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# Genetic Divergence Analysis for Quantitative Traits in Rice (*Oryza sativa* L.) Germplasm under Irrigated Conditions

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#### Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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#### ABSTRACT

The present study was with the aim of knowing genetic divergence analysis for quantitative traits among the rice genotypes for agronomical and yield traits. The experimental material consists of 36 rice genotypes sown in a Randomized Block Design with three replications and observations recorded on randomly selected five plants for 13 quantitative traits. The results indicated that the genotype Nagarjuna shows greater yield among the genotypes. The genetic parameters show high

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GCV and PCV in grain yield per plant, and according to D<sup>2</sup> analysis, the 36 genotypes were grouped into 6 clusters based on Tocher's method of cluster formation. Cluster I is the largest cluster, consisting of 29 genotypes, while Cluster II has 3 genotypes. Cluster III, IV, V, and VI were the smallest, with only a single genotype. The maximum intra-cluster distance was found in cluster II (39.19), followed by cluster I (30.15), and the maximum inter-cluster distance was shown in between cluster VI and cluster II (178.97), followed by cluster VI and cluster V (151.75). Based on cluster means, Cluster III (39.67) shows greater among 6 clusters for grain yield per plant as contributing the highest among all the quantitative characters. Hence, these traits could be focused on for selection while improving seed yield and seed quality. On the basis of their greater inter-cluster distance, high value of cluster means, and performance of the individual germplasms for the character, the germplasm could be used in improving seed yield and seed quality programs for improvement of different plant characters of the rice.

Keywords: Rice; genetic variability; heritability; genetic advance; diversity.

#### 1. INTRODUCTION

Rice belongs to the botanical genus Oryza and the grass family Poaceae. There are two main species of cultivated rice: *Oryza sativa* (Asian rice) and *Oryza glaberrima* (African rice), which have chromosome number 2n=24. It is the staple food crop for nearly four billion people around the world, providing 27% of calories in low- and medium-income countries. Almost 95% of the rice production happens in Asian countries, and nearly half of the global population consumes it.

"Rice is a cereal crop important for food security and is a staple in the diets of many billions of people worldwide. Moreover one-third of the world's population relies on rice as their main source of food. The world's demand for food production, particularly rice, is expected to rise in the ensuing decades as a result of population growth. The majority of people on the planet eat rice as their primary staple meal, and it accounts for 19% of worldwide per capita energy consumption. Rice is a nutritious cereal crop grown primarily for human consumption. It is the primary source of energy and protein, providing significant amounts of the required nutritional intakes of zinc and niacin however, rice contains extremely little calcium, iron, thiamine, and riboflavin, as well as almost no beta-carotene at all. Rice protein is biologically the richest among cereal proteins due to its high true digestibility (88%) and also contains minerals and fibre. Rice provides 50-80% of the daily caloric intake for the impoverished. Cereals, snack meals, brewed beverages, flour, oil (rice bran oil), syrup, and religious ceremonies are only a few of the applications for rice. Rice is also said to have medicinal properties and is used in many countries, including India. Compared to other crop species, rice germplasm possesses a vast

reservoir rich in genetic variation, but only a very small part has been incorporated into breeding programs, resulting in high genetic similarity in commercial rice cultivars" (Koli et al., 2022).

"The selection of parents is very important for hybridization for better crop improvement. The parents involved in the development of varieties should be divergent. The germplasm provides immense scope for wide variability. The Breeding program depends on the nature and magnitude genetic diversity present among of the genotypes. Genetic diversity among genotypes that allows for better selection in cross combinations with divergent parents offers the chance to combine different gene constellations. This also leads to promising seareaant derivatives resulting from the complementary interaction of the divergent genes from the parent plants" (Vanisri et al., 2020).

"Rice yield is a quantitatively complex property. Selecting parents solely on yield is frequently deceptive. As a result, understanding the link between yield and its contributing features is critical for plant breeders to develop an effective selection strategy in order to produce an economically viable variety. A plant breeder heavily on information concerning relies phenotypic and genotypic interactions of many economic variables while selecting and breeding different genotypes to increase yield potential. Such analysis studies also include the selection of the genetically divergent parents to obtain desirable combinations. Information about the degree and nature of genetic divergence would aid plant breeders in selecting the appropriate parent for future hybridization. Genetic divergence is an effective tool for parent selection for hybridization programs" (Koli et al., 2022).

"Mahalanobis D<sup>2</sup> used to analyze genotype differences based on quantitative variables. However, since physical attributes are heavily influenced bv environmental factors and determined by additive gene effects, relying solely on these traits for diversity analysis can produce inaccurate results (Mangudi et al., 2024). This D<sup>2</sup>analysis measures the relative contributions of various components to diversity at both inter-cluster and intra-cluster levels. Genotypes from widely divergent clusters are likely to yield heterotic combinations and greater variability in segregating generations" (Sadia et al., 2020).

#### 1.1 Objectives

- 1. To evaluate rice genotype for yield and yield attributing traits
- 2. To identify the divergent parents for future Hybridization program
- 3. To study about diversity for morphological characters genetic

#### 2. MATERIALS AND METHODS

The experiment was conducted with 36 rice genotypes in the crop research farm, Department Plant of Genetics and Breeding, Naini Agricultural Institute, Sam Higginbottom University of Agriculture, Technology, and Sciences, Prayagraj - 211007. The experimental site is situated at 98m above sea level at 25.57° N latitude and 81.56<sup>o</sup> N longitude. In These areas climate changes drastically, like the temperature may drop to 1°C at the winter and reach nearly 50°C during summer. The 36 rice genotypes are taken from Indian Institute of Rice Research (IIRR), Hyderabad. These genotypes are used to perform experiment, the conducted in Randomised Block Design with three replications with performed spacing row to row of 20cm and plant to plant of 15cm. In each replication, five randomly competitive plants were selected and examined and recorded following 13 quantitative characters: Days to 50% flowering, Days to maturity, Flag leaf length (cm), Flag leaf width (cm), Plant height (cm), Number of total tillers, Panicle length (cm), Number of Grains per panicle, Number of productive tillers, Test weight (g), Biomass (g), Harvest Index (%) and Grain yield per plant (g). Analysis of variance was carried out for all the traits, and the data were subjected multivariate analvsis following Mahalanobis's D<sup>2</sup> statistics (Mahalanobis et al., 1936). to measure the genetic divergence, followed by the clustering of genotypes based on

13 characters following Tocher's method (Kumar et al., 2017). The Analysis of variance for the experimental design was conducted following the methods outlined by Panse and Sukhatme genotypic phenotypic and The (1961). coefficients of variation were estimated using the approach proposed by Burton and de Vane (1952). According to Johnson et al., (1955) the of Phenotypic Coefficient Variation and Genotypic Coefficient of Variation were categorized as low (0-10%), moderate (10-20%), and high (>20%). Heritability (h<sup>2</sup>) is defined as the percentage of the ratio of genotypic variance (σ²g) to phenotypic variance  $(\sigma^2 p)$ , with heritability in the broad sense (h<sup>2</sup>b) estimated according to Johnson et al. (1955). Robinson et al. classified heritability into low (0 to 30%), moderate (30 to 60%), and high (> 60%). Genetic advance measures genetic improvement achieved through selection, with the genetic advance as a percentage of the mean classified as low (<10%), moderate (10-20%), and high (> 20%) according to Johnson et al. (1955). Genetic divergence analysis was performed using Mahalanobis D<sup>2</sup> statistics (1936). Genotypes were grouped into clusters using Tocher's method (Rao, 1952). Additionally, we calculated intra- and inter-cluster distances as well as the mean performances of clusters for all traits. B.Satish Chandra et al., (2022).

#### 2.1 Experimental Material

The 36 rice genotypes used in this research are mentioned in Table 1.

#### 3. RESULTS AND DISCUSSION

Analysis of Variance for different characters is presented in Table 2. The mean sum of squares shows significant differences of all the characters to all the genotypes. The variability indicates among the genotypes. That this helps with the selection of desirable. The performance of genotypes displays statistical difference and these differences with help in selection and for better crop improvement. The mean sum of squares of analysis of variance shows significant at the 1% level of significance for all 13 quantitative traits indicating the presence of genetic differences in the experimental material, suggesting the importance of the genetic variability to identify the desirable genetic makeup and provide better scope for selection and better crop improvement.

The Genetic Parameters such as Phenotypic coefficient of variance (PCV), Genotypic coefficient of variance (GCV), Heritability (h<sup>2</sup>) in

broad sense, genetic advance (GA), Genetic advance as percent of mean (GA%) for thirteen characters are explained as follows. Based on the PCV and GCV estimates. PCV is showing greater than GCV in all the characters, indicating that there is variability presence due to environment (Reddy et al., 2023). In PCV, Grain yield per plant (29.23) is showing greater among the characters, followed by number of spikelets per panicle (27.10), Biological yield per plant (20.94). In GCV Grain yield per plant (31.34) is showing greater among the characters, followed by Number of spikelets per panicle (28.66) and Biological yield per plant (23.74). Heritability is showing high range of estimates in all the characters, indicating that there is less influence of environment in the variability among the genotypes. Number of spikelets per panicle (89.40) exhibiting greater in the heritability, followed by Number of tillers per plant (88.06), Grain yield per plant (87.02). Genetic

Advancement exhibits greater in Number of spikelets per panicle (86,43%), followed by Plant height (27.96%), Days to 50% flowering (25.81%), while Flag leaf width (0.28), Number of panicles per plant (2.70%), Panicle length (2.90%) shows least among the characters. In Genetic advancement, as percent of mean shows Grain yield per plant (56.18%) exhibit greater, and followed by Number of spikelets per panicle (52.79%), Biological yield (38.03%) while the Panicle length (11.84%), and Days to maturity (14.00%) was exhibit least. All the characters showed high in heritability coupled with genetic advance as percent of mean, indicating the characters are predominantly regulated by additive gene action. As a result, the more additive genes are accumulated for the crop improvement, based on the phenotypic expression selection would be more desirable that leads to better crop improvement.

Table 1. Experimental material used in the research	Table 1.	Experimental	material	used in	the	research
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S.NO	Name of Genotypes	S.NO	Name of Genotypes
1	SAMPADA	19	DHAN-58
2	NAGARJUNA	20	NIDHI
3	SURAKSHA	21	LAGEDHAN
4	DHAN-57	22	KAUKA SEL
5	MANDHYA VIJAYA	23	NLR-145
6	JARAVA	24	NLR-40054
7	BINA-17	25	NLR-30491
8	DHAN-46	26	NLR-33641
9	DHAN-53	27	NLR-4001
10	DHAN-62	28	NLR-33359
11	BINA DHAN-10	29	JGL-3828
12	DRRH-3	30	JGL-1118
13	DHAN-54	31	JGL-3855
14	DHAN-52	32	WGL-18833
15	PHALGUNA	33	WGL-401
16	GAUTAM	34	WGL-4727
17	KRISHNA HAMSA	35	WGL-962
18	DHAN-69	36	NDR-359 (CHECK)

## Table 2. Analysis of variance of 13 traits in 36 promising rice genotypes for agronomic traits evaluated under field condition in *Kharif*, 2022

SI.No.	Source	Mean Sum of Squares (MSS)				
		Replication	Treatment	Error		
	Degrees of freedom	2	35	70		
1	Days to 50% flowering	16.4540	598.822**	36.378		
2	Days to maturity	11.5930	379.374**	47.916		
3	Plant height (cm)	51.3070	822.022**	82.939		
4	Flag leaf length (cm)	4.8970	79.485**	7.732		
5	Flag leaf width (cm)	0.0160	0.084**	0.008		
6	Panicle length (cm)	1.7920	15.282**	3.665		
7	Number of tillers per plant	1.0710	8.748**	0.378		
8	Number of panicles per plant	1.6940	7.108**	0.576		

SI.No.	Source	Mean Sum of Squares (MSS)			
		Replication	Treatment	Error	
	Degrees of freedom	2	35	70	
9	Number of Spikelets per panicle	465.9140	6140.816**	233.282	
10	Test weight (g)	3.1260	43.582**	4.06	
11	Biological yield per plant (g)	75.0920	284.547**	24.772	
12	Harvest Index (%)	15.4110	230.184**	25.276	
13	Grain yield per plant (g)	14.0320	112.167**	5.311	

Table 3. Genetic parameters of	yield and yield components	36 promising	rice genotypes for
agronomic traits	evaluated under field condition	ion in <i>Kharif</i> ,	2022

SI.No.	Characters	GCV	PCV	h2 (Broad	Genetic	Gen.Adv as % of
				Sense)	Advancement 5%	Mean 5%
1	Days to 50% flowering	14.147	15.459	83.75	25.813	26.67
2	Days to maturity	8.141	9.747	69.75	18.084	14.006
3	Plant height (cm)	12.269	14.184	74.814	27.967	21.86
4	Flag leaf length (cm)	13.191	15.174	75.57	8.758	23.622
5	Flag leaf width (cm)	13.018	14.932	76.011	0.286	23.38
6	Panicle length (cm)	8.024	11.194	51.378	2.906	11.848
7	Number of tillers per	18.729	19.958	88.064	3.229	36.206
	plant					
8	Number of panicles	17.302	19.457	79.078	2.703	31.696
	per plant					
9	Number of Spikelets	27.104	28.664	89.408	86.437	52.794
	per panicle					
10	Test weight (g)	16.213	18.543	76.442	6.537	29.2
11	Biological yield per	20.94	23.747	77.756	16.903	38.038
	plant (g)					
12	Harvest Index (%)	17.803	20.839	72.99	14.545	31.333
13	Grain yield per plant	29.237	31.341	87.025	11.469	56.185
	(g)					

#### Table 4. Grouping of 36 rice genotypes into different clusters

Cluster Group	No. of Genotypes	List of Genotypes
1 Cluster	29	DHAN - 54, DHAN - 52, DHAN - 46, DHAN - 69, DHAN - 57, SURAKSHA, GAUTAM, SAMPADA, BINA - 17, NLR - 4001, DRRH - 3, NLR -145, JGL - 3828, DHAN - 62, PHALGUNA, KRISHNA HAMSA, BINA DHAN - 10, JARAVA, WGL - 1 8833, DHAN - 53, DHAN - 58, NIDHI, WGL - 401, NLR - 30491, NLR - 40054, WGL - 962, MANDHYA VIJAYA, JGL- 3855 & NLR - 33359
2 Cluster	3	LAGEDHAN, KAUKASEL & NLR - 33641
3 Cluster	1	NAGARJUNA
4 Cluster	1	JGL - 1118
5 Cluster	1	WGL- 4727
6 Cluster	1	NDR-359 (Check)

#### Table 5. Inter and Intra cluster distances of 36 genotypes of rice

Cluster Distances						
	Cluster 1	Cluster 2	Cluster 3	Cluster 4	Cluster 5	Cluster 6
Cluster 1	30.15	78.87	66.78	51.88	50.41	109.73
Cluster 2		39.19	105.53	70.18	141.15	178.97
Cluster 3			0.00	60.19	94.24	138.27
Cluster 4				0.00	73.53	141.27
Cluster 5					0.00	151.75
Cluster 6						0.00



Fig. 1. Bar diagram depicting GCV, PCV, Heritability and Genetic advance for 13 quantitative characters of rice

	Cluster 1	Cluster 2	Cluster 3	Cluster 4	Cluster 5	Cluster 6
Days to 50% flowering	93.68	123.89	106.00	108.00	77.00	105.00
Days to maturity	126.99	142.22	139.00	143.00	126.00	131.00
Plant height (cm)	124.22	158.09	151.40	116.33	136.14	125.00
Flag leaf length (cm)	37.10	35.99	30.07	39.50	45.43	35.81
Flag leaf width (cm)	1.18	1.29	1.16	1.44	1.45	1.83
Panicle length (cm)	24.34	23.09	23.68	24.33	31.74	27.97
Number of tillers per	8.79	7.40	11.53	8.40	8.33	15.60
plant						
Number of panicles Per	8.41	7.36	11.53	7.80	7.67	14.00
plant						
Number of Spikelet per	156.00	206.87	127.00	254.27	157.47	210.60
panicle						
Test weight (g)	22.78	17.39	22.00	18.66	29.34	23.11
Biological yield per plant	41.50	56.07	69.27	56.40	50.80	51.50
(g)						
Harvest Index (%)	47.00	31.34	56.75	57.43	55.99	44.01
Grain yield per plant (g)	19.31	17.38	39.67	32.40	28.07	22.67

Table 6. Cluster mean of 13 biometrical traits of 36 rice genotypes

Table 7. Percent contribution of 13 biometrical traits of 36 rice genotypes towards genetic
divergence

SL.NO.	Source	Contribution %	Times ranked 1st
1	Days to 50% flowering	9.52	60
2	Days to maturity	9.00	57
3	Plant height (cm)	4.52	29
4	Flag leaf length (cm)	5.64	36
5	Flag leaf width (cm)	2.54	16
6	Panicle length (cm)	3.21	20
7	Number of tillers per hill	10.41	66
8	Number of panicles per hill	6.32	40
9	Number of Spikelets per panicle	5.65	36
10	Test weight (g)	8.89	56
11	Biological yield per plant (g)	6.65	42
12	Harvest Index (%)	12.00	76
13	Grain yield per plant (g)	15.65	99

Cluster analysis is performed among the 36 rice genotypes, resulting in all genotypes being divided into 6 clusters. Cluster I (29) has a greater number of genotypes, followed by cluster II (3) and remaining Clusters III, IV, V and VI contain single genotype in each cluster shown in Table 4. Based on the cluster distances Cluster II (39.19) has greater Intra cluster distances, and Cluster I (30.15) shown and remaining displays zero. According to Inter cluster distances, Cluster VI and Cluster II (178.97) have the greater among the clusters followed by Cluster VI and Cluster V (151.75), shown in Table 5. Based on the distances of clusters, best lines should be selected as parents for future hybridization, and this helps in crop improvement. Among the six clusters, the means grouped on the basis of 36 rice genotypes of every character, Cluster III (39.67) has greatest among the clusters in Grain yield per plant, followed by Cluster IV (32.40), Cluster V (28.07), Cluster VI (22.67), Cluster I (19.31), Cluster II (17.38), shown in Table 6. Based on the Percentage Contribution Grain yield per plant (15.65%) exhibits greater among the characters followed by Harvest Index (12%), Number of tillers per hill (10.41%). Considerable variability existed in the genotypes for all the characters studied. These were the genotypes with high mean values in the desirable direction, i.e., From the present investigation, it is concluded that among 36 genotypes of rice, WGL-401 showed early flowering (75 days), NLR-33359 had characters like early maturity (109 days), High Grain Yield per Plant is seen in Nagarjuna (39.6 g), Kaukasel showed high plant height (177.6 cm), and high Biomass is seen in Nagarjuna (69.2 g), and WGL-4727 (31.74 cm) showed high Panicle Length. The Highest GCV

and PCV were depicted for Grain Yield per Plant, Number of Spikelets per Panicle, Biological Yield per hill, and Number of tillers per hill. The highest Heritability was observed for the Number of Spikelets per panicle, followed by Number of tillers per hill, Grain yield per plant (g), Days to 50% flowering, Number of panicles per hill, and biological yield per plant (g). the Genetic Advance in percent of mean is high for Grain yield per plant, Number of Spikelets per panicle, biological yield per plant, Number of tillers per hill, and Number of panicles per hill. Based on D<sup>2</sup> values 36 genotypes grouped into 6 clusters. Among 6 clusters, cluster I had a greater number of genotypes (29) followed by cluster II (3), III, IV,

VI (1) genotype: each, cluster VI and V. and cluster II (178.97) followed by cluster VI and cluster V (151.75), were most diverse from each other, and the genotypes present in these clusters provide a broad spectrum of variability in segregation and may be used as parents in the future hybridization program to develop desirable genotypes for grain yield improvement in rice genotypes. The trait Grain yield per hill had maximum contribution towards to genetic divergence, followed by Harvest Index, Number of tillers per hill, Days to 50% flowering, Days to maturity, Test Weight. Therefore, all these characters should be given due consideration during selection for crop improvement.



Fig. 2. Dendrogram showing distribution of 36 rice genotypes into different clusters



Fig. 3. Inter and Intra cluster distance of 36 genotypes of ri



Fig. 4. Pie chart showing percentage contribution of 13 biometrical traits of 36 rice genotypes towards genetic divergence

#### 4. CONCLUSION

Considerable variability existed in the genotypes for all the characters studied. These were the genotypes with high mean values in desirable direction, i.e., From the present investigation, it is concluded that among 36 genotypes of rice, WGL-401 showed early flowering (75 days), NLR-33359 had characters like early maturity (109 days), High Grain Yield per Plant is seen in Nagarjuna (39.6 g), Kaukasel showed high plant height (177.6 cm), and high Biomass is seen in Nagarjuna (69.2 g); WGL-4727 (31.74 cm) showed high Panicle Length. Highest GCV and PCV were depicted for Grain Yield per Plant, Number of Spikelets per Panicle, Biological yield per plant per plant, and Number of tillers per plant. The highest Heritability was observed for Number of Spikelets per panicle, followed by Number of tillers per plant. Grain vield per plant (g), Days to 50% flowering, Number of panicles per plant, Biological yield per plant per plant (g). the Genetic Advance in percent of mean is high for Grain yield per plant, Number of Spikelets per panicle, Biological yield per plant per plant, Number of tillers per plant, Number of panicles per plant. Based on D<sup>2</sup> values, 36 genotypes grouped into 6 clusters. Among 6 clusters, cluster I had a greater number of genotypes (29) followed by cluster II (3), III, IV, V, and VI (1) genotype each. cluster VI and cluster II (178.97) followed by cluster VI and cluster V (151.75), were most diverse from each other and the genotypes present in these clusters provide a broad spectrum of variability in segregation and may be used as parents in the future hybridization program to develop desirable genotypes for grain yield improvement in rice genotypes. The trait Grain yield per plant had maximum contribution towards to genetic divergence followed by Harvest Index, Number of tillers per plant, Days to 50% flowering, Days to maturity, Test Weight. Therefore, these characters should be given due consideration during selection for crop improvement.

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#### **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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