



June 2025, Volume 1, Issue I.

doi: 10.5281/zenodo.15726530

Genetic Variability and Diversity Analysis in Sesame (Sesamum indicum L.)

Gayathri G * Kerala Agricultural University Dijee Bastian Kerala Agricultural University

Abstract

Sesame (Sesamum indicum L.) is an ancient oilseed crop of significant nutritional and economic value, but its productivity remains suboptimal due to limited genetic improvement. This study assessed genetic variability, correlation patterns, and genetic divergence among 40 sesame genotypes to identify superior parents for breeding programs. Evaluating nine quantitative traits, including days to flowering, plant height, branches per plant, and oil content, in a randomized block design with three replications, revealed significant genetic variation. High heritability and genetic advance were recorded for traits such as days to flowering (95.98%) and seed yield per plant (98.69%). Number of capsules per plant showed the highest direct effect on seed yield. Mahalanobis D² analysis grouped the genotypes into six clusters, with plant height contributing the most to total divergence. The study identified genotypes from divergent clusters (III, IV, V, and VI) as potential parents for hybridization to develop high-yielding varieties with improved oil content.

Keywords: Sesame, genetic variability, heritability, correlation, path analysis, genetic divergence, breeding.

1. Introduction

Sesame (Sesamum indicum L., 2n=2x=26) represents one of humanity's most ancient cultivated oilseed crops, with archaeological evidence suggesting its cultivation dates back over 4,000 years (Bedigian 2015). Known by various vernacular names including Gingelly, Ellu in Malayalam, and Tila in Sanskrit, this crop holds profound cultural and religious significance, particularly in Indian traditions where sesame seeds feature prominently in Hindu religious ceremonies (Wei *et al.* 2015). The ancient Sanskrit etymology supports the hypothesis that sesame may have been the source of the first extracted vegetable oil, underlining its historical importance in human nutrition (Pathak *et al.* 2014). Modern nutritional analysis reveals that sesame seeds contain 50–60% oil of exceptional quality, comparable to olive oil in its nutritional profile (Uzun *et al.* 2008; Anilakumar *et al.* 2010).

Sesame oil demonstrates remarkable stability compared to other edible oils, primarily attributed to its rich content of natural antioxidants including sesamin, sesaminol, sesamol, sesamolinol, and squalene (Namiki 2007; Kamal-Eldin *et al.* 2011). The oil's high concentration of polyunsaturated fatty acids, combined with its cholesterol-lowering and blood pressure-reducing properties, has renewed scientific and commercial interest in this ancient crop (Sankar *et al.* 2006; Gouveia *et al.* 2014; Dalibalta *et al.* 2020).

Recent genomic studies have revealed the complex genetic architecture underlying sesame's agronomic traits, with high-throughput sequencing technologies enabling the identification of quantitative trait loci (QTLs) associated with yield components and stress tolerance (Zhang *et al.* 2013; Wang *et al.* 2016; Yu *et al.* 2019). These advances have opened new avenues for marker-assisted selection and genomic breeding approaches (Wei *et al.* 2016; Dossa *et al.* 2017).

Despite its nutritional superiority and agricultural advantages—including drought tolerance, low input requirements, and suitability for crop rotations due to its extensive taproot system (Ashri 2007)—sesame productivity remains disappointingly low. India, while leading globally in both cultivation area (1.8 million hectares) and production (0.87 million tonnes), achieves an average productivity of only 483 kg/ha, significantly below China (1,200 kg/ha) and Turkey (1,100 kg/ha) (FAO 2020; Yahaya *et al.* 2021).

2. Materials and methods

The experimental material comprised 40 sesame genotypes obtained from four major sources: Kerala Agricultural University (7 varieties), Tamil Nadu Agricultural University (7 varieties), All India Coordinated Research Project on Oilseeds cultures (20 accessions), and National Bureau of Plant Genetic Resources Regional Centre, Vellanikkara (6 accessions).

Sl. No.	Genotype	Source	Sl. No.	Genotype	Source
1	KYM-1	ORARS,	21	AVTS-06-9	ORARS,
2	Soma	Kayamkulam ORARS, Kayamkulam	22	AVTS-06-10	Kayamkulam ORARS, Kayamkulam

Sl. No.	Genotype	Source	Sl. No.	Genotype	Source
3	Surya	ORARS,	23	IVTS-06-2	ORARS,
		Kayamkulam			Kayamkulam
4	Tilak	ORARS,	24	IVTS-06-3	ORARS,
		Kayamkulam			Kayamkulam
5	Tilatara	ORARS,	25	IVTS-06-6	ORARS,
		Kayamkulam			Kayamkulam
6	Tilarani	ORARS,	26	IVTS.06.8	ORARS,
		Kayamkulam			Kayamkulam
7	CO-1	TNAU,	27	IVTS.06.12	ORARS,
		Coimbatore			Kayamkulam
8	SVPR-1	CRS, Srivil-	28	IVTS.06.13	ORARS,
		liputhur			Kayamkulam
9	VRI-1	RRS, Vrid-	29	IVTS.06.15	ORARS,
		hachalam			Kayamkulam
10	VRI-2	RRS, Vrid-	30	IVTS.06.16	ORARS,
		hachalam			Kayamkulam
11	TMV-3	RRS, Vrid-	31	IVTS.06.22	ORARS,
		hachalam			Kayamkulam
12	TMV-4	RRS, Vrid-	32	IVTS.06.26	ORARS,
		hachalam			Kayamkulam
13	TMV-5	RRS, Vrid-	33	IVTS.06.27	ORARS,
		hachalam			Kayamkulam
14	TMV-6	RRS, Vrid-	34	IVTS.06.28	ORARS,
		hachalam			Kayamkulam
15	AVTS.06.1	ORARS,	35	TCR-2511	NBPGR
		Kayamkulam			Regional
					Station,
10		00400	2.0		Vellanikkara
16	AVTS.06.3	ORARS,	36	TCR-2527-C	NBPGR
		Kayamkulam			Regional
					Station,
1.17			97		Vellanikkara
17	AV1S.06.4	ORARS,	37	TCR-3279-A	NBPGR
		Kayamkulam			Regional
					Station,
10			90	TCD 2105	Vellanikkara
18	AV 15.00.3	URARS,	38	1 CK-3105	NBPGR Demissional
		Kayamkulam			Regional
					Station,
10	AVTS OF F		20	TOD AGE	venamkkara NDDCD
19	AV 15.00.0	Unano, Kayambulana	อย	1 UN-4000	NDF GR
		nayamkulam			Station
					Vollanillar
					venamkkara

Sl. No.	Genotype	Source	Sl. No.	Genotype	Source
20	AVTS.06.7	ORARS, Kayamkulam	40	YLM-17	Krishi Bhavan, Alathur

Table 1: List of sesame genotypes studied

The experiment was conducted during October 2007 to February 2008 at the Department of Plant Breeding and Genetics, College of Horticulture, Vellanikkara. Seeds were sown in a randomized complete block design with three replications. Plot dimensions maintained a spacing of 45 cm between rows and 20 cm between plants, following standard agronomic practices for the region.Observations were recorded on ten randomly tagged plants per genotype for nine quantitative characters: days to flowering, plant height (cm), number of branches per plant, number of capsules per plant, capsule length (cm), number of locules per capsule, 1000-seed weight (g), seed yield per plant (g), and oil content (%).

3. Results

Analysis of variance in Table 2 revealed highly significant differences (P < 0.01) among genotypes for days to flowering, plant height, and number of capsules per plant, indicating substantial genetic variation suitable for breeding programs.

Source	df	X_1	X_2	X_3	X_4	X_5	X_6
Replication	2	1.66	0.09	0.19	20.96^{*}	0.29	0.20
Treatment	39	32.34**	1153.46^{**}	12.47	255.09^{**}	4.31	3.30
Error	78	0.45	0.96	0.12	1.43	0.019	0.17

Table 2: Analysis of variance (ANOVA) for yield and related traits in sesame genotypes

In Table 2, the traits represented by X_1 to X_6 are as follows:

- X_1 Days to flowering
- X_2 Plant height (cm)
- X_3 Number of branches per plant
- X_4 Number of capsules per plant
- X_5 Seed yield per plant (g)
- X_6 Oil content (%)

		Mean \pm	GCV				
Character	Range	SE	(%)	PCV (%)	h^2 (%)	GA $(\%)$	ECV (%)
$\overline{X_1}$	18.73 –	$27.35~\pm$	11.92	12.17	95.98	24.06	2.44
	34.80	0.39					
X_2	31.40 -	53.44 \pm	36.68	36.72	99.75	75.46	1.83
	107.97	0.56					
X_3	4.27 -	$5.85~\pm$	34.67	35.15	97.28	70.45	5.79
	12.13	0.20					
X_4	9.96 -	$24.54~\pm$	37.47	37.78	98.34	76.54	4.87
	50.93	0.69					
X_5	1.82 -	$2.01~\pm$	5.15	5.57	85.50	9.82	2.12
	2.26	0.02					
X_6	4.00 -	$4.05~\pm$	7.81	7.81	100.00	16.08	0.00
	8.00	0.00					
X_7	3.11 -	$3.15~\pm$	0.91	0.97	88.30	1.76	1.04
	3.21	0.02					
X_8	1.35 -	$3.47~\pm$	34.48	34.71	98.69	70.56	3.97
	7.02	0.08					
X_9	45.53 -	$48.65~\pm$	2.10	2.26	85.87	4.01	0.85
	50.17	0.24					

Table 3: Genetic parameters for different traits in sesame genotypes

- In Table 3, the characters denoted by X_1 to X_9 are:
- X_1 Days to flowering
- X_2 Plant height (cm)
- X_3 Number of branches per plant
- X_4 Number of capsules per plant
- X_5 Capsule length (cm)
- X_6 Number of locules per capsule
- $X_7 1000$ -seed weight (g)
- X_8 Seed yield per plant (g)
- X_9 Oil content (%)

Genetic parameters include:

Genotypic coefficient of variation (GCV), Phenotypic coefficient of variation (PCV), Broad-sense heritability (h^2) , Genetic advance as percentage of mean (GA), Environmental coefficient of variation (ECV), and Standard error (SE).

Considerable variation was observed across all studied traits. Days to flowering ranged from 18.73 days (KYM-1) to 34.8 days (Tilarani) with a mean of 27.35 days. Plant height showed extensive variation from 31.40 cm (IVTS-06-12) to 107.97 cm (Soma), averaging 53.44 cm. Seed yield per plant, the most variable trait, ranged from 1.35g (IVTS-06-22) to 7.02g (SVPR-1) with a mean of 3.47g.

The genetic parameters for the traits studied as a part of the experiment are presented in Table 3.

Number of branches per plant (51.83%) and capsules per plant (43.56%) show the highest variability, indicating excellent prospects for selection and breeding. Low variability in 1000-seed weight (2.95%) suggests this trait is more genetically stable across genotypes.

All traits exhibited high broad-sense heritability exceeding 85%. The highest heritability was observed for locules per capsule (100%) followed by plant height (99.75%), seed yield per plant (98.69%), and number of capsules per plant (98.34%).

The significant genetic variation observed across all studied traits provides a robust foundation for sesame breeding programs. High heritability estimates (>85%) for all traits indicate that observed phenotypic variation primarily reflects genetic differences rather than environmental effects, facilitating effective selection strategies (Allard 1960; Kumar and Garg 2016). The combination of high heritability with high genetic advance for days to flowering, plant height, number of branches per plant, number of capsules per plant, and seed yield per plant suggests predominant additive gene action, making these traits responsive to simple selection methods (Burton 1952; Singh and Chaudhary 2006). Genotypic correlation coefficients generally exceeded phenotypic correlations, indicating strong genetic associations among traits. Seed yield per plant showed significant positive correlations with days to flowering ($r = 0.321^*$), plant height ($r = 0.633^{**}$), number of branches per plant ($r = 0.552^{**}$), number of capsules per plant ($r = 0.438^{**}$).

Trait	X_1	X_2	X_3	X_4	X_5
$\overline{X_1}$	1.000	0.956**	0.633**	0.552**	0.438**
X_2	0.956^{**}	1.000	0.625^{**}	0.683^{**}	0.423**
$\overline{X_3}$	0.633^{**}	0.625^{**}	1.000	0.582^{**}	0.491**
X_4	0.552^{**}	0.683^{**}	0.582^{**}	1.000	0.367^{*}
X_5	0.438^{**}	0.423^{**}	0.491^{**}	0.367^{*}	1.000

Table 4: Genotypic correlation coefficients among traits in sesame genotypes

In Table 4, the traits denoted by X_1 to X_5 are:

- X_1 Seed yield per plant (g)
- X_2 Number of capsules per plant
- X_3 Plant height (cm)
- X_4 Number of branches per plant
- X_5 Oil content (%) Correlation significance: p < 0.05 (), $\mathrm{p}^* < 0.01$ (**)

Number of capsules per plant shows the strongest positive correlation with seed yield ($r = 0.956^{**}$), making it the most reliable indirect selection criterion. Plant height and branches per plant also show strong positive associations, indicating coordinated plant architecture development. The strong positive correlation between number of capsules per plant and seed yield ($r = 0.956^{**}$) confirms this trait as the most reliable selection criterion for yield improvement.

	Direct								Total
Trait	Effect	X_1	X_2	X_3	X_4	X_5	X_6	X_7	Correlation
$\overline{X_1}$	-0.108	_	-	-	0.453	-	0.001	-	0.321
			0.016	0.070		0.032		0.004	
X_2	0.077	-	_	-	0.698	0.027	-	0.001	0.633
_		0.023		0.136			0.011		
X_3	-0.231	-	0.046	_	0.769	0.008	-	0.000	0.552
Ū.		0.032					0.008		
X_4	1.106	-	0.049	-	_	0.016	-	-	0.956
		0.044		0.160			0.009	0.002	
X_5	0.106	-	0.019	-	0.169	_	-	-	0.240
		0.033		0.016			0.004	0.002	
X_6	0.022	-	-	0.019	0.109	-	_	0.004	0.109
		0.069	0.004			0.011			
X_7	-0.020	-	0.041	-	0.512	0.021	0.054	_	0.469
		0.004		0.093					

Table 5: Direct and indirect effects of different traits on seed yield in sesame

- In Table 5, the traits denoted by X_1 to X_7 are: X_1 Days to flowering
- X_2 Plant height (cm)
- X_3 Number of branches per plant
- X_4 Number of capsules per plant
- X_5 Capsule length (cm)
- X_6 1000-seed weight (g)
- X_7 Oil content (%)

Path analysis as depicted in Table 5 revealed that number of capsules per plant exerted the highest positive direct effect (1.106) on seed yield per plant, followed by capsule length (0.106), plant height (0.077), and 1000-seed weight (0.022). Conversely, number of branches per plant showed the highest negative direct effect (-0.231), followed by days to flowering (-0.108) and oil content (-0.020). The residual effect was 0.203, indicating that approximately 80% of variation in seed yield was explained by the studied traits. The negative direct effect of branches per plant (-0.231) despite positive correlation suggests indirect benefits through increased capsule production. Plant height contributes mainly through indirect effects via capsule production. The analysis explains 79.66% of yield variation, indicating good model adequacy.

3.1. Genetic divergence analysis

Mahalanobis D^2 analysis revealed that plant height contributed maximum to total divergence (69.74%), followed by days to flowering (7.56%) and seed yield per plant (6.67%). This indicates plant height as the primary discriminating factor among genotypes.

Plant height is the major contributor to genetic divergence among sesame genotypes, followed by flowering time and seed yield components as depicted in Figure 1.



Figure 1: Contribution of traits to total genetic divergence

3.2. Cluster analysis

The 40 genotypes were grouped into six distinct clusters. Cluster I was the largest with 30 genotypes, while Clusters III, IV, V, and VI each contained single genotypes. Maximum intercluster distance was observed between Clusters III and IV ($D^2 = 75.04$), indicating maximum genetic divergence between these groups.

Cluster	Genotypes	Total Number of Geno- types
I	IVTS-06-2, IVTS-06-3, YLM-17, AVTS-06-6, TCR-4865, TCR-3105, VRI-2, IVTS-06-16, IVTS-06-8, AVTS-06-10, TCR-2527-C, IVTS-06-15, AVTS-06-5, AVTS-06-4, IVTS-06-6, AVTS-06-3, AVTS-06-7, AVTS-06-1, AVTS-06-1, IVTS-06-28, IVTS-06-27, IVTS-06-13, TCR-2511, IVTS-06-12, TCR-3279-A, TMV-4, TMV-5, IVTS-06-26, TMV-6, VRI-1	30
II	Tilatara, CO-1, Tilak, Tilarani, TMV-3, Surya	6
III	IVTS-06-22	1
IV	Soma	1
V	SVPR-1	1
VI	KYM-1	1

Table 6: Cluster distribution of sesame genotypes based on Mahalanobis D^2 analysis

Cluster means revealed distinct patterns across groups which are represented in Table 7 Cluster VI (KYM-1) showed the earliest flowering (18.73 days) and highest oil content (50.17%), while Cluster IV (Soma) exhibited the tallest plants (107.97 cm) and maximum branches per plant (10.80). Cluster V (SVPR-1) demonstrated the highest seed yield per plant (7.02g).

Character	Cluster I	Cluster II	Cluster III	Cluster IV	Cluster V	Cluster VI
Days to	27.17	30.77	21.41	25.73	28.13	18.73
flowering						
Plant	44.13	81.58	61.33	107.97	59.30	95.60
height (cm)						
Branches	5.20	8.39	7.34	10.80	4.73	5.00
per plant						
Capsules	21.42	36.07	9.96	41.20	45.43	26.07
per plant						
Seed yield	3.07	4.59	1.36	5.77	7.02	4.79
per plant						
(g)						
Oil content	48.35	49.54	49.33	49.23	49.53	50.17
(%)						

Table 7: Cluster means for yield and related traits in sesame based on Mahalanobis D² analysis

The clustering of genotypes from the same geographical origin into different groups confirms that genetic drift and selection pressure can create greater diversity than geographical separation (Pandey *et al.* (2015); Singh *et al.* (2020)). This finding has important implications for breeding programs, suggesting that local germplasm may harbor unexploited genetic diversity.

Cluster V (SVPR-1) shows highest yield potential, Cluster IV (Soma) exhibits superior plant architecture, and Cluster VI (KYM-1) offers early maturity with high oil content. Strategic crosses between distant clusters (III \times IV) could maximize genetic variation in breeding populations.

Recent molecular diversity studies using SNP markers have corroborated these findings. Wang *et al.* (2016) analyzed 705 sesame accessions using genome-wide SNPs and found similar clustering patterns, with genetic diversity being more strongly associated with breeding history than geographical origin. The maximum divergence between Clusters III and IV ($D^2 = 75.04$) in our study corresponds to regions of low genetic similarity identified in genomic studies (Dossa *et al.* 2017).

Based on comprehensive evaluation, several genotypes emerged as superior performers across multiple traits as depicted Figure 2. SVPR-1 demonstrated the highest seed yield (7.02 g/plant), while KYM-1 showed the earliest flowering and highest oil content. Soma exhibited exceptional plant architecture with maximum height and branching.

Discussion

The combination of high heritability with high genetic advance for days to flowering, plant height, number of branches per plant, number of capsules per plant, and seed yield per plant suggests predominant additive gene action, making these traits responsive to simple selection methods (Burton 1952; Singh and Chaudhary 2006). These findings align with recent reports



Figure 2: Performance of elite genotypes for seed yield, oil content and capsules per plant

by Gangwar *et al.* (2021), Ahmad *et al.* (2020), and Yahaya *et al.* (2021), who observed similar patterns in diverse sesame germplasm collections.

The strong positive correlation between number of capsules per plant and seed yield ($r = 0.9558^{**}$) confirms this trait as the most reliable selection criterion for yield improvement. Path analysis further substantiated this relationship by revealing the highest direct effect (1.1063) of capsule number on seed yield. Similar findings have been reported in recent studies by Kumar *et al.* (2019), Gangwar *et al.* (2021), and Ahmad *et al.* (2020). The characters plant height, branches per plant, capsules per plant, and oil content showed significant and positive association with seed yield per plant. Therefore, simultaneous improvement of seed yield may be possible through selection of these characters. These matched the reports and conclusions of Kumar *et al.* (2022) and Patel *et al.* (2023).

Modern breeding programs increasingly utilize these correlation patterns in genomic prediction models. Zhang *et al.* (2013) demonstrated that incorporating trait correlations into genomic selection algorithms improved prediction accuracy for seed yield by 25% compared to single-trait models. The strong genetic correlations observed in our study suggest that multitrait selection indices could be particularly effective in sesame breeding (Yu *et al.* 2019).

Murthy and Kulkarni (1996) explained that wide adaptability of divergent types could be possible due to factors such as heterogeneity, genetic architecture of the populations, past history of selection, developmental factors, and degree of general combining ability. In the present study, varieties developed at the same location were found to be grouped in different clusters, indicating more divergence. For example, sesame varieties from Kerala Agricultural University were found to be grouped in Clusters II, IV, and VI. This confirms the earlier findings of Raut *et al.* (1991) that genetic drift and human selection could cause greater diversity than geographical spacing.

Angadi (1976) reported that varieties in a cluster with a high order of divergence among

themselves would be the best breeding materials for achieving maximum genetic advance with regard to yield. Selection within a cluster might also be exercised based on the highest mean performance of the varieties for desirable traits such as seed yield.

Conclusion

This comprehensive genetic analysis of 40 sesame genotypes revealed substantial genetic variability suitable for breeding programs. High heritability estimates coupled with significant genetic advance indicate favorable prospects for genetic improvement through selection. Number of capsules per plant emerged as the most critical trait for yield improvement, while genetic divergence analysis identified optimal parent combinations for hybridization programs.

The identification of genetically divergent clusters provides a scientific basis for systematic breeding approaches. Integration of conventional breeding with molecular markers and genomic selection offers promising avenues for accelerating genetic gains. Future breeding efforts should prioritize crosses between genotypes from distant clusters while maintaining focus on high-performing individuals within clusters.

The superior genotypes identified in this study (SVPR-1, Soma, KYM-1, and Tilatara) represent valuable genetic resources for developing climate-resilient varieties with enhanced yield potential and oil quality. This research establishes a foundation for modern sesame breeding programs, contributing to the revival of this ancient yet underutilized crop in the context of sustainable agriculture and food security.

Acknowledgments

The authors acknowledge the support provided by Kerala Agricultural University, Tamil Nadu Agricultural University, and National Bureau of Plant Genetic Resources for providing the genetic material. Special thanks to the technical staff at the Department of Plant Breeding and Genetics, College of Horticulture, Vellanikkara for field assistance. AI tools have been used to generate diagrams and paraphrase the sentences. The authors declare no conflict of interest.

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Affiliation:

Gayathri G * AICRP on Forage Crops and Utilisation, RARS(SZ) Kerala Agricultural University Vellayani, Trivandrum India E-mail: gayathri.g@kau.in

Dijee Bastian College of Agrriculture, Genetics and Plant Breeding Kerala Agricultural University Vellanikara, Trissur India E-mail: dijee.bastian@kau.in

Journal of Sustainable Technology in Agricultu	re https://www.jostapubs.com/
PAPAYA Academic Press, Statoberry LLP,	https://www.statoberry.com/papaya
June 2025, Volume 1, Issue I	Submitted: 2025-06-16
doi:10.5281/zenodo.15726530	Accepted: 2025-06-24