

Journal of Experimental Agriculture International

Volume 46, Issue 9, Page 741-760, 2024; Article no.JEAI.122804 ISSN: 2457-0591 (Past name: American Journal of Experimental Agriculture, Past ISSN: 2231-0606)

# Utilizing Genetic Traits Distributions to Enhance Rice Breeding Programs: A Study of Skewness and Kurtosis in Segregating Generations

# G. S. Shai Prasanna <sup>a</sup>, J. L. Joshi <sup>b\*</sup> and Ajish Muraleedharan <sup>c</sup>

 <sup>a</sup> Division of Crop Improvement, Dhanalakshmi Srinivasan Agriculture College, Thuraiyur Road, Perambalur – 621 212, Tamil Nadu, India.
<sup>b</sup> Agricultural Research Station (ARS), Thirupathisaram-629901, Kanya Kumari, Tamil Nadu, India.
<sup>c</sup> Department of Horticulture, Agricultural College and Research Institute, Killikulam, Vallanad -628262, Tamil Nadu, India.

# Authors' contributions

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

# Article Information

DOI: https://doi.org/10.9734/jeai/2024/v46i92871

#### **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/122804

> Received: 04/07/2024 Accepted: 08/09/2024 Published: 09/09/2024

Original Research Article

# ABSTRACT

This study analyzed the genetic basis of skewness and kurtosis in segregating generations of six rice crosses. Skewness and kurtosis values were calculated for ten biometrical traits across the populations. The traits examined included plant height, number of productive tillers, panicle length, grain yield, and grain dimensions. Results showed diverse patterns of skewness and kurtosis across traits, crosses and generations. Platykurtic distributions were predominant for most traits,

**Cite as:** Prasanna, G. S. Shai, J. L. Joshi, and Ajish Muraleedharan. 2024. "Utilizing Genetic Traits Distributions to Enhance Rice Breeding Programs: A Study of Skewness and Kurtosis in Segregating Generations". Journal of Experimental Agriculture International 46 (9):741-60. https://doi.org/10.9734/jeai/2024/v46i92871.

<sup>\*</sup>Corresponding author: E-mail: jeya.joshi@gmail.com;

suggesting flatter distributions with lighter tails compared to normal. Skewness varied between positive and negative across traits and crosses. Some traits like grain L/B ratio tended to show positive skewness more frequently. Plant height often exhibited negative skewness. The diversity in distribution characteristics indicates significant genetic variability in the populations. Traits showing platykurtic distributions with negative skewness, such as plant height in several crosses, suggest duplicate epistasis and polygenic control. Positively skewed platykurtic distributions for traits like grain dimensions in some crosses indicate complementary epistasis requiring intense selection. Hence, skewness denotes the horizontal pull and kurtosis deals with vertical peaks of the segregating population. The findings provide insights into the genetic control of key rice traits and have implications for breeding strategies. The complex distribution patterns revealed offer a rich basis for selective breeding, depending on specific trait objectives.

Keywords: Skewness; kurtosis; segregating generations and rice.

# 1. INTRODUCTION

Rice genetics and breeding have seen significant advancements in recent decades. Understanding the genetic architecture of complex traits in rice is crucial for crop improvement. Traits such as yield, grain quality, and stress tolerance are typically controlled by multiple genes and are influenced by environmental factors. The study of segregating populations, particularly in early generations like F2 and F3, provides valuable insights into the genetics of these biometrical traits. Analysis of trait distributions in these populations can reveal important information about gene actions, interactions, and the number of genes controlling a trait. Rice played a very important role in the local food security and sustainable development of agriculture, in addition to their significance as genetic resource for rice genetic improvement [1].

Skewness and Kurtosis are statistical measures that provide insights into the asymmetry and flatness, tail heaviness of a frequency distribution of segregating populations in rice, these measures help to understand the nature of the data distribution and make informed decisions about analyzing different plant populations. In case of skewness, if skewness value of zero indicates a perfectly symmetrical distribution, positive skewness suggests that the distribution is skewed to the right, indicating that there may be more plants with lower values and negative skewness suggests that the distribution is skewed to the left, indicating that there may be more plants with higher values. In case of kurtosis, if kurtosis value of zero is considered normal or mesokurtic. Values higher than zero indicate heavier tails, while values lower than zero indicate lighter tails, Positive kurtosis indicates a distribution with heavy tails and a sharp peak (leptokurtic), suggesting that extreme values are more likely and negative kurtosis

indicates a distribution with light tails and a flatter peak (platykurtic), suggesting that extreme values are less likely. In this context, measures of distribution such as skewness and kurtosis offer powerful tools for genetic analysis. Skewness provides information about the asymmetry of trait distribution, while kurtosis indicates the peakedness or flatness of the distribution relative to a normal distribution. These parameters can suggest the presence of major genes, the nature of gene action (additive, dominance, or epistasis), and the relative number of genes controlling a trait. The present study aims to investigate the genetic basis of skewness and kurtosis in F2 and F3 segregating populations derived from six rice crosses. By examining these distribution properties for ten important biometrical traits, we seek to gain insights into the genetic control of these traits. This information will be valuable for designing effective breeding strategies, predicting breeding progress, and ultimately contributing to the development of improved rice varieties to meet future global food demands.

# 2. MATERIALS AND METHODS

The present study on measures of distribution was carried out at the Plant Breeding Farm, Department of Genetics and Plant Breeding, Faculty of Agriculture, Annamalai University, Chidambaram during Kharif season of 2021 and Kharif season of 2022. The seeds from the F1 population of six crosses involving five parents namely ADT 43, ADT 45, ASD 16, ASD 12 and TKM 6 were raised as F<sub>2</sub> population and seeds collected from the F2 plants were forwarded as F3 population. The F2 and F3 generations were derived from the following crosses viz., Cross 1 (ADT 43 X TKM 6), Cross 2 (ADT 43 X ASD 12), Cross 3 (ADT 45 X TKM 6), Cross 4 (ADT 45 X ASD 12), Cross 5 (ASD 16 X TKM 6) and Cross 6 (ASD 16 X ASD 12).

Meas	sures of Skewness	Measures of Kurtosis					
γ <sub>1</sub> value	Type of skewness	γ₂ value	Type of kurtosis				
γ1 is positive	Positively skewed	γ2 is positive	Leptokurtic				
γ <sub>1</sub> is negative	Negatively skewed	γ <sub>2</sub> is negative	Platykurtic				
γ <sub>1</sub> is zero	Symmetrical distribution	γ <sub>2</sub> is zero	Mesokurtic				

Table 1. Interpretation of Skewness and Kurtosis based on obtained values

#### 2.1 Karl Pearson's Measures of Skewness

Skewness refers to the symmetry of distribution., we mean the asymmetry of the distribution to compare the Skewness of different distribution we prefer to use a dimensionless number. If we have more mass to the right of the highest ordinate of the frequency curve, the curve is said to be positively skewed and it's negatively skewed of more mass is to the left of the highest ordinate. A measure of skewness is obtained as making use of the second and third moments about the mean.  $\beta$ 1 as a measure of skewness does not talk about the direction of skewness, *i.e.*, positive or negative. Because µ3 being the sum of cubes of the deviations from mean may be positive or negative but µ<sub>3</sub> is always positive. Also, µ2 being the variance always positive. Hence,  $\beta_1$  would be always positive. This drawback is removed if we calculate Karl Pearson's Gamma coefficient y1which is the square root of  $\beta_1$ . The interpretation method of skewness is presented in Table 1. A skewness values between -1 to +1 is excellent, while -2 to +2 is generally acceptable. Values beyond this suggest substantial nonnormality [2].

Karl Pearson defined the following  $\beta$  and  $\gamma$  coefficients of skewness, based upon the second and third central moments:

$$\begin{aligned} \gamma_1 &= \pm \sqrt{\beta_1} \\ &= \sqrt{\frac{\mu_3^2}{\mu_2^3}} \end{aligned}$$

whereas,

 $\mu_2$  = Second order central moment of distribution  $\mu_3$  = Third order central moment of distribution

$$\mu_2 = \frac{\sum (x - \overline{x})^2}{N}$$
$$\mu_3 = \frac{\sum (x - \overline{x})^3}{N}$$

#### 2.2 Karl Pearson's Measures of Kurtosis

Kurtosis refers to the degree of fatness or peakness in the region about the mode of a

frequency curve. For calculating the kurtosis, the second and fourth central moments of variable are used. The method of interpretation of kurtosis is presented in Table 1.

The measure of kurtosis is the value of the coefficient  $\beta_2$ , it is defined as

$$\boldsymbol{\beta}_2 = \sqrt{\frac{\boldsymbol{\mu}_4}{\boldsymbol{\mu}_2}}$$

whereas,

$$\mu_2 = \frac{\sum (x - \overline{x})^2}{N}$$
$$\mu_4 = \frac{\sum (x - \overline{x})^4}{N}$$

 $\mu_2$  = Second order central moment of distribution  $\mu_4$  = Fourth order central moment of distribution

The deviation of the  $\beta_2$  is used as a measure of kurtosis,  $\gamma_2$  is designed as

$$\gamma_2 = \beta_2 - 3$$

#### 3. RESULTS AND DISCUSSION

The study of distribution properties such as coefficient of skewness (third degree statistic) and kurtosis (fourth degree statistic) provided insight into the nature of gene action [3] and number of genes controlling the traits [4], respectively. The parameter 'd' and 'h' in the genetic expectations of skewness (-3/2 d<sup>2</sup>h) represented additive gene effects and dominance gene effects, respectively. Skewness and kurtosis were more powerful than first order and second order statistics which revealed interaction genetic effects [5].

The skewed distribution of a trait in general suggesting that the trait was under the control of non-additive gene action, especially the epistasis and influenced by environmental variables [6 and 7]. Maximizing the genetic gain in respect of these biometrical traits with positively skewed distribution required more intense selection from the existing variability [7 and 8]. Skewness was a

measure of the asymmetry of the probability distribution of a real-valued random variable about its mean. In a frequency distribution, if the peak is at left and the right tail was longer called as positively skewed and if the peak is towards the right and the left tail is longer, then it was said to be negatively skewed. Positive skewness was associated with complementary gene interactions while negative skewness was associated with duplicate (additive x additive) gene interactions. The genes controlling the trait with skewed distribution tend to be predominantly dominant irrespective of whether they had increasing or decreasing effect on the trait. Genetic gain in respect of all the mentioned traits showing negatively skewed distribution was rapid under mild selection from the existing variability.

Kurtosis was a measure of the "peakness" of the probability distribution of a real valued random variable. A normal distribution had kurtosis exactly 0, called mesokurtic. A distribution with kurtosis (< 0) was called platykurtic. A distribution with kurtosis (> 0) was called leptokurtic. The traits with leptokurtic and platykurtic distribution were controlled by fewer and large number of genes, respectively. Kurtosis was negative or close to zero in the absence of gene interaction and it was positive in the presence of gene interactions [6,5,9]. Studies on the amount of gene interaction were undoubtedly needed so as to increase the efficiency of our selection and breeding programmes. Selection intensity could be higher under complementary than under duplicate relationship. Probably, progress in improving population performance may be greater under complementary interaction than under duplicate gene interaction [5]. The inferences on the relative number of genes and nature of genetic control of different traits in F2 and F<sub>3</sub> generation of two rice crosses are discussed below,

Platykurtic and negatively skewed distribution was observed for plant height, number of productive tillers per plant, total number of tillers, panicle length, number of grains per panicle, and grain yield per plant in both  $F_2$  and  $F_3$  generation and thousand grain weight, grain length traits in  $F_2$  generation in ADT 43 X TKM 6. The traits plant height, panicle length, grain length in  $F_2$ and  $F_3$  generation and trait grain breath in  $F_3$ generation. The traits plant height, panicle length, grain length in both  $F_2$  and  $F_3$  generation and grain breath in  $F_3$  generation of ADT 43 X ASD 12. The traits plant height, number of productive tillers, total number of tillers in both  $F_2$ 

and  $F_3$  generation, grain length in  $F_2$  and panicle length in F<sub>3</sub> generation in ADT 45 X TKM 6. The traits number of productive tillers, total number of tillers, grain breath, grain yield per plant in F2 and F<sub>3</sub> generation then thousand grain weight in F<sub>2</sub> and grain length in F<sub>3</sub> generation in ADT 45 X ASD 12. The traits number of productive tillers, total number of tillers, grain breath, grain length in  $F_2$  and  $F_3$  generation then thousand grain weight, grain yield per plant in F<sub>2</sub> and plant height in F<sub>3</sub> generation in ASD 16 X TKM 6. The traits panicle length, number grains per panicle, thousand grain weight, grain breath in both F2 and F<sub>3</sub> generation and number of productive tillers, grain yield per plant in F3 generation in ASD 16 X ASD 12 which suggested the involvement of more segregating genes with majority of them having increased effects and duplicate type of epistasis. Hence, a greater number of genes segregates for these traits and the selection was not be more effective in other words, it indicated a slow selection response. Similar observations were reported by Karim et al. [10] and Nikhitha et al. [11]. The trait plant height of majority of crosses exerted a significant skewed platykurtic negatively distribution signifying the prevalence of dominance and dominance-based duplicate epistasis, specifying that more than one gene controls the trait. The negatively skewed platykurtic curve indicated that, these traits are under the control of many genes with duplicate gene action. Mild selection will help in achieving faster genetic gain in these traits.

Platykurtic and positively skewed distribution suggested the involvement of relatively large number of segregating genes with majority of them had a decreasing effects and dominance based complementary type of interaction in the inheritance of grain length in F<sub>2</sub> generation and grain length, grain breath, grain L/B ratio and thousand grain weight in F<sub>3</sub> generation of ADT 43 X TKM 6. The traits number of productive tillers, total number of tillers, number of grains per panicle, grain yield per plant in both  $F_2$  and  $F_3$ generation and grain breath, grain L/B ratio in F2 generation of ADT 43 X ASD 12. The traits number of grains per panicle, thousand grain weight, grain breath in both  $F_2$  and  $F_3$  generation then panicle length in F<sub>2</sub> and grain length, grain vield per plant in F3 generation of ADT 45 X TKM 6. The traits number of grains per panicle, panicle length in both F<sub>2</sub> and F<sub>3</sub> generation and trait panicle length in F2 generation of ADT 45 X ASD 12. The traits panicle length, plant height in F<sub>2</sub> generation and traits panicle length, number

F <sub>2</sub>	Cross 1		Cross 2		Cross 3		Cross 4		Cross 5		Cross 6	
Traits	SKW	KRT	SKW	KRT	SKW	KRT	SKW	KRT	SKW	KRT	SKW	KRT
PH	-0.49	-0.54	-0.17	-0.58	-0.06	-1.31	0.02	-1.30	0.10	-1.02	0.40	-0.22
NPT	-0.30	-0.46	0.14	-0.87	-0.12	-0.98	-0.12	-0.98	-0.49	-0.18	0.56	0.03
TNT	-0.27	-0.49	0.09	-0.87	-0.10	-1.03	-0.46	-0.69	-0.54	-0.15	0.70	0.38
PL	-0.33	-1.05	-0.13	-1.03	0.01	-0.53	0.26	-0.84	0.2	-0.49	-0.03	-0.60
NGP	-0.16	-0.78	0.12	-1.19	0.11	-0.76	0.35	-0.65	-3.49	2.57	-0.15	-0.28
TGW	-0.10	-0.20	0.68	0.55	0.26	-0.59	-0.16	-0.38	-0.07	-0.17	-0.26	-0.43
GL	-0.003	-0.53	-0.02	-0.58	-2.47	2.88	-1.59	1.64	-0.05	-0.41	0.18	-0.17
GB	0.01	-0.44	0.03	-0.41	0.017	-0.15	-0.23	-0.13	-0.15	-0.55	-0.01	-0.58
L/B R	0.84	0.76	0.51	-0.004	-2.78	0.19	-1.97	5.64	1.01	1.16	0.21	-0.93
GYP	-0.18	-0.51	0.30	-0.50	0.20	-0.33	-0.29	-0.60	-0.40	-0.20	0.46	-0.01

Table 2. Frequency distribution analysis using skewness and kurtosis values for six rice crosses in F<sub>2</sub> generation

Where, SKW – skewness, KRT – Kurtosis, PH – Plant Height, NPT – Number of Productive Tillers, TNT – Total Number of Tillers, PL – Panicle Length, NGP – Number of Grains per Panicle, TGW – Thousand Grain Weight, GL – Grain Length, GB – Grain Breath, L/B R – Length/Breath ratio and GYP – Grain Yield per Plant

Table 3 Fred	uoney distribution	analysis using skow	nose and kurtosis val	luce for six rico cross	os in E. concration
Table 5. Lieq	uency uistribution	analysis using shew	ness and kultosis va		es in 1 3 generation

F <sub>3</sub>	C	Cross 1		Cross 2		Cross 3		Cross 4		Cross 5		Cross 6	
Traits	SKW	KRT											
PH	-0.53	-0.27	-0.27	-0.96	-0.11	-1.30	-0.08	-1.25	-0.09	-1.22	0.17	-0.75	
NPT	-0.08	-0.41	0.13	-0.97	-0.42	-0.66	-0.65	-0.55	-0.19	-0.27	-0.19	-0.27	
TNT	-0.24	-0.34	0.11	-1.07	-0.67	-0.47	-0.73	-0.32	-0.32	-0.29	0.52	0.46	
PL	-0.40	-0.90	-0.30	-0.81	-0.41	-0.26	0.43	-0.75	0.61	-0.19	-0.20	-0.89	
NGP	-0.35	-0.61	0.20	-1.14	0.41	-0.89	0.48	-0.61	0.09	-1.43	-0.08	-0.58	
TGW	0.14	-0.77	-0.18	0.23	0.27	-0.41	-0.56	1.01	0.12	-0.51	-0.08	-0.41	
GL	0.16	-0.46	-0.07	-0.50	0.20	-0.43	-0.09	-0.58	-0.23	-0.16	0.55	0.45	
GB	0.32	-0.53	-0.03	-0.40	0.15	-0.63	-0.11	-0.65	-0.33	-0.31	-0.13	-0.70	
L/B R	0.43	-0.06	0.56	0.01	0.30	0.51	-4.55	3.48	1.08	1.20	0.25	-0.99	
GYP	-0.13	-0.66	0.31	-0.61	0.44	-0.11	-0.36	-0.89	0.22	-1.05	-0.04	-0.53	

Where, SKW – skewness, KRT – Kurtosis, PH – Plant Height, NPT – Number of Productive Tillers, TNT – Total Number of Tillers, PL – Panicle Length, NGP – Number of Grains per Panicle, TGW – Thousand Grain Weight, GL – Grain Length, GB – Grain Breath, L/B R – Length/Breath ratio and GYP – Grain Yield per Plant

of grains per panicle, thousand grain weight, grain yield per plant in  $F_3$  generation of ASD 16 X TKM 6. The traits plant height, grain L/B ratio in both  $F_2$  and  $F_3$  generation then traits grain length, grain yield per plant in  $F_2$  generation of ASD 16 X ASD 12. The Maximization of genetic gain in respect of the traits with positively skewed

distribution required intense selection from the existing variability [12]. In the present study, a positively skewed and platykurtic curve, suggests that these traits are governed by a large number of genes with dominance based gene interactions. Intense selection is needed for faster genetic gains in these traits.



Fig. 1. Frequency distribution of  $F_2$  generation in cross ADT 43 X TKM 6 for ten biometrical traits



Fig. 2. Frequency distribution of  $F_2$  generation in cross ADT 43 X ASD 12 for ten biometrical traits



Fig. 3. Frequency distribution of  $F_2$  generation of cross ADT 45 X TKM 6 for ten biometrical traits



Fig. 4. Frequency distribution of  $F_2$  generation of cross ADT 45 X ASD 12 for ten biometrical traits



Fig. 5. Frequency distribution of  $F_2$  generation of cross ASD 16 X TKM 6 for ten biometrical traits



Fig. 6. Frequency distribution of  $F_2$  generation of cross ASD 16 X ASD 12 for ten biometrical traits



Fig. 7. Frequency distribution of  $F_3$  generation of cross ADT 43 X TKM 6 for ten biometrical traits



Fig. 8. Frequency distribution of  $F_3$  generation of cross ADT 43 X ASD 12 for ten biometrical traits



Prasanna et al.; J. Exp. Agric. Int., vol. 46, no. 9, pp. 741-760, 2024; Article no.JEAI.122804

Fig. 9. Frequency distribution of  $F_3$  generation of cross ADT 45 X TKM 6 for ten biometrical traits



Fig. 10. Frequency distribution of  $F_3$  generation of cross ADT 45 X ASD 12 for ten biometrical traits



Fig. 11. Frequency distribution of  $F_3$  generation of cross ASD 16 X TKM 6 for ten biometrical traits



Fig. 12. Frequency distribution of  $F_3$  generation of cross ASD 16 X ASD 12 for ten biometrical traits

Leptokurtic and positively skewed distribution was noticed for grain L/B ratio in F<sub>2</sub> generation of ADT 43 X TKM 6. The traits thousand grain weight in F<sub>2</sub> and grain L/B ratio in F<sub>3</sub> generation of ADT 43 X ASD 12. The trait grain L/B ratio in F<sub>3</sub> generation of ADT 45 X TKM 6. The trait grain L/B ratio in F<sub>2</sub> and F<sub>3</sub> generation of ASD 16 X TKM 6. The traits number of productive tillers, total number of tillers in F2 and traits total number of tillers, grain length in F<sub>3</sub> generation of ASD 16 X ASD 12 which is in agreement with outcomes of Sheshaiah et al. [13], Priyanka et al. [14] and Nikhitha et al. [11] for number of productive tillers per plant. The mentioned traits indicated that involvement of fewer segregating genes with majority of these had a decreasing effects and complementary type of interaction in the inheritance of these traits. A leptokurtic and positively skewed distribution can suggest the presence of genetic heterogeneity. This may occur if there are multiple loci affecting the trait. and the alleles at these loci interact in complex ways. It might also indicate that there are some rare, high-impact alleles contributing to the trait.

Leptokurtic and negatively skewed distribution reported for the trait thousand grain weight in F<sub>3</sub> generation of ADT 43 X ASD 12. The trait grain L/B ratio in F<sub>2</sub> generation of ADT 45 X TKM 6. The traits grain length, grain L/B ratio in  $F_2$ generation and traits thousand grain weight, grain L/B ratio in F<sub>3</sub> generation of ADT 45 X ASD 12. The trait number of grains per panicle in  $F_2$ generation of ASD 16 X TKM 6. To achieve maximum genetic gain in respect to this traits, intense selection would be required. The method of using skewness and kurtosis for detecting additive epistasis is very simple and could be used for crop improvement. A leptokurtic and negatively skewed distribution might suggest that genetic constraints or selection pressures favor individuals with higher values of a trait, but there are some genetic factors that lead to lower values. This might imply that there are genetic interactions leading to some individuals having significantly lower trait values, possibly due to deleterious alleles or other factors.

In case of skewness on both generations, traits showed a mix of positive and negative skewness across the crosses. There was no consistent pattern of skewness for most traits across all crosses. Some traits, like grain L/B ratio, tended to show positive skewness more often than others. Plant height often showed negative skewness, indicating a tendency towards shorter plants. In case of kurtosis on both generations,

Platykurtic distributions were predominant for most traits in both generations. This suggests that many traits had flatter distributions with lighter tails compared to a normal distribution. Leptokurtic distributions were less common but occurred occasionally, especially for traits like grain L/B ratio and thousand grain weight. The cross-specific observations reveal that, while there were variations between crosses, no single cross consistently produced dramatically different distribution characteristics compared to the Each cross had its own unique others. combination of skewness and kurtosis values for different traits. The diversity in skewness and kurtosis values across traits, crosses, and generations indicates significant variability in the genetic populations. This variability suggests potential for selection and improvement in breeding programs.

The generation-specific trends of skewness and kurtosis shows that, there were no dramatic shifts in distribution characteristics from F2 to F3 for most traits. Some traits showed slight changes in skewness or kurtosis between generations, but these changes were not consistent across all crosses. Trait-specific observations of important traits shows that, number of tillers (both productive and total) frequently showed negative skewness. Grain yield per plant showed mixed skewness across crosses and generations. Grain L/B ratio often stood out with positive skewness and leptokurtic distributions in several crosses. The skewness and kurtosis values of F2 generation, F3 generation are presented in Table 2 and Table 3, respectively. The Figs. 1 to 6 shows the frequency distribution patten of ten observed biometrical traits of six rice crosses in F2 generation and Figs. 7 to 12 shows the frequency distribution patten of observed ten biometrical traits of six rice crosses in F<sub>3</sub> generation. Similar studies for frequency distribution using skewness and kurtosis analysis were done by Thirugnanakumar et al. [8], Kiran et al. [15], Ajay et al. [16], Hosagoudar and Shashidhar [17], Choudhary et al. [18], Nirubana et al. [19], Priyanka et al. [14], Khandappagol et al. [20], Nikhitha et al. [11], Fathima et al. [21], Shamini and Selvi [12].

# 4. CONCLUSION

In culmination, the platykurtic with negative skewness suggested duplicate epistasis, large number of segregating genes, and slow selection response for traits like plant height, number of productive tillers per plant, total number of tillers, grain yield per plant in ADT 43 X TKM 6, ADT 45 X TKM 6, ADT 45 X ASD 12, ASD 16 X TKM 6 and for number of grains per panicle, panicle length in ADT 43 X TKM 6, ASD 16 X ASD 12 crosses, indicated the polygenic control and duplicate epistasis. The platykurtic with positive skewness suggested complementary epistasis, more segregating genes with decreasing effects, and need for intense selection for traits like grain length, grain breath, grain yield per plant also showed platykurtic and positive skewness for ADT 43 X ASD 12, ADT 45 X TKM 6, ASD 16 X TKM 6, suggesting need for intense selection. The Leptokurtic suggested that, the fewer segregating genes with complementary epistasis for traits like grain L/B ratio in crosses. Traits like grain L/B ratio showed variable skewness and kurtosis across generations and crosses. indicating complex genetic control.

This analysis reveals complex distribution patterns across traits, crosses, and generations. The predominance of platykurtic distributions and the mix of positive and negative skewness values indicate diverse genetic expressions in these rice populations. This variability provides a rich basis for further selective breeding, depending on the desired traits and breeding objectives.

# DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that, NO generative Al technologies such as Large Language Models and text-to-image generators have not been used during writing or editing of this manuscript.

# **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

# REFERENCES

- 1. Prasanna GS, Joshi JL, Muraleedharan A. Distinctiveness, uniformity and stability (DUS) characterization in twenty five landraces of rice (*Oryza sativa* L.). Journal of Advances in Biology & Biotechnology. 2024;27(3):77-84.
- 2. Hair JF, Hult GTM, Ringle CM, Sarstedt M. A prime on partial least squares structure equation modelling (PLS-SEM) (3 ed.). Thousand Oaks, CA; Sage; 2022.
- 3. Fisher RA, Immer FR, Tedin O. The genetical interpretation of statistics of the

third degree in the study of quantitative inheritance. Genetics. 1932;17(2):107.

- 4. Robson DS. Applications of the k 4 statistic to genetic variance component analyses. Biometrics. 1956;12(4):433-444.
- 5. Choo TM, Reinbergs E. Analyses of skewness and kurtosis for detecting gene interaction in a doubled haploid population 1. Crop Science. 1982;22(2):231-235.
- Pooni HS, Jinks JL, Cornish MA. The causes and consequences of nonnormality in predicting the properties of recombinant inbred lines. Heredity. 1977;38(3):329-338.
- 7. Roy D. Analysis of skewness and kurtosis. Plant breeding–The Analysis and Exploitation of Variation. Narosa Publishing House. New Delhi. India, 300-304; 2000.
- Thirugnanakumar S, Narasimman R, Anandan A, Kumar NS. Studies of genetics of yield and yield component characters in F2 and F3 generations of rice (*Oryza sativa* L.). African Journal of Biotechnology. 2011;10(41):7987-7997.
- 9. Kotch GP, Ortiz R, Peloquin SJ. Genetic analysis by use of potato haploid populations. Genome. 1992;35(1):103-108.
- Karim D, Siddique MNA, Sarkar U, Hasnat Z, Sultana J. Phenotypic and genotypic correlation co-efficient of quantitative characters and character association of aromatic rice. Journal of Bioscience and Agriculture Research. 2014;1(1):34-46.
- 11. Nikhitha TC, Pushpham R, Raveendran M, Manonmani S. Genetic variability and frequency distribution studies in F2 population involving traditional variety mappillai samba. Electronic Journal of Plant Breeding. 2020;11(03):933-938.
- Shamini K, Selvi B. Assessment of frequency distribution in F3 generation of sorghum (*Sorghum bicolor* L. Moench.) for grain yield and its attributed traits; 2022.
- 13. Sheshaiah BM, Dushyantha Kumar S, Gangaprasad GN. TH, Gowda Hosagoudar Shashidhar HE. Studies on variability and frequency distribution of yield and yield related traits in F2 population of rice (Oryza sativa L.). International Journal of Current Microbiology and Applied Sciences. 2018;7(9):2048-2052.
- 14. Priyanka AR, Gnanamalar RP, 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.

- 15. Kiran KK, Rao MRG, Suresh K. Variability and frequency distribution studies in F2 population of two crosses of rice (*Oryza sativa* L.). 2013;153-159.
- 16. Ajay BC, Byregowda M, Veerakumar GN, Ganapathy KN, Meena M, Babu HP, Reena Μ. Genetic association and frequency distribution in segregating generations derived from pigeonpea crosses. Indian Journal of Genetics and Plant Breeding. 2016;76(02):181-186.
- Hosagoudar G, Shashidhar H. Studies on variability and frequency distribution of yield and yield related traits in F2 population of rice (*Oryza sativa* L.). Int. J. Curr. Microbiol. App. Sci. 2018;7(9):2048-2052.
- Choudhary AK, Haider ZA, Mishra SB, Prasad K, Bhushan S, Kumar S, Prasad JP. Assessment of genetic variability for yield and yield attributing traits in F2 and

F3 population of Rice (*Oryza sativa* L.) Cross. Current Journal of Applied Science and Technology. 2018 31(2), 1-5.

- Nirubana V, Vanniarajan C, Aananthi N, Banumathy S, Thiyageshwari S, Ramalingam J. Variability and frequency distribution studies in F2 segregating population of rice with phosphorous starvation tolerance Gene (OsPSTOL 1) introgressed. Int J Curr Microbiol App Sci. 2019;8(9):2620-2628.
- 20. 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):1630-1634.
- 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.

**Disclaimer/Publisher's Note:** The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of the publisher and/or the editor(s). This publisher and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.

© Copyright (2024): Author(s). The licensee is the journal publisher. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Peer-review history: The peer review history for this paper can be accessed here: https://www.sdiarticle5.com/review-history/122804