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Research Article

Multivariate analysis in parental lines and land races of pearl millet [*Pennisetum glaucum* (L.) R. Br.]

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Abstract

Present investigation was carried out with 31 pearl millet genotypes containing R lines, B lines and land races at Department of Millets, Tamil Nadu Agricultural University, Coimbatore during *kharif* 2019 in order to assess the genetic diversity and to compare different methods of multivariate analysis. Mahalanobi's D² cluster analysis resulted in six clusters with the highest inter cluster distance observed between cluster V and cluster VI. Cluster mean showed that cluster III and cluster I genotypes played significant influence on yield and yield attributing characters. In agglomerative hierarchical cluster (AHC) analysis, the highest inter cluster distance was observed between cluster II and cluster VI. Cluster V and cluster I represented the highest cluster mean for yield and yield component traits. Principal component analysis (PCA) resulted four principal components with eigen values more than one explaining 73.2 per cent variability. The biplot revealed five clusters with cluster I am possessing maximum number of genotypes and positively associated to most of the traits. PT 6706, PT 6709, Nattu Cumbu, Cumbu 2, PT 6676 and PT 6067 were the top-ranking genotypes upon PCA analysis with positive PC1 scores. All three multivariate analyses revealed considerable divergence in the experimental material as well as a comparable type of clustering in the diversity of R line, B line and small seeded land races and hence can be used in future breeding programmes.

Keywords: Mahalanobis' D², Agglomerative hierarchical clustering, Principal component analysis, Pearl millet

INTRODUCTION

Pearl millet [*Pennisetum glaucum* (L.) R. Br.] is a small-grained tropical C₄ cereal crop grown in most adverse agroclimatic conditions (Kumar *et al.*, 2020) and important in the lives of the poor and low-income groups (Govindaraj *et al.*, 2020). Pearl millet has a global area of approximately 340 lakh hectares and a yield of 310 lakh tonnes, ranking sixth among grains after wheat, rice, maize, barley, and sorghum (Patil *et al.*, 2020). India is the largest producer of pearl millet in the world, with a production of 97.0 lakh tonnes from 75 lakh ha

area (INDIASTAT, 2018). In India, pearl millet hybrids account for 70% of total pearl millet land area, with the remainder occupied by open-pollinated varieties (OPVs) or landraces (Patil *et al.*, 2020) indicating the importance of hybrid development in India. Pearl millet was grown on around 0.63 lakh hectares in Tamil Nadu, with productivity of 2,277 kg/ha (Statistical Handbook of Tamil Nadu, 2019). Understanding the genetics and diversity of pearl millet will help in opening the door to additional possibilities for using it as a fodder and grain

crop in today's variable environment. Broadening the genetic base is crucial for boosting genetic gain for yield by introducing diverse germplasm accessions to attain maximum heterosis. Hybrid development programme is strongly depending upon the selection of diverse seed and pollen parent to develop high yielding hybrids (Sharma *et al.*, 2020). Multivariate analysis, such as cluster analysis and principal component analysis (PCA) are the statistical procedures used to create the cluster in order to classify and identify divergent parents.

The D^2 statistics proposed by Mahalanobis is the most appropriate method for selecting morphologically divergent parents as it furnishes a measure of actual variation between any pair of populations (Malik *et al.*, 2017). Mahalanobis's generalized distance is estimated by D^2 statistics for discriminating population considering a set of parameters together rather than inferring from indices based on morphological similarities and polygenic relationship (Sankar *et al.*, 2014; Singh and Gupta, 1979; Rasitha *et al.*, 2020; Swamynatham *et al.*, 2020). Hierarchical cluster analysis is a commonly used method for forming clusters and displaying similarities and dissimilarities between pairs of genotypes, in which agglomerative hierarchical clustering were formed by grouping cases into bigger and bigger clusters until all cases are members of a single cluster. Principal components analysis (PCA) is the data reduction technique applicable to quantitative type of data and transforms multi-correlated variables into another set of uncorrelated variables (Kumar *et al.*, 2020).

The current study employed multivariate analysis, comprising cluster analysis (Agglomerative hierarchical clustering (AHC) and Mahalanobis' D^2 statistics) and principal component analysis (PCA) with the objective to examine the diversity of 31 important pearl millet lines and to make comparison of the different methods.

MATERIALS AND METHODS

The current investigation was carried out at the Department of Millets, Tamil Nadu Agricultural University, Coimbatore which lies in western agroclimatic zone of Tamil Nadu, India. This zone has an altitude in the range of 200 m to 600 m and located between 11°55' to 10°02' N latitude and 76°51' to 78°09' latitude. Field experiment was conducted during the main cropping season *kharif* (June to October) during 2019 at Department of Millets, Tamil Nadu Agricultural University, Coimbatore which is located at 11°01' to 30.7°N longitude and 76°55' to 35.0°E latitude. Total of 31 pearl millet genotypes which included 17 restores lines, 3 maintainer lines, 10 land races and an open pollinated variety Dhanashakti was utilized for the present investigation. Experimental material was laid out in randomized complete block design (RCBD) with two replications. A total of 16 quantitative traits were recorded on randomly selected five competitive plants in two replications except days to 50% spike emergence,

which was a single observation by visual assessment of group of plants in each replication on plot basis. Then, the mean data in each replication were subjected to statistical analysis.

The statistical analysis of replicated data was carried out with the help of the software WINDOSTAT ver 7.1 for D^2 statistics. D^2 statistics was originally developed by Mahalanobis (1936) and Rao (1952) suggested its application for the assessment of genetic diversity in plant breeding. The genotypes were grouped on the basis of minimum generalized distance using the Tocher's methods. Software XLSTAT was used for employing AHC (Agglomerative Hierarchical Clustering) method, where Euclidean distance between the genotypes were calculated from the unweighted pair group method using arithmetic averages (UPGMA) and PCA (Principal Component Analysis) was carried out by software XLSTAT Version 2014.5.0 for standardized mean data. Cluster diagram for AHC was analysed through software Graphical Genotypes (GGT 2.0).

RESULTS AND DISCUSSION

Mahalanobis's D^2 statistics is a technique for the assessment of genetic diversity in various breeding materials. For exploiting heterosis in hybrid development programme, it is necessary to utilize parents with maximum genetic divergence. More diverse the parents, more are the chances of pronounced heterotic effects and increased spectrum of variability in the segregating generations (Govindaraj *et al.*, 2011). Mahalanobis's D^2 analysis employed for the grouping 31 pearl millet genotypes using 16 yield and yield attributing traits resulted in six major clusters. The dendrogram for the D^2 cluster analysis using Tocher method is depicted in the **Fig. 1**. Wilk's Criterion simultaneous test of significance showed that there was highly significant difference among genotypes for all the characters. Out of six clusters, cluster I possessed maximum genotypes of 23 followed by four genotypes in cluster II. Most of R (restorer line) and B (maintainer line) lines were in cluster I and all small seeded land races like Kuttu cumbu 1, Kuttu cumbu 2 and Kuttu cumbu 3 were in cluster II. Pattern of distinct clusters and allotment of land races and breeding lines in different clusters indicated the presence of divergence between land races and lines used in experimental material. Out of three B lines, two B lines (ICMB 98222, ICMB 99222) came under cluster I and cluster VII possessed one B line (ICMB 06111). This indicated the clear differentiation between maintainer lines (Kaushik *et al.*, 2018). Among the land races, the genotypes Cumbu 1, Shoolgiri local and Nattucumbu were fell in cluster I, whereas cluster III possessed Kizikuppam local and cluster II accounted for four small seeded land races. These uneven distribution of land races to different clusters and most lines falling into few clusters suggested that land races collected from the same geographic area were not necessarily closely related and different regions did not necessarily

have different genetic background (Upadhyay and Murty, 1970; Dave and Joshi, 1995 and Govindaraj *et al.*, 2011). Genetic drift and selection under different environments could have caused greater divergence than geographical distance (Upadhyay and Murty, 1970).

Higher inter-cluster distance was observed than intra-cluster distance (**Table 1**). The maximum intra cluster distance was observed in cluster I followed by cluster II and least in cluster III, IV, V and VI as these had only one genotype. The maximum inter cluster distance was observed between cluster V and cluster VI followed by cluster III and cluster VI, cluster IV and cluster VI and minimum cluster distance was observed between cluster IV and cluster V. The higher inter-cluster distance than intra-cluster distance showed homogeneity and narrow genetic variability within a cluster (Singh and Gupta, 1979; Malik *et al.*, 2017). Lesser intra cluster distance indicated that the genotypes inside the cluster should be nearly identical in their characteristics and less divergent. More intra cluster distance may be due to degree of heterogeneity and pedigree and hence, selection will be efficient if it is based on highest mean for desirable traits (Ramya *et al.*, 2017; Kaushik *et al.*, 2018; Rasitha *et al.*, 2020). The genotypes within the clusters can be subjected to further analysis for morphological trait uniformity and to test for general combining ability to combine 4 to 10 lines to develop synthetics and composites. Maximum inter cluster distance was observed for cluster V, cluster VI and cluster III, which had PT6583, ICMB06111 and Kizikuppam local genotypes, respectively. These genotypes could be utilized for evaluation of specific combining ability (*sca*), hybrid development and to obtain good recombinants in F_2 (Govindaraj *et al.*, 2011; Sankar *et al.*, 2014; Ramya *et al.*, 2017; Rasitha *et al.*, 2020). Maximum inter cluster distance also lead to wide spectrum of variability in segregating population to operate selection (Singh *et al.*, 1981; Govindaraj *et al.*, 2011; Athoni *et al.*, 2016). It was also indicated that the genotypes with the restricted genetic divergence having relatively smaller statistical distances or falling in the same cluster were also likely

to produce desirable heterotic effects in the population resulting from crossing if they complement some major weaknesses of each other as against those involving genotypes which falling in distant clusters and possessing wide genetic divergence.

Cluster II showed highest mean for the number of productive tillers. Cluster III recorded highest mean for leaf length, flag leaf length, flag leaf width, thousand seed weight and single plant yield. Cluster IV recorded maximum mean for the traits like leaf width, number of nodes, panicle diameter total, number of grains per panicle and biological yield. Cluster V recorded highest mean for leaf sheath length, flag leaf width, number of nodes, panicle length and plant height. Cluster VI showed maximum mean for the trait days to flowering and harvest index (**Table 2**). From the results of cluster mean values, it was clear that for good yield and yield attributing traits, cluster III played prominent role and it included land race Kizikuppam local followed by the cluster I which included almost all-important R and B lines. Cluster II comprised of small seeded land races with high cluster mean for number of productive tillers and plant height with fair amount of biological yield could be used for the fodder purpose (Dave and Joshi, 1995). It was also important to note that, the cluster mean for days to 50% spike emergence was low in cluster III and cluster II, indicating early maturity enabling the cultivars to escape from terminal drought (Govindaraj *et al.*, 2011; Sumathi, *et al.*, 2016; Rasitha *et al.*, 2020).

Highest percent contribution to the total variability was due to thousand grain weight followed by single plant yield, plant height, panicle length, days to 50 % spike emergence and number of productive tillers (**Table 2**). Nearly 90 per cent of variability was contributed by thousand grain weight, single plant yield, plant height, number of nodes, biological yield, leaf width, panicle length, days to 50% spike emergence and panicle diameter indicating the opportunity of these traits for selection in the given experimental material

Table 1. Estimates of intra (diagonal bolded) and inter (non-diagonal non-bolded) cluster distances in pearl millet genotypes for yield and yield attributing traits by D² method

Clusters	Cluster I	Cluster II	Cluster III	Cluster IV	Cluster V	Cluster VI
Cluster I	11.37 (3.37)	16.54 (4.07)	13.22 (3.64)	13.99 (3.74)	13.91 (3.73)	17.03 (4.13)
Cluster II		10.71 (3.27)	19.53 (4.42)	15.22 (3.9)	15.56 (3.94)	18.73 (4.33)
Cluster III			0 (0)	19.96 (4.47)	19.29 (4.39)	20.11 (4.48)
Cluster IV				0 (0)	8.55 (2.92)	20.07 (4.48)
Cluster V					0 (0)	20.89 (4.57)
Cluster VI						0 (0)

Values in the parenthesis are "D" distance

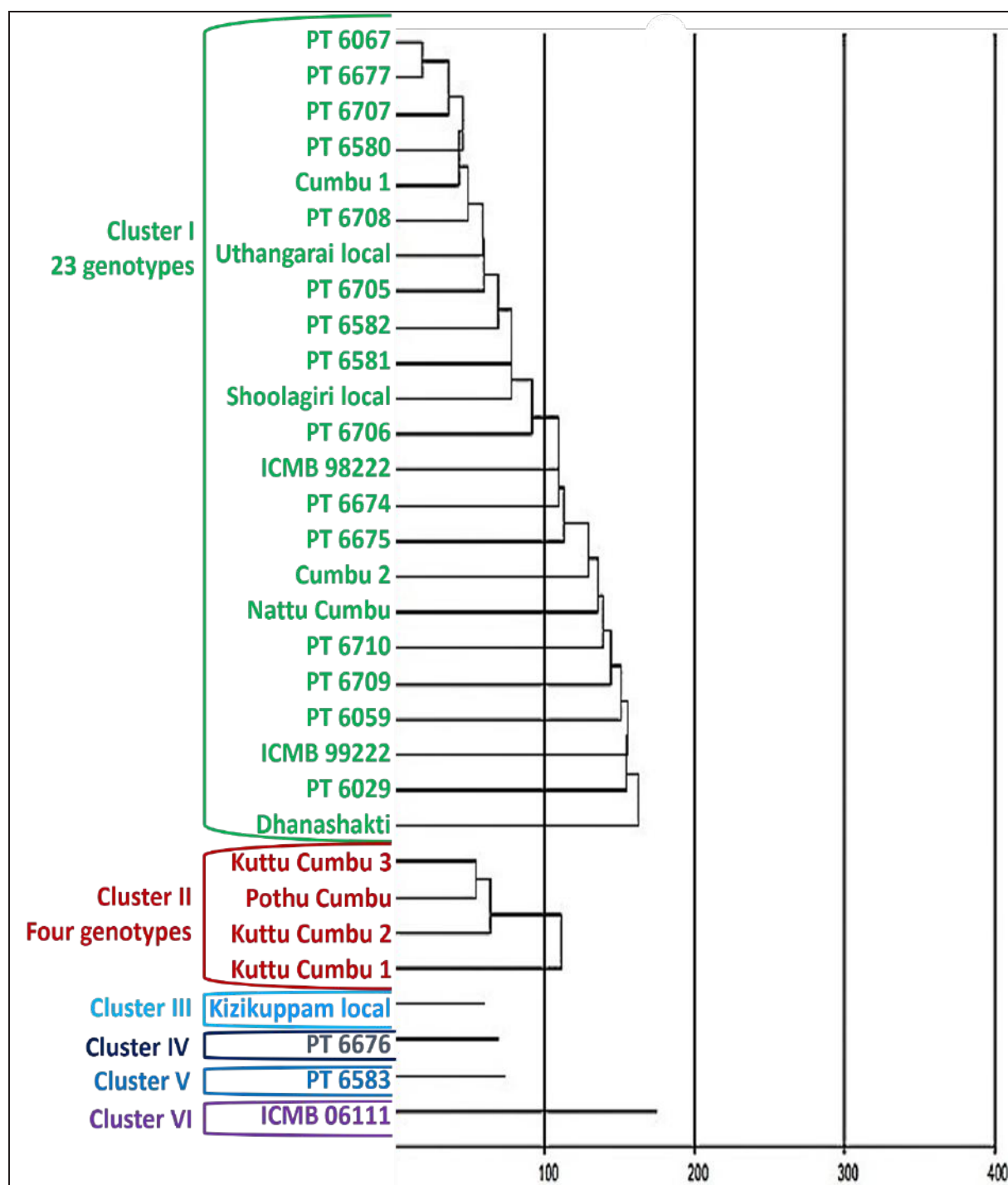


Fig. 1. Dendrogram representing the D^2 cluster analysis using Tocher method in pearl millet genotypes for yield and yield attributing traits

Table 2. Cluster mean of pearl millet genotypes and percentage contribution to total variability for yield and yield attributing traits by D² method

Cluster	DTF	LSL	LL	LW	FL	FW	NN	PL	PD	NT	PH	TGP	TSW	BY	SPY	HI
Cluster I	50.39	17.84	57.25	4.27	44.16	4.25	6.34	23.48	3.06	3.23	157.03	2739.81	10.92	75.94	51.22	70.91
Cluster II	40.25	12.19	45.13	3.09	29.13	3.26	5.92	18.50	1.72	4.83	173.66	1506.95	4.80	77.33	30.63	41.86
Cluster III	40.00	16.75	73.00	3.95	53.00	4.50	5.34	25.33	2.92	3.66	170.54	2486.54	12.20	71.50	79.00	110.84
Cluster IV	50.00	19.00	64.00	4.76	39.67	4.23	7.00	31.33	3.15	3.33	160.00	4144.96	6.10	109.64	43.12	42.90
Cluster V	52.00	23.33	48.33	4.20	43.33	4.50	7.00	31.67	3.04	2.66	177.91	1944.42	9.92	99.77	31.58	31.85
Cluster VI	58.00	14.66	34.00	3.46	25.33	3.50	4.00	15.66	2.68	3.33	83.67	2304.33	7.05	32.74	43.45	132.76
Per cent contribution	4.3	1.94	0.22	7.1	1.08	1.29	8.39	4.95	3.87	1.29	9.46	3.01	28.6	7.74	12.9	3.87

DTF- Days to 50 % spike emergence , LSL- Leaf sheath length (cm), LL- leaf length (cm), LW- Leaf width (cm), FL- Flag leaf length (cm), FW- Flag leaf width (cm), NN- Number of nodes , PL- Panicle length (cm), PD- Panicle diameter (cm), NT- Number of productive tillers , PH- plant height (cm), TGP – Total number of grains per panicle, TSW - Thousand grain weight (g), BY - Biological yield (g), SPY - Single plant yield (g) and HI - Harvest index (%)

(Shanmuganathan *et al.*, 2006; Athoni *et al.*, 2016; Rasitha *et al.*, 2020). The low contribution to genetic divergence by other characters may be due to the fact that selection towards uniformity in these characters could have caused an eroding effect on genetic diversity (Govindaraj *et al.*, 2011).

Agglomerative hierarchical cluster analysis of 16 yield and yield attributing traits in 31 pearl millet genotypes resulted in six major clusters and were depicted in dendrogram derived by AHC method (**Fig. 2**). Out of six clusters, cluster I showed the maximum genotypes of about 18, followed by five genotypes in cluster II, four genotypes in cluster VI, two genotypes in cluster V and one genotype in cluster III and cluster IV each. Six major clusters indicated the presence of divergence between the land races and lines, which could be utilized for breeding programmes in future (Reddy *et al.*, 1996; Zhang *et al.*, 2010; Kiprotich *et al.*, 2015; Yadav *et al.*, 2016; Pujar *et al.*, 2020; Sharma *et al.*, 2020; Nagendra *et al.*, 2020). The pattern of cluster analysis differentiated the R lines, B lines and small seeded land races (Kumar *et al.*, 2020). The other land races were scattered around cluster I and cluster V, indicating that the lines grouped in different clusters despite their place of development and geographical distribution and showed that geographical isolation was not directly related to genetic diversity (Murty and Tiwari 1967; Kumar *et al.*, 2020). In contradictory, clustering pattern in pearl millet depended on origin of collection, geographical origin, pedigree relation or close area of cultivation as reported by Animasaun *et al.* (2017). Chaudhary *et al.* (2015) also explained that genotypes related to the place of origin showed tendency to group in the same cluster because of dependence upon the directional selection pressure that lead to well evolved homeostatic mechanism that would favour consistency of the associated character. It showed that geographic diversity does not essentially lead to genetic diversity, the factors of original domestication and environmental

conditions at the time of development played an important role in perpetuation.

The maximum intra cluster distance was observed in cluster II followed by cluster I, cluster VI, cluster V and zero for the cluster III and cluster IV as they possessed one genotype each. The highest inter cluster distance was observed between cluster III and cluster VI, followed by cluster III and cluster VI, cluster I and cluster VI. The minimum inter cluster distance was observed between cluster I and cluster III (**Table 3**). Intra cluster distance revealed that there was a similarity within the clusters and diversity between clusters, as observed by Govindaraj *et al.* (2020). Maximum inter cluster distance (cluster II and cluster VI) explained the divergence between the genotypes of two different clusters and could be intercrossed to prepare the base population. Assessment of lines from these clusters for *per se* performance and specific combining ability to identify best lines and could help to use them as pollinators for development of hybrids. It would help in choosing the parents for evaluation of specific combining ability and selection of heterotic hybrid (Ghazy *et al.*, 2015; Kiprotich *et al.*, 2015; Sharma *et al.*, 2020).

Cluster I showed the highest mean for leaf sheath length, number of nodes and panicle length. Cluster II showed the maximum mean for the trait panicle diameter and total number of grains per panicle. Cluster III showed highest mean for the trait flag leaf width and leaf width. Cluster IV showed the highest mean for harvest index and days to 50% spike emergence. Cluster V showed the highest mean for single plant yield, leaf length, flag leaf length, thousand grain weight biological yield and single plant yield. The highest mean for number of productive tillers and plant height was shown by cluster VI (**Table 4**). Cluster mean values showed a wide range of variation for all the characters undertaken in the study. Cluster I included almost all R lines with high mean for yield

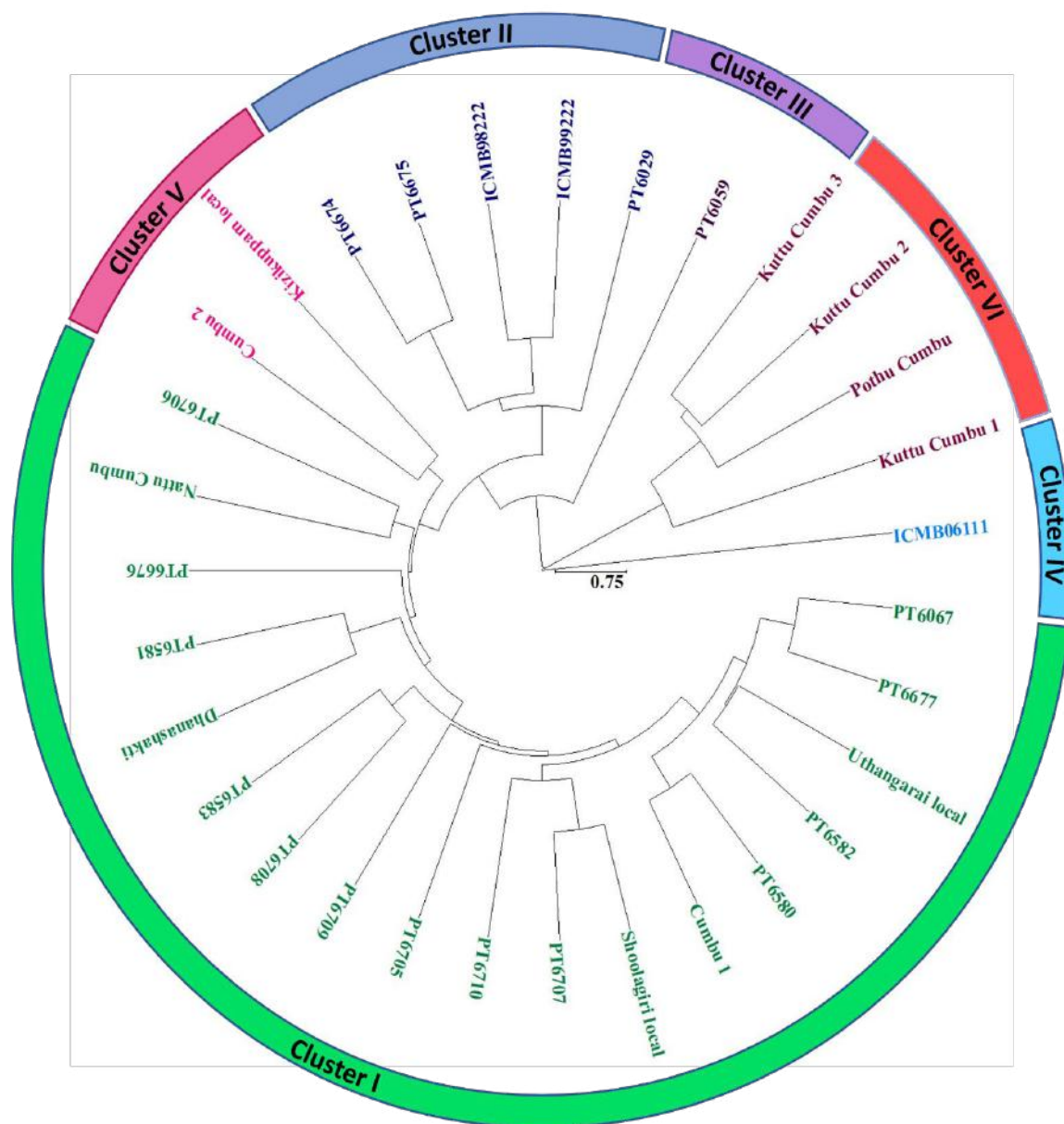


Fig. 2. Dendrogram depicting pearl millet genotypes derived by AHC method of clustering for yield and yield attributing traits

Table 3. Estimates of intra and inter cluster analysis of 31 pearl millet genotypes for yield and yield attributing traits by AHC method

Cluster	Cluster I	Cluster II	Cluster III	Cluster IV	Cluster V	Cluster VI
Cluster I	19.15	21.25	9.93	19.85	13.03	34.24
Cluster II		22.83	20.44	28.77	24.88	40.26
Cluster III			0.00	20.60	15.28	34.77
Cluster IV				0.00	15.74	28.45
Cluster V					6.76	31.78
Cluster VI						11.25

Table 4. Estimates of cluster mean of pearl millet genotypes for yield and yield attributing traits by AHC method

Cluster	DTF	LSL	LL	LW	FL	FW	NN	PL	PD	NT	PH	TGP	TSW	BY	SPY	HI
Cluster I	50.67	18.74	59.68	4.34	44.94	4.32	6.52	25.87	3.00	3.37	167.77	2678.66	10.66	83.78	54.40	68.86
Cluster II	48.20	16.90	47.29	3.97	38.07	3.73	6.40	18.67	3.29	2.67	133.29	3127.02	10.31	51.55	33.64	66.06
Cluster III	56.00	16.00	53.00	4.73	47.33	5.17	4.33	23.67	2.94	1.33	95.00	2713.09	9.89	62.04	19.20	30.94
Cluster IV	58.00	14.67	34.00	3.47	25.33	3.50	4.00	15.67	2.68	3.33	83.67	2304.33	7.05	32.73	43.45	132.73
Cluster V	46.00	15.88	69.25	4.23	52.50	4.60	5.50	22.83	3.01	4.33	169.43	2513.73	12.99	99.91	82.58	88.81
Cluster VI	40.25	12.19	45.13	3.09	29.13	3.26	5.92	18.50	1.72	4.83	173.67	1506.95	4.80	77.33	30.63	41.54

DTF - Days to 50 % spike emergence , LSL - Leaf sheath length (cm), LL - leaf length (cm), LW - Leaf width (cm), FL - Flag leaf length (cm), FW - Flag leaf width (cm), NN - Number of nodes , PL - Panicle length (cm), PD - Panicle diameter (cm), NT - Number of productive tillers , PH - plant height (cm), TGP – Total number of grains per panicle, TSW - Thousand grain weight (g), BY - Biological yield (g), SPY - Single plant yield (g) and HI - Harvest index (%)

attributing traits indicating the significance of these lines to become potential parents for yield contributing traits. Out of three B lines, two B lines (ICMB 98222 and ICMB 99222) and out of 17 R lines three R lines (PT 6675, PT6674 and PT 6029) fell under cluster II, which showed the highest mean for panicle diameter and total number of grains per panicle. These lines could be used as potential parents for obtaining the panicles of bigger size with a greater number of seeds per panicle. The highest mean for the number of productive tillers and plant height observed for cluster VI included small seeded land races which could be used for forage purpose. The genotypes with contrast mean performance from these clusters could be utilized as potential parents in the development of hybrids for harnessing heterosis (Drabo *et al.*, 2013; Kiprotich *et al.*, 2015; Nehra *et al.*, 2016; Santos *et al.*, 2017; Sharma *et al.*, 2020).

Principal component analysis is an effective approach for reducing the variability in multiple characters to the principal components with the first principal component capturing the maximum variability. The PCA based on correlation was used to study interrelationship between different characters. Principal component analysis with correlation matrix is best to determine the principal factors, as it does not require the normal distribution assumption of populations (Chaudhary *et al.*, 2015; Sharma *et al.*, 2020). PC with higher eigen values and variables with high factor loadings were considered as best representative of system attributes. In the present investigation, PCA was performed for yield and other attributes in pearl millet genotypes (Table 5 and Fig. 3). The first four principal components accounted for 73.20% of the total variability with eigen values more than one. The PC1 accounted for 37.55% of total variability followed by PC2, PC3 and PC4 exhibited 17.93%, 10.59% and 7.13% of total variability respectively. From the result it was revealed that maximum variability was spread within first four principal components where PC1 showed the highest variability among four. Hence, it was

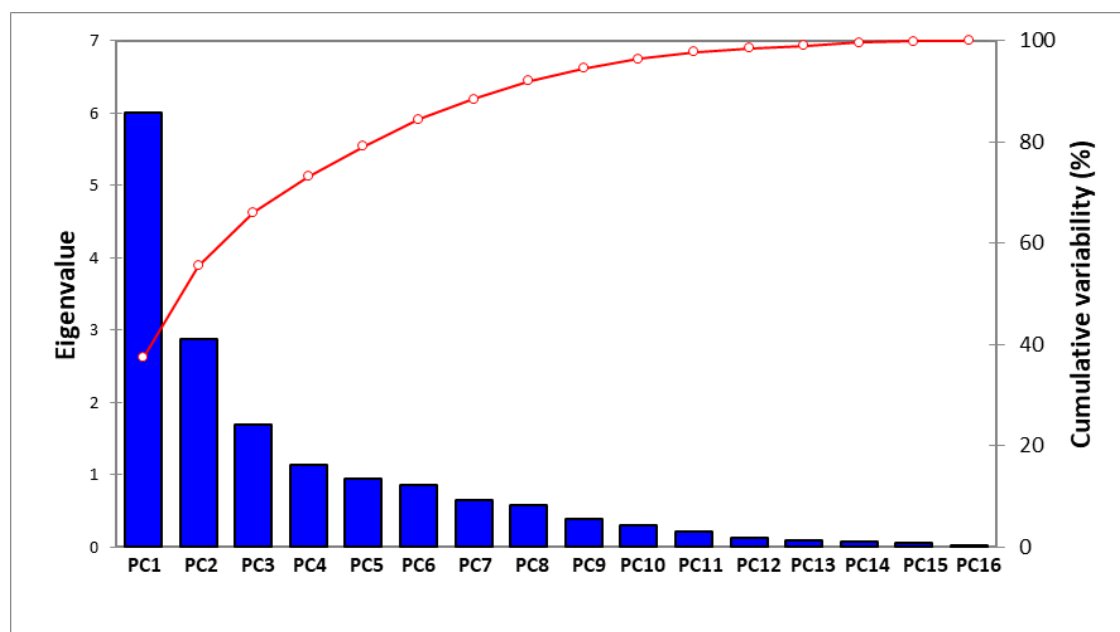
recommended to consider the characters or genotypes lying near and showing more PC1 score for catching the variability of particular trait (Ramya *et al.*, 2017; Jain and Diwan, 2021).

From the factor loading of PCA analysis, it was revealed that PC1 accounted maximum variability for most of the traits and other traits such as number of productive tillers, plant height and biological yield were captured in PC2. The harvest index and single plant yield were accounted in PC3 (Table 6). Result of factor loading of PCA analysis indicated that the maximum variability accounted by the PC1 was highly related to most of the yield attributing traits. PC2 showed maximum factor loading for number of productive tillers, plant height and biological yield, which were related to small seeded land races. Since, PC3 captured maximum variability for single plant yield and harvest index, the genotypes captured under this component can be utilized in improvement of crop for above mentioned traits. Characters which showed high positive or high negative contributed more to the diversity. The sign here indicates the relationship between variable and principal components. PC1 showed negative factor loading for number of productive tillers (-0.31) indicating the negative correlation with the trait (Ghazy *et al.*, 2015; Chaudhary *et al.*, 2015; Malik *et al.*, 2017; Rasitha *et al.*, 2020; Ramya *et al.*, 2017; Jain and Diwan, 2021).

PCA results were generally are displayed as a biplot, in which axes correspond to the new system of coordinates (Fig. 4). The direction of arrow denotes the maximum change in great quantity and the length could be related with the rate of change occur. The acute coordinate angle (<90°) between the traits or principal component axis and trait shows the positive association between these traits, whereas obtuse angle (>90°) shows negative and right angle (=90°) indicates no correlation between the traits (Govindaraj *et al.*, 2020). Most of the traits were in acute angle with the PC1 coordinates except number of productive tillers. The third quadrant did not have

Table 5. Eigen values and estimates of per cent variability accounted by the principal component analysis

	PC1	PC2	PC3	PC4	PC5	PC6	PC7	PC8	PC9	PC10	PC11	PC12	PC13	PC14	PC15	PC16
Eigenvalue	6.01	2.87	1.69	1.14	0.95	0.86	0.65	0.57	0.39	0.31	0.21	0.12	0.09	0.08	0.05	0.01
Variability (%)	37.55	17.93	10.59	7.13	5.91	5.35	4.05	3.59	2.41	1.92	1.30	0.76	0.59	0.51	0.30	0.09
Cumulative %	37.55	55.49	66.07	73.20	79.12	84.47	88.52	92.12	94.53	96.46	97.76	98.51	99.10	99.61	99.91	100.00

**Fig. 3. Scree plot showing eigen values and percentage of cumulative variability****Table 6. Factor loading of four important principal components of pearl millet genotypes for yield and yield attributing traits**

S. No.	Traits	PC1	PC2	PC3	PC4
1	DTF	0.55	-0.47	-0.09	-0.11
2	LSL	0.77	-0.11	-0.21	-0.09
3	LL	0.74	0.33	0.06	-0.08
4	LW	0.84	-0.14	-0.19	-0.18
5	FL	0.86	0.13	-0.01	-0.23
6	FW	0.73	-0.01	-0.10	-0.38
7	NN	0.41	0.40	-0.13	0.34
8	PL	0.71	0.20	-0.33	-0.20
9	PD	0.71	-0.44	0.05	0.44
10	NT	-0.31	0.75	0.31	0.04
11	PH	0.28	0.84	-0.06	-0.11
12	TGP	0.52	-0.45	0.05	0.46
13	TSW	0.69	-0.06	0.38	0.27
14	BY	0.34	0.65	-0.25	0.36
15	SPY	0.57	0.37	0.68	0.11
16	HI	0.22	-0.23	0.84	-0.31

DTF - Days to 50 % spike emergence , LSL - Leaf sheath length , LL - leaf length , LW - Leaf width , FL - Flag leaf length, FW - Flag leaf width , NN - Number of nodes , PL - Panicle length , PD - Panicle diameter, NT - Number of productive tillers , PH - plant height, TGP – Total number of grains per panicle, TSW - Thousand grain weight , BY - Biological yield , SPY - Single plant yield and HI - Harvest index

any traits. All most all the traits were in acute angle with single plant yield. The lowest acute and adjacent angle with single plant yield were observed for leaf length and number of nodes. Number of productive tillers showed obtuse angle, which indicated the negative correlation between the traits. Single plant yield showed acute angle with most of the traits indicating positive correlation and exhibited the significance of the trait selection for improvement of yield attributing characters mainly like leaf length, number of nodes, panicle length etc. which were highly correlated (Kalagare *et al.*, 2021; Sharma *et al.*, 2020 ; Pujar *et al.*, 2020).

Five different clusters were observed, where cluster I with the maximum number of genotypes possessed most of the genotypes on first quadrant followed by cluster II with 6 genotypes resided on third quadrant. Cluster I possessed most of the R lines and no B lines, whereas cluster II showed the two B lines (ICMB 98222 and ICMB 99222) and cluster IV showed one B line (ICMB 06111) on second quadrant. Cluster III possessed small seeded land races (Kuttu Cumbu 1, Kuttu Cumbu 2, Kuttu Cumbu 3 and Pothu Cumbu) on second quadrant. Biplot representing the genotypes is highly beneficial to identify the genotypes and character association and making the cluster for identification of diverse parents for utilization in development of hybrid. On the biplot, genotypes which closure to each other are similar and farthest are divergent (Sharma *et al.*, 2020). The distance between the location of any two genotypes on the score plot is indirectly proportional to the degree of similarity. Genotypes which are nearer to the origin are contributing less to the variability, while those far from the origin are extremes and mostly extremes are favourable for breeding programme (Kiprotich *et al.*, 2015). Similar genotypes formed group on the biplot which were differentiated by clusters. Results clearly indicated that the cluster formed on PC could differentiate the R lines, B lines and small seeded land races. Clusters contributing to the variability depended on how much far away the cluster formed from the origin. Cluster I and cluster II were nearer to the origin, whereas cluster III, cluster IV and cluster V were somewhat far away from the origin compared to cluster I and cluster II. Land races belonging to the cluster III contributed more to the variation (Bashir *et al.*, 2014; Nehra *et al.*, 2016; Animasaun *et al.*, 2017; Jain and Diwan, 2021).

The relationship between yield attributing traits with genotypes on first two principal components presented in the Fig.4. Among the five-clusters observed, cluster I with maximum genotypes was positively correlated with most of the traits except number of productive tillers. Most of the genotypes in the clusters I were bold seeded and possessed high single plant yield. Cluster III possessed small seeded land races and showed positive correlation with number of productive tillers, plant height and biological yield and negative correlation with days to flowering, harvest index, total number of grains per

panicle and panicle diameter. The genotypes, PT 6067, Dnanashakti, PT 6581, PT6580, Uthangarai local and Shoolagiri local were nearer to the origin. The genotypes, PT 6707, PT 6067, Dnanashakti, PT 6581, PT6580, PT 6705, Uthangarai local and Shoolagiri local were nearer to the origin indicating their stability and less variation for the characters. Cluster III with small seeded four land races showed positive relationship with number of productive tillers, plant height and biological yield and can be involved in the breeding for fodder crops. Cluster I was almost opposite the cluster II and cluster IV indicated the diversity between the clusters. Accessions from diverse group will maximize opportunities to obtain transgressive segregants as there is a higher chance from genotypes to contribute unique desirable alleles at various loci. Hence, it is recommended to use the genotypes present in cluster I, cluster II, cluster IV and cluster V to intercross among these clusters (Chaudhary *et al.*, 2015; Rasitha *et al.*, 2020; Sharma *et al.*, 2020).

List of top ten pearl millet accessions based on their PC score were arranged in a descending order of their scores (Table 7). The genotypes PT 6706, PT 6709, Nattu Cumbu, PT 6580, Cumbu 1, Cumbu 2, PT 6676, PT 6708, Kizikuppam local and PT 6067 showed maximum scores in PC1. The PC2 captured maximum score for the genotypes as Kuttu Cumbu 1, Pothu Cumbu, Cumbu 2, Kuttu Cumbu 2, PT 6580, Cumbu 1, Kizikuppam local, Kuttu Cumbu 3, PT 6706 and Dhanashakti. PC3 possessed maximum score for the genotypes like Kizikuppam local, ICMB 06111, Nattu Cumbu, Dhanashakti, Cumbu 2, PT 6705, PT 6582, PT 6581, PT 6706 and PT 6675. In present investigation, genotypes were identified through PC scores, where top ten PC1 scores were obtained by PT 6706, PT 6709, Nattu Cumbu, PT 6580, Cumbu 1, Cumbu 2, PT 6676, PT 6708, Kizikuppam local and PT 6067 in descending order. These genotypes showed overall highest mean values for the traits positively related to PC1 and had high factor loading that included most of yield and yield attributing traits. But number of productive tillers, total number of grains per panicle and plant height were related to PC2 and top score was possessed mainly by Kuttu Cumbu 1, Pothu Cumbu, Cumbu 2, Kuttu Cumbu 2, PT 6580, Cumbu 1, Kizikuppam local, Kuttu Cumbu 3, PT 6706 and Dhanashakti. These results were in concordance with the biplot determination (Fig. 4) (Chaudhary *et al.*, 2015; Sharma *et al.*, 2020). The above small seeded land races showed earliness, which is an important heat escaping attributes of pearl millet and could be used as potential parents in breeding programmes for early flowering as explained by Malik *et al.* (2017). The factor loading for single plant yield and harvest index were high for PC2 and the genotypes Kizikuppam local, ICMB 06111, Nattu Cumbu, Dhanashakti, Cumbu 2 were the top five scoring genotypes in PC2 indicating the expected significant yield improvement by these genotypes.

All three multivariate analysis (D^2 statistics, AHC

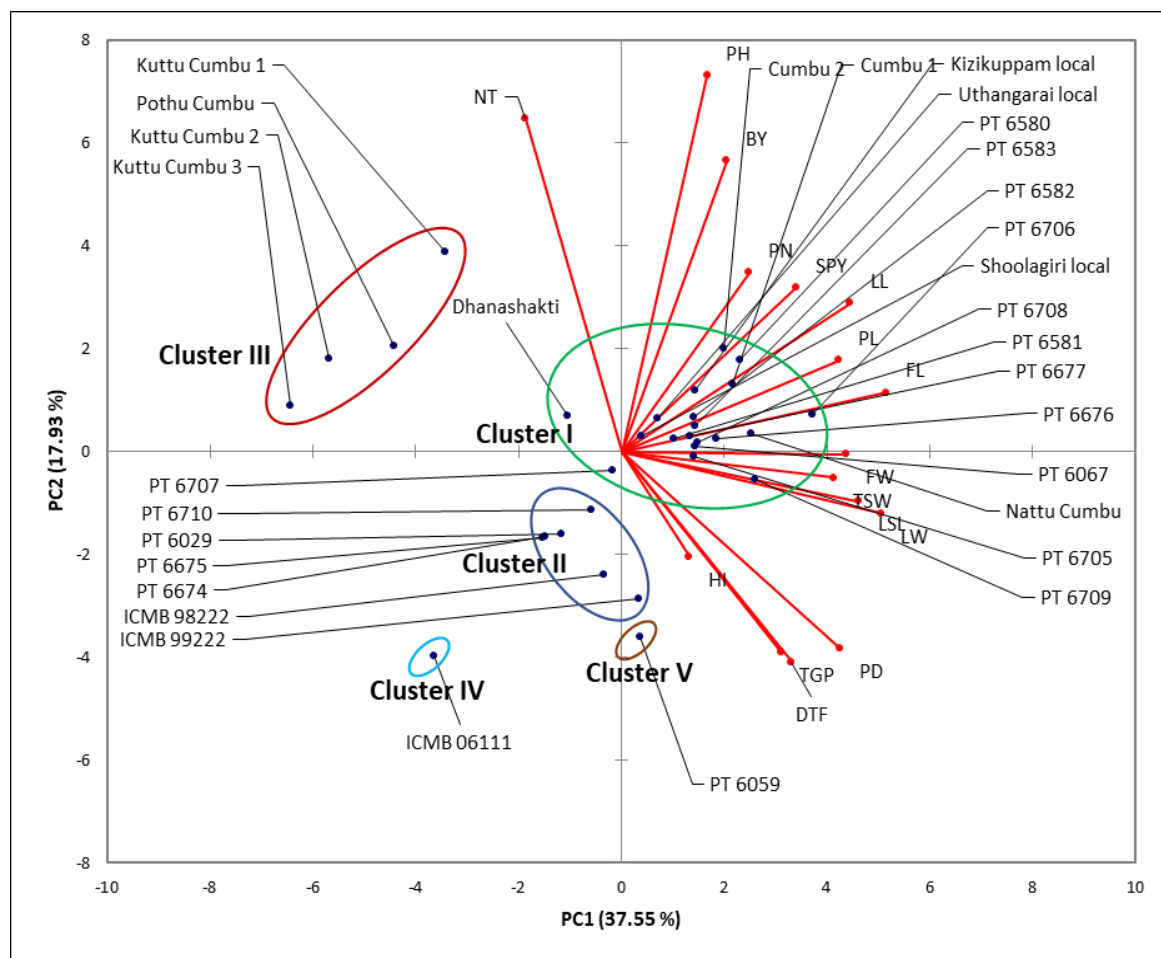


Fig. 4. Biplot representing the relationship between yield and yield attributing traits of pearl millet genotypes on first two principal components

Table 7. List of top ten pearl millet accessions based on their PC scores

Rank	PC1		PC2		PC3	
	DTF, LSL, LL, LW, FL, FW, NN, PL, PD, TGP, TSW		NT, PH		SPY, HI	
	Genotypes	Score	Genotypes	Score	Genotypes	Score
1	PT 6706	3.73	KuttuCumbu 1	3.89	Kizikuppam local	2.43
2	PT 6709	2.59	PothuCumbu	2.06	ICMB 06111	2.34
3	NattuCumbu	2.52	Cumbu 2	2.01	NattuCumbu	1.98
4	PT 6580	2.32	KuttuCumbu 2	1.80	Dhanashakti	1.95
5	Cumbu 1	2.16	PT 6580	1.79	Cumbu 2	1.70
6	Cumbu 2	1.99	Cumbu 1	1.31	PT 6705	1.16
7	PT 6676	1.85	Kizikuppam local	1.19	PT 6582	1.11
8	PT 6708	1.49	KuttuCumbu 3	0.88	PT 6581	1.08
9	Kizikuppam local	1.44	PT 6706	0.71	PT 6706	0.93
10	PT 6067	1.43	Dhanashakti	0.70	PT 6675	0.68

DTF- Days to 50 % spike emergence , LSL- Leaf sheath length, LL- leaf length , LW- Leaf width , FL- Flag leaf length , FW- Flag leaf width , NN- Number of nodes , PL- Panicle length, PD- Panicle diameter , NT- Number of productive tillers , PH- plant height , TGP – Total number of grains per panicle, TSW - Thousand grain weight , BY - Biological yield , SPY - Single plant yield and HI - Harvest index

Table 8. Identification of diverse parents and land races through different multivariate techniques

S. No.	D ² statistics	Agglomerative hierarchical clustering	Principal component analysis
1	Kuttu cumbu 1, Kuttu cumbu 2, Kuttu cumbu 3 and Pothu Cumbu	Kuttu cumbu 1, Kuttu cumbu 2, Kuttu cumbu 3 and Pothu Cumbu	Kuttu cumbu 1, Kuttu cumbu 2, Kuttu cumbu 3 and Pothu Cumbu
2	Kizikuppam local	Kizikuppam local Cumbu 2	
3	PT 6676	PT 6059	PT 6059
4	PT 6583	PT 6674, PT6675, ICMB 98222, ICMB 99222, PT 6029	PT 6674, PT6675, ICMB 98222, ICMB 99222, PT 6029, PT 6710
5	ICMB 06111	ICMB 06111	ICMB 06111

method and PCA) showed that there was a significant divergence between 31 genotypes. The cluster analysis Mahalanobis' D² statistics and Agglomerative hierarchical clustering (AHC) showed the variation in number of clusters and grouping of genotypes into different clusters. This is mainly because of the method of clustering and the type of data input for the cluster analysis. In case of Mahalanobis' D² statistics replicated data were used as input. During the testing of the significance (Wilk's Criterion simultaneous test of significance), it excluded the replication variances and consider only the determinants of error and error plus varietal variances. For transformation of the original variable to uncorrelated variable by pivotal condensation method and for calculation of D² values it uses error variance and covariance. The cluster formation by Tocher method was used where the cluster formation depends on the arbitrary value of D² which is generally approximately near to the maximum D² value between any two population (Singh and Chaudhary, 1977). In case of Agglomerative hierarchical clustering (AHC), standardized mean data was used for the calculation of distances between the mean of the different genotypes for characters. Euclidian distances (coefficient) were calculated and used for formation of dendrograms by unweighted pair group method using arithmetic averages (UPGMA). The number of clusters in AHC were decided by the value of Euclidian distance as dissimilarity coefficient in the dendrograms (Govindaraj *et al.*, 2020). Therefore, the number of clusters and grouping of genotypes into different clusters varied from one method to other method of cluster analysis. However, by making consensus it is observed that the comparable type of clustering in the diversity of R line, B line and land races. Especially the small seeded land races Kuttu cumbu 1, Kuttu cumbu 2, Kuttu cumbu 3 and Pothu Cumbu were grouped in one cluster and the B line ICMB 06111 was clearly different from other clusters (Table 8). Similar pattern of clustering was observed between AHC and PCA as the data used for analysis were mean data in both the cases (Chaudhary *et al.*, 2015; Malik *et al.*, 2017)

In the present investigation the main objective of selection of both the cluster analysis is to improve the selection criteria for the identification of diverse parents. The Mahalanobis' D² statistics is comparatively superior

as it uses the replicated data, whereas agglomerative hierarchical clustering method does not require replicated data. Principal component analysis (PCA) helps in identifying genotypes extreme for the variation which is highly useful in plant breeding (Kiprotich *et al.*, 2015). From the principal component scores it is possible to identify the genotypes contributing highest variation for several characters and genotypes are ranked accordingly. Hence, for cluster analysis, PCA has added advantage for selection of genotypes in breeding programmes.

In the present study the main aim of employing the multivariate analysis was to identify the potential parents and classify the pearl millet genotypes comprising R lines, B lines and land races. Three types of multivariate analysis viz. D² statistics, AHC method and PCA were utilized to analyse the genetic variation present in the set of genotypes. Even though each method has its own pros and cons for synthesizing the observed data and providing classificatory analysis, all the methods were compared to identify the elite genotypes with associated quantitative traits and making a selection strategy for crop improvement. In order to apply more selection pressure and to achieve highest genetic gain from the selected potential genotypes, classificatory approaches were performed. Each method has indicated the different groupings of genotypes. However, by making consensus, the most common parents which showed diverse in all three multivariate analysis included small seeded land races like Kuttu cumbu 1, Kuttu cumbu 2, Kuttu cumbu 3 and Pothu Cumbu and also the B line ICMB 06111. Highest cluster mean for single plant yield, thousand seed weight, flag leaf length and leaf length were observed for the cluster containing the genotype Kizikuppam local constantly in both Mahalanobis' D² statistics and AHC method of cluster analysis. PT 6706, PT 6709, Nattu Cumbu, Cumbu 2, PT 6676 and PT 6067 were the top-ranking genotypes upon PCA analysis with positive PC1 scores. High contribution of traits to total variability was by thousand grain weight, single plant yield, plant height and biological yield. Hence, there is much scope for selection of these traits among the genotypes studied for the exploitation of heterosis in hybrids and for obtaining broad spectrum of variation in segregating material for yield attributing traits.

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