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Genetic Diversity Analysis for Forage Yield and Quality Traits in Sorghum [*Sorghum Bicolor* **L. Moench] Germplasm**

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Authors' contributions

This work was carried out in collaboration between both authors. Both authors read and approved the final manuscript.

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ABSTRACT

This study investigated 280 sorghum germplasm lines using an augmented block design at GBPUAT Pantnagar under normal sowing conditions during the 2019 Kharif season. The observations were recorded on different yield contributing traits such as days to flowering, plant height, number of leaves; stem girth etc., quality traits such as protein content, total soluble solids, in vivo dry matter digestibility etc., and quality traits like cellulose content, silica content, and

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hemicelluloses etc. The statistical analysis for genetic diversity was done using hierarchical cluster analysis. Hierarchical cluster analysis revealed significant genetic diversity in the sorghum germplasm concerning various yield-related traits, quality traits, and biochemical traits. The 280 germplasm lines were grouped into XI distinct non-overlapping clusters.The cluster-VIII (50) consisted of highest number of genotypes whereas lowest numbers of genotypes were grouped into cluster-III (1).The maximum intra-cluster distance was observed in cluster-IV (54.652) suggesting highest genetic diversity among genotypes of this cluster in comparison to other clusters whereas minimum intra-cluster distance was observed in cluster-III (0.000).The highest inter-cluster distance was observed between clusters-III and VI (334.554)suggested distant relationship between members of these two clusters and upon crossing the members of these two clusters will give more genetic diversity in segregating generationwhereas minimum inter-cluster distance was observed betweenclusters-VIII and IX (52.512) suggested a closer relationship between these two clusters and low degree of genetic diversity among the genotypes.. Presence of substantial genetic diversity among the genotypes screened in the present study indicated that this material may serve as a good source for selecting the diverse parents for hybridization programme. In order to increase the possibility of isolating good trangressivesegregants in the segregating generations it would be logical to attempt crosses between the diverse genotypes belonging to clusters separated by large inter-cluster distances.

Keywords: Germplasm; cluster; genetic diversity; intra and inter; cluster distance.

1. INTRODUCTION

Sorghum is one of the most important and widely grown crops globally, covering an area of 41.14 million hectares with a production of approximately 58.72 million tonnes. "In India, sorghum covers 5.00 million hectares with a grain production of 4.50 million tonnes [1]. Sorghum is known by various names in Africa, such that *guinea-corn, dawa* or *sorgho* in West Africa, *durra* in the Sudan, *mshelia* in Ethiopia and Eritrea, *mtama* in East Africa, *kaffir corn* in South Africa and *amabele*or *mabele* in several countries in Southern Africa. In the Indian subcontinent, it is known as *jowar* (Hindi)*, cholam* (Tamil Nadu), *jonna* (Andhra Pradesh) and *jola* (Karnataka). Five basic races of cultivated sorghum are recognized as *Bicolor, Guinea, Kafir, Durra* and *Caudatum"* [2].

"It has extensive variability of usage such as forage sorghum, grain sorghum and sweet sorghum, providing food, fodder, feed, fuel and fiber. The crop is mainly grown in tropical and subtropical areas because of its drought tolerance capacity, and quick growing habit, good palatability where agro-climatic conditions such as rainfall, temperature and soil are variable. Much of the crop is grown in the stressprone and marginal areas of the semi-arid tropics, mainly on small holdings. In Northern Western India, it is grown for meeting the major fodder requirement of *kharif* and summer seasons. Precise information on nature and degree of genetic variability helps the plant

breeder in selecting the genetically diverse parents for the purposeful hybridization" [3]. "Genetic improvement of yield especially in selfpollinated crops depends on nature and amount of genetic diversity" [4].

"Nutritionally, among the *kharif* fodders, sorghum is a crop *par excellence* with starch (63-68%), potential of high digestibility (50-60%), dry matter (20-35%), sugars (8-17%), crude protein (7.5- 10.0%), calcium (0.53%), phosphorus (0.24%), and crude fiber (30-32%)". Sheoran et al*.* [5] "Beside the higher content of carbohydrates, it has iron (Fe) and vitamin B₃contentswhich are higher than maize and rice. It is a major staple food of many countries in Asia and Africa, sorghum is now a major feed crop in the United States, Argentina, Mexico, South Africa, and Australia". Miller and Kebede, [6].

Genetic diversity and relationship among different individuals is a prerequisite for any successful breeding programme. "Genetic diversity among accessions provides opportunities for improvement of agronomic and nutritional quality traits in crops". Huang, [7] "It aids plant breeders to characterize and classify accessions into heterotic groups". Menz et al., [8] "Genetic diversity of plants determines their potential for improved efficiency and hence their use for breeding, which eventually may result in enhanced forage production. Genetic diversity explains the genetic differences between different populations within a species or between species. The parents having more genetic diversity result into higher heterotic expression in F_1 and greater amount of genetic variability in segregating populations". Shekhawat et al., [9]. "One of the important approaches to sorghum breeding is hybridization and subsequent selection. Parents' choice is the first step in plant breeding program through hybridization. In order to obtain transgressive segregants, genetic diversity between parents is necessary" [10]. "The higher genetic diversity between parents, the higher heterosis in progeny can be observed" [4]. Estimation of genetic diversity is one of appropriate tools for parental selection in sorghum hybridization programs. Appropriate selection of the parents is essential to be used in crossing nurseries to enhance the genetic recombination for potential yield increase. In view of the above, there is need to screen the diversity of sorghum germplasm based on yield and quality parameters to find out their suitability in different breeding programmes.There is a need to make genuine efforts to assess available diversity. Hence the present investigation was conducted to estimate the magnitude of genetic diversity present among the elite sorghum genotypes.

2. MATERIALS AND METHODS

The initial research related to germplasm screening was carried out in the experimental area of Instructional Dairy Farm, Nagla, G.B. Pant University of Agriculture and Technology. Pantnagar, District U. S. Nagar, Uttarakhand during *Kharif*, 2019. The experimental material for this experiment consisted of two hundreds and eighty diverse germplasm lines of sorghum along with six checks *viz*., SSG 59-3, Pant Chari-5, Pant Chari- 6, CSV-21 F, CSH-22S, and CSV-24SS. The germplasm lines were evaluated in Augmented Block Design during *Kharif* season of 2019. The experiment was carried out in an Augmented Block Design [11-13] with each block containing 35 test entries and 6 checks which were randomly allocated in 8 blocks. All genotypes were sown on 27th July 2019 in single row of 5 meter length with a row spacing of 45 cm. All recommended agronomic practices for sorghum were followed to ensure healthy crop growth. The observations were recorded ondays to 50% flowering, days to maturity, number of leaves per plant, number of nodes, plant height (cm), leaf length (cm), leaf width (cm), leaf area (cm²), flag leaf length (cm), flag leaf width (cm), stem girth (cm), internodal length (cm), panicle length (cm), panicle width (cm), leaf:stem ratio, 1000-grains weight (gm), grain yield per plant (gm), green fodder yield per plant (gm), dry

fodder yield per plant (gm), foliar diseases zonate leaf spot and anthracnose [14] shoot fly (*Atherigonasoccata*) incidence (Dead hearts %), dry matter (%), brix %, HCN content (ppm) Hogg and Ahlagreen, [15] and Gilchrist et al. [16] protein content (%) Jeckson, [17] in-vitro dry matter disappearance (IVDMD) % Erwin and Ellinston, [18] neutral detergent fiber Van Soest, [19] acid detergent fiber (%) and cellulose (%) Van Soest, [19] acid detergent lignin (%), cellulose (%) and silica (%) Van Soest, [19]. Hierarchical cluster analysis was performed on the basis of Euclidean distance between the genotypes. Euclidean distance was calculated by using the following method:

Euclidean distance: The Euclidean distance between ith and kth accession is:

$$
D_{ik} = \Big[E_{j=1}^n \big(A_{ij} - A_{kj}\big)^2\Big]^{1/2}
$$

Where,

 D_{ik} = Euclidean distance between ith and kth accession

 A_{ii} = performance of ith accession for ith character.

 A_{ki} = performance of kth accession for jthcharacter.

 $m =$ number of accessions (I or $k = 1, 2...$ m) $n =$ number of characters $(i = 1, 2...n)$

When the similarity matrix is computed from distance function, the hierarchical clustering method begins by finding the link between the two closest genotypes [20]. Statistical analysis was conducted using Indostat software in Hyderabad**.**

3. RESULTS AND DISCUSSION

Understanding the genetic diversity of parents in hybridization programs is essential, as crosses involving genetically diverse parents are likely to produce not only high heterotic effects but also desirable transgressive segregants in later generations. The hierarchial cluster analysis discriminates genotypes in a different cluster on the basis of genetic diversity among the genotypes and thus enable breeder to select more genetically diverse parents for their crossing programme to recover desirable seggregants. The genotypes included in the same cluster may have different generations of time, different parental combinations or different generations of the same parental combinations. This proved that geographical diversity need not necessarily be related sown conditioned to the genetic diversity.

The hierarchial cluster analysis had been found to be a potent tool in quantifying the degree of divergence in germplasm. This analysis provides a measurement of relative contribution of different components on diversity both in inter cluster and intra cluster level and genotypes drawn from widely divergent clusters are likely to produce heterotic combinations and wide variability in segregating generation [21]. Among the different approaches of selecting parents, selection based on diversity has its own merit.

3.1 Distribution of Genotypes into Different Clusters

The clustering pattern of genotypes on the basis of Hierarchial cluster analysis has been presented in Table 1and Fig. 1 during *Kharif* 2019. The genotypes were grouped into XI distinct non-overlapping clusters suggesting considerable amount of genetic diversity prsesent in the experimental material. The cluster pattern of the genotypes showed nonparallelism between geographic and genetic diversity [22]. The genotypes were grouped into XI distinct non-overlapping clusters. The cluster-VIII (50) consisted of highest number of genotypes followed by cluster-IX (46), cluster-I (39), cluster-VII (36), cluster- V (35), cluster- IV (27), cluster- XI (24), cluster- II (14), cluster- X (11), cluster-VI (3) and lowest number of genotypes were grouped into cluster-III (1).

Cluster-I: This cluster consisted of thirty nine genotypes viz.,E2-2, Malwan, SSG-212, HJ-513, IS-23586, HC-171, IS-3318, SSG-222, PC-23, ICSV-702, IS-12743, CSV-10, 1890 (08BZL-01- 43-1), IS-20703-1, GP-2011-471, SSG-260, SSG-263, SSG-234, IS-1219, 9533-1, PC-1001, SPV-1752, UTFS-42, IS-9722, EJN-40,PC-1002, EJN-46, SSG-611, SMC-14, SEVS-2, B-437 (09B-RUS-04), IS-2363, IS-9162, IS-607, ICSV-111, SPV-1725, IS-6090, PSSV-61, UPFS-38 x UPFS-36. This cluster had highest cluster mean for acid detergent fiber, cellulose content, lignin content and silica content.

Cluster-II: This cluster exhibited fourteen genotypes viz., CO (FS)-29, IS-18850, SSG-21, IS-30117, IS-3353, E-159, IS-14357, IS-13566, IS-18927, IS-18844, IS-18933, SMC-8, SMC-13, and IS-28313. This cluster had high cluster mean for days to flowering, number of leaves, panicle width, green fodder yield per plant, dry fodder yield per plant, protein percent, acid detergent fiber, cellulose content and lignin content.

Cluster-III: This cluster consisted lowest number of genotype (1) i.e. IS-14241. This cluster had highest cluster mean for days to flowering, days to maturity, number of leaves, number of nodes, leaf length, flag leaf length, panicle length, panicle width, green fodder yield per plant, dry fodder yield per plant, hydrocyanic acid content, protein percent, neutral detergent fiber and hemicelluloses content.

Cluster-IV: This cluster consisted of twenty seven genotypes viz., SEVS-1, IS-4726-2, IS-2101, IS-25419-2, SMC-5, IS-25419-1, IS-1478, IS-23988, IS-5434-1, IS-6045, IS-14278-1, IS-6953, IS-7002, IS-21977, UTMC-523, IS-15008- 1, CS-3541-1, RS-673, IS-20740, IS-20782, IS-23948-1, IS-20399, JJ-1041, Pant Chari-5, IS-21622, IS-21461, and SSG-59-3. . This cluster had high cluster mean for leaf width, leaf area, flag leaf width, leaf: stem ratio, total soluble solids, protein percent, cellulose content and shoot fly incidence.

Cluster-V: This cluster had thirty five genotypes viz., UTMC-531, ESRK-7, SSG-227, CSV-14, SPV-1749, UPChari-1, EJN-37, EJN-54, SMC-2, SMC-6, Nizamabad, (SDSL-92101 x IS-3359) x Pant Chari-5, UP Chari-2, UPFS-38, IS-3359, PC-121, Pant Chari-3, IS-3199, GM-1378-1, IS-29794, GGUV-55, GMC-1422, SPV-1252, SPV-1616, SPV-1750, SST-4, SPV-1753, SRF-285, R-74(09R-AGR-26), R-77 (09R-AGR-26), RAJ-21, R-72(09R-AGR-23), UPFS-39, R-73 (09- AGR-24), and R-255 (09R-SS-26). This cluster had high cluster mean for number of leaves, leaf length, leaf width, leaf area, flag leaf length, flag leaf width, stem girth, 1000-grains weight, grain yield per plant, dry matter percent, in-vitro dry matter disappearance, silica content and zonate leaf spot.

Cluster-VI: This cluster had only three genotypes viz., CSV-19, CSV-24SS, and CHS-22SS. This cluster had high cluster mean for number of nodes, leaf width, leaf area, flag leaf width, internodal length, leaf: stem ratio, 1000 grains weight, green fodder yield, dry fodder yield per plant, total soluble solids, in-vitro dry matter disappearance, acid detergent fiber and hemicelluloses content.

Cluster-VII: This cluster consisted of thirty six genotypes viz., IS-2549-3, ICSR-93023, EJN-73, SMC-9, HC-260, GP-2011-372, E-1, ESRK-10, UPFS-35, E-25, E-105, EJN-59, ESRK-12, ESRK-16, SSG-223, 1910(08BZL-01-32-4), 1946 (08RLD-01-5-3), 1941(08RLD-01-5-3), R-72

(09R-AGR-23), EJN-58, PM-98019-2, GD-68717-1, UPFS-34, IS-14756, RAJ-16, EJN-49, EJN-68, IS-3821, HC-136, RAJ-32, EP-122, E-7, E-28, ESRK-4, EJN-39, and IS-12735. . This cluster had high cluster mean for days to flowering, days to maturity, plant height, stem girth, 1000-grains weight, grain yield per plant and anthracnose.

Cluster-VIII: This cluster was marked with highest number of genotypes (50) viz., ART-1008, UPFS-38 x IS-7002, SSG-219, SSG-256, IS-3313, SSG-244, UPFS-37 x UPMC-6, Rajasthan Local, GGUB-25, PC-23 x (SDSL-92101 x UPFS-23), EG-11, SRF-286, SL-44, SPV-462, UPFS-40, SMC-7, EJN-51, SPV-1754, UPMC-504 x UPMC-8, UTFS-48, GP-2011-44-1, UTFS-49, SMC-11, IS-15680, UPFS-36 x Pant Chari-6, IS-3821, SMC-18, HC-171, SMC-17, SSG-245, UPMC-503 x (SDSL-92101 x UPFS-23), ESRK-29, EJN-67, SSG-221, Ramkel, MP Chari, EJN-38, SSG-236, ESRK-27, SSG-241, SSG-250, SSG-224, SSG-256, SMC-3, SSG-227, SSG-243, SSG-248, SSG-234-1, SSG-253, and SSG-226. This cluster had high cluster mean for plant height, stem girth, internodal length, panicle length, panicle width and neutral detergent fiber.

Cluster-IX: This cluster had forty six genotypes viz., EJ-3, RAJ-9-1, EJ-42, C-43, RAJ-15, IS-313, Pant Chari-6, RS-29, UPFS-36(Pant Chari-7), UTMC-532, IS-3314, IS-3345, IS-3145, EA-11, IS-12956, GGUB-27, IS-699, RAJ-20, Pant Chari-5 x UPMC-512 , CSV-21F, SSV-74, SSG-304, IS-4307, SMC-12, ICSV-95119-1-2, 77113, IS-639, IS-29691, SMC-10, IS-31861, ESRK-26, IS-3359, SSG-226, SSG-225-1, UPFS-38 x SSG-59-3, SSG-225-2, IS-6193, IS-21602-1, IS-3237-2, IS-14298-1, NSSV-259, IS-14333-1, IS-18008-2, IS-22241, PSSV-49, and GMS-1338. This cluster had high cluster mean for internodal length, neutral detergent fiber, hemicelluloses content and zonate leaf spot.

Cluster-X: This cluster consisted of eleven genotypes viz., EJ-19, EJ-26, EJ-27, EJ-40, EJ-25, IS-25733, EJ-30, IS-4925, IS-33096, CSV-17, and IS-23992. This cluster had high cluster mean for leaf: stem ratio, dry matter percent, hydrocyanic acid content, total soluble solids, invitro dry matter disappearance and lignin content.

Cluster-XI: This cluster had twenty four genotypes viz., EJ-19, EJ-15, EJ-48-1, EJN-62, IS-14816, EJN-57, EJN-56, EJN-60, EJN-63, EJN-64, GGUB-36, IS-29314, GP-2011-110-1,

EJ-30, EP-135, EP-124, EJ-24, EJN-48-2, EJ-30, EJN-43, EJN-45, GP-2011-18-2, EJN-47, and EJN-52. This cluster had highest cluster mean for hydrocyanic acid content.

The pattern of distribution of genotypes in different cluster exhibited that geographical diversity was not related to genetic diversity as genotypes of same geographical region were grouped into different clusters and vice-versa Deep et al. [23] Karadi and Kajjidoni [24] Kavya et al. [25] Thant et al. [26] Umakanth et al. [27] Deep et al. [28] Kanbar et al. [29] Varaprasad and Sridhar (2020), Navya et al. [30] Sameera et al. [31] Pal et al. [32] Rohilla et al. [33] and Rathod et al*.* [34].

3.2 Average Intra and Inter Cluster Distances

The intra-cluster and inter-cluster distances were calculated to determine the genetic relationship between members of different clusters and among the individuals within a cluster. The intracluster and inter-cluster distances has been represented in Table 2. Inter-cluster distance is the main criterion for the selection of genotypes [35]. The genotypes belonging to those clusters having maximum inter-cluster distance are genetically more divergent and hybridization between these genotypes of different clusters is likely to produce wide range of variability with desirable individuals in segregating generations Damor et al. [36] Prasad et al. [37] Tesfaye [38] Ahlawat et al. [39] More et al. [40] Swamy et al. [41] Deep et al. [42] Karadi and Kajjidoni [24] Kavya et al. [25] Thant et al. [26] Umakanth et al. [27] Deep et al. [28] Kanbar et al. [29] Varaprasad and Sridhar (2020).

3.2.1 Intra-cluster distance

The maximum intra-cluster distance was observed in cluster-IV (54.652) suggesting highest genetic diversity among genotypes of this cluster in comparison to other clusters followed by cluster-VII (52.193), cluster-V (50.732), cluster-I (50.481), cluster-IX (49.583), cluster-XI (46.631), cluster-II (46.316), cluster-X (40.065), cluster-VIII (38.591), cluster-VI (30.366) whereas minimum intra-cluster distance was observed in cluster-III (0.000).

3.2.2 Inter-cluster distance

The genotypes belonging to those clusters having maximum inter-cluster distance are genetically more divergent and hybridization between these genotypes of different clusters is likely to produce wide variability with desirable individuals. The highest inter-cluster distance was observed between clusters III and VI (334.554), suggesting a distant relationship between members of these two clusters. Crossing these members is likely to result in greater genetic diversity in the segregating generation followed by clusters-III and X (321.316), clusters-III and XI (291.861), clusters-I and III (289.659), clusters-III and IV (280.335), clusters-III and IX (270.152), clusters-III and V (267.593), clusters-III and VII (260.334), clusters-III and VIII (245.298), clusters-II and III (218.755), clusters-II and VI (161.43), clusters-VI and X (137.992), clusters-VI and XI (125.337), clusters-VI and IX (116.215), clusters-I and VI (112.814), clusters-VI and VII (110.712), clusters-VI and VIII (108.318), clusters-IV and VI (95.978), clusters-II and X (92.231), clusters-V and X (91.74), clusters-VII and X (90.572), clusters-II and XI (88.733), clusters-V and VI (88.182), clusters-II and V (86.633), clusters-II and IV (84.875), clusters-II and VII (78.683), clusters-VIII and X (76.258), clusters-I and X (74.436), clusters-V and XI (74.418), clusters-I and II (74.38), clusters-II and IX (74.378), clusters-I and XI (73.742), clusters-IV and X (73.702), clusters-I and VII (72.079), clusters-III and XI (71.454), clusters-I and IX (71.197), clusters-V and IX (69.247), clusters-VII and IX (66.597), clusters-IV and VII (66.407), clusters-I and V (65.409), clusters-I and IV (64.572), clusters-X and XI (64.481), clusters-VII and XI (64.078), clusters-IV and IX (63.058), clusters-II and VIII (62.93), clusters-V and VII (62.632), clusters-VIII and XI (62.178), clusters-IX and X (62.109), clusters-IV and V (61.652), clusters-I and VIII (61.239), clusters-IX and XI (61.224), clusters-IV and VIII (58.536), clusters-VII and VIII (54.901**),**clusters V and VIII (53.071) whereas lowest inter-cluster distance was observed between clusters-VIII and IX (52.512) suggested a closer relationship between these two clusters and low degree of genetic diversity among the genotypes. Presence of substantial genetic diversity among the genotypes screened in the present study indicated that this material may serve as a good source for selecting the diverse parents for hybridization programme. In order to increase the possibility of isolating good trangressivesegregants in the segregating generations it would be logical to attempt crosses between the diverse genotypes belonging to clusters separated by large inter-cluster distances.

3.3 Cluster Mean for Different Characters

Cluster means were calculated for all the yield and quality traits along with some biochemical traits which exhibited considerable differences among the clusters. The mean performance of the clusters was used to select genetically diverse and agronomically superior genotypes under present study (Table 3, Fig. 2, Fig. 3 and Fig. 4).

The maximum cluster mean for days to flowering was observed in cluster-III (85.100) followed by cluster-II (78.400), cluster-VII (76.900), cluster-IV (67.200), cluster-I (66.700), cluster-VIII (63.300), cluster-VI (63.200), cluster-IX (63.200), cluster-XI (61.500), cluster-V (60.900) whereas minimum by cluster-X (55.000).The highest cluster mean for days to maturity was exhibited by cluster-III (147.000), cluster-VII (142.000), cluster-II (141.000), cluster-VI (133.000), cluster-IV (129.000), cluster-I (129.000), cluster-VIII (129.000), cluster-XI (126.000), cluster-IX (125.000), cluster-V (123.000) whereas lowest cluster mean for days to maturity was exhibited by cluster-X (121.000).

The maximum cluster mean for number of leaves was observed in cluster-III (20.000) followed by cluster-II (19.000), cluster-V (18.000), cluster-VII (18.000), cluster-VIII (18.000), cluster-VI (18.000), cluster-I (16.000), cluster-IX (16.000), cluster-XI (16.000), cluster-IV (15.000) whereas minimum by cluster-X (11.000).The highest cluster mean for number of nodes was exhibited by cluster-III (18.640), cluster-VI (17.620), cluster-II (17.370), cluster-V (16.560), cluster-VII (16.260), cluster-VIII (15.820), cluster-I (15.100), cluster-IX (14.890), cluster-XI (14.200), cluster-IV (14.100) whereas lowest cluster mean for number of nodes was exhibited by cluster-X (10.860).

The maximum cluster mean for plant height was observed in cluster-II (414.000) followed by cluster-VIII (405.000), cluster-VII (385.100), cluster-III (368.000), cluster-V (362.000), cluster-I (358.000), cluster-IV (350.000), cluster-IX (347.000), cluster-XI (340.000), cluster-X (298.000) whereas minimum by cluster-VI (291.000).The highest cluster mean for leaf length was exhibited by cluster-III (99.700), cluster-V (93.390), cluster-II (92.620), cluster-IV (88.020), cluster-III (86.040), cluster-I (85.370), cluster-VII (84.310), cluster-VI (84.300), cluster-IX (75.620), cluster-XI (73.790) whereas lowest cluster mean for leaf length was exhibited by cluster-X (72.170).

Table 1. Distribution of genotypes into different clusters during *Kharif* **2019**

	Cluster-I	Cluster-II	Cluster-III	Cluster-IV	Cluster-V	Cluster-VI	Cluster-VII	Cluster-VIII	Cluster-IX	Cluster-X	Cluster-XI
Cluster-I	50.481	74.38	289.659	64.572	65.409	112.814	72.079	61.239	71.197	74.436	73.742
Cluster-II		46.316	218.755	84.875	86.633	161.43	78.683	62.93	74.378	92.231	88.733
Cluster-III			0	280.335	267.593	334.554	260.334	245.298	270.152	321.316	291.861
Cluster-IV				54.652	61.562	95.978	66.407	58.536	63.058	73.702	71.454
Cluster-V					50.732	88.182	62.632	53.071	69.247	91.74	74.418
Cluster-VI						30.336	110.712	108.318	116.215	137.992	125.337
Cluster-VII							52.193	54.901	66.597	90.572	64.078
Cluster-VIII								38.591	52.512	76.258	62.178
Cluster-IX									49.583	62.109	61.224
Cluster-X										40.065	64.481
Cluster-XI											46.631

Table 2. Intra and inter cluster distances between the clusters based on hierarchical cluster analysis of sorghum germplasm during *Kharif* **2019**

Table 3. Cluster means for different characters in sorghum germplasm during *Kharif* **2019**

DF= Days to 50% flowering, DM= Days to maturity, NL= Number of leaves per plant, NN= Number of nodes, PH= Plant height (cm), LL= Leaf length (cm), LW= Leaf width (cm), LA= Leaf area (cm²), FLL= Flag leaf length (cm), FLW= Flag leaf width (cm), SG= Stem girth (cm).

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Continued											
	INL	PL	PW	∟:S∶	TGW	GYP	GFY	DFY	$DM\%$	TSS%	HCN
Cluster-I	29.581	25.906	13.021	0.332	23.401	91.245	320.271	116.739	38.683	6.721	91.898
Cluster-II	27.588	26.765	16.242	0.329	11.329	43.226	385.67	143.558	38.382	6.316	88.498
Cluster-III	20.064	37.761	21.026	0.325	13.674	49.769	917.729	472.253	35.359	7.129	112.846
Cluster-IV	30.309	21.904	10.593	0.352	23.57	91.528	331.357	116.663	35,908	12.027	91.101
Cluster-V	27.477	25.156	11.383	0.324	28.323	111.204	367.003	133.47	38.888	6.976	87.628
Cluster-VI	36.314	20.648	11.91	0.448	39.71	103.899	447.29	183.378	42.945	12.969	73.57
Cluster-VII	25.632	19.782	11.05	0.33	26.537	103.512	326.097	118.397	37.792	5.987	91.683
Cluster-VIII	30.675	29.685	15.984	0.337	21.235	83.287	334.915	118.615	37.568	5.75	92.714
Cluster-IX	32.079	24.37	12.701	0.351	21.928	85.401	296.728	106.556	37.002	9.05	85.54
Cluster-X	30.349	16.725	6.892	0.369	15.568	60.496	241.79	94.781	39.143	9.472	101.112
Cluster-XI	29.364	13.242	6.333	0.337	20.727	79.548	301.921	109.743	38.462	4.828	95.928

INL= Internodal length (cm), PL= Panicle length (cm), PW= Panicle width (cm), L:S= Leaf:stem ratio, TGW= 1000-grains weight (gm), GYP= Grain yield per plant (gm), GFY= Green fodder yield per plant (gm), DFY= Dry fodder yield per plant (gm), DM= Dry matter (%), TSS= Total soluble solids (%), HCN= HCN content (ppm)

PP= Protein content (%), IVDMD= Iin-vitro dry matter disappearance (IVDMD), NDF= Neutral detergent fiber, ADF=Acid detergent fiber (%), C= Cellulose (%), L= Lignin (%), S= Silica (%), HC= Hemicellulose, A= Anthracnose (%), ZLS= Zonate Leaf Spot, SFI= Shoot Fly Index (%)

Fig. 1. Distribution of genotypes into different clusters during *Kharif* **2019**

Fig. 2. Cluster means for DF, DM, NL, NN, PH, LL, LW, LA, FLL, FLW, and SG

Fig. 3. Cluster means for INL, PL, PW, L:S, TGW, GYP, GFY, DFY, DM%, TSS%, and HCN

Fig. 4. Cluster means for PP, IVDMD, NDF, ADF, C, L, S, HC, A, ZLS, and SFI

The maximum cluster mean for leaf width was observed in cluster-VI (11.300) followed by cluster-V (9.721), cluster-IV (9.255), cluster-VII (8.669), cluster-I (8.487), cluster-VIII (8.372), cluster-XI (7.960), cluster-IX (7.494), cluster-X (7.005), cluster-II (5.478) whereas minimum by cluster-III (4.881).The highest cluster mean for leaf area was exhibited by cluster-VI (733.000), cluster-V (559.000), cluster-IV (526.000), cluster-VII (463.000), cluster-I (457.000), cluster-VIII (455.000), cluster-XI (379.000), cluster-IX (357.000), cluster-III (333.000), cluster-X (315.000) whereas lowest cluster mean for leaf area was exhibited by cluster-II (308.000).

The maximum cluster mean for flag leaf length was observed in cluster-III (51.990) followed by cluster-V (47.970), cluster-II (46.970), cluster-IV (44.890), cluster-VIII (44.030), cluster-I (43.780), cluster-VI (43.180), cluster-VII (43.110), cluster-
IX (38.220), cluster-XI (38.030) whereas IX (38.220), cluster-XI (38.030) minimum by cluster-X (36.740).The highest cluster mean for flag leaf width was exhibited by cluster-VI (9.000), cluster-V (5.000), cluster-IV (5.000), cluster-VII (4.000), cluster-I (4.000), cluster-XI (4.000), cluster-VIII (4.000), cluster-IX (4.000), cluster-X (4.000), cluster-III (3.000) whereas lowest cluster mean for flag leaf width was exhibited by cluster-II (3.000).

The maximum cluster mean for stem girth was observed in cluster-VII (2.821) followed by cluster-VIII (2.739), cluster-V (2.667), cluster-XI (2.656), cluster-I (2.525), cluster-IV (2.403), cluster-VI (2.323), cluster-IX (2.247), cluster-X (2.220), cluster-II (2.105) whereas minimum by cluster-III (2.082).The highest cluster mean for inter-nodal length was exhibited by cluster-VI (36.300), cluster-IX (32.100), cluster-VIII (30.700), cluster-X (30.300), cluster-IV (30.300), cluster-I (29.600), cluster-XI (29.400), cluster-II (27.600), cluster-V (27.500), cluster-VII (25.600) whereas lowest cluster mean for intermodal length was exhibited by cluster-III (20.100).

The maximum cluster mean for panicle length was observed in cluster-III (38.000) followed by cluster-VIII (30.000), cluster-II (27.000), cluster-I (26.000), cluster-V (25.000), cluster-IX (24.000), cluster-IV (22.000), cluster-VI (21.000), cluster-VII (20.000), cluster-X (17.000) whereas minimum by cluster-XI (13.000).The highest cluster mean for panicle width was exhibited by cluster-III (21.000), cluster-II (16.000), cluster-VIII (16.000), cluster-I (13.000), cluster-IX (13.000), cluster-VI (12.000), cluster-V (11.000), cluster-VII (11.000), cluster-IV (11.000), cluster-X (7.000)

whereas lowest cluster mean for panicle width was exhibited by cluster-XI (6.000).

The maximum cluster mean for leaf:stem ratio was observed in cluster-VI (0.448) followed by cluster-X (0.369), cluster-IV (0.352), cluster-IX (0.351), cluster-VIII (0.337), cluster-XI (0.337), cluster-I (0.332), cluster-VII (0.330), cluster-II (0.329), cluster-III (0.325) whereas minimum by cluster-V (0.324).The highest cluster mean for 1000-grains weight was exhibited by cluster-VI (39.700), cluster-V (28.300), cluster-VII (26.500), cluster-IV (23.600), cluster-I (23.400), cluster-IX (21.900), cluster-VIII (21.200), cluster-XI (20.700), cluster-X (15.600), cluster-III (13.700) whereas lowest cluster mean for 1000-grains weight was exhibited by cluster-II (11.300).

The maximum cluster mean for grain yield per plant was observed in cluster-V (111.2) followed by cluster-VI (103.900), cluster-VII (103.500), cluster-IV (91.530), cluster-I (91.250), cluster-IX (85.400), cluster-VIII (83.290), cluster-XI (79.550), cluster-X (60.500), cluster-III (49.770) whereas minimum by cluster-II (43.250).The highest cluster mean for green fodder yield per plant was exhibited by cluster-III (917.717), cluster-VI (447.300), cluster-II (385.700), cluster-V (367.000), cluster-VIII (334.900), cluster-IV (331.400), cluster-VII (326.100), cluster-I (320.300), cluster-XI (301.900), cluster-IX (296.700) whereas lowest cluster mean for green fodder yield per plant was exhibited by cluster-X (241.800).

The maximum cluster mean for dry fodder yield was observed in cluster-III (472.000) followed by cluster-VI (183.000), cluster-II (144.000), cluster-V (133.000), cluster-VIII (119.000), cluster-VII (118.000), cluster-I (117.000), cluster-IV (117.000), cluster-XI (110.000), cluster-IX (107.000) whereas minimum by cluster-X (94.800).The highest cluster mean for dry matter percent was exhibited by cluster-VI (42.950), cluster-X (39.140), cluster-V (38.890), cluster-I (38.680), cluster-XI (38.460), cluster-II (38.380), cluster-VII (37.790), cluster-VIII (37.570), cluster-IX (37.000), cluster-IV (35.910) whereas lowest cluster mean for dry matter percent was exhibited by cluster-III (25.360).

The maximum cluster mean for total soluble solids was observed in cluster-VI (13.000) followed by cluster-IV (12.000), cluster-X (9.000), cluster-IX (9.000), cluster-III (7.000), cluster-V (7.000), cluster-I (7.000), cluster-II (6.000), cluster-VII (6.000), cluster-VIII (6.000) whereas

minimum by cluster-XI (5.000).The highest cluster mean for hydrocyanic acid content was exhibited by cluster-III (112.850), cluster-X (101.110), cluster-XI (95.928), cluster-VIII (92.714), cluster-I (91.898), cluster-VII (91.683), cluster-IV (91.101), cluster-II (88.498), cluster-V (87.628), cluster-IX (85.540) whereas lowest cluster mean for hydrocyanic acid content was exhibited by cluster-VI (73.570).

The maximum cluster mean for protein percent was observed in cluster-III (16.400) followed by cluster-II (12.800), cluster-IV (12.400), cluster-I (12.200), cluster-VIII (12.000), cluster-VII (11.900), cluster-IX (11.400), cluster-V (11.400), cluster-XI (11.100), cluster-X (10.000) whereas minimum by cluster-VI (7.250).The highest cluster mean for in-vivo dry matter digestibility was exhibited by cluster-VI (59.000), cluster-X (58.000), cluster-V (58.000), cluster-I (58.000), cluster-II (57.000), cluster-III (57.000), cluster-VIII (57.000), cluster-VII (56.000), cluster-IX (56.000), cluster-XI (56.000) whereas lowest cluster mean for in-vivo dry matter digestibility was exhibited by cluster-IV (54.000).

The maximum cluster mean for neutral detergent fiber was observed in cluster-III (60.000) followed by cluster-VIII (57.000), cluster-IX (57.000), cluster-VII (57.000), cluster-XI (56.000), cluster-V (56.000), cluster-IV (56.000), cluster-II (56.000), cluster-X (56.000), cluster-I (54.000) whereas minimum by cluster-VI (52.000).The highest cluster mean for acid detergent fiber was exhibited by cluster-I (39.730), cluster-VI (37.690), cluster-II (37.590), cluster-X (36.500), cluster-IV (36.500), cluster-V (36.280), cluster-VIII (35.660), cluster-XI (35.480), cluster-VII (35.350), cluster-IX (35.090) whereas lowest cluster mean for acid detergent fiber was exhibited by cluster-III (32.780).

The maximum cluster mean for cellulose content was observed in cluster-I (31.300) followed by cluster-II (30.400), cluster-IV (30.100), cluster-IX (29.700), cluster-VIII (29.700), cluster-V (29.700), cluster-XI (29.400), cluster-IX (29.300), cluster-VII (29.300), cluster-VI (28.400) whereas minimum by cluster-III (28.100).The highest cluster mean for lignin content was exhibited by cluster-I (7.204), cluster-X (6.484), cluster-II (6.000), cluster-IV (5.810), cluster-IX (5.498), cluster-V (5.474), cluster-XI (5.429), cluster-VII (5.370), cluster-III (5.206), cluster-VIII (5.171) whereas lowest cluster mean for lignin content was exhibited by cluster-VI (5.063).

The maximum cluster mean for silica content was observed in cluster-I (2.615) followed by cluster-IV (2.321), cluster-V (2.307), cluster-X (2.183), cluster-II (2.180), cluster-VI (2.156), cluster-XI (2.060), cluster-VIII (2.048), cluster-VII (1.968), cluster-IX (1.817) whereas minimum by cluster-III (1.303).The highest cluster mean for hemicelluloses content was exhibited by cluster-III (30.100), cluster-IX (25.400), cluster-VI (25.100), cluster-VII (24.700), cluster-VIII (24.600), cluster-XI (24.000), cluster-V (23.200), cluster-IV (22.600), cluster-II (22.000), cluster-X (21.900) whereas lowest cluster mean for hemicelluloses content was exhibited by cluster-I (17.800).

The maximum cluster mean for anthracnose was observed in cluster-XI (65.56) followed by cluster-III (27.910), cluster-VII (27.250), cluster-I (24.060), cluster-VIII (21.870), cluster-IX (20.300), cluster-V (17.430), cluster-IV (15.140), cluster-II (15.020), cluster-X (14.680) whereas minimum by cluster-VI (8.406).The highest cluster mean for zonate leaf spot was exhibited by cluster-IV (11.000), cluster-V (11.000), cluster-IX (10.000), cluster-XI (8.000), cluster-VI (8.000), cluster-VII (8.000), cluster-VIII (7.000), cluster-I (7.000), cluster-II (4.000), cluster-X (10.300) whereas lowest cluster mean for zonate leaf spot was exhibited by cluster-III (2.000).The maximum cluster mean for shoot fly incidence was observed in cluster-XI (36.169) followed by cluster-VII (35.928), cluster-IV (23.360), cluster-I (21.052), cluster-V (18.010), cluster-VI (17.823), cluster-X (17.543), cluster-IX (16.746), cluster-VIII (15.240), cluster-II (11.199) whereas minimum by cluster-III (5.656).

Classification of the germplasm in to divergent groups based on inter cluster distances, per se performance and selection of parents from diverse clusters was reported in several studies Kumar et al*.* [43] Rahman et al*.* (2015),Usha and Rekha [44] Doijad et al. [45] Jain and Patel [46] Damor et al. [47] Prasad et al. [37] Tesfaye [38] Ahlawat et al. [48] More et al. [40] Swamy et al. [41] Deep et al. [23] Karadi and Kajjidoni [24] Kavya et al. [25] Thant et al. [26].

Crosses suggesting parents belonging to most divergent clusters would be expected to manifest maximum heterosis and also wide variability of genetic architecture [26,28]. These results of our present study areare somewhat in accordance with the findings of, Varaprasad and Sridhar (2020), Navya et al. [30] Sameera et al. [31] Pal et al. [32] Rohilla et al. [49] and Rathod et al*.* [50] Raghavendra et al. [51].

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4. SUMMARY AND CONCLUSION

It can be summarized and concluded from the above discussion that there is a presence of huge amount of genetic variability in the material under investigation as seven different clusters were obtained and intra cluster distance were found to be lesser than the inter cluster distances. The genotypes were grouped into XI distinct non-overlapping clusters.The cluster-VIII (50) consisted of highest number of genotypes whereas lowest numbers of genotypes were grouped into cluster-III (1).The maximum intracluster distance was observed in cluster-IV (54.652) suggesting highest genetic diversity among genotypes of this cluster in comparison to other clusters whereas minimum intra-cluster distance was observed in cluster-III (0.000).The clusters with high intra-cluster distances suggested that genotypes in these clusters were more genetic diverse than the genotypes in other clusters with low intra-cluster distances. Low intra-cluster distance suggested a closer relationship between these two clusters and low degree of genetic diversity among the genotypes whereas high intra cluster distance represented high amount of genetic diversity among members of same cluster. The highest inter-cluster distance was observed between clusters-III and VI (334.554)suggested distant relationship between members of these two clusters and upon crossing the members of these two clusters will give more genetic diversity in segregating generationwhereas minimum inter-cluster distance was observed between clusters-VIII and IX (52.512) suggested a closer relationship between these two clusters and low degree of genetic diversity among the genotypes.Presence of substantial genetic diversity among the genotypes screened in the present study indicated that this material may serve as a good source for selecting the diverse parents for hybridization programme. To increase the likelihood of isolating desirable transgressive

segregants in segregating generations, it would be logical to attempt crosses between diverse genotypes from clusters separated by large intercluster distances.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of manuscripts.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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