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Marker Assisted Selection in Advanced Backcross Population of Rice Variety, MTU1010 for Bacterial Blight and Blast Resistance

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Authors' contributions

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

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ABSTRACT

The present study was aimed for identification of the bacterial blight and blast resistant lines in the advanced back cross population of MTU-1010 using marker assisted foreground selection and also for evaluation of agro-morphological characters. The M-16-59 introgressed line (derived from an intercrossing of BC2F1 plants of MTU-1010 x GPP2 and MTU-1010 x NLR 145) is developed under ongoing DBTDBSRR subproject -IV possessing *Xa21, xa13*, *Pi1* and *Pi54* resistance genes having broad spectrum resistance to bacterial blight and blast is used as a donor parent and MTU-1010 was used as a recurrent parent for the back cross. Before attempting the backcross both the parents were verified for the target genes along with the original donors GPP2 and NLR 145 using gene specific/linked molecular markers *viz*., *xa13-promo* for *xa13* gene, *pTA248* for *Xa21* gene, RM 224 for *Pi1* and *Pi54 MAS* for *Pi*54. under the F1 generation (equivalent to BC3F1 because two backcrosses were completed earlier), 120 plants were screened, and 16 plants were confirmed for xa13, Xa21, Pi1, and Pi54 genes under heterozygous conditions. These confirmed plants were assessed for agro-morphological characteristics such as yield, grain features, and plant type. The results showed that the confirmed heterozygous plants were comparable to MTU-1010 and passed to the next round of selection and evaluation. The $BC_1F_1-198th$ plant (96.8%) was selfed to produce BC_1F_2 population. Twenty five plants were advanced to BC_1F_3 generation based on BB resistance and above other phenotypic characters. The selected BC_1F_3 progenies were screened for blast and BB resistance. Four BC1F³ progenies with four target genes (*xa13xa13Xa21Xa21Pi54Pi54Pi1Pi1*) showed very high level of resistance to both the diseases. $BC_1F_2-198-52^{nd}$ line found similar to MTU1010 with respect to yield and yield related characters besides showing resistance to both BB and blast.Improved lines of MTU1010 can be advanced for multi-location testing under All India Coordinated Rice Improvement Project (AICRIP) for their evaluation and possible release for the benefit of rice farmers.

1. INTRODUCTION

"Rice is the staple food for more than half of the world's population, and global rice demand is estimated to rise from 6.76 × 108 t in 2010 to 8.52 × 108 t in 2035" (Khush, 2013). "To produce 1.76 \times 108 t additional rice, it is needed to increase the yield and also minimize the yield loss caused by various diseases and insect pests. Among the biotic stresses, bacterial blight (BB) and blast are important diseases that results in significant yield reduction worldwide. BB is caused by a bacterium, *Xanthomonas oryzae pv. oryzae*, which is a serious problem in irrigated and shallow lowland conditions in India causing yield losses ranging from 74 to 81% based on severity of the disease" (Srinivasan et al., 2005). "Rice blast disease, caused by *Magnaporthe oryzae,* is one of the most serious diseases of rice. While it is present nearly everywhere rice is grown, blast is more of a

problem in the temperate flooded and tropical upland cropping systems, marked by cooler climates" (Scardaci et al., 1997). "In Andhra Pradesh and Telangana yield losses are very high especially in Nellore, West Godavari and Rangareddy districts" Rajarajeswari (Rajarajeswari et al., 2006). "Breeding and the development of resistant cultivars carrying major resistance (R) genes have been the most effective and economical strategy to control BB disease to have a neutral effect on the environment" (Huang et al., 1997, Singh et al., 2001) and (Jena and Mackill, 2008). "However, cultivars undergo rapid breakdown in their resistance mainly by the emergence of new pathotypes, due to the high instability in the genome of the pathogen" (Dean et al., 2005). "Therefore, bringing together multiple genes conferring resistance to more than one pathotype into one genetic background is necessary for durable resistance. However, conventional

Keywords: Bacterial leaf blight; blast; molecular markers; foreground selection; background selection; recurrent parent genome; molecular breeding; MTU1010; xa13; Xa21; Pi54 and Pi1 genes.

breeding methods to improve rice cultivars for BB resistance have not found much success" (Shin et al., 2011).

To date, at least 40 BB resistance (Kim et al., 2015) "genes conferring host resistance against various strains of *Xanthomonas oryzae pv. oryzae (Xoo* have been identified. Out of this 40, 29 dominant and 11 recessive genes have been identified and registered" (Ranjith. Ellur K et al., 2015). "Using MAS breeding approaches three or more BB genes, like *xa5, xa13, Xa21* have been successfully pyramided in diverse elite rice varieties like IR64, PR106, Pusa Basmati 1, Lalat, Tapaswini, Swarna, IR64 and Samba Mahsuri" (Sundaram et al., 2014). "Using the gene pyramid approach, a three-gene combination appeared to be the most effective with *Xa21* contributing the largest component of resistance" (Pradhan et ai., 2015). "Globally, 100 rice blast major resistance genes (R-genes) have been identified (Devanna et al*.,* 2014)), out of which 19 blast resistance genes have been cloned and over 50 major rice blast R genes have been mapped" (Hayashi et al., 2005, Chen et al., 2006). "The most of identified blast R genes were found in a cluster on chromosome 6, 11 and 12 (Yang et al*.,* 2008). Recently the *Pi1* leaf blast resistance gene has been introgressed into the D521 line derived from the donor line BL122" (Fu et al., 2012).

"M-16-59, a gene stacked line developed from the ongoing DBTDBSRR sub project IV funded by DBT, Government of India at Institute of Biotechnology possessed two BB resistance genes (*xa13* and *Xa21*) and two blast resistance genes (*Pi 54* and *Pi1*). This line was developed by intercrossing of BC_2F_1 plants of MTU-1010 x GPP2 and MTU-1010 x NLR 145. M-16-59 is carrying the four genes with 85% recurrent parent genome (MTU 1010) and was recovered from the ICF₂ segregating population. Though this introgressed line is having four genes with 85% recovery it still resembles its donor" (Zheng et al., 2016). So, in the present study an attempt was made to in progress BB and blast resistant genes *xa13, Xa21, Pi54 and Pi1* to further increase the resistance and also to improve the recovery of MTU-1010. Resistance genes linked/gene specific molecular markers were used for foreground selection while polymorphic primer pairs that are spread all over rice genome were used for background selection to carry out MAS. In our study MTU-1010 is used as recurrent parent and M-16-59 is used as donor parent.

2. MATERIALS AND METHODS

2.1 Plant Material

Cottondora Sannalu (MTU 1010) is an elite mega rice variety derived from the cross, Krishnaveni/ IR64 possessing short duration, high yielding ability and long slender grain quality was used as recurrent parent. M-16-59, an introgressed MTU1010 line possessing four [two BB (xa13 and *Xa21*) and two Blast (*Pi54* and *Pi1*)] biotic stress resistance genes with 85% MTU-1010 genome was used as the donor parent in the present study. The nucleus seed of MTU-1010 and NLR145 were obtained from APRRI, Maruteru respectively, while GPP2 seed which was used for positive check control were obtained from ICAR-IIRR, Rajendranagar, Hyderabad.

2.2 Molecular Marker Analysis

MTU-1010 and M-16-59 were verified for the target genes along with the original donors GPP2 and NLR145. This verification was carried out using gene specific molecular markers (Table 1) viz., xa13-promo, pTA248, Pi54 MAS for *xa13, Xa21, Pi54* genes respectively. While a gene linked marker RM224 was used for *Pi1* gene.

2.3 Genomic SSR Markers for Background Selection

Parental polymorphism survey between donor and recurrent parents was carried out by using 354 genomic SSR markers covering all the twelve chromosomes, selected from gramene data base [\(www.gramene.org\)](http://www.gramene.org/) (Fig. 1). Polymorphic markers were used for background selection in F_1 , BC₁F₁ and BC₁F₂ generations. In the earlier study of Aruna Kumari 2013, 616 SSR markers were tested covering all twelve chromosomes, while 108 markers showed polymorphism between recurrent parent (MTU-1010) and the two donor parents (GPP2 and NLR 145) during the development of M-16-59, an introgressed line of MTU-1010 carrying *xa13, Xa21, Pi54* and *Pi1* genes.

2.4 Generation of F¹ Material

Staggered sowings were taken up with 7 days interval to obtain synchrony between MTU-1010 and donor parent M-16-59 in order to make crosses. Twenty-five days old seedlings were transplanted in two row plots with a spacing of 20 x 20 cm. Fertilizer application, inter cultivation,

water management and plant protection
measures were adopted as per the measures were adopted as per the recommendation of PJTSAU. F_1 material was generated by making cross between MTU-1010 and M-16-59 during wet season, 2014 at ARI, Rajendranagar.

2.5 Generation of BC1F1, BC1F² and BC1F³ along with Parent Material

F1 seeds were raised at ARI, Rajendranagar during dry season, 2014-15. DNA was isolated from F_1 plants and were verified for the four target genes *xa13, Xa21, Pi54* and *Pi1* using the foreground markers viz., *pTA248, xa13 prom, Pi54-MAS* and RM224, respectively. The one true hybrid (*Xa13xa13, Xa21xa21, Pi54pi54, Pi1pi1*) plant was backcrossed with MTU-1010 (using the F_1 as male parent and MTU-1010 as female parent) to generate BC_1F_1 seeds. After foreground and background selection in BC_1F_1 generation, the BC_1F_1 plant with high recurrent parent genome (RPG) was selfed to generate BC₁F₂ generation. Foreground selection was carried out in 1060 BC1F² population. Background analysis was carried out in 20 selected BC₁F₂ plants. In addition to twenty selected four and three gene BC_1F_2 plants, five plants with two gene combination were also selected based on their resistance to BB and selfed to generate BC_1F_3 generation.

2.6 DNA Extraction and PCR Analysis

DNA was isolated from the leaf samples according to Zheng et al. (1996). The quality and quantity of DNA was estimated in 0.8% agarose gel using 500ug/ml lamda (ë) Hind III DNA (New England Biolabs) as reference standard. PCR was carried out to detect the presence of four genes. PCR and gel electrophoresis protocols recommended by Sundaram et al. (2008) and Ramkumar et al. (2011) were adopted for marker-assisted selection of target genes *xa13, Xa21, Pi54* and *Pi1,* respectively.

2.7 Evaluation of Agro-morphological Characters

The F_1 plants were transplanted in the main field at a spacing of 20 cm \times 15 cm along with the donor and recurrent parents. Standard agronomic practices were followed to raise a healthy crop and agro-morphological characters like days to 50% flowering, days to maturity; plant height (cm), number of productive panicles per plant, panicle weight (g), panicle length (cm), grain yield per plant (g), 1000 grain weight (g) and grain type were recorded.

2.8 Evaluation of BC1F2 and BC1F³ Progenies for Agro-Morphological Parameters

The BC_1F_2 plants showing homozygosity for 2, 3 and 4 target genes were advanced to $BC₁F₃$ generation. 25 $BC₁F₃$ progenies along with MTU-1010 were grown during wet season, 2016 at ARI, Rajendranagar. The phenotypic data was recorded on twenty five BC_1F_2 plants possessing four target genes *viz., xa13xa13,Xa21Xa21, Pi54Pi54,Pi1Pi1* (4 plants), 3 genes (16 plants) and 2 genes (5 plants) in different combinations in homozygous condition for Days to 50% flowering (DFF), Plant height (cm) and Grain type were recorded along with the recurrent parent MTU-1010. In BC_1F_3 the material was raised in Randomized Block Design (RBD) with two replications. Each progeny was planted in 3 rows with a spacing of 20 X 15 cm. A healthy crop was raised by following standard agronomic practices recommended by PJTASU. Data was collected from five randomly selected plants from each replication for Days of 50% flowering (DFF), Plant height (cm), No. of panicles per plant, Number of filled grains per panicles, Panicle length (cm), Grain yield per plant (g), Thousand seed weight (g) and Grain type. Data on DUS characters *viz.,* Basal Leaf : Sheath Color, Leaf : Auricles, Leaf : Anthocyanin Colouration of auricles, Leaf : Shape of ligule, Leaf : color of ligule, Flag Leaf : Attitude of blade (Early observation), Time of heading (50% of the plants with panicles), Lemma : Anthocyanin coloration of area below apex, Stem length (excluding panicles; excluding floating rice), Stem : Anthocyanin coloration of nodes, Panicle : Length of main axis, Flag Leaf : Attitude of blade (late observation), Panicle : Curvature of main axis, Spikelet : Color of tip of lemma, Panicle : Awns, Panicle : Attitude of branches, Panicle : Exsertion, Sterile lemma : Color, Leaf : Senescence, Panicle : Presence of secondary branch, Lemma and palea color was collected in comparison with recurrent parent. The data on DUS characters was recorded as per the guidelines (Subbaet al., 2013).

2.9 Statistical Analysis

The data collected from BC_1F_3 progenies, which was raised in RBD design was subjected to analysis of variance (ANOVA), by using OPSTAT version 9.1 software.

3. RESULTS

In the present study to improve the recurrent parent genome recovery and development of resistant lines against bacterial blight and blast of MTU-1010 Marker assisted breeding has been successfully applied (Hari et al., 2013, Khanna et al., 2015) as MAS saves time and offers a very simple efficient and accurate method (Singh et al., 2012).

Verification of the Parents for the Resistance Genes Using Gene Specific/Linked Markers: Verification of the parents for the resistance genes using gene specific/ linked polymorphic markers is an important prerequisite before starting marker-assisted backcross breeding. A marker which is monomorphic bears no value in selection work because this type of marker cannot distinguish the two parental genotypes viz. MTU-1010, the recurrent or recipient parent and M-16-59, the donor parent of the MABC program. A total of 4 primers specific to *Xa21,*

xa13 BB genes and *Pi1, Pi54* blast genes were surveyed for finding out polymorphic markers and all of them were found as polymorphic.

The results (Fig. 2) revealed that 500bp resistance allele of *xa13* gene was amplified with *xa13 promoter* primer in M-16-59 line. This band was exactly identical to the band that was amplified in the check material, GPP2. (Fig. 3) The marker *pTA248* amplified 900bp resistance allele in M-16-59 line, which was similar with that of GPP2 confirming that the resistant parent was carrying *Xa21* gene. Magar et al*. (2014)*, Hajira Shaik et al*. (2014)* and Balachiranjeevi et al*. (2015)* also utilized *xa13 promoter* and *pTA248* primers for validation of parents and foreground analysis in backcross derived population. In the similar way 250bp resistance allele of *Pi54* gene was amplified with *Pi54-MAS* in M-16-59 line and is similar to NLR145 which is used as the positive check control and also for *Pi1* gene in M-16-59 line the band is obtained at 150bp a resistant allele, when amplified with RM224, which is identical to that of original donor NLR145. These results confirmed that M-16-59 line was carrying *xa13*, *Xa21, Pi54* and *Pi1* genes. Jamal-oddin et al*., (2015)* used the same primers *Pi54-MAS* and RM224 for *Pi54* and *Pi1* genes in MAS.

Fig. 1. Parental polymorphism between recurrent parent and donors with genomic SSR markers

M: 50 bp LadderP: MTU1010 D1: GPP2

D 2: NLR145

Red color ring indicates parental monomorphism with primer RM124 and yellow color rings indicate parental polymorphism with primer RM19629 and RM19472

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Fig. 2. Foreground analysis for confirmation of F1s using *xa13-prom* **for** *xa13* **gene** *Note: Lane M: 50 bp ladder, Lane p: Recipient MTU1010, Lane D:Donar M-16-59, 1-12 are F1 plants in heterozygous condition*

Fig. 3. Foreground analysis for confirmation of F1s using *PTA248* **for** *Xa21* **gene** *Note: Lane M: 50 bp ladder, Lane p : Recipient MTU1010, Lane D:Donar M-16-59, 1-9 are F1 plants in heterozygous condition*

4. DISCUSSION AND CONCLUSION

4.1 Foreground Selection of BC1F¹ Progeny

"Among 257 BC_1F_1 plants, 6 BC_1F_1 plants showed the presence of all four genes *xa13*, *Xa21, Pi54* and *Pi1* in heterozygous condition. This result is similar to previous reports on the successful utilization of MABC to transfer BB and Blast resistance genes into several elite rice varieties" (Denget al., 2012), (Suh et al*.,* 2013); (Win et al., 2013, Dash et al., 2016, Abhilash et al., 2016). "In the present study, MABC clearly overcome the obstacles when breeding for biotic resistance by conventional breeding method and demonstrated that MABC is generally an effective strategy for genes or QTL pyramiding. Identification of positive heterozygous plants for *xa13*, *Xa21, Pi54 and Pi1* genes in BC1F¹ generation is very difficult and time consuming job, if done, based on phenotype based selection alone. Hence molecular markers used in the present study allowed precise selection of positive plants for four genes" (Ribaut et al., 1998). "MAS is particularly useful for identification of heterozygous individuals for recessive genes like *xa13*. In the absence of

marker, identifying backcross plants that have this type of recessive genes would require progeny testing, which is an addition of one more generation study and cumbersome too" (Sundaram et al., 2008). "Like any other genetic markers, the PCR based DNA markers used in the present study (i.e. *xa13-prom, pTA248, Pi54 MAS* and RM224) are located very near to/within *xa13*, *Xa21, Pi54 and Pi1* genes" (Sundaram et al., 2011, Ronald et al., 1992, Ramkumaret al., 2011). Hence these markers can be used to complement classical breeding techniques in order to select segregating plants at early stage based on the DNA marker genotype rather than waiting to observe the phenotypic disease screening (i.e. rice blast and bacterial blight).

4.2 Background Selection of BC1F¹ Progeny

The percent recurrent parent genome recovery observed in this study was identical to that of Sundaram et al*. 920080* and also with Hasan et al*.* (2015). Background screening with RM polymorphic SSR markers is shown in Figs. 4 and 5. The present BC_1F_1 is equal to BC_4F_1 as three backcrosses were completed earlier. In our study, we could recover 96.8% RPG as per the expectation of 96.875% of RPG in fourth backcross generation. Balachiranjeevi et al*.,* 2015 utilized marker assisted backcross breeding for recovering the plants with three biotic resistance (*xa13, Xa21* and *Pi54*) genes with maximum recurrent parent genome of DRR17B.

4.3 Foreground Selection of BC1F² Population

Foreground selection of the BC_1F_2 population was carried out using PCR based gene specific and gene linked markers for the target genes. A total of 1060 BC1F2 plants were screened for homozygosity of all four target resistance genes. This BC_1F_2 population exhibited donor parent type, heterozygote and recurrent parent type alleles for all four markers.

Identification of homozygous BC_1F_2 plants is very important because if the selected BC₁F₂ plants contain one or more of the target genes in heterozygous condition, they will segregate in next generation. The plants carrying the donor parent alleles were selected. In the present study, only homozygous plants with the desirable gene combinations (i.e.*xa13xa13Xa21Xa21Pi54Pi54Pi1Pi1)* were

selected for further advancement and evaluation.

4.4 Background Selection of BC1F2 Gene Positive Plants

At BC_1F_2 generation (BC_1F_2 is equal to BC_4F_2 as three backcrosses completed earlier), the recovery of RPG was observed to be nearly equivalent to the theoretically expected value of 96.8%. In BC1F2 generation the recurrent parent genome recovery percentage was ranged between 96.0 and 97.8%. Identification of plants carrying more than one target gene with desired recurrent parent genome is extremely difficult through conventional breeding. In our study, we could identify the plants with four and three gene combinations with more than 96 % RPG, which is impossible through conventional breeding which were developed through the incorporation of the blast resistance *Pi-7(t), Pi-d(t)1* and *Pir2-3(t)* genes and qLN2 QTL into the MR263 background using an MABC breeding approach (Ahmed et al., (2016) ly, MR263-BR-3, MR263- BR-4, MR263-BR-13 and MR263-BR-26). They used simple sequence repeat (SSR) markers RM5961 and RM263 (linked to the blast resistance genes and QTL) for foreground

selection and a collection of 65 polymorphic SSR markers for background selection in backcrossed and selfed generations. Background analysis (BC2F⁴ generation) revealed the highest rate of recurrent parent genome recovery of 96.1% in MR263-BR-4-3 and 94.3% in MR263-BR-3-2. In a similar study Tanweer et al., (2015) introgressed blast resistance genes (Putative *Pib* and *Pi-54*) into elite rice cultivar MR219 through Marker-Assisted Selection. For background selection they used a total of 72 polymorphic markers. The minimum recovery of the recurrent parent genome in an improved line was 94% and the maximum recovery in an improved line was 97.5%. The percentage of chromosome segments derived from PongsuSeribu 2 was 2.5% and remained constant in all of the advanced improved lines. The average proportions of the recurrent parent genome in all 15 improved lines were 96.17%, showing the maximum similarity observed at the phenotypic level with the recurrent parent. Similarly,. (Basavaraj et al., 2010, Basavaraj et al., 2009) carried out marker assisted background selection in the 10 best BC_2F_5 families of Pusa6B and PRR78 using 74 STMS markers polymorphic between Pusa6B and Pusal46 and 54 STMS markers polymorphic between PRR78 and Pusa1460. They recovered the recurrent parent genome ranging from 85.14 to 97.30% and 87.04 to 92.81% in the 10 selected BC₂F₅ families of Pusa6B and PRR78 respectively. Rajpurohit et al. (2011) also tested 209 rice SSR markers for background selection out of which a set of 95 markers showed polymorphism between the parents Type 3 Basmati and PR106-P2. Sixteen BC₂F₃ progenies with nearly Type 3 Basmati seeds were finally selected for background profiling using 95 SSR and 12 ISSR markers. On the basis of SSR markers, these lines showed background recovery from 81.57% (41-3-40) to 92 10% (29-1-35). Pyramid line 29-1-35 recovered maximum recurrent parent genome (92.0%) followed by line 31-4-2 with RPG (91.05%) (Divya et al., 2015, Harlan et al., 1922, Hospital et al., 2001, Hospital et al., 1992, Kauffman et al., 1973).

4.5 Foreground Selection of BC1F³ Progenies

Foreground selection was carried out in BC_1F_3 progenies using PCR based SSR markers Figs. 6,7,8,9. Selected BC1F3 progenies were screened for BB resistance along with resistant donor parent, GPP2 and recurrent parent, MTU1010 using a virulent isolate of *Xoo (DX-020)* Fig. 10. Twenty five BC1F3 progenies were screened for Blast resistance at Indian Institute of Rice Research (IIRR), Rajendranagar, Hyderabad. Local isolates of *Magnaportheoryzae* from IIRR, were used to screen the donor,

recurrent parents along withBC₁F₃progenies under in vivo conditions following uniform blast nursery (UBN) method Fig. 11 (Khushet al., 1997, Khush et al., 1990, Lalitha et al., 2013, Manish et al., 2013, Pandey et al., 2012, Singh et al., 2011).

Table 1. Resistance and susceptible allele sizes of target gene specific / linked markers

Gene	Markers	Resistance allele	Susceptible allele	Reference
xa13	xa13-prom	500 _{bp}	250bp	Sundaramet al., (2008)
Xa21	PTA248	900 _{bp}	650 _{bp}	Ronald et al., (1992)]
Pi54	Pi54 MAS	200bp	350bp	Ramkumaret al., (2011)
Pi1	RM224	130bp	150 _{bp}	Hittalmaniet al., 2000

Fig. 4. Background analysis for recurrent parent genome recovery of F1 plants using RM 19629

Fig. 5. Background analysis for recurrent parent genome recovery of F1 plants using RM 22552 Note: 1to 9 equals to $F_1 - 5$, $F_1 - 17$, $F_1 - 19$, $F_1 - 27$, $F_1 - 51$, $F_1 - 63$, $F_1 - 65$, $F_1 - 69$, $F_1 - 77$ plants, respectively *The plants possessing MTU1010 allele were selected*

Representation of graphical genotype of selected BC1F² plants in the genomic region around *xa13* **(on Chromosome 8),** *Xa21, Pi54* **and** *Pi1* **(on Chromosome 11) based on analysis with parental polymorphic SSR markers**

Chromosome – 8 Chromosome – 11

Note: R – Recipient (MTU1010), D- Donor (GPP2- *xa13* **and** *Xa21***) / (NLR145-** *Pi54* **and** *Pi1***), 1: BC1F2-198-317, 2: BC1F2-198-52, 3: BC1F2-198-581, 4: BC1F2-198-620**

Fig. 6. Foreground analysis for confirmation of BC1F3s using *xa13-prom* **for** *xa13* **gene** *Note: Lane M: 50 bp ladder, Lane p: Recipient MTU1010, Lane D:Donar M-16-59, 1-25 are BC¹ F3 plants*

Fig. 7. Foreground analysis for confirmation of BC1F3s using *Pi54 MAS* **for** *Pi54* **gene** *Note: Lane M: 50 bp ladder, Lane p: Recipient MTU1010, Lane D:Donar M-16-59, 1-25 are BC¹ F3 plants*

Fig. 8. Foreground analysis for confirmation of BC1F3s using *PTA248* **for** *Xa21* **gene** *Note: Lane M: 50 bp ladder, Lane p: Recipient MTU1010, Lane D:Donar M-16-59, 1-25 are BC1 F3 plant*

Fig. 9. Foreground analysis for confirmation of BC1F3s using RM224 for *Pi1* **gene** *Note: Lane M: 50 bp ladder, Lane p: Recipient MTU1010, Lane D:Donar M-16-59, 1-25 are BC¹ F3 plants*

Fig. 10. Field level screening of BC1F³ progenies against BB resistance with IIRR isolate *(DX-020)***. Arrows indicates the plant showing**

Fig. 11. Blast nursery screening of BC1F3 progenies at IIRR, Rajendranagar, Hyderabad

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

We hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of this manuscript.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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