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Genetic Variability, Heritability and Genetic Advance in F₂ Segregating Population of Cross RNR-15048 x Dokra-Dokri in Rice (*Oryza sativa* L.)

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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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Original Research Article

ABSTRACT

The present experiment was conducted to study the variability, heritability and genetic advance in F_2 segregating population derived from the cross, RNR-15048 x Dokra-Dokri, evaluated in augmented design with two checks at ICAR-IIRR, Hyderabad during the *Kharif*-2022. Analysis of variance revealed the presence of significant differences for all eleven traits examined, indicating the presence of genetic variation between the individuals studied. Higher values of Genotypic

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Coefficient of Variation (GCV) and Phenotypic Coefficient of Variation (PCV) were observed for panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, and single plant yield. Least difference was observed between PCV and GCV value for almost all the traits indicating the little influence by the environment and possibility for genetic improvement of these traits through selection. The traits plant height, panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, spikelet fertility, thousand grain weight and single plant yield recorded high heritability (>60%) associated with high genetic advance as percent of mean (>20) resulting in no environmental influence on the expression and selection for improvement of such characters could be worthwhile. High heritability coupled with moderate genetic advance as percent of mean was observed for panicle length propounding non-additive gene action in their inheritance suggesting heterosis breeding could be useful for improving these traits. Overall high PCV and GCV coupled with high heritability and high genetic advance as percent of mean was noticed for panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, and single plant yield suggesting significant amount of variability with predominance of additive type of gene action. Hence, direct phenotypic selection.

Keywords: Variability; F₂ segregating population; heritability; additive gene action; heterosis; phenotypic coefficient of variation (PCV); genotypic coefficient of variation (GCV).

1. INTRODUCTION

"Rice (Oryza sativa L.) is a major staple food grain crop and carbohydrate-dietary source of more than 50 per cent of the world's population" [1]. "More than 90% of the world's rice is produced and consumed in Southeast Asia and tropical Latin America" [2]. "Despite the fact that India generates 22% of the world's rice supply, population growth in India is expected to jeopardize food security by 2050, putting pressure on future rice demand" [3] "The production per unit area needs to be increased from 2.4 tonnes per ha to 3.4 tonnes per ha assuming the rice area under plough remains at same level [4]. "Although significant improvement in rice production and productivity, was achieved through green revolution, yield plateau limited the further improvement sideways affected by various biotic and abiotic factors. Hence, guaranteeing food security in future is a big challenging task, particularly, for rice breeders in India as rice is pivotal for food and nutritional security in the country. Rice varieties with productivity levels and elevated flexible agronomic features are required for breaking the existing yield threshold for which hybridization is one of the highly adapted and practically feasible solutions for creation of desired variability and achieving high productivity levels" [5].

"Heritability measures a character's transmissibility from one generation to another" [6]. "Presence of Genetic variation among grain yield and yield attributes is the basic requisite to be considered during selection for the successful yield improvement through breeding. High segregation and recombination are observed in segregating populations which can be imposed selection of superior progenies" for [7]. "Coefficient of variation is the measure of variability present in the traits of which genotypic coefficient of variation (GCV) projects heritable variability and phenotypic coefficient of variation (PCV) assesses the effect of environment on the genotype. Differences between PCV and GCV indicates environmental effects on traits viz... small differences indicate low genetic difference, and large differences indicate high environmental effects. Hence small differences between GCV and PCV must be considered for yield improvement in further breeding programs" [8]. "Genetic advance measures the difference between the mean genotypic values of selected population and the original population from which these were selected. Heritability estimates along with genetic advance are more precise in predicting the gain under selection" [9]. "The genetics of traits can be better understood using third- and fourth-degree statistics i.e., skewness and kurtosis in segregating generations" [10]. Hence the present study was undertaken to determine variability, heritability, genetic advance as a percent of the mean in F₂ segregating generation which could be useful in selection and further improvement.

2. MATERIALS AND METHODS

The experiment was carried out at Indian Council of Agricultural Research -Indian Institute of Rice Research (ICAR-IIRR), Rajendranagar, Hyderabad, during *kharif* 2022 with Augmented Randomized Complete Block Design (ARCBD) consisting of 105 F₂ segregating population derived from cross RNR-15048 x Dokra-Dokri presented in Table 1. The experimental area is divided into five blocks where each block was assigned with 24 entries, that included 22 F2'S and 2 checks RNR-15048 and Dokra-Dokri. Prior determining individuals in each block, two checks were initially randomized. The seeds were direct sown in the nursery beds and twenty-five days old seedlings were transplanted in the main field with a spacing of 20 x 15 cm. All necessary intercultural operations, weeding and plant protection measures were taken to ensure uniform and healthy crop stand as per the agronomic practices of rice cultivation. Eleven traits, namely, Plant Height (cm) at maturity stage, Panicle length (PL) (cm), Total Number of Tillers per plant (TNT), No. of Productive Tillers per plant (NPT), Panicle weight (PW), No. of filled grains per panicle (NFG), No. of unfilled grains per panicle (NUFG), Total No. of grains per panicle (TGP), Spikelet fertility (SF) (%), 1000 grain weight (TGW)(gm), and Single plant yield (SPY) (gm) were recorded in every individual F₂ plant.

2.1 Statistical Analysis

The genetic variability parameters, namely, Genotypic and Phenotypic Coefficient of Variation were computed as outlined by Burton [11] while estimates of Heritability was calculated using the formula given by Allard [12] and genetic advance as percentage of mean was calculated by adopting the formula given by Johnson *et al.*1955 [9] by using the R software version 1.4.1717.

3. RESULTS AND DISCUSSION

3.1 Mean Performance

The F_2 data revealed that variance due to treatment (ignoring blocks) was highly significant for all the characters except total number of tillers per plant (TNT) and productive tillers (NPT) revealing that F_2 segregating population under study had enough variation showing the inherent genetic differences between the individuals given in Table-2. The Plant height had a range of variance 86.0 cm to 194.00 cm, with an average of 147.92 cm which is in agreement with Gupta et al. [13]. The average panicle length in the F₂ population studied was noticed to be 28.29 cm, the lowest panicle length was 19.61 cm and the highest was 32.89 cm, similar results were observed by Manojkumar et al. [14]. "Total number of tillers per plant ranged from 4.30 to 18.30 with an average of 9.01 whereas the average number of productive tillers per plant was 7.66, with a range of 4.00 to 14.50. The average spikelet fertility was 71.09 per cent, with the lowest spikelet fertility value being 32.02 per cent and the greatest spikelet fertility value being 96.77 per cent". [15] The average TGW was 24.94 gm, with the lowest and the highest of 12.45 and 39.75 g respectively. Similar variation in F2 were reported by Khandappagol et al. [16]. Single plant yield had a mean value of 27.19 g with minimum and maximum of 8.53 and 56.04 g respectively which is in agreement with the reports of Manojkumar et al. [14].

3.2 PCV and GCV

"The development of a plant characteristic is greatly influenced by genetic variability and heritability. Prior knowledge of population variability estimates and the heritable elements of a trait are helpful tools in any breeding programme. It would be more challenging to enhance a character through selection unless a significant amount of the variation is heritable. Breeding would benefit from knowledge of the character's heritability apart from its phenotypic and genotypic coefficient of variation. In order to compare the variability seen across various traits, the coefficient of variation expressed at the phenotypic and genotypic levels was used" [15].

The results of PCV and GCV are presented in Table 3 indicated high to moderate values for almost all the traits further higher PCV value was observed compared to GCV value, showing the effect of environment. Greater difference between PCV and GCV values was recorded in Total number of tillers (TNT) followed by number of productive tillers (NPT) specifying

Table 1. Details of the parents of the intra-specific cross studied in the present investigation

Parents	Details
RNR-15048	Short slender grain type with low glycemic index (51.72), good cooking quality, test weight of 12.9 g and resistant to blast developed from the cross between MTU1010 and JGL 3855.
Dokra-Dokri	Longest and boldest grain with 14 mm long, high in protein (6.06%) low in fiber with high test weight of 40.5 g.

Source	Treatment	Check	Test	Test vs Check	. Block	Std Error	Std. Deviation	CV	Residuals
Df	106	1	104	1	4				4
PH	927.99 **	10497.60 **	844.03 **	89.97 ^{ns}	17.00 ^{ns}	2.87	29.66	3.1	21.10
PL	6.44 **	62.20 **	5.93 **	3.18 **	1.39 *	0.25	2.55	1.26	0.13
TNT	6.25 ^{ns}	22.50 *	4.47 ^{ns}	174.86 **	2.65 ^{ns}	0.23	2.39	17.82	2.75
NPT	5.16 ^{ns}	25.60 *	3.30 ^{ns}	178.30 **	0.75 ^{ns}	0.19	1.99	17.08	1.85
PW	3.05 **	30.35 **	2.81 *	0.09 ^{ns}	0.23 ^{ns}	0.16	1.68	9.54	0.21
NFG	4175.13 **	41731.60 **	3716.55 **	14311.10 **	186.35 ^{ns}	6	62.04	6.26	119.35
NUFG	940.63 **	722.50 *	857.46 **	9808.63 **	76.15 ^{ns}	2.79	28.9	12.33	60.75
TGP	3939.68 **	53436.10 **	3497.56 **	423.97 *	42.10 ^{ns}	5.8	59.97	2.63	39.10
SF	205.29 **	4.44 ^{ns}	188.63 **	2138.70 **	15.54 ^{ns}	1.32	13.65	4.64	11.20
TGW	26.83 **	1863.22 **	9.31 **	12.75 **	0.56 ^{ns}	0.35	3.57	2.77	0.48
SPY	151.74 ***	19.91 ^{ns}	135.23 **	2000.30 **	8.97 ^{ns}	1.12	11.63	8.36	5.56

Table 2. Analysis of variance for 11 characters of F₂ segregating population

Significant values are as follows: ns- Non-significant, * Significant at 5%, ** Significant at 1%; PH – Plant height (cm); PL – Panicle length (cm); TNT- Total number of tillers; NPT-Number of productive tillers; PW-Panicle weight (g); NFG-Number of filled grains; NUFG- No. of Unfilled grains; TGP- Total grains per panicle; SF – Spikelet fertility (%); TGW- Thousand grain weight (g) and SPY- Single plant yield (g).

Table 3. Magnitude of variability, heritability, and genetic advance for different characters of F₂ segregating population during *kharif*, 2022

Trait	Mean	Min	Max	GCV (%)	PCV (%)	h ²(%)	GA (%)	GAM (at 50%)	Skewness	Kurtosis
PH	147.92	86.00	194.00	19.39	19.64	97.50	58.44	39.51	-0.70 **	2.11 **
PL	28.29	19.61	32.89	8.52	8.61	97.86	4.92	17.38	-0.71 **	3.66 ^{ns}
TNT	9.01	4.30	18.30	14.58	23.49	38.55	1.68	18.68	0.64 **	4.22 *
NPT	7.66	4.00	14.50	15.73	23.72	44.00	1.65	21.53	0.73 **	3.92 ^{ns}
PW	4.78	1.23	9.21	33.73	35.06	92.58	3.20	66.96	-0.07 ^{ns}	2.46 ^{ns}
NFG	171.75	48.60	347.60	34.92	35.50	96.79	121.73	70.88	0.11 ^{ns}	2.62 ^{ns}
NUFG	65.46	6.80	159.80	43.12	44.73	92.92	56.13	85.74	0.51 *	3.13 ^{ns}
TGP	237.21	105.40	384.90	24.79	24.93	98.88	120.64	50.86	-0.04 ^{ns}	2.33 ^{ns}
SF	71.09	32.02	96.77	18.74	19.32	94.06	26.65	37.49	-0.58 *	2.90 ^{ns}
TGW	24.94	12.45	39.75	11.92	12.23	94.85	5.97	23.94	0.25 ^{ns}	5.91 **
SPY	27.19	8.53	56.04	41.89	42.77	95.89	23.00	84.62	0.64 **	2.81 ^{ns}

GCV - Genotypic coefficient of variation; PCV – Phenotypic coefficient of variation; h² – Heritability; GA – Genetic advance; GAM- Genetic advance as percent of mean; PH – Plant height (cm); PL – Panicle length (cm); TNT- Total number of tillers; NPT-Number of productive tillers; PW-Panicle weight (g); NFG-Number of filled grains; NUFG- No. of Unfilled grains; TGP- Total grains per panicle; SF – Spikelet fertility (%); TGW- Thousand grain weight (g) and SPY- Single plant yield (g). higher influence of environment on these traits. resulting in low heritability values. However, panicle length, total grains per panicle, thousand grain weiaht recorded minimum variation between GCV and PCV values, presented in Fig.1 indicating lesser influence of environment resulting in high heritability values (\geq 95 percent). Similar results were reported by Vijayakumari et al. [17] for panicle length and Gupta et al. [13] for thousand grain weight. Higher values of PCV and GCV were observed for panicle weight (35.06, 33.73) number of filled grains (35.50, 34.92), number of unfilled grains (44.73, 43.12), total grains per panicle (24.93, 24.79), and single plant yield (42.77, 41.89). These results were consistent with Dileep Kumar et al. [18], Sravani et al. [19] for above traits; Borah et al. [20], Manojkumar et al. [14] for filled grains and total arains per panicle; Kujur et al. [21], Abdul Fiyaz et al. [22] for single plant yield. High PCV but moderate GCV values were observed for total number of tillers (23.49, 14.58) and number of productive tillers (23.72, 15.73). Moderate values of PCV and GCV were observed for plant height (19.64,19.39), spikelet fertility (19.32,18.74) and TGW (12.23, 11.92). Similar results were observed by Sravani et al. [19] for above three traits; vijayakumari et al. [17] for spikelet fertility; Abdul Fiyaz et al. [22] and Faysal et al. [23] for TGW suggesting greater influence of genetic factors on these traits and relatively less influence of the environment. Low values of PCV and GCV were observed for panicle length (8.61, 8.52) similar values of low PCV and GCV were

reported by Gnaneswari et al. [24] Yumkhaibam et al. [25].

3.3 Heritability (h²), and Genetic Advance (GA)

The coefficient of variation doesn't offer the full scope of heritable variation. When selection parameters heritability in conjunction with genetic advance are followed greater degree of accuracy is possible that are more helpful in predicting the genetic gain under selection than heritability estimates alone. However, it is not necessary that a character showing high heritability will also exhibit high genetic advance Johnson *et al.* [11].

The traits plant height, panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, spikelet fertility, thousand grain weight and single plant yield recorded high heritability (>60%) associated with high genetic advance as percent of mean (>20) presented in Table 3 and Fig. 2,3. Similar results were obtained by Dileep Kumar et al. [18], Sravani et al. [19], vijayakumari et al. [17] for above traits; Manojkumar et al. [14] for panicle length, number of filled grains per panicle, total grains per panicle and test weight; Gupta et al. [13] for plant height, single plant yield; Abdul Fiyaz et al. [22] for total grains per panicle and test weight resulting in no environmental influence on the expression and selection for improvement of such characters could be worthwhile.



Fig. 1. PCV and GCV for different traits in F₂ Population of Rice



Fig. 2. Heritability (h²) for different traits in F₂ Population of Rice



Fig 3. Genetic advance over mean for different traits in F₂ population of rice

High heritability coupled with moderate genetic advance as percent of mean was observed for panicle length earlier reported by Lakshmi *et al.* [26], Gupta *et al.* [13], Borah *et al.* [20] showing greater role of non-additive gene action in their inheritance suggesting heterosis breeding for improving these traits. Moderate heritability and genetic advance as percent of mean was

observed for total number of tillers per plant which is in conformity with the results of Singh *et al.* [27]. Moderate heritability associated with high genetic advance as percent of mean was observed for number of productive tillers per plant indicating the presence of both additive and non-additive gene effects and considerable effect of the environment. Overall high GCV and PCV coupled with high heritability and high genetic advance as percent of mean was recorded for panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, and single plant yield. Similar findings were reported by Shivani et al. [28], Bagudam et al. [29], Manojkumar et al. [14] for filled grains per panicle, total grains per panicle; Bhargava et al. [30] for total grains per panicle; suggesting significant amount of variability with predominance of additive type of gene action. Hence, direct phenotypic selection would be effective for improvement of these traits even in early generations.

4. CONCLUSION

The present study showed presence of adequate variability in F₂ segregating population that could be effectively manipulated using appropriate breeding techniques and program to develop improved varieties. Narrow range of difference between GCV and PCV was observed for panicle weight, number of filled grains, number of unfilled grains, total grains per panicle and single plant yield indicating little influence of environment on the expression of these characters. High GCV, PCV, heritability and genetic advance as a percent of mean were observed for panicle weight, number of filled grains, number of unfilled grains, total grains per panicle, and single plant vield indicating the predominance of additive gene action and the possibility of direct phenotypic selection through these traits even in early generations.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

- 1. Khush GS. What it will take to feed 5.0 billion rice consumers in 2030. Plant molecular biology. 2005;59(1):1-6.
- 2. Fiyaz RA, Shivani D, Chaithanya K, Mounika K, Chiranjeevi M, Laha GS,

Sundaram RM. Genetic improvement of rice for bacterial blight resistance: Present status and future prospects. Rice Science. 2022;29(2):118-132.

- Fathima MA, Geetha S, Amudha K, Uma D. Genetic variability, frequency distribution and association analysis in ADT (R) 48 x Kavuni derived F2 population of rice (*Oryza sativa* L.). Electronic Journal of Plant Breeding. 2021;12(3):659-666.
- Central Rice Research Institute (CRRI) -Indian Council of Agricultural Research. Cuttack (Odisha). -VISION; 2050. Available:http://www.crri.nic.in/ebookcrrivisi on 2050_final_16 Jan13.pdf.
- 5. Ahmed MI, Siddiq EA. Rice. In: Banga, S.S and Banga, S.K. eds. Hybrid cultivar development, Narosa Publishing House, New Delhi, India. 1998;221-256.
- Satheesh kumar P, Saravanan K. Genetic variability, correlation and path analysis in rice (*Oryza Sativa* L.). Int. J. Current. Res. 2012;4:82–85.
- Priyanka A, Gnanamalar R, Banumathy S, Senthil N, Hemalatha G. Genetic variability and frequency distribution studies in F2 segregating generation of rice. Electronic Journal of Plant Breeding. 2019;10(3):988-994.
- 8. Tuhina-Khatun M, Hanafi MM, Yusop RM, Wong M, Salleh FM, Ferdous J. Genetic variation, heritability, and diversity analysis of upland rice (*Oryza sativa* L.) genotypes based on quantitative traits. BioMed research international. 2015.
- Johnson HW, Robinson HF, Comstock RE. Estimates of genetic and environmental variability of Soybeans. Agronomy Journal. 1955;47:314-318.
- Savitha P, Usha kumara R. Genetic Variability Studies in F₂ and F₃ Segregating Generations for Yield and its Components in Rice (*Oryza sativa* L.). Indian Journal of Science and Technology. 2015;8(17).
- 11. Burton GW. Quantitative inheritance in grasses. In: Proc. 6th Inter Grassland Congr. 1952;1: 277–283.
- Allard RW. Principles of Plant Breeding. 2nd ed. John Wiley and Sons. Inc; c. 1960;485.
- 13. Gupta P, Parikh M, Tandekar K. Genetic variability assessment in accessions of rice (*Oryza sativa* L.). 2023;12(3):4721-4725.
- Manojkumar D, SrinivasT, Rao LS, Suneetha Y, Sundaram RM, Kumari VP. Study of genetic variability and trait associations in F2 Population of YH3 x

AKDRMS 21-54 intra-specific cross of rice. The Pharma Innovation Journal. 2022;11(9):1735-1742.

- Manojkumar D, Srinivas T, Rao LS, Suneetha Y, Sundaram RM, Kumari VP. Study of genetic variability and trait associations in F2 Population of YH3 x AKDRMS 21-54 intra-specific cross of rice.
- Khandappagol M, Rajanna MP, Savita SK. Variability and frequency distribution studies in F₂ population of two crosses involving traditional varieties of rice (*Oryza sativa* L). Journal of Pharmacognosy and Phytochemistry. 2019;8(1):1630-4.
- Vijayakumari M, Pillai MA, Senguttuvel P, Saravanan S, Sheela J. Assessment of genetic variability and correlation studies in direct seeded aerobic rice ecosystem. 2022;11(8):398-403.
- Dileep Kumar GD, Abdul Fiyaz R, Viswanatha KP, subbarao LV, Chimote VP, Raghuwanshi KS et al. Assessment of Genetic Variability Parameters Among the F₂ Population of a Cross Between Jaya × Isogenic Line of MTU1010 for Yield and its Component Traits in Rice. Journal of Rice Research. 2022;15(2):118-122.
- 19. Sravani B, Anuradha C, Sundaram RM, Supriya K, Gandhi N, Fiyaz RA. Genetic Variability, Genetic Heritability and Advance in 3K Rice (Oryza sativa L.) Genotypes. International Journal of Environment and Climate Change. 2022;12(11):2259-2265.
- Borah N, Pratim P, Behera BV, Sarma RN. Genetic variability, heritability and genetic advance of aromatic rice for yield and its components. In Biol. Forum-Int. J. 2023;15(5):1516-1522.
- Kujur VK, Abhinav Sao MK, Tiwari A. Genetic variability, heritability and association analyses for yield and related characters in rice germplasm (*Oryza sativa* L.). The Pharma Innovation Journal. 2023;12(4):2236-2240.
- 22. Abdul Fiyaz R, Ramya KT, Chikkalingaiah AB, Gireesh C, RS K. Genetic variability, correlation and path coefficient analysis studies in rice (*Oryza sativa* L.) under

alkaline soil condition. Electronic Journal of Plant Breeding. 2011;2(4):531-537.

- 23. Faysal ASM, Ali L, Azam MG, Sarker U, Ercisli S, Golokhvast KS, Marc RA. Genetic variability, character association, and path coefficient analysis in transplant Aman rice genotypes. Plants. 2022;11(21):2952.
- Gnaneswari VM, Krishnan V, Anandhan T, Vengadessan V, Nadaradja S. Tamilzharasi M. Assessment of genetic variability and diversity analysis in medium duration rice accessions. Electronic Journal of Plant Breeding. 2023;14(1):329-335.
- 25. Yumkhaibam RS, Ram M, Mehandi S. Prakash S. Assessment of genetic variability for yield and yield contributing traits in rice (*Oryza sativa* L.).2023;12(6):1816-1820
- 26. Lakshmi MS, Suneetha Y, Srinivas T. Genetic variability, correlation and path analysis for grain yield and yield components in rice genotypes. Journal of Pharmacognosy and Phytochemistry. 2021;10(1):1369-1372.
- Singh SK, Singh CM, Lal GM. Assessment of genetic variability for yield and its component characters in rice (*Oryza sativa* L.). Research in Plant Biology. 2011;1(4): 73-76.
- Shivani D, Cheralu C, Neeraja CN, Shankar VG. Genetic Variability Studies in Swarna X Type 3 RIL Population of Rice (*Oryza sativa* L.). International Journal of Pure and Applied Bioscience. 2018;6(2):384-392.
- Bagudam R, Eswari KB, Badri J, Rao PR. Correlation and path analysis for yield and its component traits in NPT core set of rice (*Oryza sativa* L.). International Journal of Current Microbiology Applied Sciences. 2018;7(9):97-108.
- Bhargava K, Shivani D, Pushpavalli SNCVL, Sundaram RM, Beulah P, Senguttuvel P. Genetic variability, correlation and path coefficient analysis in segregating population of rice. Electronic Journal of Plant Breeding. 2021;12 (2):549-555.

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