

Principal component analysis for yield and quality traits of blackgram

(*Vigna mungo* (L.) Hepper)

Abstract

The study consists of fifty-nine blackgram genotypes, which were evaluated for fourteen quantitative and qualitative traits. In order to determine the relationship and diversity among the blackgram genotypes taken for study. A field experiment was conducted at the Regional Agricultural Research Station, Lam, Guntur district, Andhra Pradesh state during *Kharif*, 2019. Principal component analysis for various yield-contributing traits was done to evaluate diversity and some quantitative and qualitative traits that had more effects on diversity. PCA results revealed that four of the five principal components had eigen values greater than one. The first five components obtained from principal component analysis (PC 1 to 5) accounted for about 76.73% of the total variation for fourteen quantitative and qualitative traits. Out of total principal components, PC 1, PC 2, PC 3, PC 4 and PC 5 were retained with values of 35.42%, 14.85%, 11.14%, 8.75% and 6.56%, respectively. The results of 2D and 3D scatter diagrams revealed LBG 904, LBG 752 and TU 94-2 genotypes to be the most diverse. Utilizing these diverse genotypes as parents in hybridization suggests obtaining desirable transgressive segregants towards the development of high yields with nutritional quality. The clustering of blackgram genotypes based on the yield and quality-attributing traits would be helpful in identifying the appropriate genotypes for effective utilisation in upcoming breeding programmes. The outcomes of principal component analysis revealed that wide genetic variability occurs between these blackgram genotypes and proposed their potential value in blackgram yield and quality improvement.

Key words : Principal component analysis, genetic divergence, yield, quality, blackgram.

1.INTRODUCTION :

Black gramme (*Vigna mungo* (L.) Hepper) with a chromosome number of $2n = 22$, belonging to the family Leguminosae and subfamily Papilionaceae, is a self-pollinating, short-duration, and widely cultivated seed legume [1]. *Vigna mungo* var. *silvestris* is the progenitor of blackgram [2]. India is the primary centre of origin [3]. Blackgram seed is a rich source of protein, fibre, several vitamins and essential minerals such as calcium and iron [4]. Blackgram is the best source of protein for vegetarians [5]. India is the largest producer and consumer of blackgram.

In India, about 3060 thousand tonnes of blackgram are produced annually from about 5602 thousand hectares of area, with an average productivity of 546 kg per hectare, while in Andhra Pradesh, about 310.56 thousand tonnes of blackgram are produced annually from about 318 thousand hectares of area, with an average productivity of around 977 kg per hectare. [6].

In any hybridization programme, genetic diversity is a prerequisite for desirable recombination [7]. Assessment of the nature and extent of genetic variability for qualitative and quantitative traits within the blackgram genotype is necessary for crop improvement in terms of crop yield and quality. Principal component analysis (PCA) allows not only the natural grouping of the genotypes but is also a precise indicator of genotype differences. The main advantage of using principal component analysis is that each genotype can be assigned to one group only. PCA has been used to identify redundancy among genotypes with similar traits and their elimination [8]. The present investigation was undertaken in this context to study the nature and magnitude of genetic diversity among 59 blackgram genotypes for yield, yield component, and quality traits using principal component analysis (PCA).

2. MATERIAL AND METHODS

The experimental material consisted of 59 blackgram genotypes obtained from IIPR, Kanpur and from the MULLaRP scheme of the Regional Agricultural Research Station, Lam, Guntur, Andhra Pradesh state. Details of the genotypes studied in the present investigation are presented in Table 1. All the 59 genotypes were sown during *Kharif* 2019–20 Regional Agricultural Research Station, Lam, Guntur, University of ANGRAU, AP. The experiment was laid out in an augmented design without replication with five blocks and four check varieties. Four check varieties are randomised in each block. The plot size was 2 rows of 3 metres each, and the spacing maintained was 30 x 10 cm. The observations were recorded on traits viz., days to flowering, plant height (cm), branches per plant, cluster per plant, pods per plant, pod length (cm), seeds per pod, days to maturity, 100 seed weight (gm), seed yield per plant (gm), harvest index (%), protein content (%), iron content (mg/100g) and zinc content (mg/100g). The observations are recorded on ten randomly selected plants from the middle of the row, avoiding the plants from the border, and they are tagged. Observations on test weight, days to 50% flowering, days to maturity, and all the quality parameters, viz., protein, iron, and zinc, were recorded on a plot basis. Total variance among the 55 genotypes and 4 check entries was separated into different sources ('genotypes + check entries', "genotypes", "check entries" and "genotypes vs check entries") using augmented design [9] presented in Table 2. Principal component analysis was carried out using the software Window Stat Version 8.5.

Table 1. Details of the blackgram genotypes studied and their sources

Sl. N	Genotype	Source	Sl. N	Genotype	Source
1	KU 96-7	CSA,Kanpur	31	PU 1501	GBPU A&T, Pantnagar
2	MBG 1070	ARS, Madhira	32	OBG 102	OUAT, Bhubaneswar
3	LBG 918	RARS, Lam	33	TBG 129	RARS, Tirupati
4	IPU 17-1	IIPR, Kanpur	34	LBG 776	RARS, Lam
5	DBGV 16	UAS, Dharwad	35	WBU 108	PORS, Berhampore
6	OBG 103	OUAT, Bhubaneswar	36	KPU1720-140	ARS, Kota
7	DKU 90	CSK HPKV, Palampur	37	LBG 709	RARS, Lam
8	Uttara	IIPR, Kanpur	38	TU 50	BARC, Mumbai
9	VBG 09-005	NPRC, Pudukkottai	39	LBG 868	RARS, Lam
10	KPU 52-87	ARS, Kota	40	TU 40	BARC, Mumbai
11	PU 31	GBPUAT, Pantnagar	41	MU 52	MSSC Ltd, Akola
12	KU 17-04	CSAU, Kanpur	42	RU 03-22-4	IGKV, Raipur
13	DKU 116	Dhaulakuan	43	KUG 818	PAU, Ludhiana
14	CO 5	NPRC, Vamban	44	VBG 12-110	NPRC, Vamban
15	GJU 1509	SDAU, S.K nagar	45	NUL 242	Nirmal seed
16	LBG 854	RARS, Lam	46	ADT 5	TNAU, Aduthurai
17	VBG 17-026	NPRC, Vamban	47	ADT6	TNAU, Aduthurai
18	VBN -5	NPRC, Vamban	48	VBG 17-029	NPRC, Vamban
19	OBG 41	OUAT, Bhubaneswar	49	OBG 101	OUAT, Bhubaneswar
20	VBG 12-062	NPRC, Vamban	50	IPU 11-6	IIPR, Kanpur
21	LBG -623	RARS, Lam	51	IPU 1702	IIPR, Kanpur
22	TU 44	BARC, Mumbai	52	LBG 972	RARS, Lam
23	ADBG 13023	TNAU, Aduthurai	53	LBG 885	RARS, Lam
24	AKU 1608	PDKV, Akola	54	LBG 883	RARS, Lam
25	IPU 12-5	IIPR, Kanpur	55	LBG 880	RARS, Lam
26	VBG 13-003	NPRC, Vamban	56	LBG 787	RARS, Lam
27	LBG 904	RARS, Lam	57	IPU 2-43	IIPR, Kanpur
28	SBC 50	RARS, Shillongani	58	LBG 752	RARS, Lam
29	TJU 134	BARC, Mumbai	59	TU 94-2	BARC, Mumbai
30	PU 1541	GBPU A&T, Pantnagar			

Table.2. Analysis of variance for quantitative and qualitative characters studied in 59 genotypes of blackgram (*Vignamungo*(L.) Hepper)

Source of Variatio	d.f	DM	DF	PH (cm)	NBP	NCP	NPP	PL(cm)
Mean Sum of Squares								
Block	4	0.178	0.5	1.721	0.013	0.19	1.487	0.011
Entries	58	13.663***	4.392 ***	33.509***	0.206 ***	3.375***	33.293***	0.248***
Checks	3	16.183***	8.600 ***	37.869***	0.114 **	27.618***	230.007***	0.616***
Varieties	54	7.267***	3.521 ***	28.559***	0.183 ***	1.785***	14.047***	0.210***
Checks vs. Varieties	1	351.494***	38.838 ***	287.714***	1.723 ***	16.484***	482.466***	1.167***
Error	12	0.141	0.433	0.876	0.016	0.06	0.879	0.006

* Significant at 5% level ** Significant at 1% level*** Significant at 0.1% level

Source of Variation	d.f	NSP	100-SW (gm)	HI (%)	Protein (%)	Iron content (mg/100g)	Zinc content (mg/100g)	SYPP (gm)
Mean Sum of Squares								
Block	4	0.058**	0.02	2.998**	0.055	0.021	0.007	0.325
Entries	58	0.247***	0.116***	19.104***	2.933***	0.962 ***	0.302***	4.089***
Checks	3	0.813***	0.254***	86.157***	10.651***	0.194 ***	1.159***	21.847***
Varieties	54	0.209***	0.105***	14.688***	2.549***	1.009 ***	0.231***	2.302***
Checks vs. Varieties	1	0.561***	0.279***	56.417***	0.465	0.739 ***	1.532***	47.299***
Error	12	0.008	0.007	0.35	0.154	0.01	0.008	0.216

* Significant at 5% level ** Significant at 1% level*** Significant at 0.1% level

DM- Days to maturity, **DF**- Days to 50% flowering, **PH**-Plant height, **NBP**-Number of branches per plant, **NCP**- Number of clusters per plant, **NPP**- Number of pods per plant, **PL**-Pod length, **NSP**- Number of seeds per pod, **100-SW**-100 Seed weight, **HI**- harvest Index, **SYPP**-Seed yield per plant.

3.RESULTS AND DISCUSSION :

In the present study, PCA was estimated for fourteen traits of fifty-nine genotypes of Blackgram. The PC1 contributed 35.423% towards variability. Characters *viz.*, seed yield per plant (0.414), number of pods per plant (0.373) and harvest index (0.379) explained the maximum variance in PC1. The second axis (PC 2) contributed 14.853% variability, and variation at this axis is because of the following traits: days to maturity (0.433), plant height (0.354) and iron content. With loading of days to 50% flowering (0.291), days to maturity (0.209), and number of branches per plant (0.167), PC 3 contributed 11.146% of variation. The fourth principal component (PC 4) contributed 8.750 percent of the total variability. This axis showed positive loadings for zinc content (0.316), number of branches per plant (0.297) and days to 50% flowering (0.148) and the

fifth principal component (PC 5) was characterised by 6.563 percent contributed towards the total variability. This axis showed positive loadings for the number of seeds per pod (0.338), pod length (0.301) and days to maturity (0.205).

The cumulative variability percentage for the first component is 35.423, while it is 50.276 for PC 2, 61.422 for PC 3, 70.173 for PC 4 and 76.736 for PC 5 (Table 3). The PCA scores for 59 blackgram genotypes in the first three principal components were computed and were considered as three axes, X, Y, and Z, and the squared genotypes from these three axes were calculated (Table 4). The pattern of spread of the genotypes in these clusters was detected to be at random with no reference to geographical diversity, as genotypes from different geographical regions were clustered in the same as well as different clusters. The PCA scores for 59 genotypes were plotted in the graph to get the 2D (PCA I as X axis and PCA II as Y axis) and 3D (PCA I as X axis, PCA II as Y axis, and PCA III as Z axis) scatter diagrams (Fig. 1 and Fig. 2).

The diverse genotypes numbered 27 (LBG 904), 58 (LBG 752), and 59 (TU 94-2), which are far away from other genotypes in the 2 dimensional and 3 dimensional diagrams (Fig 1 & 2), may be used as parents in hybridization to exploit the transgressive segregants.

Usage of PCA for getting 2D and 3D digrams and to understand the genetic diversity was earlier used in various crops. In finger millet, [10] discovered that eigen values greater than one accounted for 76.41 percent of the cumulative variance among the first three axes. In cotton, [11] discovered 81.99 percent variability among six principal components, while in mungbean, [12] discovered 78 percent total variance among five principal components for identifying promising parents and producing superior segregants in subsequent generations.

Table 3. Eigen value, per cent variance and percent cumulative variance for five principal components (PCs) and factor loading between PCs and traits studied in blackgram (*Vigna mungo*(L.) Hepper)

Canonical Roots Analysis (P. C. A.)						
	Components	PC1	PC2	PC3	PC4	PC5
	Eigene Value (Root)	4.95926	2.07946	1.56049	1.22509	0.91885
	% Var. Exp.	35.42328	14.8533	11.14639	8.75061	6.56322
	Cum. Var. Exp.	35.42328	50.27658	61.42296	70.17357	76.73679
	Characters	PC1	PC2	PC3	PC4	PC5
1	Days to 50% Flowering	0.27311	0.27958	0.29113	0.14814	0.09769
2	Days to maturity	0.2272	0.43322	0.20978	0.00721	0.20514
3	Plsnt height (cm)	0.22832	0.35434	0.15533	-0.03623	0.05019
4	Number of branches per plant	0.26926	0.27798	0.16774	0.29795	0.09285
5	Number of clusters per plant	0.32718	-0.39255	0.03246	0.12817	0.02131
6	Number of pods per plant	0.37304	-0.32604	0.07726	0.09448	-0.02828
7	Pod length (cm)	0.19888	-0.03188	-0.52202	-0.01192	0.30183
8	Number of seeds per pod	0.29012	0.02819	-0.36248	-0.17266	0.3382
9	100-Seed weight (g)	0.17746	0.15724	0.07679	-0.42969	-0.66854
10	Harvest Index (%)	0.37911	-0.23543	0.0894	-0.04726	-0.13137
11	Protein content (%)	0.09716	0.12295	-0.10932	-0.71973	0.15591
12	Iron content (mg/100g)	0.10224	0.32065	-0.41532	0.15348	-0.15554
13	Zinc content (mg/100g)	0.11112	0.17104	-0.4559	0.3168	-0.45407
14	Seed Yield Per plant (g)	0.41446	-0.20438	0.04332	-0.06169	-0.10775

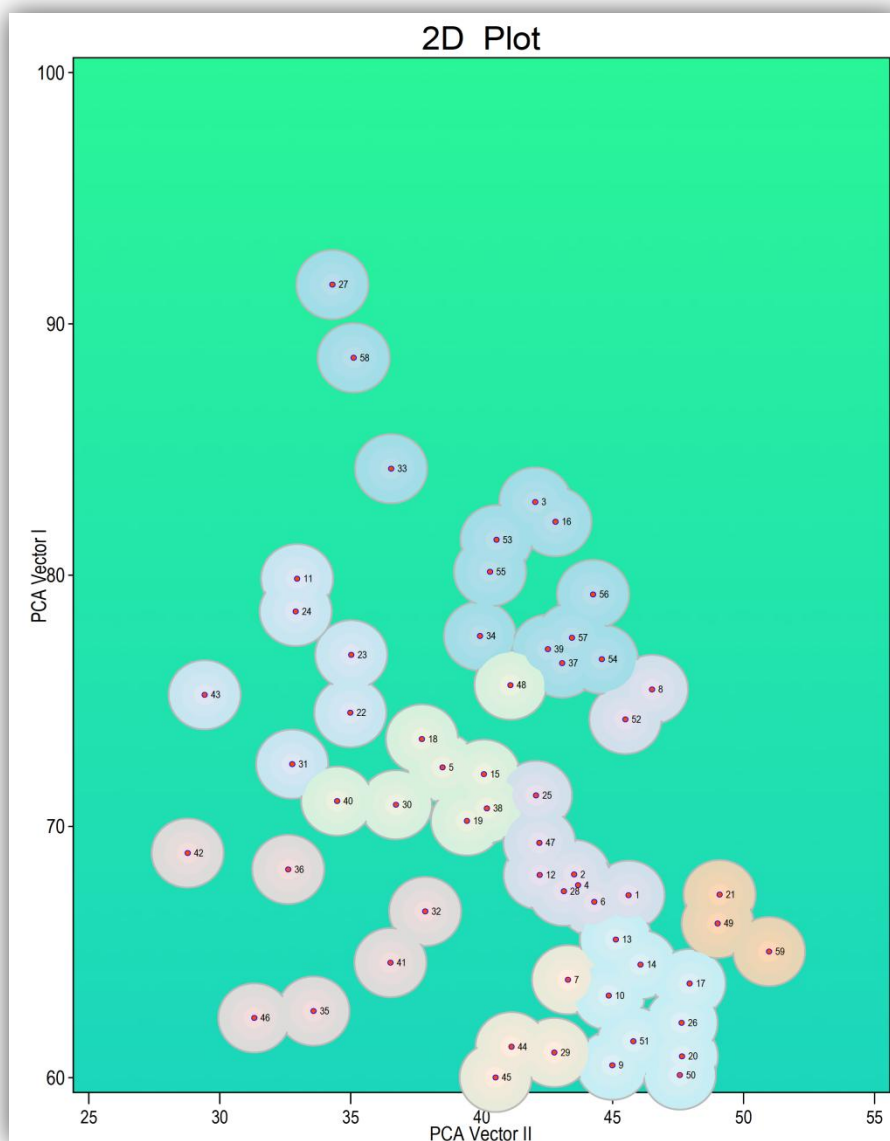


Fig 1 Two dimensional graph showing relative positions of 59 blackgram genotypes based on PCA score

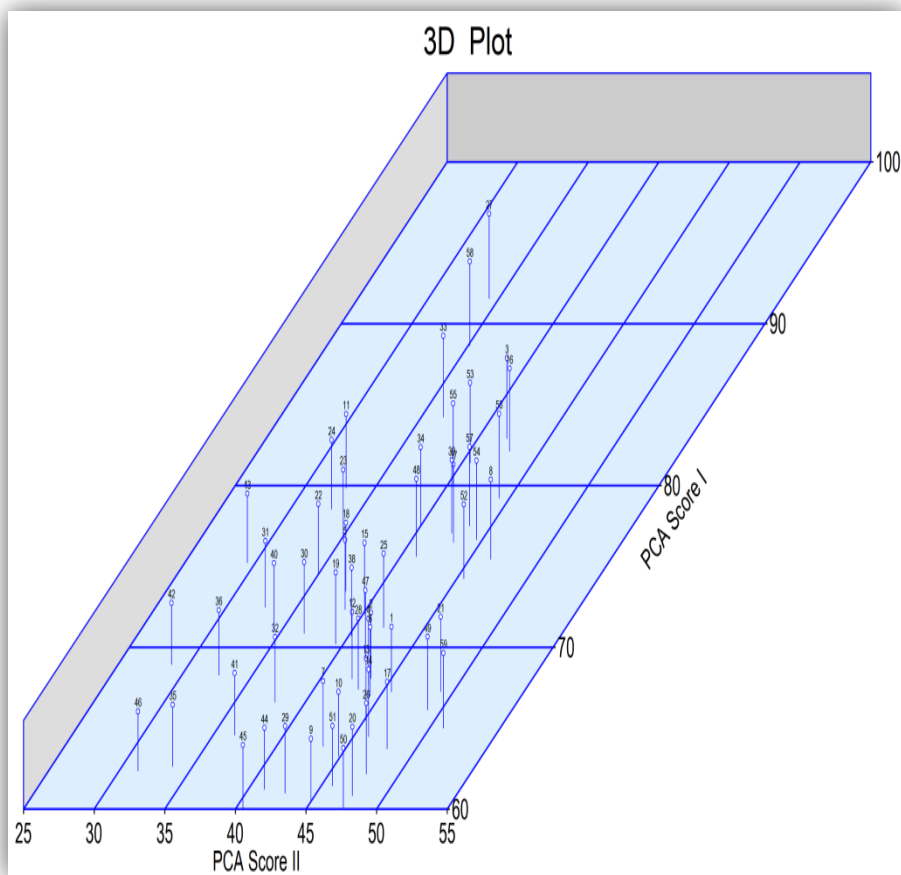


Fig 2: Three dimensional graph showing relative positions of 59 blackgram genotypes based on PCA scores

Table 4. The PCA scores of genotypes of 59 genotypes of blackgram (*Vigna mungo* (L.) Hepper)

		PCA I	PCA II	PCA III			PCA I	PCA II	PCA III
Sl. N	Genotype	X Vector	Y Vector	Z Vector	Sl.N	Genotype	X Vector	Y Vector	Z Vector
1	KU 96-7	67.258	45.608	24.878	31	PU 1501	72.475	32.769	25.393
2	MBG 1070	68.084	43.533	25.197	32	OBG 102	66.615	37.847	24.953
3	LBG 918	82.908	42.048	30.78	33	TBG 129	84.233	36.546	31.122
4	IPU 17-1	67.656	43.68	25.455	34	LBG 776	77.575	39.937	29.713
5	DBGV 16	72.344	38.51	26.59	35	WBU 108	62.646	33.58	23.636
6	OBG 103	66.993	44.303	26.466	36	KPU1720-140	68.284	32.614	24.874
7	DKU 90	63.891	43.296	24.968	37	LBG 709	76.493	43.074	29.774
8	Uttara	75.447	46.506	30.513	38	TU 50	70.712	40.198	26.063
9	VBG 09-005	60.486	44.99	23.94	39	LBG 868	77.049	42.532	27.985
10	KPU 52-87	63.263	44.855	24.741	40	TU 40	71.005	34.483	26.012
11	PU 31	79.856	32.953	28.255	41	MU 52	64.572	36.519	23.778
12	KU 17-04	68.064	42.215	25.552	42	RU 03-22-4	68.937	28.775	23.605
13	DKU 116	65.492	45.125	23.768	43	KUG 818	75.232	29.423	26.446
14	CO 5	64.494	46.066	25.711	44	VBG 12-110	61.227	41.14	23.534
15	GU 1509	72.076	40.088	27.142	45	NUL 242	60.002	40.53	24.539
16	LBG 854	82.122	42.82	31.723	46	ADT 5	62.375	31.32	22.704
17	VBG 17-026	63.743	47.94	25.517	47	ADT6	69.341	42.203	25.988
18	VBN -5	73.474	37.719	26.144	48	VBG 17-029	75.618	41.1	29.654
19	OBG 41	70.214	39.44	27.367	49	OBG 101	66.132	49.011	28.065
20	VBG 12-062	60.845	47.651	26.278	50	IPU 11-6	60.098	47.568	22.765
21	LBG -623	67.281	49.084	28.553	51	IPU 1702	61.443	45.792	22.879
22	TU 44	74.522	34.986	26.853	52	LBG 972	74.255	45.487	28.396
23	ADBG 13023	76.824	35.02	25.773	53	LBG 885	81.405	40.568	30.618
24	AKU 1608	78.55	32.9	26.408	54	LBG 883	76.652	44.589	30.278
25	IPU 12-5	71.231	42.078	28.246	55	LBG 880	80.132	40.321	30.641
26	VBG 13-003	62.177	47.635	27.096	56	LBG 787A©	79.229	44.254	32.258
27	LBG 904	91.561	34.303	32.467	57	IPU 2-43A©	77.506	43.449	30.243
28	SBC 50	67.415	43.141	27.107	58	LBG 752A©	88.647	35.114	32.227
29	TJU 134	60.994	42.774	25.62	59	TU 94-2A©	65.014	50.973	28.735
30	PU 1541	70.858	36.728	27.314					

On the basis of PCA-based clustering, 59 genotypes were divided into 7 clusters, with cluster 1 (24 genotypes) having the most genotypes, followed by cluster 2 (23 genotypes), cluster 3 (7 genotypes), cluster 5 (2 genotypes), and clusters 4, 6, and 7 being solitary groupings that were chosen to be more diverse than the other clusters (Table 5). Table 6 shows the inter- and intra-cluster distances between different genotypes. Cluster 2 (142.28) had the highest intra-cluster distance, followed by cluster 3 (141.96), whereas clusters 5 and 6 (1396.70) had the highest inter-cluster distance, followed by cluster 3 and 5 (1172.96). This finding suggests that genotypes from clusters separated by a large statistical distance should be used in future hybridization programmes.

Table 5. Clustering pattern by Tocher's method in 59 genotypes of blackgram (*Vignamungo* (L.) Hepper)

Characters group	No.of genotype	List of genotypes
1 Cluster	24	MBG 1070, OBG 103, KU 96-7, IPJ 17-1, SBC 50, ADT6, IPU 12-5, KU 17-04, CO 5, DKU 116, DKU 90, TU 50 ,VBG 17-026 ,OBG 41, GJU 1509,VBG 12-110,VBG 13-003,VBG 12-062,KPU 52-87, VBG 09-005,OBG 101,LBG -623,IPU 11-6, IPU 1702
2 Cluster	23	LBG 918, LBG 854, LBG 880, LBG 885, TBG 129, LBG 776, IPJ 2-43, VBG 17-029, LBG 883, LBG 787, LBG 709, LBG 868, LBG 972, Uttara, VBN -5, PU 31, TU 44, PU 1541, ADBG 13023, DBGV 16, AKU 1608, PU 1501, TU 40
3 Cluster	7	TJU 134, NUL 242, OBG 102, MU 52, WBJ 108, ADT 5, KPU 1720-140
4 Cluster	1	KUG 818
5 Cluster	2	LBG 904, LBG 752
6 Cluster	1	TU 94-2
7 Cluster	1	RU 03-22-4

Table 6. Average intra and inter-cluster distances among seven clusters (Tocher's method) of 59 blackgram (*Vignamungo* (L.) genotypes

Cluster Distances							
Cluster	1	2	3	4	5	6	7
1	108.84	337.15	280.52	477.92	1044.34	203.41	453.02
2		142.28	462.80	225.15	372.31	515.40	422.06
3			141.96	352.14	1172.90	478.91	347.40
4				0.00	442.76	798.15	112.52
5					42.67	1396.70	867.30
6						0.00	828.23
7							0.00

* Diagonal values are intra-cluster distances. Off diagonal values are inter-cluster distances.

4. CONCLUSION:

This result suggests that genotypes in clusters that are separated by a high statistical distance should be utilised in potential hybridization programmes. The diverse genotypes numbered 27 (LBG 904), 58 (LBG 752), and 59 (TU-94-2) which are far away from other genotypes in the 2D and 3D dimension diagrams (Fig 1 and 2) and clusters 5 (LBG 904, LBG 752), and cluster 6 (TU-94-2) had the highest inter-cluster distance and may be used as parents in hybridization to exploit the transgressive segregants.

5. ACKNOWLEDGEMENT

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COMPETING INTERESTS DISCLAIMER:

Authors have declared that no competing interests exist. The products used for this research are commonly and predominantly use products in our area of research and country. There is absolutely no conflict of interest between the authors and producers of the products because we do not intend to use these products as an avenue for any litigation but for the advancement of knowledge.

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