



Determining the Genetic Divergence for Biometrical Attributes in Cluster Bean (*Cyamopsis tetragonoloba* (L.) Taub. Accessions through Mahanalobis D^2 Statistics

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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

Aims: Genetic divergence plays a crucial part in determining the selection of genotypes for any breeding programme in order to generate superior segregants. Assessing the local or related genotypes for divergence, imposes a necessity to conduct experiments on specific agro-ecological situations to breed specific varieties suitable for vegetable and gum purpose.

Study Design: Randomized Block Design

Place and Duration of Study: An elaborate study was performed in January to April 2022 at the field in Poothurai village of Villupuram District, Tamil Nadu to evaluate and screen the cluster bean germplasms.

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Methodology: A total of 102 genotypes (including Pusa Navbahar - a check variety) procured from various regions of the country were utilized. Observations were recorded for growth attributes (plant height; number of branches per plant; days to maturity), flowering and pod characters (days to 50 % flowering; number of clusters per plant; number of pods per cluster; pod length; pod girth), yield parameters (pod weight; number of pods per plant; total pod yield per plant), seed characters (number of seeds per pod; hundred seed weight; total seed yield per plant) and physiological components (guar gum; protein and fibre) on five randomly selected plants.

Results: It divulged a paramount of diversity with the genotypes being grouped into 14 clusters based on 17 parameters. Seven clusters had more than 2 genotypes, and the remaining were solitary. Yield parameters such as total per plant yield (22.20%), pods per cluster (15.30%), and clusters per plant (14.80%) contributed to the highest percentage diversity. The distance between intra clusters ranges from 0.00 to 2031.87. The maximum distance between clusters observed is between cluster V and cluster IX (15980.81).

Conclusion: Of the 102 genotypes, Cluster XIV having the sole member Pusa Navbahar performed better for six out of seventeen parameters observed for the mean of clusters. With respect to the above results obtained and *per se* performance of the landraces, selection can be made to obtain superior cluster bean recombinants.

Keywords: Cluster bean; multivariate statistic; genetic diversity; Mahalanobis D^2 ; yield attributes.

1. Introduction

Cluster bean, an underexploited leguminous vegetable belonging to Leguminaceae family is highly drought tolerant, well adapted to poor and erratic climatic conditions (Santhosha et al., 2017). It is a potential vegetable grown for its tender pods that are nutritionally affluent containing good sources of energy, protein, fats, carbohydrates, vitamins, calcium and iron (Choyal et al., 2017). It has also gained attention for its fine endospermic gum, making it a potential cash crop for both industrial and pharmaceutical purposes (Kgasudi et al., 2020). India holds a superlative share of 80 % of total gum production in the world. Owing to its multi-purpose nature, it is used as a forage crop and for fixing the atmospheric nitrogen thereby improving soil fertility (Remzeena et al., 2021). According Gopala Krishnan et al., 2011 to, Adak guar, which is produced primarily for its gum content and as fodder in the arid regions of North Western India, has been found to act as a bridge between the wild and domesticated ones. The cluster bean under cultivation shows typical signs of domestication syndrome, including dense growing habit, indehiscent edible pods, seed dormancy loss, cylindrical and large seeds.

Assessing the local or related genotypes for divergence, imposes a necessity to conduct experiments on specific agro-ecological situations to breed specific varieties suitable for vegetable and gum purpose. It is vital to breed for deviations that are suited to specific agro-ecological environments for seed and gum

purposes in order to assess local or related genotypes and figure out divergence. With very low divergence, only modest results can be achieved. The breeder will need to increase germplasm enrichment or may have to resort to increased divergence through mutations, polyploidy breeding and hybridization programme (Wankhade et al., 2017). Genetic diversity of elite germplasm opens up an array of possibilities in determining better parents and generating superior hybrids. While planning the breeding goals, having an understanding of the genetic divergence of different parameters, especially those that impact quantitative and qualitative traits, would be significant. Mahalanobis statistics indicates a scale of magnitude for the divergence between the two genotypes that are being assessed (Mahalanobis, 1936). Hence an analysis was done out to determine the degree of genetic diversity between 102 accessions for 17 biometric traits.

2. Materials and Methods

An elaborate study was performed in January to April 2022 at the field in Poothurai village of Villupuram District, Tamil Nadu to evaluate and screen the cluster bean germplasms. The experimental plot is situated at 11.9667° N and 79.7596° E with 23 MSL altitude. A total of 102 genotypes (including Pusa Navbahar - a check variety) procured from various regions of the country were utilized. The accessions were evaluated under Randomized Block Design (RBD) planted at a spacing of 45 cm between the ridges and 15 cm between the plants. The

genotypes were planted in individual rows of 5 m length. Optimum management practices as given by Tamil Nadu Agricultural University were followed during its entire growing period.

Observations were recorded for growth attributes (plant height; number of branches per plant; days to maturity), flowering and pod characters (days to 50 % flowering; number of clusters per plant; number of pods per cluster; pod length; pod girth), yield parameters (pod weight; number of pods per plant; total pod yield per plant), seed characters (number of seeds per pod; hundred seed weight; total seed yield per plant) and physiological components (guar gum; protein and fibre) on five randomly selected plants. The multivariate tool, Mahankali's D^2 statistic Rao, (1952) was implemented to estimate genetic divergence in cluster bean landraces. Tocher's method was used to determine the group constellation. Other genetic divergence parameters like percentage contribution of individual traits towards divergence, average intra and inter cluster distances, cluster mean values were assessed.

3. Results and Discussion

Diversity at genetic level provides greater scope for exploiting the variation among the genotypes making it a pre-requisite in any breeding studies. D^2 analysis is a widely used multivariate statistic to measure genetic diversity across a set of genotypes. The evaluation of 102 genotypes including the check variety showed significant variations for growth, flowering, yield and physiological parameters. Analysis of variance (ANOVA) exhibited highly positive significant differences in all genotypes for all parameters studied both at 1 % and 5% level of significance, suggesting substantial amount of diversity among the genotypes.

3.1 Group Constellations of Cluster Bean Accessions

With respect to the relative magnitude of genetic distance, the genotypes were clustered into 14 groups (Table 1) indicating vast amount of variation among the genotypes. This large grouping was also reported by Manivannan and Anandakumar, 2013. Out of fourteen clusters, cluster I was the largest (54) followed by cluster II (13) and cluster VI (8) genotypes. Cluster III, cluster IV, Cluster V and Cluster VII had 4, 7, 6 and 3 accessions respectively. The remaining seven clusters from Cluster VIII to Cluster XIV

were solitary. Cluster VII comprising of 7 genotypes are those that are highly recommended for vegetable purpose. Cluster XIV had only one genotype, Pusa Navbar, a high yielding vegetable genotype. Results complementary to this was reported by Kumar et al., (2014). Formation of solitary clusters may be a result of extreme selection or prevention in gene flow Wankhade et al. (2017) making them extraordinary contributors of one or more characters (Choyal et al., 2017). The results also revealed that agro-morphological grouping of genotypes is not necessarily correlated with their geographic origin. This was in accordance with that of Sultan et al., (2012). Through application of clustering technique, it is feasible to compare all population pairs that could possibly exist inside a given group precisely, leading to authentic crosses.

3.2 Inter and Intra Cluster Distances

The D^2 values obtained after grouping the genotypes into fourteen clusters indicated the presence of appreciable amount of genetic diversity (Table 2). The distance among the clusters ranged from 0.00 to 2031.87. This was found to be maximum in Cluster I (2031.87) followed by Cluster VI (1697.01) and Cluster III (1600.01). The presence of wide intra cluster distance indicates tremendous scope in exchange of genetic materials among the accessions (Mishra et al., 2019). Seven clusters which were single member groups had intra-cluster distance values as 0. The inter cluster distance revealed maximum distance between Cluster V and Cluster IX (15980.81), closely followed by Cluster XI and Cluster XII (15105.61), Cluster XII and Cluster IV (14953.97), Cluster III and Cluster XII (14375.62) and Cluster IX and Cluster XI (13869.07). The distance between Cluster II and Cluster X was found to be least with 2852.03. The larger the magnitude of distance between the clusters, the more the genetic diversity. The narrow distance values indicated that the genotypes of the particular cluster were close to the other cluster pair (Vishnoi et al., 2017). The fact that the distance between clusters is greater than the distance within clusters suggests that there is significant variation between the genotypes. For obtaining pertinent segregants, genotypes from clusters demonstrating large inter-cluster distances must be selected. This indication helps in the development of heterotic hybrids. Similar records were also observed by Pathak et al., 2010, Rai and Dharmatti, 2013.

Table 1. Cluster bean accessions being grouped into different clusters

Cluster	Genotype Code (CT)	Genotype Name
I (54)	1 3 4 6 7 9 12 14 16 17 18 19 20 21 27 28 29 30 32 34 37 39 43 45 47 48 51 52 54 57 58 59 60 61 62 63 64 66 67 68 69 70 71 77 78 79 81 82 85 87 88 93 95 96	IC 28283, IC 34235, IC 34458, IC 39998, IC 40023, IC 40073 IC 40188, IC 102825, IC 102867, IC 116531, IC 116532, IC 116535, IC 116535, IC 116537, IC 116543, IC 116571, IC 116572, IC 116574 IC 116576, IC 116601 IC 116607, IC 116614, IC 116625, IC 116643, IC 116687 IC 116723, IC 116724, RGC 1002, RGC 1038, RGC 471, RGC 1017, RGC 986, RGC 1053, RGC 1003, RGC 197, HGS 898, HG 75, HGS 16, HGS 894, HGS 182, HGS 885, HG 258, HG 365, HG 832, GAUG 980, GAUG 14, GAUG 005, Kandan Bahar, NS 662, NCB Nandhini, Desi Guar, Naveen, Guar Kranti, Karnataka Local, Indore local
II (13)	2 5 8 10 11 13 15 22 23 26 46 84 92	IC 33704, IC 39993, IC 40040, IC 40161, IC 40182, IC 4050 BIC 102825, IC 102828, IC 116547, IC 116554, IC 116570, IC 116700, IC Anjali, Goma Manjari
III (4)	33 55 56 97	IC 116605, RGC 1066, RGC 936, Telangana Local
IV (7)	86 90 94 98 99 100 101	Yanam Local, Sivapuri Local, MDU 1, Thiruvanamalai Local, Chegalpattu Local, Pusa Sadabahar, Orthanad Local
V (6)	35 36 40 41 44 49	IC 116609, IC 116612, IC 116626, IC 116630, IC 116645, IC 116758
VI (8)	42 53 65 72 73 76 83 89	IC 116642, RGC 1055, HGS 100, HG 884, HG 870, GAUG 13, Selection 51, Ankur Rani
VII (3)	74 75 80	GAUG 982, GAUG 003, GAUG 004
VIII	24	IC 116559
IX	25	IC 116568
X	31	IC 116580
XI	38	IC 116620
XII	50	IC 116883
XIII	91	Lakshmi
XIV	102	Pusa Navbahar

Table 2. Intra-cluster (diagonal) and Inter-cluster distances among the fourteen clusters

Cluster	I	II	III	IV	V	VI	VII	VIII	IX	X	XI	XII	XIII	XIV
I	2031.87 (45.08)	4030.00 (63.48)	3053.91 (55.26)	4355.82 (66.00)	7921.47 (89.00)	3250.49 (57.01)	3986.36 (63.14)	5942.73 (77.09)	8771.79 (93.66)	3517.20 (59.31)	7153.90 (84.58)	9285.96 (96.36)	4386.92 (66.23)	7893.17 (88.84)
II		1517.21 (38.95)	5918.54 (76.93)	3773.53 (61.43)	10620.55 (103.06)	6681.11 (81.74)	5109.33 (71.48)	5216.22 (72.22)	3463.60 (58.85)	2852.03 (53.40)	8550.09 (92.47)	5487.69 (74.08)	5332.08 (73.02)	5333.55 (73.03)
III			1600.01 (40.00)	4014.72 (63.36)	13348.62 (115.54)	6518.15 (80.74)	6647.81 (81.53)	11366.32 (106.61)	10323.95 (101.61)	5224.71 (72.28)	9875.72 (99.38)	14375.62 (119.90)	8670.51 (93.12)	8553.62 (92.49)
IV				1376.26 (37.10)	12591.36 (112.21)	8119.64 (90.11)	5507.64 (74.21)	9284.38 (96.36)	6747.84 (82.15)	3385.08 (58.18)	8545.50 (92.44)	12257.32 (110.71)	6052.02 (77.79)	2654.38 (51.52)
V					1525.59	4625.84	3626.86	6586.28	15980.81	8035.86	3853.72	12079.53	6311.24	15316.70

Cluster	I	II	III	IV	V	VI	VII	VIII	IX	X	XI	XII	XIII	XIV
					(39.06)	(68.01)	(60.22)	(81.16)	(126.42)	(89.64)	(62.08)	(109.91)	(79.44)	(123.76)
VI						1697.01 (41.19)	4070.22 (63.80)	4039.71 (63.56)	12205.45 (110.48)	5044.04 (71.02)	7327.15 (85.60)	8763.19 (93.61)	3230.89 (56.84)	11738.85 (108.35)
VII							1479.36 (38.46)	5534.67 (74.40)	8943.22 (94.57)	3572.40 (59.77)	3111.91 (55.78)	9710.50 (98.54)	5310.23 (72.87)	8815.02 (93.89)
VIII								0.00 (0.00)	8449.88 (91.92)	4404.74 (66.37)	10820.49 (104.02)	5226.89 (72.30)	3329.40 (57.70)	9983.73 (99.92)
IX									0.00 (0.00)	6829.30 (82.64)	13869.07 (117.77)	4098.29 (64.02)	10135.71 (100.68)	8423.90 (91.78)
X										0.00 (0.00)	6523.29 (80.77)	9361.05 (96.75)	5188.05 (72.03)	4257.83 (65.25)
XI											0.00 (0.00)	15105.61 (122.90)	8668.27 (93.10)	10903.83 (104.42)
XII												0.00 (0.00)	7095.09 (84.23)	14953.97 (122.29)
XIII													0.00 (0.00)	7045.16 (83.94)
XIV														0.00 (0.00)

Table 3. Cluster means for different quantitative and qualitative characters

Parameters	I	II	III	IV	V	VI	VII	VIII	IX	X	XI	XII	XIII	XIV
PH	90.13	88.38	99.19	110.23	96.33	95.64	92.55	89.19	115.10	128.27	77.62	88.48	80.98	131.01
NOB	4.04	3.43	0.00	0.00	9.16	8.09	6.97	9.20	3.13	2.80	4.26	10.25	8.86	0.00
DUR	104.31	113.46	97.67	109.44	107.57	95.96	120.32	106.79	103.05	102.89	100.20	106.85	115.53	122.57
DFPF	40.50	41.70	39.49	40.53	44.97	40.07	42.33	48.26	46.50	42.50	45.20	45.50	41.35	41.40
NCP	16.87	14.52	13.19	19.44	45.26	21.65	28.34	14.21	12.00	21.52	57.25	10.80	17.00	19.26
NPPC	3.91	7.40	3.59	6.00	2.69	3.19	3.58	6.63	12.35	4.93	2.14	12.08	4.65	6.29
PL	5.75	5.69	6.12	9.93	4.97	6.15	5.10	5.87	5.18	8.13	4.69	4.39	11.34	13.60
PG	6.60	6.65	5.70	6.27	6.77	6.86	7.03	6.50	5.20	6.70	7.40	7.30	6.77	6.51
PW	1.49	1.80	1.30	1.73	1.71	1.28	1.11	1.30	1.11	1.61	1.50	0.92	1.83	2.86
NPPP	57.65	91.72	44.63	93.04	95.64	58.49	88.03	89.21	136.20	90.09	125.21	118.46	75.54	115.35
TPYPP	80.10	136.48	51.36	139.04	134.48	69.41	92.82	116.30	147.88	141.54	190.39	105.48	134.32	256.32
NSPP	6.59	6.34	6.73	8.34	7.14	6.74	6.83	7.87	8.91	4.80	7.00	6.15	8.62	9.59
HSW	2.89	2.99	2.85	3.22	2.81	2.98	3.18	3.60	3.03	2.66	2.51	2.72	2.78	3.40

Parameters	I	II	III	IV	V	VI	VII	VIII	IX	X	XI	XII	XIII	XIV
TSYPP	8.83	11.02	6.87	9.22	9.30	10.27	9.70	19.46	10.02	16.25	5.10	8.67	9.92	14.53
GUAR	24.33	26.32	25.73	21.73	24.59	26.29	21.57	27.23	31.62	30.61	32.60	31.12	22.10	25.61
C. PROTEIN	16.91	14.18	17.76	17.66	18.00	16.22	14.75	15.03	18.02	17.53	16.50	14.32	16.10	21.60
C. FIBRE	5.11	4.92	5.35	4.60	5.48	5.09	5.51	5.32	5.03	4.80	5.39	5.13	5.02	5.62

*PH: Plant Height; NOB: Number of primary branches; DUR: Duration; DFPF: Days to 50% flowering; NCPP: Number of clusters plant⁻¹; NPPC: Number of pods cluster⁻¹; PL: Pod length; PG: Pod girth; PW: Pod weight; TNPPP: Total number of pods plant⁻¹; TPYPP: Total pod yield plant⁻¹; NSPP: Number of seeds plant⁻¹; HSW: Hundred seed weight; TSYPP: Total seed yield plant⁻¹, Guar Gum; C. Protein: Crude Protein; C. Fibre: Crude Fibre

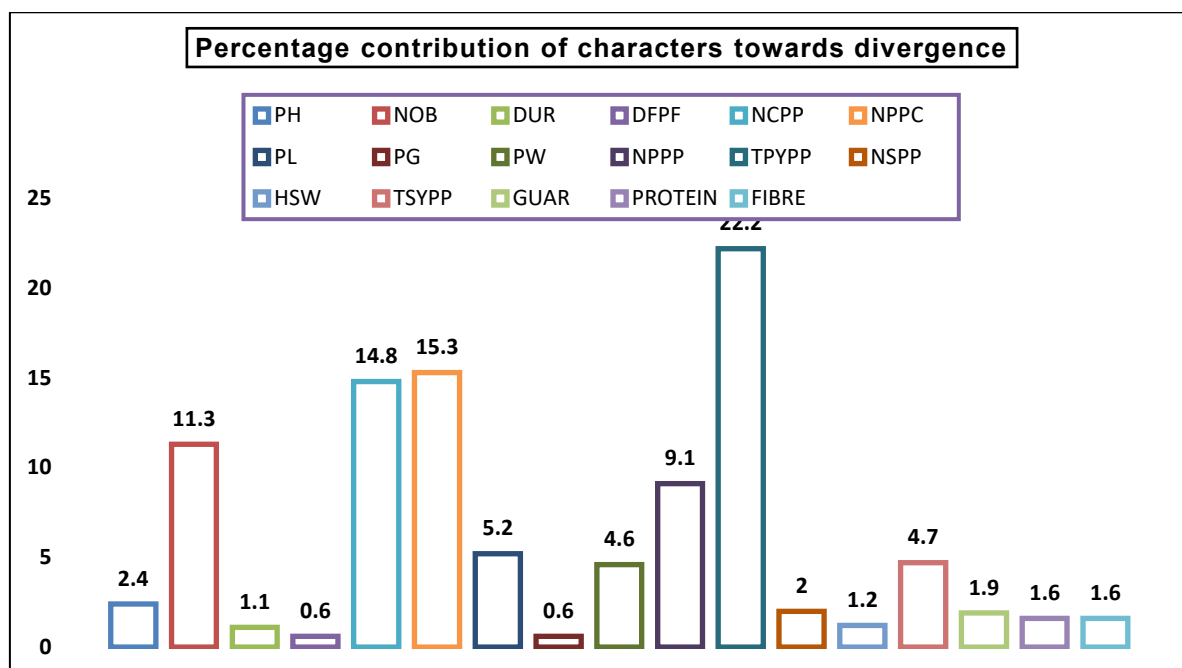


Fig. 1. Percentage contribution of biometrical parameters towards divergence

PH: Plant Height; NOB: Number of branches per plant; DUR: Duration; DFPF: Days to 50% flowering; NCPP: Number of clusters per plant; NPPC: Number of pods per cluster; PL: Pod length; PG: Pod girth; PW: Pod weight; NPPP: Number of pods per plant; TPYPP: Total pod yield per plant; NSPP: Number of seeds per plant; HSW: Hundred seed weight; TSYPP: Total seed yield per plant, GG: Guar Gum; PR: Protein; FB: Fibre

3.3 Cluster Means of Biometrical Traits

In accordance with the clustering pattern, the mean parameters were calculated and presented in Table (3). Height of the plant recorded highest values in cluster XIV (131.01) and the lowest in cluster XI (77.62). In case of number of primary branches, the top value was recorded in cluster XII (10.25), whereas cluster III, IV and XIV attributed to zero. The earliest group to have completed its duration belonged to cluster VI (95.96), whereas cluster XIV had the longest duration of 122.57 days. Cluster III (39.49) was the earliest to produce 50 % of its flowers, while cluster VIII required 48.26 days. Cluster XI had the highest cluster mean of 57.25 clusters per plant, while cluster XII had the lowest cluster mean of 10.80 clusters per plant. With respect to the number of pods per cluster, the maximum mean values were observed in cluster IX (12.35) and the minimum being in cluster XI (2.14). The cluster mean value from cluster XIV (13.60) conformed superlative for pod length and cluster XII (4.39) were the smallest. Pod girth documented maximum and minimum values at cluster XI (7.40) and cluster IX (5.20) respectively. The pod weight was highest in cluster XIV (2.86) and lowest in cluster XII (0.92). Cluster IX (136.20) adhered to maximal number of pods per plant, whereas the minimal values

were evident in cluster III (44.63). The total pod yield per plant was documented to be superlative in cluster XIV (256.32) and minimum in cluster III (51.36).

Similarly, the number of seeds per pods were highest in cluster XIV (9.59) and recorded lowest in cluster X (4.80). For hundred seed weight and total seed yield per plot, it was evident from table (3) the maximum values were recorded in cluster VIII (3.60 and 19.46) and least values in cluster XI (2.51 and 5.10) respectively.

The values for physiological parameters were recorded for guar gum, protein and fiber. The highest value for guar gum was recorded in cluster XI (32.60) and the lowest from cluster VII (21.57). In case of protein, cluster XIV showed superlative values with 21.60 % and the least being in cluster II (14.18). Minimal values being the positive aspect was recorded in cluster IV (4.60) and maximum value being in cluster XIV (5.62). Cluster XIV, comprising of the solitary member Pusa Navbahar, had maximum cluster mean values for six out of seventeen parameters which include plant height, pod length, pod weight, total pod yield per plant, number of seeds per pod and protein content showing its superiority over other clusters. The present findings are in accordance with those of (Kumar

et al., 2014, Manivannan and Anandakumar, 2013).

3.4 Percentage Contribution of Characters towards Divergence

In addition to genetic divergence, genotype and character performance with the greatest potential for divergence should be prioritized. The method employed by Nadarajan and Gunasekaran, (2012) on ranking of genotypes by mean and critical difference for all traits, rather than their magnitude of contribution was found to be more effective. For the ranking of genotypes, this method depends on the contribution of characteristics to genetic diversity. Consequently, the combined ranking identifies superior genotypes for subsequent use as possible parents for a hybridization undertaking.

The contribution of characters towards divergence is presented in Figure (1). Total pod yield per plant contributed to maximum divergence with 22.20 %, followed by number of pods per cluster (15.30 %), number of clusters per plant (14.80 %), number of primary branches (11.30 %) and number of pods per plant (9.10 %). Similar findings on maximum contribution from pod yield was also observed by Remzeena et al., (2021), Choyal et al., (2017). Nadarajan and Gunasekaran, (2012) reported on number of pods per cluster being a major contributor of divergence. The characters viz., total pod yield per plant, number of pods per cluster, number of clusters per plant, number of primary branches and number of pods per plant were major contributors of divergence owing to 72.7 % of total divergence. Similar reports on diversity were given by Sushmitha et al. 2025, Khatoon et al., 2024 in cluster bean. Importance should be given for these characters while selecting genetically diverse genotypes.

4. Conclusion

The investigation put forth the paramount availability of genetic divergence in cluster bean accessions collected from various regions of the country. During any selection process, further emphasis must be given to those characters that contribute to maximum divergence. Most divergent clusters having high D^2 values in cross combination proves to provide greater manifestation of heterosis. Genotypes for suitable characters can be selected based on their cluster mean values i.e. lowest cluster mean values for duration and days to 50% flowering and higher mean values for yield aspects. In the event that the desired traits are not available, the

implementation of a hybridization program to take advantage of the hybrid vigor potential in cluster beans helps to bridge this gap, which in turn also helps to expand the genetic diversity of cluster bean production.

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 the writing or editing of this manuscript.

Competing interests

Authors have declared that they have no known competing financial interests or non-financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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