

*International Journal of Environment and Climate Change*

*12(11): 3612-3622, 2022; Article no.IJECC.93476 ISSN: 2581-8627 (Past name: British Journal of Environment & Climate Change, Past ISSN: 2231–4784)* 

# **Assessment of Variability and Genetic Diversity for Elite Rice (***Oryza sativa* **L.) Genotypes of Telangana and Andhra Pradesh**

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#### *Authors' contributions*

*This work was carried out in collaboration among all authors. Authors RH, DB, SV and VGS designed experiment and guided at different stages of research. Author RH performed experiment and data analysis. Authors RH, KB and APB wrote the first draft of manuscript. Authors MMG and SS helped in literature searches and manuscript improvisation. All the authors read and approved the final manuscript.*

#### *Article Information*

DOI: 10.9734/IJECC/2022/v12i111409

#### **Open Peer Review History:**

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: https://www.sdiarticle5.com/review-history/93476

*Original Research Article*

*Received 11 September 2022 Accepted 02 November 2022 Published 04 November 2022*

## **ABSTRACT**

The current investigation was undertaken with eighteen elite rice genotypes to study the genetic diversity, variability, heritability and genetic advance for yield and yield attributing traits. The ANOVA revealed the significant variability among the genotypes for the traits studied. 1000 seed weight is

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having highest phenotypic coeffcient variation and genotypic coeffcient variation (24.882 & 24.835) followed by grain yield per plant (20.315 & 19.523 respectively). Heritability estimates of yield attributing traits ranged from 58.3% (Panicle length) to 99.6% (1000 seed weight). High heritability along with higher genetic advance observed for 1000 seed weight, numbers of productive tillers, plant height and grain yield per plant indicating that selection for these traits would be effective. The diversity analysis revealed that genotypes were grouped into four clusters, cluster I was the largest comprising 14 genotypes and cluster II with two genotypes followed by Cluster III and IV with one variety each indicating a high degree of divergence among the genotypes.

*Keywords: Cluster analysis; genetic diversity; GCV; heritability; D<sup>2</sup>statistics.*

# **1. INTRODUCTION**

"Rice (*Oryza sativa L.*) is one of the most important leading cereal grain produced, consumed and traded in the world after wheat. It is the staple food crop for more than half of the world human populations. India has shown a tremendous increase in its rice production over the past few decades and this trend has continued in recent years. In India, it is grown on around 45.5 m.ha area with an estimated production of 124.0 MT during 2022-23 crop seasons" (Anonymous) [1]. Land scarcity, climate change and narrow genetic base of breeding materials are further reducing the yield in Asian countries. Yield enhancement is the vital breeding objective in any rice breeding programmes and knowledge on the nature and magnitude of genetic variation governing the inheritance of quantitative characters like yield and its characteristics is essential for effective genetic improvement. A critical analysis of the genetic variability parameters, namely, genotypic coefficient of variability (GCV), phenotypic coefficient of variability (PCV), heritability and genetic advance for different traits of economic importance is a major pre-requisite for any plant breeder to work with crop improvement programmes.

"The genetic diversity available in the crop determines the success of any crop improvement programme. Genetic diversity determines the inherent potential of a cross for heterosis and the frequency of the desirable recombinants in the advanced generation. So, genetic distance plays a vital role, as optimum parental diversity is required to obtain superior genotypes in segregating populations" [2]. Genetic diversity is very effective to predict potential genetic gain. A hybridization programme involving genetically diverse parents belonging to different clusters would provide an opportunity for bringing together gene constellations of diverse nature, promising hybrid derivatives resulted probably due to the complementary interaction of divergent genes in parents. The assessment of divergence in the genotypes is indispensable to know the spectrum of diversity. Improvement in grain yield is normally accomplished through the involvement of genetically diverse parents in breeding programmes. "Mahalanobis's  $D^2$ statistics [3] have been used in several crops, for grouping genotypes into different clusters based on the magnitude of diversity". It is a multivariate statistical tool used to estimate the diversity and cluster the genotypes based on genetic divergence [4]. "Further, it also considers the contribution of each trait towards total diversity. Hence, provides the idea about the total genetic divergence among the genotypes with the estimates of inter-cluster and intra-cluster distance" [5]. "This will assist in identifying diverse parents for hybridization programmes by estimating divergence at intervarietal, species and subspecies levels in classifying problems in crop plants" [6,7,8]. Hence in the current study, eighteen elite rice genotypes are evaluated to assess the genetic variability and nature and magnitude of genetic diversity for further utilization in breeding programmes.

# **2. MATERIALS AND METHODS**

## **2.1 Plant Material**

The present study, comprised of eighteen elite rice genotypes includes medium slender and long slender genotypes, which were grown commercially in Telangana and Andra Pradesh India, have high market demand and are in the regular seed chain. Details of the genotypes along with the name of the research centers where the selected genotypes were developed are furnished in Table 1.

<b>Genotypes</b>	Name of the research center
JGL 1798, JGL 3855, JGL 11470, JGL 18047,	Regional Agricultural Research Station,
JGL 11118, JGL 17004 and JGL 24423	Polasa, Jagtial
<b>KNM 118</b>	Agricultural Research Station, Kunaram,
	Karimnagar District
RNR 15048, Tellahamsa, RNR 2458 and	Rice Research Unit, Rajendranagar,
RNR 2465	Hyderabad
WGL 44, Ramappa and WGL 32100	Regional Agricultural Research Station,
	Warangal
BPT 5204 and MTU 1010	Acharya N. G. Ranga Agriculture University,
	Bapatla and Maruteru
<b>KPS 2874</b>	Agricultural Research Station, Kamapasagar,
	Nalgonda District

**Table 1. List of rice genotypes and research centers where they developed and released**

#### **2.2 Field Experiments**

The Nucleus seeds of selected genotypes obtained from the respective research stations were raised at Seed Research and Technology Center (SRTC), Professor Jayashankar Telangana State Agricultural University, Hyderabad, India (17.3226° N, 78.4172° E and 559 meter above MSL) during Kharif, 2018-19. Thirty days old seedlings of each variety were transplanted in 4 rows of 4 meters length with a spacing of 30 cm between rows and 20 cm between plants in a randomized complete block design (RCBD) with three replications. All the necessary precautions and management practices were adopted to maintain a uniform plant population. The observation was recorded for yield and yield attributing traits, namely, days to 50 per cent flowering, productive tillers per plant, 1000 grain weight, plant height, grains per panicle, panicle length, days to maturity and seed yield per plant (PPVFRA 2001) [9]. The observations on plant height, panicle length, productive tillers per plant and number of filled grains per panicle were recorded from ten randomly selected plants for each entry in each replication. However, days to 50 per cent flowering, days to maturity and grain/seed yield were recorded on plot basis. In contrast, observations for 1000-grain weight were obtained from a random grain sample drawn from each plot in each entry and replication.

## **2.3 Data Analysis**

The data obtained was subjected to standard statistical procedures. Analysis of variance (ANOVA) is used to test whether significant difference among the treatments or not, by following the procedure of randomized complete block design (RCBD) analysis [10]. Genotypic and phenotypic coefficients of variation were computed following the methodology outlined by Burton [11], while the estimates of heritability and genetic advance were obtained as per the procedures outlined by Burton and Devane [12] and Johnson et al. [13] respectively. The  $D^2$ statistics was used to assess the genetic diversity among the 18 elite rice genotypes for eight yield attributing traits, and the procedure was given by C R Rao [14]. The genotypes were grouped into several different clusters by Tocher's method described by Rao (1952) based on the genetic divergence.

## **3. RESULTS AND DISCUSSION**

For the crop to be properly described, assessed, and enhanced, genetic variation within the crop is a crucial component. The genotypic variability of any crop is directly correlated with its improvement. Eighteen elite rice genotypes were assessed in light of the potential threat of genetic erosion and the unrestrained introduction of new variations. All the rice genotypes have shown substantial magnitudes of variance for all the investigated variables are expressed in Table 2. This has demonstrated the hypothesis of this experiment.

# **3.1 Morphological Variability**

The mean values of eight morphological features examined for eighteen elite genotypes were shown in Table 1. The rice genotype JGL 11470 (82.7 cm) was found to be the shortest, and KNM 118 to be the tallest (99.1 cm). Abbasi et al. [15] suggested that reducing plant height could increase the plants' resilience to lodging and lessen the significant yield losses inflicted by this attribute. Genotype Tellahamsa showed less productive tillers per plant (8), whereas KNM 118

$\overline{\mathsf{SI}}$ .	<b>Genotypes</b>	Days to 50%	Days to	<b>Plant</b>	<b>Productive</b>	<b>Panicle</b>	Grains/	1000 grain	<b>Seed</b>
no	name	flowering	maturity	height(cm)	tillers	length(cm)	panicle	weight (g)	yield/plant (g)
1	<b>BPT 5204</b>	112	142	83.6	10.2	20.1	214	13.2	31.3
2	<b>JGL 11118</b>	88	118	95.7	10.4	21.6	233	15.2	36.1
3	<b>JGL 11470</b>	104	134	82.7	9.7	20.5	279	14.3	35.5
4	<b>JGL 17004</b>	72	103	83.7	9.5	20.5	166	13.5	32.6
5	<b>JGL 1798</b>	90	120	94.3	8.8	20.1	185	13.7	27.6
6	<b>JGL 18047</b>	85	120	88.2	10.3	20.7	191	23.1	49.4
	<b>JGL 24423</b>	89	121	95.5	9.9	22.0	236	23.5	49.1
8	<b>JGL 3855</b>	99	129	90.0	10.9	20.6	201	13.7	46.4
9	<b>KNM 118</b>	91	121	99.2	11.0	21.0	182	21.5	48.1
10	<b>KPS 2874</b>	101	131	91.4	10.1	19.9	256	13.5	53.5
11	<b>MTU 1010</b>	92	121	86.9	9.6	19.5	183	23.9	36.5
12	Ramappa	95	123	98.0	9.9	20.8	231	14.9	44.3
13	<b>RNR 15048</b>	93	123	94.5	10.5	21.8	256	12.5	35.1
14	<b>RNR 2458</b>	93	129	90.5	10.1	20.3	227	14.7	43.2
15	<b>RNR 2465</b>	100	130	89.3	10.2	21.8	207	14.5	47.1
16	<b>Tellahams</b>	90	121	90.1	8.0	20.8	174	21.5	37.6
	a								
17	<b>WGL 32100</b>	101	132	97.4	9.4	20.7	176	13.3	33.0
18	<b>WGL 44</b>	105	132	89.1	9.9	21.9	198	12.5	29.5
<b>Mean</b>		94	125	91.1	9.9	20.8	211	16.3	39.8
Range		72-112	103-142	82.7-99.2	$8.0 - 11.0$	19.5-22.0	166-279	12.5-23.9	27.6-53.5
S.E.		0.773	0.9	0.47	0.266	0.29	10.1	0.44	1.29
C.V.		1.4	1.2	0.9	4.6	2.4	8.3	4.7	5.6
C.D. 5%		2.22	2.62	1.36	0.76	0.84	29.31	1.28	3.71
C.D. 1%		2.98	3.51	1.83	1.02	1.13	39.35	1.73	4.98

**Table 2. Mean performance for yield and yield attributing traits of elite rice genotypes**

has shown the highest (11). Rice grain yield and biomass output are closely related to the number of productive tillers [16]. It is demonstrating its value in programmes to boost yield. The greater number of grains per panicle (279) were observed in JGL 18047; on the other hand, genotype JGL 17004 produced lowest number of grains per panicle. The grain yield per plant varies from 27.6 g (JGL 1798) to 53.5g (KPS 2874) with average of 39.8g. The highest 23.9g of 1000 seed weight were recorded in the MTU 1010 and lowest was observed for (12.5g) genotype WGL44.

The days to maturity to vary from 103 days to 142 days among the studied genotypes. The rice genotype JGL17004 recorded early flowering (72days) and maturity (103 days), whereas genotype BPT5204 locally called samba mashuri showed late flowering (105 days) and maturity(142days), having very long vegetative phase. In the current study, the panicle length was found to vary from 19.5 cm (MTU 1010) to 22.0 cm (JGL 18047), with a mean value of 20.70 cm across the genotypes panel. Singh et al, [17] suggested, when choosing parents for a rice crop development programme, one factor to take into consideration is the association between various attributes and the manifestation of genetic variance.

#### **3.2 Genetic Variability, Heritability and Genetic Advance**

ANOVA showed the significant difference for the yield and yield attributing traits among the genotypes (Table 3). A Wide range of variability was observed for days to 50 percent flowering (72-112days), days to maturity (103-142 days), plant height (82.7-99.2cm), number of productive tillers per plant (8.0-11.0), panicle length (19.5- 22.0cm), grain per panicle (166-279), 1000 seed weight (12.5-23.9gm), grain yield per plant (27.6- 53.5gm). The character plant height shown the

more variability among the genotypes indicates the presence dwarf and tall genotypes. Plant height showed greater variation among genotypes, indicating the presence of dwarf and tall genotypes. The variability in 1000 seed weight, grains per panicle, productive tillers per plant, days to 50% flowering, and days to maturity has been demonstrated. The greater magnitude of genetic variability in the population provides good opportunities for the selection of traits to obtain the elite genotypes [18].

The magnitude of the difference between GCV and PCV indicates the extent of environmental influence on a particular trait (Asante et al. [19]). In the current study, PCV was a little greater in magnitude than the corresponding GCV for all the traits studied (Table 4), which showed little influence of environment on the expression of these traits. Similar results are delineated by Rashid et al. [20] and Gyawali et al. [21], Lalitha et al. [18]. PCV ranges from 3.686 to 24.882 and GCV ranges from 2.814 to 24.835. The 1000 seed weight is having highest PCV and GCV (24.882 & 24.835) followed by grain yield per plant (20.315 & 19.523 respectively) indicates that selection for these traits would be effective. The magnitude of difference between PCV and GCV is nearly negligible for the 1000 seed weight and plant height indicating the limited role of the environment in the expression of these traits. The panicle length (2.814) and days to maturity (4.034) have low GCV, which indicates the little scope for improvement of these traits. Similar results were reported by Lalitha et al. [18], Kumar et al. [22] and Srujana et al. [23]. However, days to 50% flowering, plant height, number of productive tillers and grains per panicle showed moderate levels of GCV which argues the considerable scope for genetic improvement of these traits. This result is in line with the reports of Tiwari et al. [24].





*\*& \*\*significance at 5 & 1per cent probability levels, respectively*

<b>Characters</b>	Range		Mean	<b>PCV</b>	<b>GCV</b>	<b>ECV</b>	$H_{\text{bs}}^2$	<b>GA</b>
	Mini.	Maxi.					(%)	
Days to 50 percent	72	112	94	6.551	6.281	1.862	91.9	15.89
flowering								
Days to maturity	103	142	125	4.470	4.034	1.926	81.4	9.611
Plant height (cm)	82.7	99.2	91.1	9.449	9.342	1.418	97.7	24.384
Number of productive	8.0	11.0	9.9	9.830	8.952	4.060	82.9	21.523
tillers per plant								
Panicle length (cm)	19.5	22.0	20.8	3.686	2.814	2.381	58.3	5.671
Grain per panicle	166	279	211	15.619	12.554	9.292	64.4	26.639
1000 seed weight $(g)$	12.5	23.9	16.3	24.882	24.835	1.517	99.6	65.444
Grain yield per plant	27.6	53.5	39.8	20.315	19.523	5.619	92.4	49.530
(g)								

**Table 4. Estimation of Variability, Heritability, and Genetic advance as percent mean of mean for yield attributing traits in rice genotypes**

*PCV-Phenotypic coefficient variation, GCV- Genotypic coefficient variation, ECV -environment coefficient variation, H 2 bs (%) - Heritability in broad sense, GA-Genetic advance as% of Mean (%)*

The variability should be heritable for selection to be efficient (Lalitha et al. [18]. The heritability estimates of yield attributing traits ranged from 58.3% (Panicle length) to 99.6% (1000 seed weight) (Table 4). 1000 seed weight, plant height, grain yield per plant and days to 50% flowering showed high (>90%) heritability compared to other traits. Hossain et al, [25] also reported similar results for days to 50% flowering and panicle length, Mohsin et al. [26] for yield traits of wheat. High heritability indicates the predominance of the genetic component in the expression of phenotype and hence the selection is effective for high heritable traits.

The genetic advance as per-cent of the mean indicates the genetic gain through selection, which is the prime indication of the effectiveness and success of any selection programme. In the current study, the genetic advance as percent of the mean ranged from 5.67% to 65.44%. Genetic advance is higher (>20%) for 1000 seed weight (65.44%), grain yield per plant (49.53%), grains per panicle (26.63%), plant height (24.38%) and number of productive tillers per plant (21.52%), which shows the presence of additive gene action for this traits. Therefore, selection is effective for the improvement of these traits. The heritability accompanying genetic advance will give a reliable picture of the quantum of genetic enhancement made through phenotypic selection. Higher heritability along with higher genetic advance observed for 1000 seed weight, number of productive tillers, plant height and grain yield per plant renders the selection more effective. Also, it showed the presence of greater additive gene action in the inheritance of these

traits. A similar result was reported by Nandan et al. [27] for panicle weight and single plant yield.

## **3.3 Genetic Diversity**

The rice genotypes were grouped into four clusters (Fig. 1) based on  $D<sup>2</sup>$  values such that the genotypes belonging to the same cluster had a smaller divergence than those belonging to different clusters. The distribution of genotypes into four clusters is shown in Table 5. Out of four clusters, cluster I was the largest comprising 14 genotypes and cluster II with two genotypes (BPT 5204, JGL 11470) followed by Cluster III (MTU 1010) and IV (JGL 17004) with one genotype each indicating a high degree of divergence among the genotypes. Similar results of grouping the genotypes into four clusters was reported by Biswash et al. [28] while studying 15 modern aman genotypes of rice based on the  $D<sup>2</sup>$ values (*O. sativa* L.) and Soe, [29] and Rathan et al. [5] reported that the rice genotypes were clustered into seven and five clusters respectively. The average intra and inter cluster  $D^2$  values can be computed from the cluster diagram where the statistical distances among the 18 genotypes were exhibited (Table 6).

The maximum intra cluster distance was observed in cluster I (72.53) with 14 genotypes, followed by cluster II (26.04) with two genotypes, indicating the existence of some genetic divergence among the accessions within clusters. Intra cluster distance for clusters III and IV was zero, since it contains only one genotype each. Rathan et al. [5] reported similar results among the 24 genotypes of rice studied. From the inter cluster  $D^2$  values of the four clusters, it



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#### **Fig. 1. Cluster Diagram of 18 elite rice genotypes by Tocher method**

*1- JGL1798, 2- JGL3855, 3- JGL11470, 4- JGL18047, 5- JGL11118, 6- JGL17004, 7- JGL24423, 8- KNM118, 9 - RNR15048, 10- Tellahamsa, 11- RNR2458, 12- RNR2465, 13- WGL44, 14-Ramapa, 15- WGL32100, 16- BPT5204, 17- MTU1010, 18- KPS2874*

can be seen that the highest divergence was noticed between clusters II and IV (284.47) followed by clusters I and IV (184.86), clusters II and I (177.1), clusters II and III (151.56) and clusters III and IV (149.24), while the lowest was noticed between clusters I and III (125.02). It is assumed that the maximum amount of heterosis would be manifested in cross combinations of the parents belonging to divergent cluster like clusters II (BPT5204, JGL11470), clusters III

(MTU1010) and clusters IV (JGL17004). Similar results were observed by Rajesh et al. [30] in 29 land- races of rice using Mahalanobis's  $D^2$ statistics for eight quantitative characters. The maximum inter cluster distance was recorded between cluster II (BPT5204, JGL11470) and cluster IV (JGL17004), these genotypes may serve as potential donors for future hybridization programmes. The objective is set for high heterosis not only for yield but also for other characters. The greater the distance between two clusters, the wider the genetic diversity between the accessions. The cluster mean revealed the existence of a considerable difference in the mean values for yield attributing traits (Table 7). The distribution of highest and lowest mean values for different characters in distinct clusters indicated the traits contributing to the total divergence. The percent contributions of each trait towards genetic divergence were presented in Fig. 2. The results showed that the contribution of plant height was highest towards genetic divergence (32.68%), followed by 1000 seed weight (31.37%), Days to 50 percent flowering (20.92%), Grain yield per plant (12.42%) The plant height, 1000 seed weight, days to 50% flowering and grain yield per plant contributed 97.39% towards total divergence. Therefore, these characters should be given importance during parental selection in each clusters for hybridization. Similar results were reported by Prasad et al. [31] and Rathan et al. [5] in 37 and 24 accessions of rice (*Oryza. sativa L*.) respectively.

**Table 5. The grouping of 18 rice genotypes into different clusters by D² analysis**

<b>Clusters</b>	<b>No. of Entries</b>	Genotypes
	14	JGL 11118, JGL 1798, JGL 18047, JGL 24423, JGL 3855, KNM 118,
		KPS 2874, Ramappa, RNR 15048, RNR 2458, RNR 2465,
		Tellahamsa, WGL 32100, WGL 44
		BPT 5204. JGL 11470
Ш		<b>MTU 1010</b>
IV		<b>JGL 17004</b>







**Fig. 2. The relative contribution of different characters to genetic diversity**

<b>Clusters</b>	<b>DFF</b>	DM	PН	PL	<b>PTP</b>	GPP	TSW	GYP	
Cluster I	94.83	125.10	93.08	21.03	9.97	211	16.30	41.48	
Cluster II	108.00	137.83	83.13	20.32	10	246.33	13.80	33.46	
Cluster III	92.33	121.33	86.86	19.58	9.60	183.33	23.93	36.50	
Cluster IV	72.40	102.83	83.67	20.47	9.50	166.00	13.50	32.63	

**Table 7. Mean values of eight yield attributing characters of 18 rice genotypes grouped in four clusters**

*DFF- Days to 50% flowering, DM- Days to maturity, PH- Plant height, PL- Panicle length, PTP- Productive tillers per plant, GPP- Grains per panicle, TSW- Thousand seed weight, GYP- Grain yield per plant*

#### **4. CONCLUSION**

The ANOVA indicated the significant differences among the rice genotypes for the traits studied. The 1000 seed weight, grain yield per plant, grain yield per panicle, numbers of productive tillers and plant height have high PCV and GCV along with greater heritability and genetic advance. So, these traits can show greater effectiveness during selection. Further, the high variability existed between the genotypes in different clusters for yield and yield attributes. Among all the characters, 1000 seed weight, plant height, days to 50% flowering and grain yield per plant contributed maximum towards genetic divergence. The maximum inter-cluster distance was noticed between cluster II and IV, cluster I and IV, cluster II and I, cluster II and III. So, the crosses involving elite genotypes BPT5204, JGL11470, MTU1010, and JGL17004 would certainly be rewarding to isolate desirable recombinants during the hybridization programme. This research aids in the identification of diverse parents for rice breeding programs.

## **ACKNOWLEDGEMENT**

The first author is thankful to the Indian Council of Agricultural Research (ICAR), New Delhi for the grant of Junior Research Fellowship (JRF) for the master's degree programme. The authors are thankful to Professor Jayashankar Telangana State Agricultural University, Hyderabad, for providing financial and other facilities to carry out the research work.

# **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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