							
47		Ι	II	III	IV	V	VI		
	Plant height (cm)	114.14	117.20	133.07	102.50	107.67	118.33		
48	Number of primary branches per plant	2.36	2.57	3.10	1.98	2.08	2.73		
	Days to first flowering	42.01	41.33	40.33	44.83	42.40	42.93		
40	Days to first picking	48.77	47.33	48.00	54.17	49.67	50.67		
49	Number of seeds per fruit	46.16	39.33	56.67	63.00	44.00	28.33		
	Number of ridges per fruit	5.04	5.00	5.00	6.67	5.13	5.00		
50	Fruit length (cm)	11.17	14.27	12.00	10.33	13.17	10.90		
	Fruit diameter (cm)	1.62	1.63	1.81	1.70	1.69	1.46		
- 1	YVMV incidence (%)	27.32	9.33	12.33	27.71	37.39	51.67		
21	Average fruit weight (g)	11.41	12.91	12.10	11.06	15.63	11.35		
	Number of fruits per plant	22.76	25.00	26.00	21.32	20.67	22.33		
52	Fruit yield per plant (g)	268.92	319.33	312.33	248.00	287.67	266.00		

**Table 7.** Mean intra and inter cluster distance  $(D^2)$  among six clusters of okra.

				<i>. . . .</i>		
Cluster	Cluster I	Cluster II	Cluster III	Cluster IV	Cluster V	Cluster VI
Cluster I	13.42	23.07	28.07	29.70	23.63	48.80
Cluster II		0.00	14.02	48.75	40.40	90.76
Cluster III			0.00	37.75	59.44	111.03
Cluster IV				7.47	45.24	90.10
Cluster V					0.00	33.34
Cluster VI						0.00

**Table 8.** Percent contribution of studied phenotypic attributes towards total genetic divergence in okra.

Traits	Times ranked first	% Contribution
Plant height	14	4.67
Number of primary branches per plants	23	7.67
Days to first flowering	2	0.67
Days to first picking	0	0.00
Number of seeds per fruits	59	19.67
Number of ridges per fruits	30	10.00
Fruit length (cm)	8	2.67
Fruit diameter (cm)	2	0.67
YVMV incidence (%)	134	44.67
Average fruit weight (g)	18	6.00
Number of fruits per plants	7	2.33
Fruit yield per plant (g)	3	1.00



Fig. 1. Clustering pattern of 25 okra genotypes selected for the study.



**Fig-2.** Intra and inter cluster distance  $(D^2)$  among six clusters of selected 25 okra genotypes.