Tagging microsatellite marker to a blast resistance gene in the irrigated rice cultivar Cica-8

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Abstract - The rice cultivar Cica-8 exhibit differential reaction to several pathotypes of Magnaporthe oryzae. The objective of the present investigation was to determine the number of alleles involved in the expression of resistance to leaf blast and identify microsatellite markers linked to these alleles. A cross between cultivar Metica-1 and Cica-8 susceptible and resistant, respectively, to pathotype IB-1 (Py1049) was made to obtain F₁, F₂, BC₁:₁ and BC₁:₂ progenies. Greenhouse tests for leaf blast reaction showed that resistance is controlled by a monogenic dominant gene. For testing microsatellite markers, DNA of both resistant and susceptible parents and F₁ and F₂ populations was extracted. As expected for single dominant gene the F₂ populations segregated at a ratio of 3:1. Of the 11 microsatellite markers tested, one marker RM 7102 was found to be closely linked to the resistant allele at a distance of 2.7 cM, in the cultivar Cica-8 to pathotype IB-1.

Key words: Oryza sativa, Pyricularia oryzae, DNA markers, marker assisted selection.

INTRODUCTION

Rice blast caused by Magnaporthe oryzae B. Couch [anamorph - Pyricularia oryzae Cavra] ranks first in order of economic importance in Brazil. The grain yield losses caused by rice blast are worldwide. In China, over 3.8 million hectares were affected in 1993 causing losses of approximately 1.1 million tons, whereas in Japan the yield losses were estimated to vary between 20 to 100% (Khush and Jena 2009). In Brazil, the losses were reported to be as high as 100% in susceptible upland rice cultivars under upland conditions (Prabhu et al. 2009). Even though there are no reliable estimates of losses in irrigated rice, they are considered significant, because the farmers resort to intensive fungicide spraying to control blast. Breeding for blast resistance is the most economic means of reducing losses in yield as well as in grain quality.

Identification of resistance genes with molecular markers is the basic requisite for marker-assisted selection (MAS) in resistance breeding. Several DNA markers closely linked to agronomic traits have been developed for indirect selection for traits of interest in segregating populations (Caixeta et al. 2009). The indirect selection is highly effective due to the absence of confounding effect of environment and also pyramiding of genes for disease resistance, which is difficult through the use of conventional plant breeding methods (Gupta and Varshney 2000). The availability of different molecular markers permits characterization of genes of interest. Nearly 60 blast resistance genes have been identified following conventional genetic analysis of resistant donors using specific pathotypes and molecular analysis techniques (Khush and Jena 2009). In rice, molecular markers have been successfully used to pyramid bacterial blight resistance genes into isogenic lines (Huang et al. 1997) and three blast resistance genes in a single genotype (Hittalmani et al. 2000).

Among the PCR-based markers (RAPD₃, SCAR₃, CAPS, STS, STMS, AFLP₃ etc.) microsatellites have attracted attention, because they are hypervariable, abundant and well distributed throughout the rice genome and are easily available through the published linkage map or in public data base and permit differentiation of homo and heterozygote individuals (McCouch et al. 2002, Caixeta et al. 2009). The speed, reliability and cost effectiveness make microsatellite analysis on acrylamide gel an attractive tool for MAS in blast resistance breeding (Fuentes et al. 2008).

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Korean cultivar Suweon 365 carries three major resistance genes, \textit{Pi}18, \textit{Pi}21\,(t), and \textit{Pi}22\,(t), against Korean isolates KI-313, KJ-101, and KJ-201, respectively, using microsatellite markers in an \textit{F}_2 population derived from a cross of cultivars Suweon and Chucheongbyeo (Ahn et al. 2000). A resistance gene, \textit{Pi-CO39}, corresponding to a virulence gene \textit{Avr1-CO39}, was identified in cultivar CO39 through linkage analysis in a \textit{F}_2 population from a cross between cultivars CO39 and 51583, with microsatellite and RFLP markers (Chauhan et al. 2002). Microsatellite markers (SSR) were used for mapping the gene \textit{Pi-k}, that confer resistance to blast races in Himalayan region of north east India (Sharma et al. 2005). In Colombia, of the 24 microsatellite sequences tested, six showed polymorphism and two markers RM1233 and RM224 were linked to the gene \textit{Pi-1\,(t)} in the same position (0.0 cm). Furthermore, microsatellite analysis of advanced rice breeding lines with genetic background showed that all known sources of blast resistance carry the specific \textit{Pi-1\,(t)} allele (Fuentes et al. 2008).

The cultivar Cica-8 shows resistance to several Brazilian blast fungus races. To effectively use its resistance gene in a rice breeding program, it is important to tag the resistance gene or genes present in this cultivar with molecular markers. The present investigation was undertaken to determine the number of alleles in the expression of rice blast resistance and identify microsatellite markers linked to these genes.

**MATERIAL AND METHODS**

**Plant material for inheritance studies**

The populations of \textit{F}_1, \textit{F}_2 and BC were obtained from a cross between irrigated rice cultivar Metica-1 and cultivar Cica-8, susceptible and resistant, respectively, to pathotype IB-1, in Embrapa Rice and Beans Research Center, Santo Antônio de Goiás, GO. A part of \textit{F}_1 seed (600) was used for backcross with susceptible parent Metica-1 (BC\_1\_t) and with resistant parent Cica-8 (BC\_2\_t) to obtain 400 seeds of each backcross and the remaining \textit{F}_2 plants were self-pollinated to generate 1400 \textit{F}_2 progeny seeds.

**Inoculation and evaluation**

The test material was planted in plastic trays (30 cm x 15 cm x 10 cm) containing 5 kg of soil fertilized with 5 g of NPK 5-30-15, 2 g of ammonium sulfate and 1 g of zinc sulfate. The seed was sown in eight 10 cm along rows at the rate of 15 seeds per row. The plants after germination were thinned to maintain 10 plants per row. Thirty plants of each parent Metica-1 and Cica-8, 20 plants of \textit{F}_1, 200 plants of \textit{F}_2, 100 plants of each backcross were planted in 12 trays.

The monosporic isolate Py 1049 of \textit{M. oryzae} originated from cultivar Metica-1 was used for inoculation and evaluation of leaf blast reaction. The isolate was identified as pathotype IB-1 based on previous inoculation testes on eight standard international differentials. Initially the pure culture was multiplied in Petri plates containing culture medium PDA (potato-dextrose-agar). For sporulation, mycelial discs were grown on oat meal agar medium in plates and incubated at 27 °C, for 10 days under continuous fluorescent day light. The aerial mycelium was scrapped with a sterilized glass rod under aseptic conditions. The plates were later kept under fluorescent day light for 48 hours with lids open but covered with sterilized cheese cloth. Inoculum was prepared by flooding culture plates with distilled water and conidia were dislodged with a paint brush. The conidial suspension along with the scrapped mycelium was filtered through double layer of cheese cloth. The final conidial suspension was adjusted to 3x10^6 conida mL^{-1} using haemocytometer.

Twenty one day-old plants with three fully opened leaves were inoculated with spores suspension on the leaves, until run-off, using a Devilbis automizer connected to an air compressor. The inoculated plants were kept in moist chamber for 24 h at temperatures ranging from 19 °C to 21°C. Subsequently they were transferred to greenhouse benches where the mean temperature was 28 °C and humidity >80%.

The plants were evaluated after seven days for leaf blast reaction using a 0-9 scale according to IRRI (1996). The disease ratings 5, 7 and 9 were considered susceptible or compatible and 0, 1 and 3 resistant or incompatible. The frequency of resistant and susceptible plants in \textit{F}_1 and backcross progeny was assessed and chi-square (x^2) analysis was done for goodness of fit (Strickberger 1971).

**Identification of microsatellite markers linked to resistance gene**

The rice leaves were macerated in liquid nitrogen and DNA extracted according to Doyle and Doyle (1987). The DNA concentration was adjusted to 10 ng \mu L^{-1} and has been estimated visually utilizing the standard DNA \textit{lambda} at concentrations of 50 ng \mu L^{-1}, 100 ng \mu L^{-1} and 200 ng \mu L^{-1}. The microsatellites utilized are shown in Table 1.

<table>
<thead>
<tr>
<th>Marker</th>
<th>Expected size (bp)</th>
<th>Gene</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>9871.T7E2b</td>
<td>642</td>
<td>\textit{Pi5}</td>
<td>Suh et al. (2009)</td>
</tr>
<tr>
<td>yca72</td>
<td>635</td>
<td>\textit{Pia}</td>
<td>Suh et al. (2009)</td>
</tr>
<tr>
<td>RM224</td>
<td>122</td>
<td>\textit{Pi1}</td>
<td>Jia and Moldenhauer (2010)</td>
</tr>
<tr>
<td>OSR32</td>
<td>247</td>
<td>\textit{Pita}</td>
<td>Li et al. (2008)</td>
</tr>
<tr>
<td>RM7102</td>
<td>170</td>
<td>\textit{Pi41}</td>
<td>Yang et al. (2009)</td>
</tr>
<tr>
<td>RM6836</td>
<td>178</td>
<td>\textit{Piz}</td>
<td>Fjellstrom et al. (2006)</td>
</tr>
<tr>
<td>pBA14</td>
<td>480</td>
<td>\textit{Pig}</td>
<td>Liu et al. (2002)</td>
</tr>
<tr>
<td>RM144</td>
<td>254</td>
<td>\textit{Pi_k}</td>
<td>Jia and Moldenhauer (2010)</td>
</tr>
<tr>
<td>RM28130</td>
<td>176</td>
<td>\textit{Pi40}</td>
<td>Yang et al. (2009)</td>
</tr>
<tr>
<td>RM1261</td>
<td>167</td>
<td>\textit{Pi41}</td>
<td>Yang et al. (2009)</td>
</tr>
<tr>
<td>JJ817</td>
<td>1450</td>
<td>\textit{Pi5}</td>
<td>Suh et al. (2009)</td>
</tr>
</tbody>
</table>
Each polymerase reaction was conducted in a final volume of 40 μL containing genomic DNA, buffer (100 mM Tris-HCl, pH 8.8 and 500 mM KCl), MgCl₂ (25 mM), dNTP (2 mM of each dATP, dGTP, dCTP e dTTP), marker (forward and reverse) and Taq DNA polymerase. The enzymatic amplification was performed in a thermocycler (ESCO Swift MaxPro) with amplification program specified for each marker. For microsatellite markers utilized in this study the reaction was processed as shown in Table 2.

Table 2. Procedure for each microsatellite marker utilized

<table>
<thead>
<tr>
<th>Marker</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>9871.T7E2b</td>
<td>4 minutes at 95 ºC, 35 cycles for 30 seconds at 95 ºC, 30 seconds at 55 ºC and 1 minute at 72 ºC and 10 minutes at 72 ºC</td>
</tr>
<tr>
<td>yca72</td>
<td>4 minutes at 95 ºC, 35 cycles for 30 seconds at 95 ºC, 30 seconds at 55 ºC and 1 minute at 72 ºC and 10 minutes at 72 ºC</td>
</tr>
<tr>
<td>RM224</td>
<td>1 minute at 94 ºC, 40 cycles for 30 seconds at 94 ºC, 30 seconds at 55 ºC and 30 seconds at 72 ºC and 10 minutes at 72 ºC</td>
</tr>
<tr>
<td>OSR32</td>
<td>5 minutes at 94 ºC, 35 cycles for 1 minute at 94 ºC, 50 seconds at 55 ºC and 1 minute at 72 ºC and 5 minute at 72 ºC</td>
</tr>
<tr>
<td>RM7102</td>
<td>3 minutes at 95 ºC, 29 cycles for 30 seconds at 95 ºC, 30 seconds at 55 ºC and 30 seconds at 72 ºC and 7 minute at 72 ºC</td>
</tr>
<tr>
<td>RM6836</td>
<td>3 minutes at 95 ºC, 35 cycles for 30 seconds at 95 ºC, 30 seconds at 55 ºC and 1 minute at 72 ºC and 10 minutes at 72 ºC</td>
</tr>
<tr>
<td>pBA14</td>
<td>4 minutes at 94 ºC, 35 cycles for 45 seconds at 94 ºC, 45 seconds at 55 ºC and 1 minute at 72 ºC and 5 minutes at 72 ºC</td>
</tr>
<tr>
<td>RM144</td>
<td>5 minutes at 94 ºC, 35 cycles for 1 minute at 94 ºC, 1 minute at 55 ºC and 2 minute at 72 ºC and 7 minute at 72 ºC</td>
</tr>
<tr>
<td>RM28130</td>
<td>3 minutes at 94 ºC, 35 cycles for 30 seconds at 94 ºC, 30 seconds at 55 ºC and 1 minute at 72 ºC and 7 minute at 72 ºC</td>
</tr>
<tr>
<td>RM1261</td>
<td>5 minutes at 94 ºC, 35 cycles for 1 minute at 94 ºC, 1 minute at 55 ºC e 2 minute at 72 ºC and 7 minute at 72 ºC</td>
</tr>
<tr>
<td>JJ817</td>
<td>4 minutes at 95 ºC, 35 cycles for 30 seconds at 95 ºC, 30 seconds at 55 ºC and 1 minute at 72 ºC and 10 minutes at 72 ºC</td>
</tr>
</tbody>
</table>

The amplification products were separated in polyacrylamide gel (6%). The images were obtained utilizing scanner HP Scanjet 3770. For the identification of markers linked to resistance genes to isolate Py 1049 in the test populations, evaluation and selection was done in the following three steps: In the first step 11 microsatellite primers (Table 1) were tested and only those that were capable of detecting polymorphism among parents were selected for second step. In the second step, only the primers capable of amplifying the same standard polymorphism in parents and bulks were selected for third step. The bulks of F₁, F₂ resistant and F₂ susceptible, each contained DNA (10 ng) of seven individuals of each population. In the final step, the above selected primers were utilized for amplification of reactions that contained DNA with each individual component of bulks previously described. Each primer selected participated in the following reaction amplification: a) resistant parent (1 reaction); b) susceptible parent (1 reaction); c) seven individuals of F₁ population (seven reactions); d) seven individuals of resistant F₂, and; e) seven individuals of susceptible F₂ (seven reactions).

Analysis

For estimating the distance between microsatellite marker and resistance gene, 200 individual F₂ populations were evaluated phenotypically and genotypically. The distance was estimated utilizing the software MAP-MAKER III with one minimum lod score of 3.0 and Kosambi function of mapping (Lander et al. 1987).

RESULTS AND DISCUSSION

Genetic analysis of resistance of the cultivar Cica-8 was conducted by inoculating F₁, F₂, BC₁ and BC₂ populations obtained from a cross between cultivars Metica-1 and Cica-8 with the isolate Py 1049 under greenhouse conditions. The observed and expected ratios are presented in Table 3. The F₁ plants showed resistant reaction whereas F₂ population segregated at a ratio of 3:1 resistant and susceptible plants indicating that the resistance to pathotype IB-1 in the cultivar Cica-8 is dominant and monogenic. These results were confirmed by the expected segregation ratio of 1:1 resistant and susceptible plants of backcross population to susceptible parent (BC₁) and resistant reaction of plants of backcross (BC₂) with resistant parent.

Table 3. Segregation of F₁, F₂, BC₁, BC₂ populations of a cross between Metica-1 and Cica-8, susceptible and resistant, respectively, to pathotype IB-1 of Magnaporthe oryzae

<table>
<thead>
<tr>
<th>Parents/ generation</th>
<th>No. of plants</th>
<th>Observed data</th>
<th>Expected ratio</th>
<th>χ²</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>R</td>
<td>S</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metica-1</td>
<td>30</td>
<td>0</td>
<td>30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cica-8</td>
<td>30</td>
<td>30</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F₁</td>
<td>20</td>
<td>20</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>F₂</td>
<td>200</td>
<td>137</td>
<td>63</td>
<td>3:1</td>
<td>4.167</td>
</tr>
<tr>
<td>BC₁</td>
<td>82</td>
<td>31</td>
<td>51</td>
<td>1:1</td>
<td>4.402</td>
</tr>
<tr>
<td>BC₂</td>
<td>100</td>
<td>79</td>
<td>21</td>
<td>1:0</td>
<td>4.203</td>
</tr>
</tbody>
</table>

1 Backcross to susceptible parent Metica-1
2 Backcross to resistant parent Cica-8
3 Expected ratio resistant to susceptible
4 Significant at 5% probability

Based on studies on genetic resistance, in five improved and four traditional cultivars using three rice blast isolates, Yu et al (1986) showed that resistance is controlled by one or two dominant genes. The analysis of isogenic lines with single resistance genes showed that the resistance is controlled by independent dominant genes (Mackill and Bonman 1992).
In Brazil, the inheritance studies conducted with 12 different crosses, Filippi and Prabhu (1996) observed that the resistance to the predominant races IB-1 and IB-9 was dominant and controlled by one or three genes. Nunes et al. (2007) showed that the resistance in irrigated rice cultivars Taim and BRS Firmeza is controlled by one dominant gene. The resistance in Cica-8 is controlled by a pair of dominant alleles and segregated independently confirming the majority of the earlier studies on inheritance of resistance conducted in Brazil and elsewhere.

Of all 11 markers tested, four markers, RM224, RM7102, RM1261 and yca72 were polymorphic and showed amplification differentiating the resistant and susceptible parent (Figure 1). These four polymorphic markers were tested in

![Figure 1. Amplification pattern observed in denatured polyacrylamide gel (6%) for all primers tested. From left to right: M: Ladder 10 bp; P1: susceptible parent (Metica-1) to isolate Py 1049 and P2: resistance donor parent (Cica-8) to the isolate Py 1049.](image-url)
bulk populations of segregating F$_2$ resistant, F$_2$ susceptible and F$_1$ plants (Figure 2). It was possible to detect the same amplification of bulks, observed in parents, in three markers (RM224, RM1261 and RM7102). Only one marker RM7102 showed standard amplification and expected polymorphism or in other words the amplification observed in resistant parent was repeated in resistant F$_1$ and F$_2$ bulks. The polymorphic bands with 170 and 190 base pairs were tested in seven individual components of each bulk.

It was possible to observe in individuals F$_1$, to F$_2$, tested with marker RM7102, the presence of two bands one on the top similar to that observed in parent Cica-8, resistant to isolate Py 1049, with 190 base pairs (bp), and one on the bottom with 170 pb, similar to that observed in susceptible cultivar Metica-1. This pattern of amplification is in accord with its heterozygote condition (Figure 2). The same amplification pattern for microsatellite marker RM7102 and individual resistant F$_2$ population (F$_2$R$_1$ to F$_2$R$_7$) was obtained. All showed similar bands indicating that these individuals are also heterozygotes. The individual susceptible F$_2$ plants showed only inferior band (170 pb), similar to the susceptible parent Metica-1. This amplification pattern suggest that the top band (190 pb) is associated with resistance to isolate Py 1049, observed in this population, and the bottom band is possibly a recessive allele, associated with the susceptibility inherited of the parent Metica-1 (Figure 2). It was verified that this marker is linked to the resistance gene at a distance of 2.7 cM (Figure 3).

**Figure 2.** Amplification pattern observed in denatured polyacrylamide gel (6%) for primers RM224, RM7102, RM1261 and yca72 (no amplification) in bulks F$_1$ and F$_2$ resistant and susceptible populations. From left to right: M: Ladder 10 bp; P1: susceptible parent (Metica-1) to isolate Py 1049; P2: resistance donor parent (Cica-8) to the isolate Py 1049; BH: F$_1$ bulk population; BFR: F$_2$ resistant bulk; BFS: F$_2$ susceptible bulk.
Figure 3. Amplification pattern observed in denatured polyacrylamide gel (6%) in individual F₁, F₂ resistant and F₂ susceptible populations by marker RM7102. From left to right: M: Ladder 10 bp; 3A, 3B and 3C: numbers 1 to 200 of F₂ resistant and susceptible plants.

Yang et al. (2009), working on identification and mapping of gene Pi4₁, utilized resistant cultivar 93-11 and susceptible cultivar Nipponbare, and F₂ population derived from this cross, to challenge with two blast isolates (CHL724 e CHL743). They showed different responses to resistance and susceptibility in these parents. For this study, they utilized 180 microsatellite markers and obtained seven markers linked to the gene Pi4₁, RM7102 being one of them.

The markers RM144, RM224 and yca72 were related to resistance genes Pi-k, Pi1 and Pi₄, respectively (Suh et al. 2009, Jia and Moldenhauer 2010), indicating that these genes are not associated with resistance in cultivars Metica-1 and Cica-8. The marker RM1261 (Yang et al. 2009) was polymorphic to parents but not to F₁, F₂R and F₂S bulks. This marker is related to gene Pi4₁, but did not show association to resistance allele of the cultivar Cica-8. It can be inferred that Pi4₁ and the allele identified in Cica-8 are similar but show some differences among them.

Bryan et al. (2000) reported that change in single aminoacid differentiate resistant and susceptible alleles in gene Pi-ta, because they modify the proteins that will be codified by gene. According to Jia et al. (2000) the substitution of single aminoacid in leucine-rich domain of gene Pi-ta or in gene AVR-Pita results in loss of resistance in plant. These results demonstrated that even with little alterations in aminoacids of genes the resistance and/or avirulence are capable of inducing variability and loss of resistance. The gene Pita is localized in a stable region close to centromere (Zhou et al. 2007). This gene codifies cytoplasmic protein NBS-LRR, following the mold of majority of cloned genes. The same is capable of producing 12 different proteins, which are effective against 10 pathotypes. On the other hand, the avirulence allele AVR-Pita is localized in one unstable telomeric region of the chromosome and the protein has been shown to be frequently altered under field conditions. The high specificity of pathotype IB-1 to cultivar Cica-8 has been reported (Araujo and Prabhu 2004) and resistance until today is highly stable. However, investigations concerning alteration of AVR gene frequency in a P. oryzae population have been done for many years around the world. But we are still looking for complementary strategies of disease management.

The microsatellite marker RM7102 which is tightly linked (2.7 cM) to the blast resistance gene, to IB-1 pathotype, in cultivar Cica-8 in rice, showed accuracy of 98% in identifying the resistant plants in segregating populations and thus provide means to conduct marker assisted selection in a rice breeding program. This information is also useful for the strategies of pyramiding different major resistance genes and gene rotation, which are important components in integrated pest management.

ACKNOWLEDGEMENTS

The authors acknowledge the support received by Embrapa Rice and Bean, by granting equipment usage and CAPES, for financial support to the principal author.
Marcadores microsatélites ligados a gene de resistência à brusone na cultivar de arroz irrigado Cica-8

Resumo - A cultivar de arroz Cica-8 apresenta reação diferencial a vários patótipos de Magnaporthe oryzae. Este trabalho teve como objetivo determinar o número de alelos envolvidos na expressão de resistência do arroz à brusone e identificar marcadores microsatélites ligados a estes alelos. Foi realizado um cruzamento entre as cultivares Metica-1 e Cica-8, resistente e suscetível, respectivamente, ao patótipo IB-1 (Py1049) para obtenção de progêniis F\textsubscript{1}, F\textsubscript{2} RC\textsubscript{1} e RC\textsubscript{2}. Testes fenotípicos realizados em casa de vegetação para reação do brusone nas folhas mostrou que a resistência é controlada por um gene dominante. Para testar marcadores microsatélites selecionados, foram feitas extrações de DNA de cada genitor e das populações F\textsubscript{1} e F\textsubscript{2}. Como esperado para um gene dominante, a população F\textsubscript{2} segregou em razão 3:1. Entre 11 marcadores microsatélites testados, um marcador, RM7102, encontrou-se ligado ao alelo de resistência à distância de 2.7 cM na cultivar Cica-8 para o patótipo IB-1.

Palavras-chave: Oryza sativa, Pyricularia oryzae, marcadores moleculares, seleção assistida.

REFERENCES


