

Improvement of bacterial blight resistance of the popular variety, Nellore Mahsuri, NLR34449 through marker-assisted breeding

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Abstract

To combat the dreaded diseases in rice like bacterial blight and blast, host plant resistance has been advocated as a sustainable method. Through the present study, we have successfully incorporated three major bacterial blight (BB) resistance genes *viz.*, *Xa21*, *xa13* and *xa5* into NLR3449, a high yielding, blast resistant, fine-grain type popular rice variety through marker-assisted backcross breeding. Foreground selection was carried out using PCR based, gene-specific markers *viz.*, pTA248 (*Xa21*), xa13prom (*xa13*) and xa5FM (*xa5*) at each generation of backcrossing, while 127 polymorphic SSR markers spanning on 12 chromosomes were used for background selection and backcrossing was limited to two rounds. At BC₂F₁ generation, a single plant (NLR-87-10) with 89.9% recovery and possessing all the three bacterial blight resistance genes was forwarded to BC₂F₂ generation. A solitary BC₂F₂ plant *viz.*, NLR-87-10-106 possessing all the three resistance genes and >90% genome recovery was identified and advanced through selfing till BC₂F₄ generation by adopting pedigree method. Three best lines at BC₂F₄ lines, possessing high level of resistance against bacterial blight and blast and equivalent or superior to NLR 34449 in terms of yield, grain quality and agro-morphological traits have been identified and advanced for multi-location trials.

Introduction

Rice is an important food crop that serves as a major carbohydrate source for nearly half of the world's population (Nguyen and Ferrero 2006; Sundaram et al. 2008 and Pradhan et al. 2015). It is grown throughout the year in a variety of agro-ecosystems in India, which include irrigated, rain-fed, deep-water and hills (Sundaram et al. 2009). We need to produce at least 40% more rice by 2030 to feed a growing population and there are many challenges in meeting the food security requirements for the future (Khush 2005). This target has to be met amidst the scarcity of natural resources and adverse effects from a rapidly changing climate and with less use of chemicals and under constant battle against new emerging pathogens and pests (Pradhan et al. 2015). Rice is affected by many diseases and among them, bacterial blight (BB) caused by *Xanthomonas oryzae* pv. *oryzae* (*Xoo*) (Ishiyama, 1922) is known to cause yield losses ranging from 74%-81% (Srinivasan and Gnanamanickam 2005), when the disease occurs in severe form. BB is particularly severe in the irrigated and rain-fed lowland ecosystems (Sundaram et al. 2008). The disease occurs in the host plant at seedling, vegetative and reproductive stages, but infection at the tillering stage causes severe blighting of leaves resulting in yield loss (Shivalingaiah and Umesha 2011). Chemical control for BB is cost intensive and not effective as successful bactericidal chemicals are not available (Devadath 1989). Thus, the exploitation of disease resistant cultivars is the most effective and eco-friendly approach for reducing yield losses due to BB (Sundaram et al. 2008). At least, 45 BB resistant genes (Neelam et al. 2019; Sundaram et al. 2014; Suk-man Kim et al. 2018) have been identified so far from the primary gene pool of rice and pyramiding two or more resistance genes has been advocated to enhance the durability of resistance (Sundaram et al. 2008).

The rice variety, NLR 34449, popularly known as Nellore Mahsuri, was released from Rice Research Station, Acharya N.G. Ranga Agricultural University, Nellore, Andhra Pradesh, India. It is popular among the rice farmers of South India, as it is high yielding, possesses highly desirable medium slender (MS) grain type, has excellent blast resistance, is of short duration (120 days), possesses non-lodging plant type and is highly amenable for machine harvesting. It is highly preferred rice by the farmers of Andhra Pradesh and other South Indian states (<http://www.kvknellore-angrau.org/index.php>). Despite its high yield and blast resistance, the variety is highly susceptible to bacterial blight disease, which limits its production and adoption significantly. Keeping these points in view, through the present study, we have transferred three major bacterial blight resistance conferring genes, *viz.*, *Xa21*, *xa13* and *xa5* from elite, fine grain, BB resistant variety Improved Samba Mahsuri (ISM) into the genetic background of NLR 34449 through marker-assisted backcross breeding (MABB).

Material And Methods:

Plant material:

NLR 34449 (popularly known as Nellore Mahsuri), a high yielding, medium slender, blast resistant and short duration variety (120-125 days) developed from the cross IR 72/BPT 5204 in the year 2009 was used as the recurrent parent. Improved Samba Mahsuri (ISM), a high yielding, medium-slender grain type, bacterial blight (BB) resistant variety developed by ICAR-Indian Institute of Rice Research (ICAR-IIRR), Hyderabad, India and CSIR-Centre for Cellular and Molecular Biology (CSIR-CCMB), Hyderabad, India, possessing three major bacterial blight (BB) resistance genes, *viz.*, *Xa21*, *xa13* and *xa5* (Sundaram et al. 2008) was used as the donor parent for bacterial blight resistance.

Marker-assisted backcross breeding strategy:

NLR 34449 was crossed with Improved Samba Mahsuri during dry season 2016-17. F₁s developed from the above mentioned cross were confirmed for their heterozygosity using target resistance gene specific markers *viz.*, pTA248 specific for *Xa21* (Ronald et al. 1992), xa13prom, specific for *xa13* (Hajira et al. 2016) and xa5FM specific for *xa5* (Hajira et al. 2016). True F₁s thus identified were then backcrossed with the recurrent parent, NLR 34449 to generate BC₁F₁s. They were subjected for foreground selection using gene-specific markers and positive plants (*i.e.* plants which are heterozygous for all the three target BB resistance genes) were then analyzed for the recurrent parent genome recovery through background selection using a set of 127 SSR parental polymorphic markers evenly distributed across the 12 rice chromosomes (listed in Supplementary Table 1B) to identify those plants with maximum recurrent parent genome recovery (RPG). A single BC₁F₁ plant positive for all three target genes and with maximum recurrent parent genome recovery was selected and backcrossed with NLR 34449 to generate BC₂F₁s. Marker-assisted foreground and background selection was repeated among the BC₂F₁ plants and a single BC₂F₁ plant possessing the target resistance genes and also with maximum recurrent genome recovery were selfed to develop BC₂F₂s. Finally, a single BC₂F₂ plant possessing *Xa21*, *xa13* and *xa5* in homozygous condition along with maximum RPG and closely resembling NLR 34449 (based on morphological features) was identified and advanced through pedigree method of breeding till BC₂F₄ (Figure 1). The extent of donor parent segment introgression among the selected backcross derived plants at BC₂F₄ generation was assessed utilizing the software tool, Graphical Genotype V 2.0 (Van Berloo, 1999) and the BC₂F_{4.5} lines were also further evaluated for bacterial blight resistance and key agro-morphological traits.

Phenotypic screening for BB and blast resistance:

Bacterial blight screening:

Thirty-day old seedlings of the improved breeding lines of NLR 34449 at BC₂F₄ generation were transplanted in the main field of experimental farm of ICAR-IIRR located at Rajendranagar, Hyderabad, India to assess their resistance against bacterial blight during wet season 2019. IX0-20, a local virulent isolate (collected from Hyderabad, Telangana State, India) of *Xanthomonas oryzae* pv. *oryzae* was used for screening of bacterial blight resistance of the breeding lines. Inoculation was done at maximum tillering stage following leaf clip method of Kauffman et al. 1973 by clipping the leaf tip (about 1 to 2cm) of the uppermost leaf with scissors dipped into the inoculum. Symptoms were measured at 15 days after inoculation based on the IRRI SES (IRRI, 2013) for their resistance/susceptibility.

Screening for blast resistance:

During dry season 2019-20, gene pyramided lines possessing *Xa21*, *xa13* and *xa5* at BC₂F₅ generation were analysed for their resistance against blast in uniform blast nursery along with the susceptible (HR12) and resistant (C101A51 and Tetep) checks using a local isolate, SPI-40 of the blast pathogen, *Magnaporthea oryzae* as per the protocol described in Mohan et al. (2011). After 15 days of inoculation, the lines were scored for their resistance/ susceptibility to the disease based on IRRI-SES scale, 1996 (IRRI, 1996).

Screening of the improved lines for agro-morphological characters:

During dry season 2020, thirty-day old seedlings of the recurrent parent NLR 34449 and donor parent ISM along with five best backcross derived lines at BC₂F₅ generation, possessing bacterial blight resistant genes in the genetic background of NLR 34449 were transplanted in the main field of experimental farm of ICAR-IIRR located at Rajendranagar, Hyderabad, India at a spacing of 15 X 20cm in 2 m² plots in three replications and the field was applied with the recommended dose of NPK fertilizers (@ 220:70:80kg/ha). Phenotypic data was collected for the selected plants for key agro-morphological traits, *viz.*, plant height (cm), days to 50% flowering (DFF), panicle length (cm), number of productive tillers per plant, panicle exertion, grain yield per 33 plants (*i.e.* per m²; in g), number of grains per panicle, grain type, L/B ratio, 1000-grain weight (g) among 33 plants for each replication (n = 3) as explained in Abhilash et al. (2016). The data was statistically analysed as per the procedure described in Freeman et al. (1978). Least Significance Difference (LSD) values at 5 to 7 percent level of significance and Coefficient of variation (CV) were calculated using standard errors of mean (S. Em. ±) using Microsoft Excel package. The software package, R studio (R Core, 2016) was used for analysis of variance (ANOVA) to determine significant variation among the improved breeding lines of NLR 34449.

Results:

Marker-assisted introgression of *Xa21*, *xa13* and *xa5* into the genetic background of NLR 34449

Ninety-five F_1 plants, which were derived from the cross NLR 34449 X ISM were confirmed for their hybridity with respect to the three target BB resistance genes *i.e.*, *Xa21*, *xa13* and *xa5* with the help of gene-specific markers *viz.*, pTA248, xa13prom and xa5FM, respectively during wet season 2017. A total of 80 F_1 plants were found to be heterozygous for all the three target BB resistant genes. These plants (*i.e.* *Xa21xa21 xa13xa13 xa5xa5*) were used as pollen donors to develop BC_1F_1 s. Among 175 BC_1F_1 s, 21 plants were found to be heterozygous for all the three target BB resistant genes (*i.e.* *Xa21*, *xa13* and *xa5*). These 21 plants were analyzed to assess the recovery of the recurrent parent genome using a set of 127 parental polymorphic primers (Supplementary table 1A, 1B) and the analysis revealed that a single BC_1F_1 plant (NLR-87) had the highest recurrent parent genome (RPG) recovery of 78.5%. The plant, NLR-87 was then used as a pollen donor to produce BC_2F_1 s. A total of 105 BC_2F_1 plants were raised and evaluated through foreground selection with gene-specific markers and thirteen were found to be positive for all the three target BB resistant genes and they were subjected for background genome recovery analysis with parental polymorphic SSR markers and a single plant (NLR-87-10) with maximum recurrent parent genome re recovery (89.9%) was identified. This plant was selfed to produce BC_2F_2 plants. A total of 425 BC_2F_2 plants, which were generated by selfing the selected BC_2F_1 plant (NLR-87-10), were grown in the field. All of them were subjected to phenotypic screening with a local isolate of *Xoo*, IX0-20 at maximum tillering stage to screen for their resistance against BB. A total of 359 plants were found to be resistant.

When the phenotypically resistant plants were subjected for foreground selection with the help gene-specific markers, a total of five plants were identified to be homozygous for all the three target resistant genes (Figure 2). A solitary plant (NLR-87-10-106) which was homozygous resistant to all the three target resistant genes and possessed with maximum recovery of NLR34449 genome (94.6%) was identified and was advanced further through pedigree method of breeding to BC_2F_4 generation. The details of number of plants screened and selected in each generation of backcrossing were given in Table 1. A set of five promising BC_2F_4 lines, *viz.*, (NLR-87-10-106-40-50, NLR-87-10-106-41-51, NLR-87-10-106-42-52, NLR-87-10-106-43-53, NLR-87-10-106-44-54) identified to be identical to recurrent parent NLR 34449 with respect to morphological traits and these lines were subjected for analysis of their resistance against bacterial blight, blast, yield and agro-morphological traits.

Table 1: Details of number of plants confirmed for foreground and background selection among the backcross population

S. No	Generation	Total no. of plants analysed	Total no. positive plants to all three target genes (<i>Xa21</i> , <i>xa13</i> , <i>xa5</i>)	% of Recurrent Parent Genome (RPG) recovery	Selected plant with maximum RPG%
1	F_1	95	11	-	
2	BC_1F_1	175	20	78.5%	NLR-87
3	BC_2F_1	105	13	89.9%	NLR-87-10
4	BC_2F_2	318	5	94.6%	NLR-87-10-106
5	BC_2F_4			96%	(i) NLR-87-10-106-40-50 (ii) NLR-87-10-106-41-51 (iii) NLR-87-10-106-42-52 (iv) NLR-87-10-106-43-53 (v) NLR-87-10-106-44-54

#The true F_1 s obtained from the cross between Nellore Mahsuri (NLR34449) x Improved Samba Mahsuri were backcrossed with Nellore Mahsuri. The best BC_1F_1 plant was identified through foreground and background selection was backcrossed with Nellore Mahsuri to generate BC_2F_1 s. The best BC_2F_1 plant (identified through foreground and background selection) was selfed to produce

BC₂F₂s. Selected homozygous BC₂F₂ plant with maximum RPG% was selfed and advanced through pedigree method of breeding to BC₂F₄ generation.

The five selected BC₂F₄ lines were further analysed to assess the extent of linkage drag around the target bacterial blight resistant genes *i.e.*, *Xa21*, *xa13* and *xa5*. This analysis revealed that for *Xa21* gene located on Chr. 11L, a segment of 0.3 Mb was introgressed at both proximal and distal ends from the donor parent genome in the best BC₂F₄ plant (NLR-87-10-106-40-50); thus, in total, a segment of 0.6Mb was introgressed from the donor parent with respect to the genomic region in the vicinity of *Xa21* (Figure 3A). With respect to *xa13* gene located on Chr. 8L, a segment of 0.5 Mb was introgressed at proximal end while another segment of 0.8 Mb was introgressed at distal end from the donor parent genome in the best BC₂F₄ plant (NLR-87-10-106-40-50); thus, in total, a segment of 1.3Mb was introgressed from the donor parent with respect to the genomic region in the vicinity of *xa13* (Figure 3B). With respect to *xa5* gene located on Chr. 5S, a segment of 0.2 Mb was introgressed at proximal end, while 0.3 Mb introgressed at distal end from the donor parent genome in the best BC₂F₄ plant (NLR-87-10-106-40-50); thus, in total, a segment of 0.5Mb was introgressed from the donor parent with respect to the genomic region in the vicinity of *xa5* (Figure 3C). All the five selected introgressed BC₂F₄ lines were subjected to screening for bacterial blight and blast resistance and also evaluated for their agro-morphological traits.

Assessment of BB and Blast resistance in the improved lines of NLR 34449:

The donor parent Improved Samba Mahsuri (ISM), RPBio- Patho- 1 and RPBio- Patho- 2 were observed to be highly resistant to the disease with a lesion length ranging from 0.0 to 1.7 ± 0.3 (Score 1). The recurrent parent, NLR 34449 was observed to be highly susceptible to the disease with a lesion length of 8.7 ± 0.0 (Score 9) when screened with IXO-20. All the five selected BC₂F₄ were observed to be resistant to BB disease, showing a lesion length of 0-1 cm, which was similar to the donor parent (*i.e.* ISM) (Fig 4A, Table 2). With respect to blast disease, the susceptible check, HR12 and the parent, donor ISM were highly susceptible to blast, with a score of 9, while the resistant check, Tetep and the recurrent parent, NLR 34449 were found to be resistant to the disease with a score of 1 and 3, respectively. The selected BC₂F₅ lines were observed to show resistant reaction against blast disease with a disease score of 2-4 (Fig 4B, Table 2)

Table 2: Reaction of selected improved lines of NLR 34449 after inoculation with bacterial blight and blast pathogen

Parents and checks	Reaction against BB		Reaction against Blast	
	IXO-20		SP140	
	Score	R/S	Score	R/S
ISM	0.0 ± 0.0	R	9	S
NLR 34449	8.7 ± 0.0	S	3	R
RPBio- Patho 1	1.3 ± 0.3	R	2	R
RPBio- Patho 2	1.7 ± 0.3	R	1	R
TN1	8.7 ± 0.3	S	-	-
HR12	-	-	9	S
NLR-86-10-106-40-50	1	R	2	R
NLR-86-10-106-41-51	1	R	3	R
NLR-86-10-106-42-52	1	R	3	R
NLR-86-10-106-43-53	1	R	4	R
NLR-86-10-106-44-54	1	R	3	R

#R- Resistant; S-Susceptible

Evaluation of Agro-morphological traits and data analysis in the improved lines of NLR 34449

Among the improved lines, BC₂F₄ plant, NLR-87-10-106-44-54 displayed better attributes with respect to most of the agro-morphological traits when compared to the recurrent parent, Nellore mahsuri. All the introgressed lines were slightly taller than the recurrent parent, Nellore Mahsuri and shorter than donor parent, Improved Samba Mahsuri, except a single line NLR-87-10-106-42-52 (75.5 ± 0.9 cm) which was observed to be slightly shorter than recurrent parent (Table 3). The introgressed lines were found to perform equivalent or better than both the recurrent as well as donor parents in terms of panicle length, number of grains per panicle, thousand grain weight and grain yield per plant with good panicle exertion (Figure 5, Table 3). Two improved lines, NLR-87-10-106-40-50 and NLR-87-10-106-42-52 recorded grain yield per plant (24.1 ± 0.6 g) equivalent to the recurrent parent. The analysis of variance for ten agro-morphological traits among the selected BC₂F₄ lines and the recurrent parent, Nellore Mahsuri revealed that the mean sum of squares due to treatments (<0.01) were highly significant for all the traits under study, indicating significant variability among the lines as compared to Nellore Mahsuri (Table 3). For genetic parameters such as genotypic coefficient of variance (GCV) and phenotypic coefficient of variance (PCV), lower values (0-10%) was recorded in days to fifty percent flowering (DFF), plant height (PH), panicle length (PL), number of grains per panicle (NGP), thousand grain weight and L/B ratio and moderate values (10-20%) for productive tillers, grain yield per plant. Values for heritability in broad sense was noticed moderate (>50%) in days to 50% flowering (DFF), plant height, panicle length, number of grains per panicle, L/B ratio and to be high for remaining traits (>60%). For Genetic advance in % of mean (GAM) lower values were (0-10%) observed in days to 50% flowering (DFF), plant height, panicle length, L/B ratio and moderate (10-20%) in number of grains per panicle, thousand seed weight and higher (>20%) with respect to the remaining traits (Table 3).

Table 3: Evaluation of agro-morphological characters in the improved lines along with parents under field conditions

S. No.	Plant identity	Days to 50% flowering (DFF)	Mean plant height (cm)	No. of productive panicles/plant	Panicle length (cm)	Number of grains per panicle	Grain Yield per plant (g)	1000 seed weight (g)	L/B ratio	Grain type	Panicle exertion
1	NLR 34449	95.0 ± 0.3	77.0 ± 0.3	13.7 ± 0.3	22.0 ± 0.6	242.0 ± 1.5	24.0 ± 0.3	14.1 ± 0.6	2.70 ± 0.00	MS	FE
2	RPBio-226	101.0 ± 1.2	84.0 ± 0.6	12.3 ± 0.9	19.2 ± 0.6	225.0 ± 1.2	17.1 ± 1.7	12.5 ± 0.7	2.70 ± 0.00	MS	PE
3	NLR-87-10-106-40-50	97.3 ± 0.3	77.7 ± 0.4	17.0 ± 0.0	22.5 ± 0.6	268.0 ± 1.2	24.1 ± 0.2	14.3 ± 0.2	2.70 ± 0.00	MS	FE
4	NLR-87-10-106-41-51	101.0 ± 0.6	78.2 ± 0.9	16.0 ± 0.6	22.1 ± 0.3	273.0 ± 0.9	24.7 ± 0.4	14.2 ± 0.2	2.70 ± 0.00	MS	FE
5	NLR-87-10-106-42-52	99.0 ± 0.6	75.5 ± 0.9	16.3 ± 0.3	22.5 ± 0.3	272.0 ± 0.9	24.1 ± 0.6	14.5 ± 0.1	2.70 ± 0.00	MS	FE
6	NLR-87-10-106-43-53	99.0 ± 0.3	77.7 ± 0.6	16.7 ± 0.9	23.0 ± 0.4	263.0 ± 1.5	26.8 ± 0.5	15.0 ± 0.1	2.70 ± 0.00	MS	FE
7	NLR-87-10-106-44-54	94.0 ± 0.9	79.0 ± 0.6	17.7 ± 0.3	23.8 ± 0.3	280.0 ± 0.9	28.8 ± 0.3	16.1 ± 0.1	2.70 ± 0.00	MS	FE
	Mean	97	78.04	15.67	22.15	260	24.23	14.37	2.68		
	CV (%)	2.7	3.7	6.5	5.45	5.61	5.50	5.08	2.61		
	F _{cal} value	3.14**	3.5**	10.63**	4.36**	5.56**	21.94**	6.42**	3.02**		
	GCV	2.27	3.36	11.75	5.77	6.91	14.53	6.83	2.14		
	PCV	3.51	4.99	13.46	7.94	8.9	15.53	8.51	3.38		
	h ² b(%)	41.69	45.37	76.25	52.84	60	87.47	64.36	40.25		
	LSD@5%	4.67	5.12	1.83	2.15	26.01	2.37	1.3	0.12		
	GAM@5%	3.02	4.66	21.14	8.64	11.06	27.99	11.28	2.8		

#CV – Coefficient of Variation, PCV – Phenotypic Coefficient of Variance, GCV – Genotypic Coefficient of Variance, h² b - broad sense of Heritability, f (Cal) – F test calculated; LSD – Least Significant Difference; GAM – Genetic advance in % of mean; MS – Medium Slender; FE – Full Exserted, and PE – Partially Exserted.

Discussion:

Rice yield and productivity is drastically reduced due to two major diseases, *viz.*, bacterial blight (BB) and blast (Ou 1985; Srinivas prasad et al. 2009). Development of improved versions of crop varieties through introgression of resistance genes with the help of molecular markers has been well demonstrated with respect to blast disease (Ashkani et al. 2014; Hari et al. 2013; Manu et al. 2012; Madhavi et al. 2013, 2016; Rekha et al. 2018) and BB management (Joseph et al. 2004; Sundaram et al. 2008, 2009, 2011). Till date, at least 45 BB resistance genes (Kim, 2018; Kumari Neelam et al. 2019; Sundaram et al. 2014) have been identified. Among them, the major, dominant, broad spectrum bacterial blight resistance gene, *Xa21* (originally derived from *O. longistaminata*) located on Chr. 11, a major recessive gene located on chromosome 8, *xa13* another major recessive gene located on chromosome 5, *xa5* are known to confer broad spectrum and durable resistance against the disease (Lalitha et al. 2013; Pradhan et al. 2015; Ramalingam et al. 2017; Rekha et al. 2018; Sundaram et al. 2008, 2009).

NLR 34449, popularly known among farmers as Nellore Mahsuri, was released from Rice Research Station, Acharya N.G. Ranga Agricultural University (ANGRAU), Nellore, Andhra Pradesh, India. This variety is popular among the farmers, as it is high yielding,

semi-dwarf in plant stature, blast resistant, early maturing (120 days) and is non-lodging type (suitable for machine harvesting). It is highly preferred by rice farmers in the states of Andhra Pradesh, Telangana and other South Indian states (<http://www.kvknellore-angrau.org/index.php>). Further, it possesses medium-slender grain type, which is highly desirable among the rice consumers in South India and varieties with medium-slender grain type like Samba Mahsuri, Sona Mahsuri etc. are getting increasingly popular across India. As NLR 34449 is highly susceptible to bacterial blight disease, we attempted to improve NLR 34449 for durable bacterial blight resistance by using marker assisted introgression of three bacterial blight resistance genes, *Xa21*, *xa13* and *xa5*.

In breeding programmes, molecular markers that are tightly linked to target genes/QTLs are very useful to overcome the limitations of conventional phenotype based breeding and improve the selection efficiency of target genes or traits in backcross breeding programmes (Jena and Mackill, 2008). This is especially through 'foreground selection' where markers are used alone or in combination with phenotypic screening in selection for the target trait(s) (Hospital and Charcosset, 1997). In the present study, the co-dominant markers, pTA248 (Ronald et al. 1992), *xa13*prom and *xa5*FM (Hajira et al. 2016) were used to screen for the presence of three bacterial blight resistance genes, viz., *Xa21*, *xa13* and *xa5* respectively and all the three markers are reported to be functional markers or very closely linked for the respective genes, with no chance or very little chance for recombination (Hajira et al. 2016). These markers were also used by Ramalingam et al. (2017) and Rekha et al. (2018). The strategy of combining phenotyping with genotyping for BB resistance in the later generation of backcrossing/selfing has been reported earlier by Hari et al. (2011; 2013), wherein a stringent phenotypic screening initially followed by selective genotyping of resistant plants has been reported to save time and resources. This is because; marker-assisted selection was done with only plants which exhibited phenotypic resistance against the targeted stress. Moreover, the segregation pattern with respect to bacterial blight resistance in phenotypic screening in F₂ generation was observed to be 55:9 as the target genes include two recessive genes (viz., *xa13* and *xa5*). These plants were advanced for evaluation of their disease resistance, agro-morphological traits at BC₂F₄ generations

Marker-assisted backcross breeding (MABB) strategy is very helpful in reducing the number of backcrosses needed for complete recovery of the recurrent parent genome (RPG). Even though there are reports which suggest that a minimum of three to four backcrosses are required for near complete recovery of recurrent parent genome (Bai et al. 2006; Hasan et al. 2015a; Sundaram et al. 2008; Sundaram et al. 2009), our recent experience and that of other studies suggest that by adopting a stringent MABB strategy, the number of backcrosses can be limited to just two and still we can maximize the recurrent parent genome recovery (Abhilash et al. 2017; Basavaraj et al. 2010; Singh et al. 2001, Miah et al. 2015; Rekha et al. 2018, Swathi et al. 2019).

Marker-assisted background selection in the early backcross generations has been advocated for quick recovery of the recurrent parent genome (Chen et al. 2001 and Joseph et al. 2004). Polymorphic microsatellite markers are usually utilized for background selection in order to assess the recovery of the recurrent parent genome and also to shorten the number of backcross generations (Hospital and Charcosset, 1997) to estimate the amount of recurrent parent genome contribution. Similar study has been done by (Abhilash kumar et al. 2015; Anila et al. 2014; Balachiranjeevi et al. 2015; Bhaskar et al. 2015; Mahadevaswamy et al. 2018; Rekha et al. 2018).

The donor parent and recurrent parent used in the present study are having the same background (i.e. Samba Mahsuri); hence backcrossing was restricted to two rounds. The recurrent parent genome (RPG) recovery was 78.5% (plant # NLR-87) in the best BC₁F₁ plant, while the value was 88.5% (plant # NLR-87-10) in the best BC₂F₁ plant. In BC₂F₂ generation, the RPG was observed to be 94.6% (line # NLR-87-10-106) and 96% (line # NLR-87-10-106-44-54) in the selected best BC₂F₄ lines. The estimation of linkage drag of donor segment in the best BC₂F₄ lines (plant # NLR-87-10-106-44-54) using graphical genotyping analysis (GGT) revealed that the extent of donor parent genome is restricted to 0.6 Mb with respect to *Xa21* on chromosome 11, 1.3 Mb with respect to *xa13* on chromosome 8 and 0.5 Mb with respect to *xa5* on chromosome 5, highlighting very high recovery of the genome of Nellore Mahsuri genome (up to 94.6%) at BC₂F₄ (NLR-87-10-106). Similar results were observed in earlier studies where *Xa21*, *xa13* and *xa5* genes were introgressed into Samba Mahsuri, Jalmagna, wherein a recovery of 97% and 94.6% respectively was documented (Pradhan et al. 2015; Sundaram et al. 2008, respectively). Few more studies also reported similar observations (Abhilash kumar et al. 2016, 2017; Balachiranjeevi et al. 2015, 2018; Basavaraj et al. 2010; Fahim Ahmed et al. 2015, Hasan et al. 2015b; Rekha et al. 2018; Shanti et al. 2012; Swathi et al. 2019).

Phenotypic screening of the improved versions of the NLR 34449 against bacterial blight disease revealed that all the lines are resistant to the disease with a score of 1. Further, they were also resistant to blast with a score of 2–4 (Fig. 4, Table 2). The results are in agreement with earlier reports, wherein the bacterial blight resistance genes, *Xa21*, *xa13* and *xa5* were deployed either singly or in

combination (Dokku et al. 2013a & 2013b; Gitishree and Rao, 2015; Huang et al. 1997; Sanchez et al. 2000; Singh et al. 2001; Sundaram et al. 2008; Yoshimura et al. 1995) and the derived lines showed resistance against the disease. The objective of the present study was to develop pyramided lines of NLR3449 with durable resistance against BB. In earlier studies, introgressed lines possessing two or more genes against bacterial blight were observed to possess higher level of resistance, possibly due to quantitative complementation (Sanchez et al. 2000; Sundaram et al. 2008). Cultivation of breeding lines of NLR34449 possessing the three resistance genes in bacterial blight endemic areas may be beneficial to farmers and also the resistance is expected to be more durable, as three resistance genes with distinct functions (Sundaram et al. 2009) have been deployed. There have been various reports, wherein a single major blast resistance gene likes *Pi1* or *Pi2* have shown the desired level of resistance against the disease (Fu et al. 2012). Similar to the recurrent parent, NLR 34449, which is highly resistant to blast disease, all the improved lines showed resistance against blast disease. Our preliminary analysis has indicated the possibility of presence of the major blast resistance gene, *Pi1* in NLR34449, based on marker analysis (data not shown) and resistance in the breeding lines could be due to the presence of *Pi1*. The results observed in the present study with respect to resistance against BB and blast are in correspondence with the results obtained in earlier reports (Abhilash et al. 2016; Balachiranjeevi et al. 2015; Hari et al. 2013; Rekha et al. 2018), wherein no negative interaction was observed between blast and BB resistance genes. The improved breeding lines with BB resistance using NLR 34449 as recurrent parent are expected to show high level of resistance against both the diseases. These two diseases are known to be widely prevalent and limiting the rice production in the Indian state of Andhra Pradesh (also known as Rice Bowl of South India) and also other Indian States (Aruna kumari et al. 2016).

Most of the improved lines were observed to be equivalent to NLR 34449 in terms of grain yield and other agromorphological characters and some of them were found to be superior to NLR 34449 with respect to yield (Table 3) and all the lines were showing good panicle exertion with plant height equivalent to NLR 34449 and hence are non-lodging (Fig. 5). Further all the selected improved lines, possessing bacterial blight and blast resistance were observed to be retaining the medium slender grain type and fine-grain quality attributes of the NLR 34449 (Fig. 5). It can be expected that the bacterial blight and blast resistant lines of NLR 34449 developed through present study may useful for the farmers to get the same price in the market, similar to NLR 34449 as they are identical in terms of grain quality and yield, plus added advantage of resistance against two major diseases. One of the significant achievement of this study is the complete recovery of the yield and yield related traits along with grain quality features in the improved lines of NLR 34449 possessing bacterial blight and blast resistance. Combining phenotype-based selection with MAS in the present study was helpful in not only recovering good features of the recurrent parent, but also helpful in selection of superior lines (with better yield and panicle exertion) as compared to the recurrent parent, NLR34449.

Conclusion:

The improved versions of Nellore mahsuri possessing BB and blast resistance, developed in the present study may offer a distinct advantage to farmers of NLR34449, whose fields are affected by both bacterial blight and blast. Further, Cultivation of such improved backcross derived lines of NLR34449 possessing resistance against bacterial blight and blast could help to improve rice production in the disease endemic areas in many states of India, wherein fine-grain type varieties like NLR34449, Samba Mahsuri, HMT Sona etc. are preferred. Further, the improved lines of NLR 34449 developed in this study can also be used as donors to transfer BB and blast resistance into other genetic backgrounds as they possess high yield and medium-slender grain type. Among the improved lines of NLR 34449, the backcross derived line # NLR-87-10-106-44-54 possessing *Xa21*, *xa13* and *xa5* showed very good phenotype and better panicle exertion along with high yield and diseases resistance and hence it has been identified as one of promising lines (Table 3). This line is being nominated for AICRIP trials and state trials for possible commercial release to farmers.

Abbreviations

NLR 34449: Nellore Mahsuri; ISM: Improved Samba Mahsuri; BB: Bacterial Blight; MABB: Marker-Assisted Backcross Breeding; ICAR-IIRR: Indian Council of Agriculture Research- Indian Institute of Rice Research; CSIR-CCMB: Council of Scientific & Industrial Research-Centre for Cellular & Molecular Biology; ANGRAU: Acharya N.G. Ranga Agricultural University; MS: Medium Slender; RPG: Recurrent Parent Genome; GCV: Genotypic Coefficient of Variance; PCV: Phenotypic Coefficient of Variance ; DFF: Days to Fifty percent Flowering; PH: Plant Height; PL: Panicle Length; NGP: Number of Grains per Panicle; S. Em.: standard errors of mean; CV: Coefficient of Variance; GAM: Genetic advance in % of mean

Declarations

Ethical approval and consent to participate:

Not applicable

Consent for Publication:

Not applicable

Availability of data and materials:

The data and information generated from the study are available in our laboratory as hard and soft copies and can be shared based on request.

Competing interests:

The authors declare that they have no competing interests.

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Contribution of authors:

AD carried out the present work and developed the improved versions of NLR 34449 with BB & blast resistance and drafted the manuscript; PV, BSM, LGS, SLV, SRM provided the infrastructural facilities for developing the improved versions of NLR 34449, RMS conceptualized and supervised the experiments and critically reviewed the manuscript; RG constantly helped in developing the improved versions of NLR 34449; DT, PE, KM, SRK, SP, HG, KRR, DMA, HSK, LB, AM, PK, MBM LGS, PMS helped with crossings, field selections and data collection for three consecutive seasons. All authors read and approved the final manuscript.

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Figures

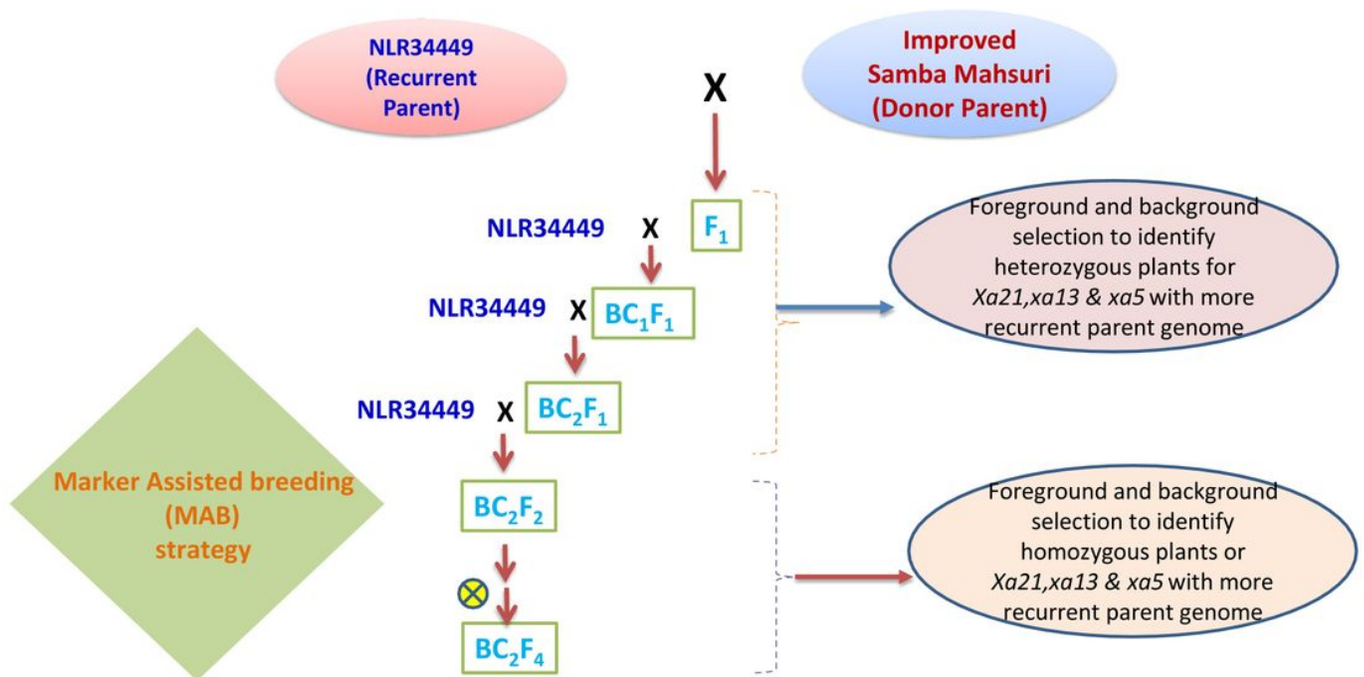


Figure 1

Outline of the crossing scheme. Nellore Mahsuri (NLR 34449) was crossed with Improved Samba Mahsuri (ISM) possessing bacterial blight resistant genes Xa21, xa13, xa5 in order to build bacterial blight resistance through step-wise, marker assisted backcross breeding strategy

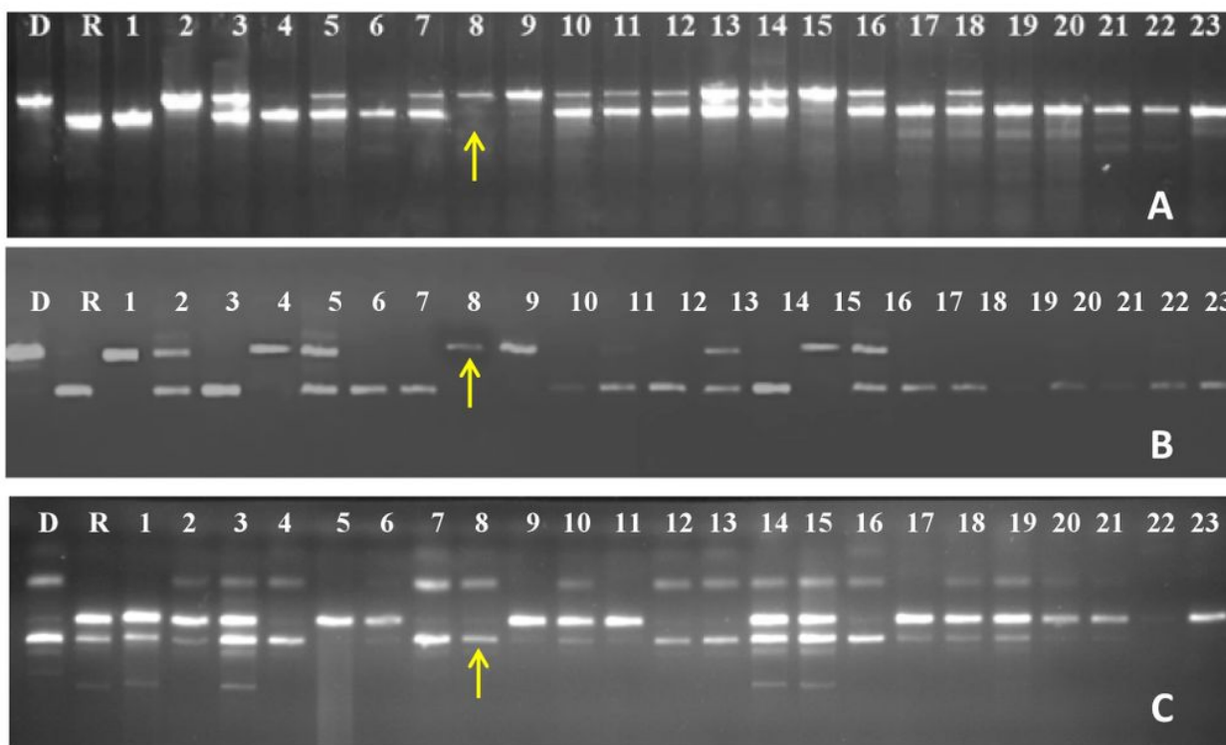


Figure 2

Identification of homozygous BC2F2 plants possessing Xa21+xa13+xa5. (A) Analysis of Xa21 gene using pTA248 marker (B) Analysis of xa13 gene using xa13prom marker (C) Analysis of xa5 gene using xa5FM marker.; R-Recurrent parent NLR34449; D- Donor parent ISM; 1-25 – test samples. Arrows indicate a positive plant possessing all the three genes in homozygous condition.

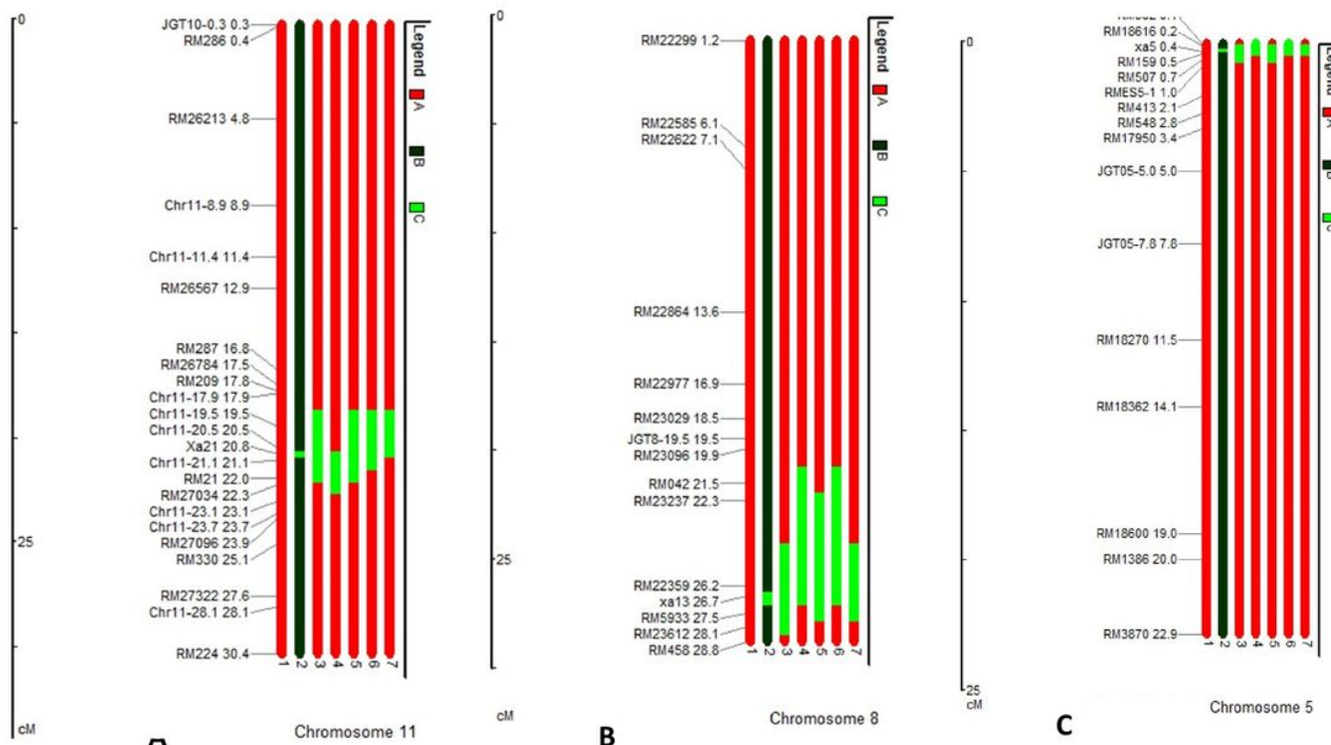


Figure 3

Analysis of donor genome introgression in the chromosomal region around the BB resistance genes, Xa21, xa13 and xa5 in the selected BC2F4 plants through graphical genotype analysis. A: Extent of background genome recovery in the genomic regions in the vicinity of Xa21 on Chromosome 11, and B: Extent of background genome recovery in the genomic regions in the vicinity of xa13 on Chromosome 8, and C: Extent of background genome recovery in the genomic region in the vicinity of xa5 on Chromosome 5

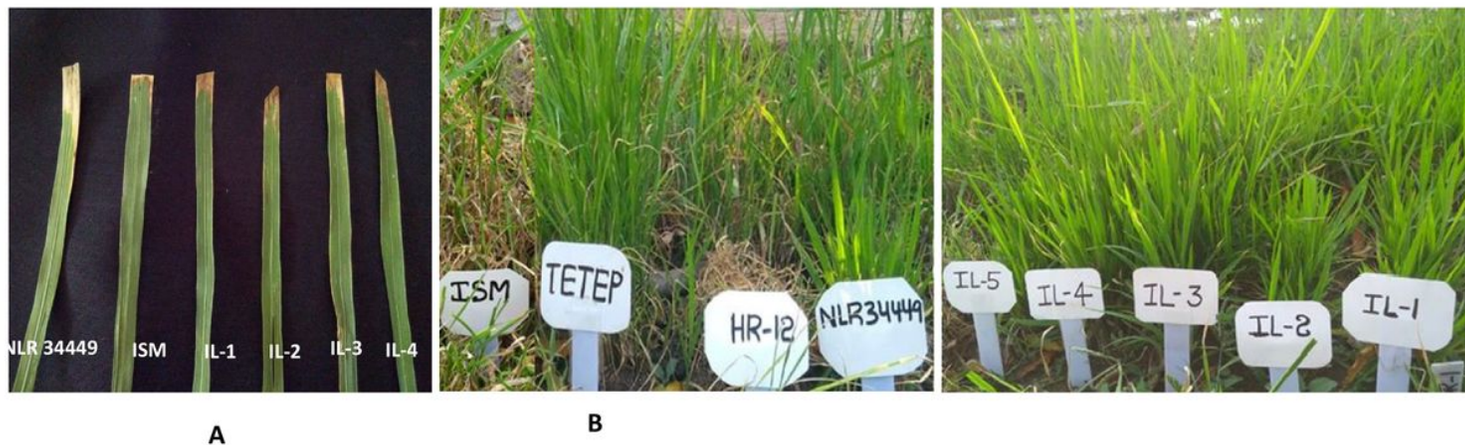


Figure 4

Phenotypic screening of NLR 34449 derived backcross lines for their resistance against bacterial blight and blast; A: Screening of the improved lines against bacterial blight disease. NLR 34449 (Susceptible check), ISM (Resistant check); IL-1 to IL-4- selected BC2F4 lines of NLR 34449. B: Screening against blast disease in Uniform Blast Nursery. Tetep, NLR 34449 (Resistant checks); ISM, HR 12 (Susceptible checks), IL-1 to IL-5: Improved lines of NLR 34449 at BC2F5 generation.

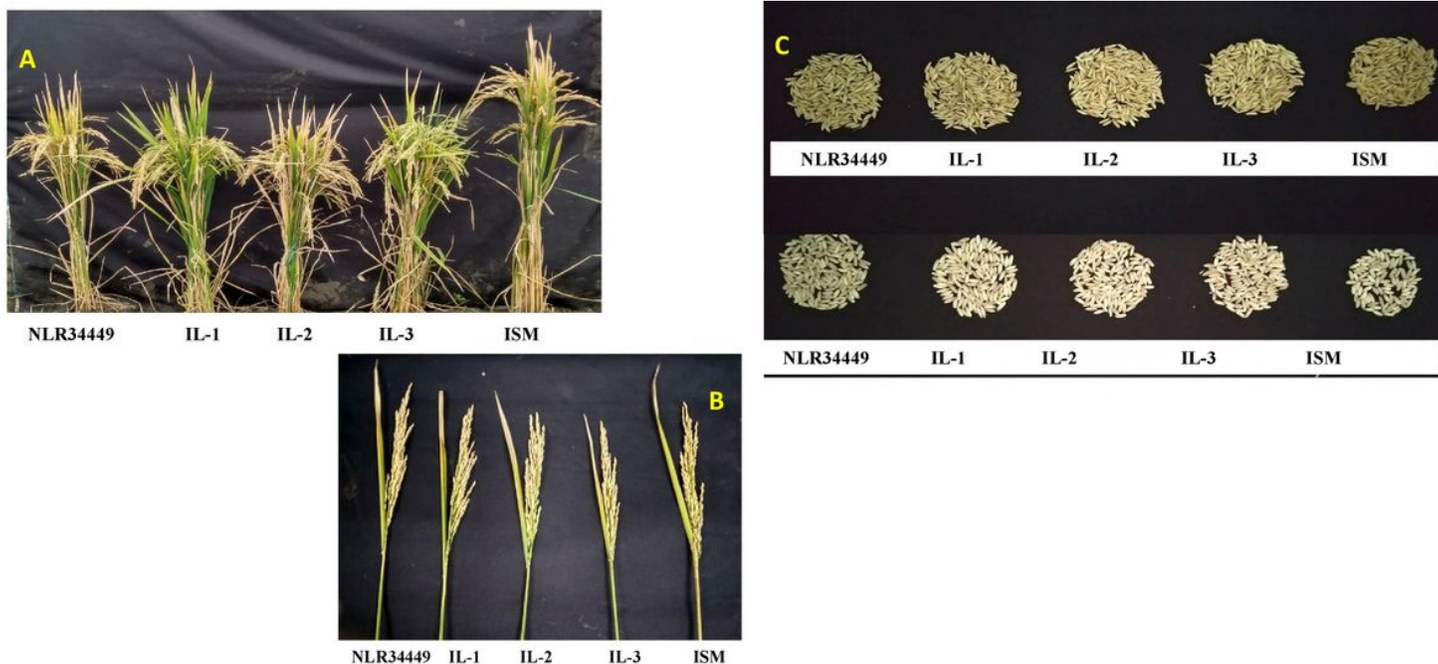


Figure 5

Improved BB and blast resistant lines of NLR 34449. (A) The best Bacterial blight (BB) and Blast Resistant Introgression Lines (ILs) of NLR 34449 with high yield (B) The best Bacterial blight (BB) and Blast resistant panicles with high grain number (C) Improved Bacterial blight (BB) and blast lines grain number and grain quality

Supplementary Files

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- [Supplemetarytables.docx](#)