

Morpho-Molecular Characterization of Rice Genotypes for Resistance to Bacterial Leaf Blight (BLB)

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ABSTRACT

Bacterial leaf blight (BLB) is one of the most devastating diseases of rice (*Oryza sativa L.*), causing substantial yield losses and posing a serious threat to food and livelihood security across rice-dependent regions of Asia and Africa. In this study, 71 rice genotypes developed through crosses among elite and improved lines, were evaluated for bacterial leaf blight (BLB) resistance using artificial clip inoculation at maximum tillering stage, with resistant (Improved Samba Mahsuri) and susceptible (Taichung Native-1, Krishnaveni) checks, under field conditions at Bapatla and Maruteru, Andhra Pradesh, India. Phenotypic screening identified nine genotypes exhibiting disease reaction towards resistance (disease scores 1–3) at both sites. Molecular screening for five BLB resistance (R) genes, *Xa21*, *xa13*, *xa5*, *Xa4*, and *Xa2*, revealed BPT-3170 carried four R genes (*xa13+xa5+Xa4+Xa2*), while eight genotypes had two genes, and 30 genotypes carried one gene. Phylogenetic analysis using 14 R gene-linked markers grouped the genotypes into three major clusters. BPT-3170 exhibited phenotypic resistance along with multiple R genes, indicating its potential to confer broad spectrum resistance and can serve as a valuable donor in BLB resistance breeding. The study also revealed the breakdown of single-gene resistance and low frequencies of *xa5*, *xa13*, and *Xa21*. These findings highlight the importance of pyramiding multiple R genes to achieve durable resistance against BLB.

Keywords: Bacterial leaf blight, Microsatellite markers, R genes, Genetic diversity.

INTRODUCTION

Rice (*Oryza sativa L.*) is a vital staple crop for billions of people worldwide and serves as a cornerstone of global food security and nutrition. Its productivity is challenged by as many as

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27 60 known rice diseases (Ou, 1972), with bacterial leaf blight (BLB) caused by *Xanthomonas*
28 *oryzae* pv. *oryzae* (*Xoo*), being one of the most destructive. BLB leads to yield losses ranging
29 from 20% to 50% (Singh *et al.*, 2011), with severity depending on the rice cultivar,
30 environmental conditions, and management practices. *Xoo* causes wilt, yellowing, and death
31 of rice plants. This disease has become a major concern, particularly in Asian countries, due to
32 the congenial climatic conditions that contribute to frequent epidemics and the lack of effective
33 control measures, making the use of resistant cultivars as the only reliable management
34 strategy. However, the durability of resistance is often compromised due to the rapid evolution
35 of *Xoo* races under selection pressure, necessitating continuous efforts to explore and identify
36 new resistant resources (Xia *et al.*, 2012).

37 The *Oryza* base repository currently lists 44 resistance genes (*Xa1* to *Xa44*) conferring
38 varying levels of resistance to diverse *Xoo* strains, highlighting the genetic complexity of the
39 pathogen-host interaction. In this study, the genotypes are screened for the resistance genes
40 *Xa21*, *xa13*, *xa5*, *Xa4*, and *Xa2*, exhibiting complementary resistance mechanisms. Gene *Xa21*,
41 identified in *Oryza longistaminata*, provides broad-spectrum resistance to diverse *Xoo* strains
42 worldwide. *xa13*, found in the variety BJ1, confers race specific resistance. *xa5* imparts
43 resistance to *Xoo* isolates from India and Nepal at all growth stages (Adhikari *et al.*, 1995). *Xa4*
44 is known for its durable resistance and is widely employed in commercial breeding programs.
45 (Ma *et al.*, 1999 and Sun *et al.*, 2003). *Xa2*, identified in cultivars *Tetep* and *Rantai Emas II*,
46 offers strong resistance across diverse backgrounds (Sakaguchi, 1967).

47 We screened 70 diverse rice genotypes for resistance to BLB using a combination of
48 phenotypic screening through artificial inoculation and molecular marker analysis. By
49 integrating these complementary approaches, this study aims to identify robust resistant
50 genotypes that can serve as valuable donors in breeding programs, ultimately contributing to
51 the development of durable BLB-resistant rice varieties for sustainable rice production.

52 MATERIALS AND METHODS

54 Plant Material

55 The experimental material comprised of 74 genotypes with Improved Samba Mahsuri (ISM)
56 as BLB resistant check, Taichung Native-1 and Krishnaveni as BLB susceptible checks and,
57 Samba Mahsuri as yield check. These genotypes are developed and provided by ARS, Bapatla,
58 Andhra Pradesh, India. Their pedigree is presented in Table 1.

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Table 1. List and Pedigree of the Genotypes used in the study.

| S. NO | GENOTYPE | PEDIGREE | S. NO | GENOTYPE | PEDIGREE |
|-------|----------|------------------------------|-------|--|---|
| 1. | BPT-1235 | SARBARMATI/W-12708 | 38. | BPT-3129 | BPT-5204/MTU-1075 |
| 2. | BPT-2231 | BPT-4358/IR-64 | 39. | BPT-3130 | BPT-5204/MTU-1075 |
| 3. | BPT-2295 | BPT-1768/NLR-33641 | 40. | BPT-3133 | BPT-5204/MTU-1001 |
| 4. | BPT-2411 | BPT-5204/BPT-4358 | 41. | BPT-3135 | BPT-5204/MTU-1001 |
| 5. | BPT-2595 | MUTANT OF BPT-2270 | 42. | BPT-3136 | RP-BIO-226/IRGC-48493 |
| 6. | BPT-2620 | MTU-1061/N-22 | 43. | BPT-3137 | RP-BIO-226/IRGC-48493 |
| 7. | BPT-2677 | MTU-2077/AJAY/MTU-2077 | 44. | BPT-3145 | RP-BIO-226/IRGC-48493 |
| 8. | BPT-2764 | MUTANT OF BPT-2270 | 45. | BPT-3147 | B-95-1/RPHR-1005/B-95-1 |
| 9. | BPT-2766 | BPT-2270/NLR-145 | 46. | BPT-3148 | RP-BIO-226/IRGC-23385/NIDHI/MTU-1081 |
| 10. | BPT-2776 | BPT-2231/NLR-145 | 47. | BPT-3150 | RP-BIO-226/JARAVA |
| 11. | BPT-2782 | NLR-145/MTU-2077 | 48. | BPT-3151 | RP-BIO-226/JARAVA |
| 12. | BPT-2808 | BPT-2231/NLR-145 | 49. | BPT-3159 | CULTURE-0910023/JP-BIO-226/CULTURE-0910023-8/BPT-5204/TETEP |
| 13. | BPT-2824 | MTU-7029/NLR-34449 | 50. | BPT-3164 | B-95-1/RPHR-1005/B-95-1 |
| 14. | BPT-2846 | MTU-1061/IR-78585-64-2-4-3 | 51. | BPT-3168 | MTU-7029/IRGC-18195/MTU-1081 |
| 15. | BPT-2848 | SWARNA/IRGC-18195/MTU-1081 | 52. | BPT-3170 | RP-BIO-226/JARAVA |
| 16. | BPT-2849 | NLR-34449/MTU-5249 | 53. | BPT-3172 | RP-BIO-226/IRGC-48493 |
| 17. | BPT-2854 | MTU-1061/IR-78585-64-2-4-3 | 54. | BPT-3178 | CULTURE-01120305/CULTURE-0910025-7 |
| 18. | BPT-2863 | MTU-2077/NLR-34449 | 55. | BPT-3208 | NLR-34449/ANNADA/NLR-34449 |
| 19. | BPT-2950 | NLR-34449/BM-71 | 56. | BPT-3244 | BPT-5204/RAMAPPA |
| 20. | BPT-2954 | NLR-34449/ANNADA/NLR-34449 | 57. | BPT-3260 | MTU-7029/IRGC-18195/MTU-1081 |
| 21. | BPT-2958 | BPT-5204/IR-50 | 58. | BPT-3261 | MTU-7029/IRGC-18195/MTU-1081 |
| 22. | BPT-3032 | BPT-5204/IR-50 | 59. | BPT-3262 | MTU-7029/IRGC-18195/MTU-1081 |
| 23. | BPT-3033 | BPT-5204/MTU-1075 | 60. | BPT-3263 | MTU-7029/IRGC-18195/MTU-1081 |
| 24. | BPT-3061 | BPT-1768/NLR-145 | 61. | BPT-3264 | CULTURE-01120305/CULTURE-0910025-7 |
| 25. | BPT-3068 | NLR-34449/RAMAPPA | 62. | BPT-3269 | RP-BIO-226/IRGC-23385/NIDHI/MTU-1081 |
| 26. | BPT-3074 | BPT-5204/MTU-1075 | 63. | BPT-3270 | RP-BIO-226/IRGC-23385/NIDHI/MTU-1081 |
| 27. | BPT-3081 | BPT-5204/MTU-1075 | 64. | BPT-3274 | BPT-5204/BPT-2605 |
| 28. | BPT-3086 | BPT-2270/IR-64/MTU-1081 | 65. | BPT-3275 | CULTURE-01120305/CULTURE-0910025-7 |
| 29. | BPT-3092 | NLR-34449/ANNADA/NLR-34449 | 66. | BPT-3276 | CULTURE-01120305/CULTURE-0910025-7 |
| 30. | BPT-3095 | MTU-5249/IR-50 | 67. | BPT-3277 | BPT-5204/O.-LONGISTAMINATA/B-95-1/SWARNA-SUB1 |
| 31. | BPT-3111 | MTU-7029/IRGC-18195/MTU-1081 | 68. | BPT-3279 | RP-BIO-226/JARAVA |
| 32. | BPT-3113 | BPT-2270/NLR-145 | 69. | BPT-3291 | SONA/MAHSURI |
| 33. | BPT-3114 | BPT-2270/NLR-145 | 70. | BPT-4358 | SONA-MAHSURI/ARC-6650 |
| 34. | BPT-3115 | BPT-2270/NLR-145 | 71. | SAMBA MAHSURI (BPT-5204) | GEB-24/TN1/MAHSURI |
| 35. | BPT-3118 | JGL-3855/RAMAPPA | 72. | IMPROVED SAMBA MAHSURI (ISM) RP BIO-226 | MAS FROM BPT-5204 AND SS1113 |
| 36. | BPT-3120 | JGL-3855/ANNADA | 73. | KRISHNAVENI (MTU-2077) | SOWBHAGYA/ARC-5984 |
| 37. | BPT-3121 | BPT-3291/RAMAPPA | 74. | TAICHUNG NATIVE-1 (TN1) | DWARF CHOW WU GEN/TSAI YUAN CHUNJ |

62 Pathogen Inoculation and Screening of Germplasm

Screening for BLB was carried out in wet season of 2020, at two locations in Andhra Pradesh, India, namely Agricultural College Farm, Bapatla ($15^{\circ} 54' 29.88''$ N; $80^{\circ} 28' 7.3092''$ E) and RARS, Maruteru ($15^{\circ} 59' 12.984''$ N; $80^{\circ} 6' 7.848''$ E). The rice genotypes were grown under irrigated conditions in a block with each entry planted in two rows of 2 m length, adopting a spacing of 20 x 15 cm. To ensure strong disease pressure, the block was flanked by two border rows of the susceptible check Taichung Native-1. Inoculum was prepared from BLB-infected

leaves collected from local fields. The leaves were cut into 1 cm pieces, and surface sterilized with 1% sodium hypochlorite. Smaller leaf bits of 5 x 5 mm size were placed in test tube having sterile distilled water for 15 to 20 min, to allow the bacteria to ooze out of the leaf tissue. A loopful of bacterial suspension was streaked onto nutrient agar (NA) media plates and incubated at $27 \pm 1^{\circ}\text{C}$ for 3 days. Single yellow, round and smooth margin, non-flat, mucous colonies were picked and purified on fresh NA plates. For field inoculation, the two days old *Xoo* cultures were suspended in sterile distilled water and adjusted to $\sim 10^8 \text{ CFU ml}^{-1}$ (OD at 600 = 0.5) using spectrophotometer. Plants were inoculated at 75 DAS by leaf clipping method (Kauffman *et al.*, 1973). Disease severity was recorded three weeks post-inoculation and scored based on IRRI SES, 2013 (Table 2).

$$\text{Per cent diseased leaf area} = \frac{\text{Total lesion length}}{\text{Total leaf area}} \times 100$$

Table 2. Disease scoring scale for bacterial leaf blight in rice as per SES, IRRI, (2013).

| Scale | Diseased Leaf Area (%) | Description |
|-------|------------------------|-----------------------------|
| 1 | 1-5 | Resistant (R) |
| 3 | 6-12 | Moderately Resistant (MR) |
| 5 | 13-25 | Moderately Susceptible (MS) |
| 7 | 26-50 | Susceptible (S) |
| 9 | 51-100 | Highly susceptible (HS) |

Identification of R Genes

DNA was isolated from young leaves of all the genotypes using the modified Cetyl Tri Methyl Ammonium Bromide (CTAB) protocol adapted from Dellaporta *et al.* (1983). A total of 16 molecular markers, previously reported, were used to screen R genes (Table 3). PCR amplification was carried out in a 14 μl reaction mixture, consisting of 3 μl of DNA (50 ng/ μl), 1.50 μl of 10X Taq buffer, 0.35 μl of dNTPs (2.5 mM), 0.75 μl of each forward and reverse primers (10 pmol), 7.40 μl of double-distilled water, and 0.25 μl of Taq polymerase (5 U/ μl). The thermal cycler was set with an initial denaturation at 94°C for 5 minutes, followed by 35 cycles of denaturation at 94°C for 40 seconds, annealing at 55°C for 40 seconds, and extension at 72°C for 1 minute, and final extension at 72°C for 10 minutes. PCR products were separated on 3% agarose gel, stained with ethidium bromide by gel electrophoresis. Distinct and unambiguous polymorphic bands were scored against a standard 100-bp DNA ladder.

The allelic data was subjected to estimation of genetic distances among the genotypes using DARwin v6.0 (Perrier and Jacquemond, 2006). For each SSR marker genetic diversity parameters, including the total number of alleles (Na), the effective number of alleles (Ne) were calculated using POPGENE version 1.32. (Yeh *et al.*, 2000) and polymorphic information content (PIC) was analysed using the following formula

$$\text{PIC} = 1 - \sum_{i=1}^n P_{ij}^2$$

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Table 3. Details of the markers linked to BLB resistance.

| S. No | Locus name | Linked gene | Primer | Chromosome number | Reference |
|-------|------------|---------------|--|-------------------|-------------------------------|
| 1. | RM349 | Xa2 | F: TTGCCATTGCGTGGAGGCG R: GTCCATCATCCCTATGGTCG | 4 | Patel <i>et al.</i> (2014) |
| 2. | RM317 | Xa2 | F: CATACTTACCAAGTTCACCGCC R: CTGGAGAGTGTCAAGCTAGTTGA | 4 | He <i>et al.</i> (2006) |
| 3. | RM144 | Xa4 | F: TGCCCTGGCGCAAATTGATCC R: GCTAGAGGAGATCAGATGGTAGTGCATG | 11 | Patel <i>et al.</i> (2014) |
| 4. | RM224 | Xa4 | F: ATCGATCGATCTTCACGAGG R: TGCTATAAAAGGCATTCGGG | 11 | Chen <i>et al.</i> (1997) |
| 5. | RM39 | xa5 | F: GCCTCTCTCGTCTCCTTCCT R: AATTCAAACATGCGGTGGC | 5 | Subudhi <i>et al.</i> (2006) |
| 6. | RM13 | xa5 | F: TCCAACATGGCAAGAGACAG R: GGTGGCATTGCAATTCCAG | 5 | Panaud <i>et al.</i> (1996) |
| 7. | RM164 | xa5 | F: TCTTGCCCGTCACTGCAGATATCC R: GCAGCCCTAATGCTACAATTCTTC | 5 | Amgai <i>et al.</i> (2015) |
| 8. | RM533 | xa8 | F: AAAGGCCGTACCTTGCCTTCC R: AGCTAGGGATCCATCCTCCAACC | 7 | Patel <i>et al.</i> (2014) |
| 9. | RM254 | Xa10 | F: AGCCCCGAATAATCCACCT R: CTGGAGGAGCATTTGGTAGC | 11 | Chen <i>et al.</i> (1997) |
| 10. | RM5509 | Xa33 | F: GATGATCCATGCTTGGCC R: TTCCAGCAGAAAGAACAGC | 6 | Korinsak <i>et al.</i> (2009) |
| 11. | RM30 | Xa33 | F: TGGGGTGGTTAGGCATCGTC R: CCTCACACACGACACGAGC | 6 | Korinsak <i>et al.</i> (2009) |
| 12. | RM206 | Xa10/Xa4 | F: CCCATGCGTTAACTATTCT R: CGTTCCATCGATCCGTATGG | 11 | Panaud <i>et al.</i> (1996) |
| 13. | RM167 | Xa4/Xa10/Xa21 | F: GATCCAGCGTGAGGAACACGT R: AGTCCGACCACAAGGTGCGTTGTC | 11 | Wu and Tanksley 1993 |
| 14. | Xa5FM | xa5 | SF: GTCTGGAATTGCTCGCGTTCG SR: TGGTAAAGTAGATACTTATCAAACCTGGA RF: AGCTCGCCATTCAAGTTCTGAG RR: TGACTTGGTTCTCAAGGCTT | - | Hajira <i>et al.</i> (2016) |
| 15. | Xa13prom | xa13 | F: GGCCATGGCTCAGTGTATT R: GAGCTCCAGCTCTCCAAATG | 8 | Hajira <i>et al.</i> (2016) |
| 16. | pTA248 | Xa21 | F: AGACCGGAAGGGTGGTCCCGGA R: AGACCGGTAATCGAAGATGAAA | 11 | Hajira <i>et al.</i> (2016) |

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104 RESULTS

105 Morphological Screening for BLB

106 The genotype's response to BLB artificial disease screening conducted at Bapatla and
107 Maruteru are presented in Table 4. Comparative analysis revealed that, out of 74 lines
108 (including checks) 37 performed similarly at both locations. For the remaining genotypes, the
109 disease scores at Maruteru exceeded those at Bapatla. Specifically, at Bapatla 7 genotypes
110 showed resistance, 12 moderately resistance, 45 moderately susceptible and 7 susceptible. At
111 Maruteru 9 genotypes showed moderately resistance, 32 moderately susceptible and 28
112 susceptible and 2 highly susceptible. Notably, 9 genotypes consistently exhibited their disease
113 reaction towards resistance (with a score of 1-3) at both the locations, while 52 genotypes
114 showed susceptibility (with score 5-9).

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117 **Table 4.** Phenotypic and genotypic screening for bacterial leaf blight resistance in 74 rice
 118 genotypes.

| S. No. | Genotypes | Disease reaction at | | | | | Xa21 | xa13 | xa5 | Xa4 | Xa2 | Total R genes expressed | Positive compatibility |
|-----------|-----------|---------------------|----------|----|----|----|------|------|-----|-----|-----|-------------------------------|---------------------------|
| | | Bapatla | Maruteru | | | | | | | | | | |
| 1. | BPT-1235 | S | S | -- | -- | -- | ++ | -- | -- | -- | -- | 1 | X |
| 2. | BPT-2231 | MS | S | -- | -- | -- | ++ | -- | -- | -- | -- | 1 | X |
| 3. | BPT-2295 | S | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 4. | BPT-2411 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 5. | BPT-2595 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 6. | BPT-2620 | MS | MS | -- | -- | -- | -- | -- | -- | ++ | -- | 1 | X |
| 7. | BPT-2677 | MS | MS | 0 | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 8. | BPT-2764 | S | S | -- | -- | -- | -- | -- | -- | -- | ++ | 1 | X |
| 9. | BPT-2766 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 10. | BPT-2776 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 11. | BPT-2782 | S | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 12. | BPT-2808 | MS | MS | -- | -- | -- | -- | -- | -- | -- | ++ | 1 | X |
| 13. | BPT-2824 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 14. | BPT-2846 | MR | MR | 0 | -- | -- | -- | -- | -- | -- | -- | 0 | X |
| 15. | BPT-2848 | R | MS | -- | -- | + | ++ | ++ | ++ | ++ | ++ | 2 | X |
| 16. | BPT-2849 | MS | S | -- | -- | -- | ++ | ++ | ++ | -- | -- | 1 | X |
| 17. | BPT-2854 | MS | HS | -- | -- | -- | ++ | ++ | ++ | -- | -- | 1 | X |
| 18. | BPT-2863 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 19. | BPT-2950 | MS | S | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 20. | BPT-2954 | MS | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 21. | BPT-2958 | MR | MR | -- | 0 | 0 | ++ | ++ | ++ | ++ | ++ | 1 | ✓ |
| 22. | BPT-3032 | MR | MS | -- | 0 | -- | -- | ++ | ++ | ++ | ++ | 1 | X |
| 23. | BPT-3033 | MS | S | 0 | 0 | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 24. | BPT-3061 | MR | MS | 0 | + | + | + | + | + | + | + | 0 | X |
| 25. | BPT-3068 | R | MR | + | + | + | + | + | + | + | ++ | 2 | ✓ |
| 26. | BPT-3074 | R | MR | 0 | -- | -- | -- | -- | -- | ++ | ++ | 1 | ✓ |
| 27. | BPT-3081 | MS | MS | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 28. | BPT-3086 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 29. | BPT-3092 | R | MR | -- | -- | -- | -- | -- | -- | ++ | ++ | 1 | ✓ |
| 30. | BPT-3095 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 31. | BPT-3111 | MS | MS | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 32. | BPT-3113 | MS | HS | 0 | + | + | + | + | + | + | + | 1 | X |
| 33. | BPT-3114 | MS | S | -- | -- | 0 | 0 | 0 | 0 | 0 | 0 | 1 | X |
| 34. | BPT-3115 | MR | S | -- | -- | -- | -- | -- | -- | -- | -- | 0 | X |
| 35. | BPT-3118 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 36. | BPT-3120 | MS | MS | 0 | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 37. | BPT-3121 | MS | S | -- | -- | -- | -- | -- | -- | ++ | ++ | 2 | X |
| 38. | BPT-3129 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 39. | BPT-3130 | MS | MS | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 40. | BPT-3133 | MS | MS | 0 | 0 | -- | -- | -- | ++ | ++ | ++ | 2 | X |
| 41. | BPT-3135 | R | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | X |
| 42. | BPT-3136 | MR | MS | -- | ++ | -- | -- | -- | -- | -- | ++ | 2 | X |
| 43. | BPT-3137 | R | MR | -- | -- | -- | -- | -- | ++ | ++ | ++ | 2 | ✓ |
| 44. | BPT-3145 | MS | S | -- | -- | -- | ++ | ++ | ++ | ++ | ++ | 2 | X |
| 45. | BPT-3147 | MS | MS | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 46. | BPT-3148 | MS | S | -- | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 47. | BPT-3150 | S | S | -- | -- | -- | + | + | + | + | + | 0 | ✓ |
| 48. | BPT-3151 | MS | MS | -- | + | -- | -- | -- | -- | -- | ++ | 1 | X |
| 49. | BPT-3159 | MR | MR | + | -- | -- | -- | -- | ++ | ++ | ++ | 2 | ✓ |
| 50. | BPT-3164 | MS | MS | -- | -- | -- | -- | -- | -- | -- | -- | 0 | ✓ |
| 51. | BPT-3168 | MS | MS | -- | 0 | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 52. | BPT-3170 | R | MR | -- | ++ | ++ | ++ | ++ | ++ | ++ | ++ | 4 | ✓ |
| 53. | BPT-3172 | MR | MS | 0 | -- | -- | -- | -- | ++ | ++ | ++ | 1 | X |
| 54. | BPT-3178 | MR | MR | -- | 0 | -- | -- | -- | -- | ++ | ++ | 1 | ✓ |

| S. No. | Genotypes | Disease reaction at | | <i>Xa21</i> | <i>xa13</i> | <i>xa5</i> | <i>Xa4</i> | <i>Xa2</i> | Total R genes expressed | Positive compatibility |
|-----------|----------------------|---------------------|----------|-------------|-------------|------------|------------|------------|-------------------------------|---------------------------|
| | | Bapatla | Maruteru | | | | | | | |
| 55. | BPT-3208 | MS | S | -- | -- | -- | ++ | -- | 1 | X |
| 56. | BPT-3244 | MS | S | -- | 0 | -- | -- | -- | 0 | ✓ |
| 57. | BPT-3260 | S | S | -- | -- | -- | -- | -- | 0 | ✓ |
| 58. | BPT-3261 | MR | S | ++ | -- | -- | -- | -- | 1 | X |
| 59. | BPT-3262 | MR | MS | -- | -- | -- | -- | -- | 0 | X |
| 60. | BPT-3263 | MS | MS | -- | + | -- | -- | -- | 0 | ✓ |
| 61. | BPT-3264 | MR | MS | -- | -- | -- | ++ | -- | 1 | X |
| 62. | BPT-3269 | MS | MS | -- | -- | -- | ++ | -- | 1 | X |
| 63. | BPT-3270 | MS | MS | -- | -- | -- | -- | -- | 0 | ✓ |
| 64. | BPT-3274 | MS | MS | -- | -- | -- | -- | -- | 0 | ✓ |
| 65. | BPT-3275 | S | S | -- | -- | -- | -- | -- | 0 | ✓ |
| 66. | BPT-3276 | MS | MS | -- | -- | -- | ++ | -- | 1 | X |
| 67. | BPT-3277 | MS | S | 0 | 0 | -- | ++ | -- | 1 | X |
| 68. | BPT-3279 | MS | S | -- | -- | -- | ++ | -- | 1 | X |
| 69. | BPT-3291 | MS | MS | -- | -- | -- | -- | -- | 0 | ✓ |
| 70. | BPT-4358 | MS | S | -- | -- | -- | -- | -- | 0 | ✓ |
| 71. | Samba Mahsuri | MS | MS | -- | -- | -- | -- | -- | 0 | ✓ |
| 72. | ISM | R | R | ++ | ++ | ++ | -- | -- | 3 | ✓ |
| 73. | Krishnaveni | S | S | -- | -- | -- | -- | -- | 0 | ✓ |
| 74. | Taichung Native-1 | HS | HS | -- | -- | -- | -- | -- | 0 | ✓ |
| | | TOTAL | | 4 | 3 | 3 | 31 | 12 | 53 | 38 |

119 R- Resistant; MR- Moderately Resistant; MS- Moderately Susceptible; S- Susceptible; HS- Highly Susceptible;
 120 ++ Resistant; +- Heterozygous; -- Susceptible; 0- Null allele; ✓ - Positively compatible; X- Positively not
 121 compatible

122

123 Molecular Characterization of Rice

124 Among the 16 markers used in this study, two SSR markers *i.e.*, RM144 and RM13 markers
 125 were not amplified, hence these markers were excluded from the analysis.

126

127 Polymorphism and Marker Efficiency

128 Analysis of the 74 rice genotypes with 14 polymorphic SSR markers revealed 47 alleles,
 129 averaging 3.22 per locus (Table 5). Alleles per locus ranged from 2 to 7, with effective allele
 130 counts from 1.05 (RM167) to 2.58 (RM39) with an average of 2.03. Polymorphism information
 131 content (PIC) ranged from 0.06 (RM167) to 0.65 (RM349, RM39), averaging 0.48. Eight were
 132 deemed highly informative, with PIC value exceeding 0.5. The maximum and the minimum
 133 allele sizes observed markers was 982 bp (pTA248) and 130 bp (RM206) respectively.

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Table 5. Genetic diversity parameters of 14 BLB resistance linked markers.

| S.No. | SSR marker | Na | Ne | PIC | Amplicon size range (bp) |
|-------|--------------------|-------------|-------------|-------------|--------------------------|
| 1. | RM349 | 7 | 2.57 | 0.65 | 190-871 |
| 2. | RM317 | 4 | 1.98 | 0.48 | 154-174 |
| 3. | RM224 | 3 | 2.55 | 0.61 | 129-163 |
| 4. | RM39 | 3 | 2.58 | 0.65 | 120-820 |
| 5. | RM164 | 4 | 1.52 | 0.33 | 240-290 |
| 6. | RM533 | 2 | 1.62 | 0.39 | 250-270 |
| 7. | RM254 | 3 | 1.82 | 0.46 | 140-170 |
| 8. | RM5509 | 3 | 2.30 | 0.57 | 270-290 |
| 9. | RM30 | 4 | 2.19 | 0.54 | 200-250 |
| 10. | RM206 | 3 | 2.56 | 0.61 | 130-170 |
| 11. | RM167 | 2 | 1.05 | 0.06 | 250-260 |
| 12. | <i>xa13</i> Prom | 2 | 1.16 | 0.18 | 270-470 |
| 13. | <i>xa5</i> FM-SR | 3 | 2.18 | 0.55 | 134-424 |
| 14. | <i>Xa21</i> pTA248 | 4 | 2.51 | 0.61 | 639-982 |
| | Maximum | 7 | 2.58 | 0.65 | 982 |
| | Minimum | 2 | 1.05 | 0.06 | 130 |
| | MEAN | 3.31 | 2.03 | 0.48 | |

141 Na- Number of alleles; Ne- Number of effective alleles; PIC- Polymorphic
142 information content; bp- base pairs

143

Marker-Assisted Selection (MAS) for BLB Resistance Genes

144 Molecular data for the genes *Xa21*, *xa13*, *xa5*, *Xa4*, and *Xa2* are presented in the Table 4.

145

***Xa21*-Linked STS Marker Analysis for BLB Resistance**

146 The presence of *Xa21* gene in germplasm was detected by the STS marker pTA248, which
147 amplified four alleles of 982bp, 737bp, 715bp and 639bp. The 982bp fragment, associated with
148 resistance, was observed in the positive control, ISM, whereas the 715bp fragment was detected
149 in the negative control, Taichung Native-1. Other variants, 737bp and 639bp, were also linked
150 to susceptibility. Among the 71 genotypes, three genotypes namely, BPT-3068, BPT-3159 and
151 BPT-3261 showed the 982bp amplicon, indicative of the presence of the *Xa21* gene. Notably,
152 BPT-3261 possessed the gene in a homozygous condition (982bp), while BPT-3068 and BPT-
153 3159 displayed the gene in a heterozygous state (982bp and 737bp). The remaining genotypes
154 produced amplicons of 737bp, 715bp, or 639bp, confirming the absence of the *Xa21* resistance
155 gene. The amplification pattern of pTA248 marker was represented in the (Fig. 1).

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158

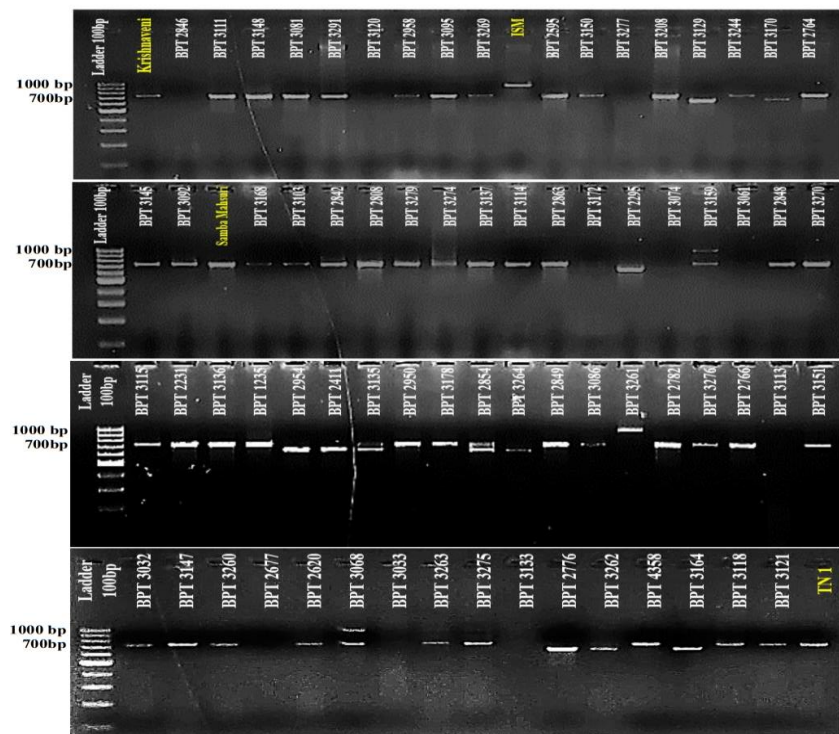


Fig. 1. Expression of *Xa21* gene amplification using marker pTA248.

Xa13-Linked STS Marker Analysis for BLB Resistance

163 The *xa13* gene in germplasm was detected by the marker *xa13*prom. The positive control,
164 ISM, showed a 470 bp fragment, while the negative control, Taichung Native-1, exhibited a
165 270 bp fragment. BPT-3136 and BPT-3170 produced amplicon of 470bp, carried *xa13* in
166 homozygous condition. The genotypes BPT-3061, BPT-3068, BPT-3113, BPT-3151 and BPT-
167 3263 produced heterozygous bands of 470bp and 270bp, where *xa13* the recessive gene was
168 not expressed. While, remaining 64 genotypes produced amplicon of 270bp (similar to
169 Taichung Native-1).

Xa5-Linked STS Marker Analysis for BLB Resistance

The STS marker *xa5FM-SR* linked to recessive *xa5* gene amplified 424bp (common), 134bp (resistance specific) and 313bp (susceptibility specific) fragments. Genotypes BPT-3145 and BPT-3170, showed of 424bp and 134bp fragments (similar to the ISM) indicating the presence of the resistant gene. BPT-2848, BPT-3061, BPT-3068, and BPT-3150 were heterozygous, producing 424 bp, 134 bp, and 313 bp fragments. The remaining 65 genotypes produced only 313bp and 134bp bands.

181 **Xa4-linked SSR marker analysis for BLB resistance**

182 The Xa4 gene was identified using the marker RM224, with resistant genotypes producing
183 a 160bp fragment and susceptible genotypes a 150bp fragment (Panwar *et al.*, 2018; Chen *et*
184 *al.* 1997). In this study, 31 genotypes carried Xa4, producing a 160bp amplicon.

185
186 **Xa2-Linked SSR Marker Analysis for BLB Resistance**

187 The Xa2 gene was detected using the SSR marker RM317. with a 154bp DNA fragment
188 indicating resistance (Hasan *et al.*, 2020; He *et al.*, 2006). In this study, 12 genotypes namely
189 BPT-2620, BPT-2764, BPT-2808, BPT-2848, BPT-3068, BPT-3121, BPT-3133, BPT-3136,
190 BPT-3137, BPT-3151, BPT-3170 and BPT-3178 amplified the154bp fragment, and considered
191 to possess Xa2.

192
193 **R Genes Expressed**

194 A total of 53 R genes were expressed from all the cultivars, for the five genes *Xa21*, *xa13*,
195 *xa5*, *Xa4* and *Xa2* studied. The gene *Xa4* was found to be most frequent, carried by 31
196 genotypes, followed by *Xa2* (12) and *Xa21* (3), while *xa5* and *xa13* were found in only two
197 genotypes. BPT-3170 possessed combination of four multiple resistance genes
198 (*xa13+xa5+Xa4+Xa2*). Two gene combinations were found in eight genotypes, namely BPT-
199 2848, BPT-3068, BPT-3121, BPT-3133, BPT-3136, BPT-3137, BPT-3145 and BPT-3159.
200 Single genes were detected in 30 genotypes, and, 32 genotypes did not express any R genes.

201
202 **Compatibility between Genotypic and Phenotypic Expression of BLB Resistance**

203 Positive compatibility between phenotypic and genotypic expression of BLB resistance was
204 observed in 38 rice lines, including checks of which nine exhibited resistance (score 1-3), and
205 29 showed susceptibility (score 5-9) (Table 4). Among the resistant genotypes namely, BPT-
206 3068, BPT-3074, BPT-3092, BPT-3137, BPT-3170, BPT-2846, BPT-2958, BPT-3159, and
207 BPT-3178, all except BPT-2846 showed positive compatibility with R gene expression. BPT-
208 3170 carried four resistance genes (*xa13+xa5+Xa4+Xa2*), while BPT-3068 (*Xa21+Xa2*), BPT-
209 3137 (*Xa4+Xa2*), and BPT-3159 (*Xa21+Xa4*) carried two genes. BPT-3074 (*Xa4*), BPT-2958
210 (*Xa4*), BPT-3178 (*Xa2*), and BPT-3092 (*Xa4*) carried a single gene.

211
212 **Molecular Diversity Analysis**

213 The phylogenetic tree was constructed using 14 markers linked to BLB resistance based on
214 neighbour-joining method (Fig. 2). The genotypes were grouped into three major clusters

(Table. 6). Cluster I, containing 32 genotypes, was further subdivided into four sub-clusters (IA, IB, IC, ID) and possessed the highest number of resistance genes (21), of which only 3 were positively compatible (Tables 7, and 8). Cluster II, the smallest with 7 genotypes and resistant check, was divided into sub-clusters IIA and IIB. Sub-cluster IIA contains exclusively resistant genotypes exhibiting multiple resistance genes namely, BPT-3068, BPT-3170 and resistant check ISM. This cluster expressed 11 R genes, 9 of which were positively compatible. Cluster III was the largest with 32 genotypes mostly showing susceptibility reaction (with score 5-9) including both the susceptible checks (Taichung Native-1 and Krishnaveni). It was divided into four sub-clusters (IIIA, IIIB, IIIC, IIID) and possessed highest number of R genes (21) with only 5 being positively compatible.

225

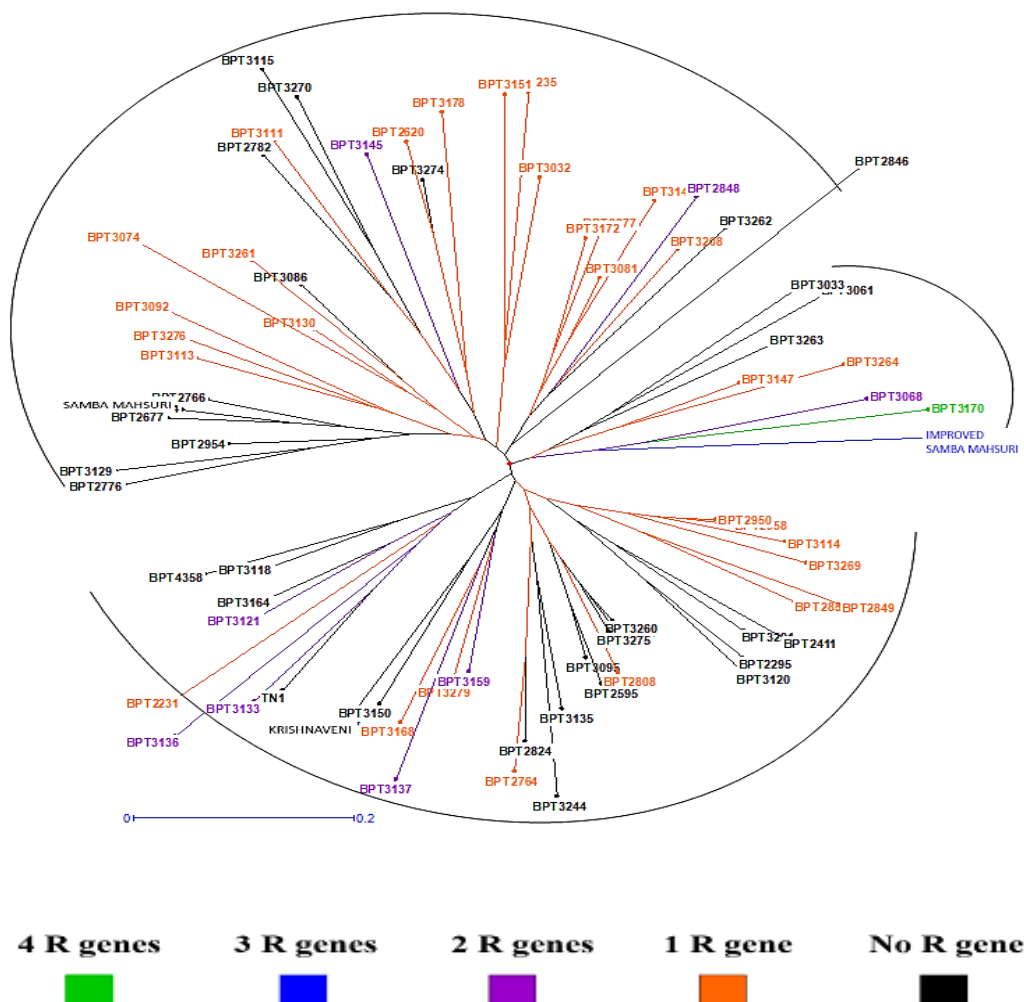


Fig. 2. Phylogenetic tree constructed based on BLB linked markers representing resistance genes.

232

Table 6. Grouping of genotypes into different clusters based on molecular diversity.

| Name of the cluster | Name of the sub-cluster | Number of Genotypes | Name of the Genotypes |
|---------------------|-------------------------|---------------------|---|
| I | IA | 13 | BPT-2776, BPT-3129, BPT-2954, BPT-2677, Samba Mahsuri, BPT-2766, BPT-3113, BPT-3276, BPT-3092, BPT-3074, BPT-3130, BPT-3261, BPT-3086 |
| | IB | 8 | BPT-2782, BPT-3111, BPT-3115, BPT-3270, BPT-3145, BPT-2620, BPT-3274, BPT-3178. |
| | IC | 3 | BPT-3151, BPT-1235, BPT-3032 |
| | ID | 9 | BPT-3172, BPT-3277, BPT-3081, BPT-3148, BPT-2848, BPT-3208, BPT-3262, BPT-2846 |
| II | IIA | 5 | BPT-3033, BPT-3061, BPT-3263, BPT-3147, BPT-3264 |
| | IIB | 2 | BPT-3068, BPT-3170, ISM |
| III | IIIA | 10 | BPT-2950, BPT-2958, BPT-3114, BPT-3269, BPT-2849, BPT-2854, BPT-2411, BPT-3291, BPT-2295, BPT-3120 |
| | IIIB | 10 | BPT-3260, BPT-2863, BPT-3275, BPT-2808, BPT-2595, BPT-3095, BPT-3135, BPT-3244, BPT-2824, BPT-2764 |
| | IIIC | 5 | BPT-3159, BPT-3279, BPT-3137, BPT-3168, BPT-3150, Krishnaveni |
| | IID | 7 | Taichung Native-1 , BPT-3133, BPT-3136, BPT-2231, BPT-3121, BPT-3164, BPT-3118, BPT-4358 |

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Table 7. Clustering of R genes for bacterial leaf blight resistance.

| Cluster No | Xa21 | xa13 | xa5 | Xa4 | Xa2 | Total |
|--------------------|----------|----------|----------|-----------|-----------|-----------|
| Cluster I | 1 | - | 1 | 15 | 4 | 21 |
| Cluster II | 2 | 2 | 2 | 3 | 2 | 11 |
| Cluster III | 1 | 1 | - | 13 | 6 | 21 |
| Total | 4 | 3 | 3 | 31 | 12 | 53 |

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Table 8. Clustering of positively compatible R genes for bacterial leaf blight based on phenotypic and genotypic marker data.

| Cluster No | Xa21 | xa13 | xa5 | Xa4 | Xa2 | Total |
|--------------------|----------|----------|----------|----------|----------|-----------|
| Cluster I | - | - | - | 2 | 1 | 3 |
| Cluster II | 2 | 2 | 2 | 1 | 2 | 9 |
| Cluster III | 1 | - | - | 3 | 1 | 5 |
| Total | 3 | 2 | 2 | 6 | 4 | 17 |

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239

DISCUSSION

This study evaluated the genetic resistance to bacterial leaf blight (BLB) in 71 rice genotypes developed from diverse elite and improved breeding lines across multilocation and revealed significant variability in disease response. Disease severity was higher at Maruteru, a known BLB hotspot, compared to Bapatla, likely due to Maruteru's favorable environment for pathogen spread. This variation can be attributed to the polygenic nature of BLB resistance, governed by over 40 R genes, and strongly influenced by environment. Genotypes showed a wide range of responses, from resistant to highly susceptible, consistent with findings by Rahman *et al.* (2017), Qudsia *et al.* (2019), and Majumder *et al.* (2019). Using 14 gene-linked markers, 47 alleles including rare and null alleles were detected across 74 rice genotypes. Rare alleles identified with markers RM349, RM30, and RM164 likely result from structural variations or mutational events (Victoria *et al.*, 2007). The

251 informativeness of SSR markers in assessing genetic diversity aligns with Ashiba *et al.* (2020),
252 and Khan *et al.* (2015).

253 Molecular characterization of five resistance genes *Xa21*, *xa13*, *xa5*, *Xa4* and *Xa2* was
254 conducted with ISM (*Xa21+xa13+xa5*) as a positive control. BPT-3170 carrying
255 *xa13+xa5+Xa4+Xa2* may confer broad spectrum resistance and serve as a valuable donor for
256 BLB resistance breeding. BPT-3061 and BPT-3068 possess the recessive resistance genes *xa13*
257 and *xa5* in heterozygous condition, requiring homozygosity for expression. Advancing these
258 lines through selfing can facilitate allele fixation and enhance resistance to BLB.

259 Among the 74 genotypes screened, 38 showed concordance between phenotypic and
260 genotypic resistance to BLB, while 36 showed discrepancies. The genotypes showing resistant
261 reaction without detected R genes may harbor additional resistance genes not included in this
262 study. Conversely, the susceptibility observed in some *Xa4*-carrying genotypes suggests
263 pathogen adaptation, likely driven by wide spread use of *Xa4*-based cultivars in India and
264 Southeast Asia (Ma *et al.*, 1999 and Sun *et al.*, 2003). This study shows that genotypic
265 screening correlates with phenotypic screening in most genotypes and it also underscore the
266 importance of combining genotypic and phenotypic data for accurate resistance evaluation.

267 Cluster analysis based on 14 R-gene-linked markers grouped genotypes into three major
268 clusters, with the resistant check (ISM) and susceptible checks (Taichung Native-1 and
269 Krishnaveni) occupying distinct clusters. The eight genotypes, BPT-3068, BPT-3074, BPT-
270 3092, BPT-3137, BPT-3170, BPT-2958, BPT-3159 and BPT-3178 that exhibited phenotypic
271 disease reaction towards resistance (score 1-3) for BLB at both the locations and demonstrated
272 positive R gene compatibility were distributed across clusters, highlighting their genetic
273 diversity. These resistant genotypes are promising resources for developing durable BLB-
274 resistant varieties. Similar studies on phylogeny analysis in rice using SSR markers were
275 reported by Ashiba *et al.* (2020), Khan *et al.* (2015) and Khannetah *et al.* (2021).

276 CONCLUSIONS

277 This study assessed genetic resistance to bacterial leaf blight (BLB) in 71 rice genotypes
278 through multilocation screening and molecular characterization. BPT-3170, exhibiting
279 phenotypic resistance and carrying multiple resistance genes (*xa13+xa5+Xa4+Xa2*), shows
280 strong potential as a donor for broad-spectrum BLB resistance. Additionally, genotypes BPT-
281 3068, BPT-3074, BPT-3092, BPT-3137, BPT-3170, BPT-2958, BPT-3159, and BPT-3178
282 showed consistent phenotypic resistance (disease score 1-3) across locations and positive

284 compatibility with R-gene expression, making them valuable genetic resources for BLB
285 resistance breeding. The findings underscore the ineffectiveness of single gene resistance and
286 emphasizes the need for pyramiding multiple R genes to achieve durable and broad-spectrum
287 resistance. Deploying diverse R-gene combinations in elite cultivars will enhance the resilience
288 of rice crops against evolving BLB pathotypes, ensuring sustainable production and food
289 security particularly in areas where BLB is a persistent threat to rice cultivation.

290

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294

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