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Identification of Good Restorer Lines through Molecular Confirmation of *Rf3* and *Rf4* Genes in Rice Hybrids

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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

Pollen fertility can be restored by nuclear-encoded genes known as fertility restorer (*Rf*) genes. The initial stage in creating high-yielding heterotic hybrids involves the identification of restorers capable of effectively restoring the fertility of CMS lines. Hybrid rice breeding programs have proven effective in achieving these goals by utilizing cytoplasmic male sterility (CMS) systems in combination with restorer genes. Among these, *Rf3* and *Rf4* genes play critical roles in restoring male fertility and are essential components of successful hybrid rice breeding. In this study, we aimed to investigate the molecular conformation of *Rf3* and *Rf4* genes in a set of restorer lines and

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hybrids in rice. DNA was isolated from 10 restorer lines and 30 rice hybrids, two checks and KMR3R were used for comparison for positive alleles. Two functional markers, RM SF21-5 for *Rf3* and RMS-PRR-9-1 for *Rf4*, were employed to identify the allelic status of the fertility restorer genes. Results revealed that among the 42 rice entries screened, five rice hybrids lacked both *Rf3* and *Rf4* genes and Four rice restorer lines and four hybrids were found to possess both *Rf3* and *Rf4* genes, making them valuable for hybrid breeding programs. Conversely, five hybrids lacked restorer genes and were unsuitable for such breeding programs. These findings establish *Rf3* and *Rf4* as major fertility restoring genes in rice, consistently contributing to complete fertility restoration. The genotyping results provide valuable insights for selecting appropriate parental lines and restorers in hybrid rice breeding programs. The presence or absence of these fertility-restoring genes plays a crucial role in developing high-yielding and productive rice varieties through effective breeding strategies.

Keywords: Rice; Rf3; Rf4; fertility restoration; male sterility.

1. INTRODUCTION

Rice (Oryza sativa) is a crucial crop worldwide, providing sustenance to a significant portion of the global population, it is the most widely grain globallv. consumed food and its consumption is projected to rise by 3% to reach 108 million tonnes, as per the USDA's report in total 2021 The global production [1]. of rice for the period 2021-2022 is estimated to be approximately 515.05 million tonnes, contributing 126,500 with India metric tonnes in the same vear (source: https://www.worldagriculturalproduction.com/crop s/rice.aspx).

A Hybrid rice breeding programs have greatly contributed to achieving high-yielding and superior quality rice varieties. These programs rely on the exploitation of cytoplasmic male sterility (CMS) systems in combination with restorer genes to restore male fertility, enabling successful hybridization. Identification of restorer lines (that restore the fertility of CMS lines) is the foremost step for superior-yielding heterotic rice hybrids [2]. Among the restorer genes, *Rf3* and *Rf4* play pivotal roles in the restoration of male fertility in CMS lines, making them essential components of hybrid rice breeding.

Understanding the molecular conformation of *Rf3* and *Rf4* genes in restorer lines is of paramount importance to ensure their efficacy in hybrid rice production. Molecular markers have emerged as powerful tools to aid in the identification and characterization of these genes, facilitating efficient selection of superior restorer lines by breeders.

Several studies have been conducted to unravel the molecular basis and genetic variation of *Rf3* and *Rf4* genes in rice. In rice, there have been a total of 17 alleles identified for fertility restoration. All of these alleles, except rf17, are dominant in rice. At least two specific genes, Rf3 (located on Rf4 chromosome and (located 1) on chromosome 10), are known to be responsible controlling fertility restoration of for WA cytoplasm in rice [3,4]. According to the researcher [5] developed gene-based functional markers, namely RMS-PPR9-1 for Rf4 and RMS SF21-5 for Rf3. These markers serve as valuable tools for identifying and studying the Rf4 and Rf3 genes. For instance, Li [6] conducted a comprehensive analysis of the genetic diversity and molecular characterization of Rf3 and Rf4 genes in a diverse panel of restorer lines. Their study revealed multiple allelic variants of Rf3 and Rf4 genes, emphasizing the significance of considering allelic diversity for effective hybrid rice breeding. Furthermore, the development of advanced molecular techniques, such as polymerase chain reaction (PCR) and DNA sequencing, has facilitated the creation of specific markers for detecting the presence of Rf3 and Rf4 genes in rice lines. Zhou [7] successfullv developed allele-specific PCR markers for Rf3 and Rf4 genes, allowing rapid and accurate screening of restorers in breeding programs. In addition to genetic diversity, the molecular conformation of Rf3 and Rf4 genes has also been linked to functional characteristics. Liu [8] conducted a study focusing on the molecular structure and function of the Rf3 gene in rice. Their research revealed that Rf3 encodes a protein with a pentatricopeptide repeat (PPR) domain, which plays a critical role in RNA binding and processing.

The main aim of this study is to identify good restorer lines which are having high restoring ability by confirming *Rf3* and *Rf4* genes through molecular studies.

2. MATERIALS AND METHODS

Plant material: The present study was conducted at Regional Agricultural Research Station. Warangal. Professor Javashankar Telangana State Agricultural University (PJTSAU) during rabi, 2023 with an objective of molecular confirmation of Restores lines for presence of two major fertility restorer genes i.e. Rf3 and Rf4 through functional markers (Table 1).

Marker assisted selection for fertility restorer genes: DNA was isolated from the 10 restorer lines and 30 rice hybrids along with two checks by following the protocol of Zheng et al., [9] and potential restorer KMR3R is used as a reference to compare the positive alleles. The PCR based SSR marker RM SF21-5 and RMS PRR 9-1 were used to identify the allelic status with respect to two major fertility restorer genes i.e. Rf3 and Rf4. PCR was performed using 1 U of Taq DNA polymerase (Fermentas, Lithuania) and 1x PCR buffer (Genei, India) in 10-µl reaction volume with a thermal profile of 94 °C for 5 min (initial denaturation), followed by 35 cvcles of denaturation at 94°C for 30s, annealing at 55°C for 30s, extension at 72°C for 1 min and a final extension of 7min at 72°C. The amplified product of Rf3 and Rf4 were electrophoretically resolved on a 3.5% Seakem LE® agarose gel (Lonza, USA) containing 0.5 mg/ml of ethidium bromide in 0.5x TBE buffer and visualized under UV.

The primer RM-SF21-5 for Rf3 produced a positive band at 172bp indicating the presence of the restorer allele, while a negative band appeared at 127bp indicating the absence of the restorer allele with the parent KMR3R. On the other hand, the functional marker RM SF21-5 for Rf3 showed positive alleles at 172bp and negative alleles at 127bp.

3. RESULTS AND DISCUSSION

A total of 30 rice hybrids and 10 restorer lines including two checks were screened to determine

the presence or absence of fertility restoration genes, namely *Rf3* and *Rf4*. The screening process involved the use of specific primers i, e. RM-SF-21-5 for *Rf3* and RMS-PRR-9-1 for *Rf4*.The results representing the screening of *Rf3* and *Rf4* are depicted in Table 2.

Molecular confirmation of *Rf3* gene by using RMS-SF-21-5 functional marker: The 30 rice hybrids and 10 restorers including two checks were screened for the presence of Rf3 gene by using RM SF21-5 functional marker, The results revealed that seven restorers (JGL24502, JGL36147, JGL36172, JGL38156, JGL36199, WGL1272, JR70) and 10 hybrids (CMS59AXJGL36147. CMS59AXJGL36172. CMS59AXJGL38156. CMS52AXJGL36172. CMS52AXJGL38156. CMS52AXJGL36199. CMS52AXJR70. CMS64XJGL38156. CMS64AXJGL36199, CMS64AXJR70) showed the prescence of Rf3 with similar banding with KMR-3.

On the other hand, 3 restorer lines (JGL24355, JGL24440. JGL35149), 17 hybrids (CMS59AXJGL24355,CMS59AXJGL24502,CMS 59AXJGL24440,CMS59AXJGL35149,CMS59AX JGL36199,CMS59AXWGL1272,CMS59AXJR70, CMS52AXJGL24355,CMS52AXJGL24440,CMS 52AXJGL35149,CMS52AXJGL36147,CMS52AX WGL1272,CMS64AXJGL24355,CMS64AXJGL2 4502,CMS64AXJGL36147,CMS64AXJGL36172, CMS64AXWGL1272) and 2 checks (US312,KRH4) showed the abscence of Rf3 gene. Moreover, three rice hybrids were found to heterozygosity exhibit for both alleles. These identified hybrids were as CMS52AXJGL24502, CMS64AXJGL24440, and CMS64AXJGL35149.

The Lane number M represents 100bp ladder, P represents positive control (KMR-3R), while the numbers from 1 to 42written on the top of gels represents list of rice entries used for present study and details were given in Table 1.



Fig. 1. Molecular confirmation of Rf3 gene by using RM SF21-5functional Marker

Table 1. List of markers used in the present study for screening of fertility restorer genes

S. No	Primer	Seque	nce	Chromosome	Annealing temperature °C	Reference
1.	RM SF 21-5	F	GAGTTGGGGGTCGAGAAATC	10	55°C	Pranathi <i>et al</i> .,
	(Rf3)	R	CGTACGTGCGGCTAGGATCAA			[5]
2.	RMS PRR 9-1	F	GAGTTTTGAATAGATTTACGTGTGGA	1	55°C	Pranathi et al.,
	(Rf4)	R	AGTGTCCAGATTCGTAGTAATGC			[5]

Table 2. Details of rice hybrids and their genotyping results by using functional markers for the present investigation

S. No.	Name of the rice entries	Rf3 gene: RM SF 21-5 marker	Rf4 gene: RMS PRR-9-1 marker
PARENT	KMR3R	RR	RR
Restorer Lines			
1.	JGL24355	rr	RR
2.	JGL24502	RR	rr
3.	JGL24440	rr	RR
4.	JGL35149	rr	RR
5.	JGL36147	RR	RR
6.	JGL36172	RR	RR
7.	JGL38156	RR	rr
8.	JGL36199	RR	rr
9.	WGL1272	RR	RR
_10.	JR 70	RR	RR
Hybrids			
11.	CMS59AXJGL24355	rr	RR
12.	CMS59AXJGL24502	rr	rr
13.	CMS59AXJGL24440	rr	Rr
14.	CMS59AXJGL35149	rr	rr
15.	CMS59AXJGL36147	RR	RR
16.	CMS59AXJGL36172	RR	RR
17.	CMS59AXJGL38156	RR	rr
18.	CMS59AXJGL36199	rr	RR
19.	CMS59AXWGL1272	rr	RR
20.	CMS59AXJR70	rr	RR
21.	CMS52XJGL24355	rr	rr

22.	CMS52XJGL24502	Rr	rr		
23.	CMS52XJGL24440	rr	Rr		
24.	CMS52XJGL35149	rr	rr		
25.	CMS52XJGL36147	rr	RR		
26.	CMS52XJGL36172	RR	RR		
27.	CMS52XJGL38156	RR	Rr		
28.	CMS52XJGL36199	RR	rr		
29.	CMS52XWGL1272	rr	rr		
30.	CMS52XJR70	RR	RR		
31.	CMS64XJGL24355	rr	RR		
32.	CMS64XJGL24502	rr	RR		
33.	CMS64XJGL24440	Rr	RR		
34.	CMS64XJGL35149	Rr	Rr		
35.	CMS64XJGL36147	rr	Rr		
36.	CMS64XJGL36172	rr	RR		
37.	CMS64XJGL38156	RR	rr		
38.	CMS64XJGL36199	RR	rr		
39.	CMS64XWGL1272	rr	rr		
40.	CMS64XJR70	RR	rr		
Checks					
41.	US312	rr	RR		
42.	KRH4	rr	RR		

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Fig. 2. Molecular confirmation of Rf4 gene by using RMS PRR9-1 functional Marker

M P 1 2 3 4 5 6 7 8 9 10 11 12 13 14 1516 17 18 19 M P 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 3

Molecular confirmation of Rf4 gene by using RMS PRR 9-1 functional marker: The 30 rice hybrids 10 restorer lines including two checks. when screened for the presence of Rf4 gene by using RMS PRR 9-1 functional marker, The results revealed that thereare7restorerlines(JGL24440,JGL35149,JGL 36147, JGL36172, JGL24355, WGL1272, JR70) 13 hybrids(CMS59AXJG24355,CMS59AXJGL36147 ,CMS59AXJGL36172,CMS59AXJGL36199,CMS 59AXWGL1272.CMS59AXJR70.CMS52AXJGL3 6147.CMS52AXJGL36172.CMS52AXJR70.CMS 64AXJGL24355,CMS64AXJGL24502,CMS64AX JGL24440,CMS64AXJGL36172)and2checks(US 312,KRH4) showed the prescence of Rf4 with similar banding with KMR-3.

Out of the 42 rice entries screened, four rice restorer lines (JGL36147. JGL36172. WGL1272.JR70) and hvbrids four (CMS59AXJGL36147, CMS59AXJGL36172. CMS52AXJGL36172) CMS52AXJR70, were found to carry both major fertility restorer genes, Rf3 and Rf4. Conversely, we also observed five rice hvbrids (CMS59AXJGL24502, CMS59AXJGL35149, CMS52AXJGL24355, CMS52AXJGL35149, and CMS52AXWGL1272) that lacked both major fertility restorer genes, Rf3 and Rf4.

The Lane number M represents 100bp ladder, P represents positive control (KMR-3R), while the numbers from 1 to 42 written on the top of gels represents list of rice entries used for present study and details were given in Table 2.

Researchers have extensively studied and confirmed the effectiveness of Rf3 and Rf4 markers in fertility restoration through various studies. For instance, Shidenur [10] conducted screening on 310 NPT lines to assess fertility restoration using DRRM-Rf3-5, DRRM-Rf3-10, and functional markers RMS-SF21-5, RM6100, RMS-PPR9-1. Similarly, Pranathi and [5] undertook screening to distinguish between 120 restorers and 44 non-restorers based on fertility restoring ability. They also developed functional markers for Rf3 and Rf4 to aid in their research. Venkanna [2] conducted a study focusing on the fertility restoration of CMS-WA lines, which is mainly governed by two independent and dominant nuclear fertility restoring genes, Rf3 and Rf4. In their study, they aimed to genotype 25 rice genotypes to determine the presence of Rf3 and Rf4 genes using functional markers.

4. CONCLUSION

Based on the genotyping results for fertility restoration, it was observed that the four rice restorer lines (JGL36147, JGL36172, WGL1272, JR70) and four hybrids (CMS59AXJGL36147, CMS59AXJGL36172, CMS52AXJGL36172, CMS52AXJR70) possess both *Rf3* and *Rf4* genes. Among ten restorer lines used in this study, three restorer lines have capability of restoring fertility in rice hybrids. Hence these three restorer lines can be used in future hybrid breeding programmes.

5. FUTURE SCOPE

The future scope of this study lies in expanding the investigation to a larger and more diverse set of rice genotypes to further validate the role of Rf3 and Rf4 genes in fertility restoration. Additionally, exploring other potential restorer and incorporating advanced genes technologies aenomic can enhance the precision and efficiency of hybrid rice breeding programs.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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