

## Parasexual Cycle Provides Genetic Segregants Equivalent to Sexual Progeny in the Rice Blast Fungus *Magnaporthe oryzae*

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### Abstract

We demonstrated that the segregation ratio of avirulence and virulence traits on rice cultivars, Hattan 3 and line K59-1, and the mating types of parasexual recombinants were consistent with those of sexual progeny in the rice blast fungus *Magnaporthe oryzae*. Our results indicated that the genetic characters of parasexual recombinants segregated in a recombination manner similar to sexual events.

**Discipline:** Plant disease

**Additional key word:** *Pyricularia oryzae*

### Introduction

Blast caused by *Magnaporthe oryzae* B. Couch (anamorph *Pyricularia oryzae* Cavara), previously known as *Magnaporthe grisea* (Hebert) Barr<sup>4</sup>, is the most serious disease of rice and limits rice production worldwide. Resistant cultivars are usually effective for controlling the disease, although newly developed resistant cultivars have been affected within several years after their introduction. The breakdown of their blast resistance results from the occurrence of pathogenic variants (races). Sexual mating, mutation, and parasexual recombination are thought to be factors affecting diversification and variability of the pathogen. Since ascospore production from a cross between Japanese field isolates had never been reported, mutation and parasexual recombination are likely the two major causes of variation in the pathogenicity of *M. oryzae*.

The rice cultivar specificity in this fungus has a gene-for-gene relationship<sup>7</sup>, and avirulence genes have been reported<sup>11,12,19,23,24</sup>. While numerous studies on the parasexual cycle have been conducted in the rice blast fungus<sup>3,5,6,8,13,14,22,25</sup>, no genetic analyses have been performed on parasexual recombinants that included avirulence genes. Therefore, we carried out a genetic study after obtaining parasexual recombinants from co-cultures of

two different antibiotic-resistant isolates of the fungus: isolate Y90-71BI derived from introducing plasmid pBARKS1, a bialaphos (BI)-resistance vector that contains the Ignite/Basta-resistance (*bar*) gene<sup>1,16</sup>, into isolate Y90-71, and isolate 3514-R-2BS derived from introducing plasmid pBF101, a blasticidin S (BS)-resistance vector that contains the blasticidin S deaminase gene (*BSD*)<sup>9</sup>, into isolate 3514-R-2<sup>15</sup>. We revealed that the segregation of avirulence to virulence (*avr/vir*) in the progeny derived from crossing Y90-71 and 3514-R-2 on Hattan 3<sup>24</sup> and line K59-1 (unpublished data) was in a 1:1 ratio. The segregation of *avr/vir* in the parasexual recombinants (BI-BS-resistant parasexual recombinants) was consistent with a 1:1 ratio on line K59-1, but not on Hattan 3<sup>15</sup>. We postulated that the segregation of *avr/vir* of the BI-BS-resistant parasexual recombinants on Hattan 3 was consistent with that of sexual progeny and that the antibiotic-resistance genes and genes for virulence on Hattan 3 might be linked as a result of transformation. To eliminate the confounding effect of linkage, in this study, we compared the segregation of *avr/vir* in the BI-BS-resistant parasexual recombinants with that in the BI-BS-resistant sexual progeny derived from crossing Y90-71BI and 3514-R-2BS on Hattan 3 and line K59-1. In addition, we examined the segregation of the mating types in the BI-BS-resistant sexual progeny derived from crossing Y90-71BI and 3514-R-2BS.

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## Materials and methods

### 1. *Magnaporthe oryzae* isolates and plant materials

*Magnaporthe oryzae* isolates and plant materials used in this study are listed in Table 1.

### 2. Crosses of blast isolates

Y90-71BI and 3514-R-2BS were crossed following the procedure described by Yaegashi<sup>21</sup> with modifications. Perithecia resulting from crossing were crushed with a scalpel to release ascospores, which were transferred to potato dextrose agar (PDA) containing BI (800 µg/ml) and BS (100 µg/ml) with a pipette to obtain BI-BS-resistant sexual progeny. After incubation for 24 to 36 h, singly germinated ascospores were separated with a micro-manipulator and transferred to PDA.

### 3. Mating type determination

Mating type was determined by crossing *M. oryzae* isolates with tester isolates, Y93-164a-1 (MAT1-1, race 132) and Y93-245c-2 (MAT1-2, race 137), as described previously<sup>15</sup>.

### 4. Pathogenicity test

The pathogenicity of the BI-BS-resistant sexual progeny was determined using a previously described method<sup>15</sup>.

## Results

### 1. BI-BS-resistant sexual progeny

Seventy sexual progeny resistant to BI and BS were obtained. All grew on PDA containing both antibiotics.

### 2. Segregation of avirulence/virulence in BI-BS-resistant sexual progeny on Hattan 3 and line K59-1

Y90-71 has *Avr-Hattan 3* and was avirulent on Hattan 3, but virulent on line K59-1 carrying *Pit*. In contrast, 3514-R-2 has *Avr-Pit* and was virulent on Hattan 3, but avirulent on line K59-1. Inoculating Hattan 3 with the 70 BI-BS-resistant sexual progeny derived from the cross between Y90-71BI and 3514-R-2BS yielded 21 avirulent and 49 virulent isolates (Table 2). Inoculating line K59-1 with the 70 BI-BS-resistant sexual progeny yielded 43 avirulent and 27 virulent isolates. Inoculating Hattan 3

**Table 1.** *Magnaporthe oryzae* isolates and plant materials used in this study

Isolates, plant materials	Relevant properties	References
<i>Magnaporthe oryzae</i> isolates		
Y90-71BI	BI <sup>r</sup> , <i>Avr-Hattan 3</i> , MAT1-1, race 102	Noguchi et al. <sup>15</sup>
3514-R-2BS	BS <sup>r</sup> , <i>Avr-Pit</i> , MAT1-2, race 136	Noguchi et al. <sup>15</sup>
Y93-164a-1	MAT1-1, race 132	Noguchi et al. <sup>15</sup>
Y93-245c-2	MAT1-2, race 137	Noguchi et al. <sup>15</sup>
BI-BS-resistant parasexual recombinants	Parasexual recombinants derived from co-cultures of Y90-71BI and 3514-R-2BS BI <sup>r</sup> , BS <sup>r</sup>	Noguchi et al. <sup>15</sup>
Plant materials		
<i>Oryza sativa</i>		
Hattan 3	Japonica-type cultivar, <i>Pik-s</i>	Yasuda et al. <sup>24</sup>
Line K59-1	One of the F <sub>3</sub> lines from the cross between K59 and Norin 3, <i>Pit</i>	Noguchi et al. <sup>15</sup>

**Table 2.** Segregation of avirulence to virulence on Hattan 3 and line K59-1 in BI-BS-resistant parasexual recombinants and in BI-BS-resistant sexual progeny

Rice plants	Sexual progeny		Parasexual recombinants <sup>a)</sup>		$\chi^2$ value <sup>b)</sup>	P value <sup>c)</sup>
	A	V	A	V		
Hattan 3	21	49	12	37	0.47	0.70–0.50
Line K59-1	43	27	24	25	1.82	0.20–0.10

Pathogenicity was assayed 6 to 7 days after spraying conidial suspension on Hattan 3 and K59-1 seedlings<sup>15,24</sup> (A: avirulence, V: virulence). a): Noguchi et al.<sup>15</sup>. b), c): Expected ratio, the ratio of avirulent to virulent in the BI-BS-resistant parasexual recombinants on Hattan 3 and line K59-1 to that in the BI-BS-resistant sexual progeny.

**Table 3. Segregation of mating types of BI-BS-resistant parasexual recombinants and BI-BS-resistant sexual progeny**

	Mating type <sup>a)</sup>		$\chi^2$ value <sup>b)</sup>	P value <sup>c)</sup>
	MAT1-1	MAT1-2		
Sexual progeny	26	34	0.002	0.98–0.95
Parasexual recombinants <sup>d)</sup>	21	28		

a): Mating types were determined by crossing with tester isolates. b), c): Expected ratio, the ratio of the mating type in the BI-BS-resistant parasexual recombinants to that in the BI-BS-resistant sexual progeny. d): Noguchi et al.<sup>15</sup>

with the 49 BI-BS-resistant parasexual recombinants yielded 12 avirulent and 37 virulent isolates<sup>15</sup>. The chi-square value of 0.47 and P value of 0.70–0.50 indicated that the segregation of avirulence and virulence in the isolates fitted that in the BI-BS-resistant sexual progeny. Inoculating line K59-1 with the 49 BI-BS-resistant parasexual recombinants yielded 24 avirulent and 25 virulent isolates<sup>15</sup>. The chi-square value of 1.82 and P value of 0.20–0.10 indicated that the ratio of avirulent to virulent in the BI-BS-resistant parasexual recombinants on line K59-1 was comparable with that in the BI-BS-resistant sexual progeny. The segregation ratio of avirulence and virulence on Hattan 3 and line K59-1 in the BI-BS-resistant parasexual recombinants was similar to that in the BI-BS-resistant progeny.

### 3. Segregation of mating type of BI-BS-resistant sexual progeny of *M. oryzae*

Sixty of the 70 BI-BS-resistant sexual progeny were used for mating type determination. Of the BI-BS-resistant sexual progeny, 26 were MAT1-1 and 34 were MAT1-2 (Table 3). Of the parasexual isolates, 21 were MAT1-1 and 28 were MAT1-2<sup>15</sup>. The chi-square value of 0.002 and P value of 0.98–0.95 indicated that the segregation of the mating type in the BI-BS-resistant parasexual recombinants fitted that in the BI-BS-resistant sexual progeny.

## Discussion

We demonstrated that the segregation ratio of the genetic character on pathogenicity and the mating type of parasexual recombinants were consistent with those of sexual progeny in the rice blast fungus *M. oryzae*. Heterokaryosis and a parasexual cycle have been reported in other filamentous fungi<sup>3,5,6,8,13,14,17,18,22,25</sup>, and parasexual recombination has proven to be a valuable tool in genetic analyses of filamentous fungi, particularly imperfect fungi. For example, the placement of adenine, lysine and white conidium gene markers in the same linkage group in *Aspergillus niger* was revealed through the parasexual cycle<sup>10</sup>. The steps in the parasexual cycle are anastomosis, heterokaryosis, diploidization, and subsequently hap-

loidization. New combinations of genes arise through haploidization, when the genes are in different linkage groups or mitotic crossing-over occurs<sup>10</sup>. Therefore, parasexual recombination may give rise to the segregation of genotypes in *M. oryzae* in a recombination manner similar to sexual events<sup>15</sup>. The segregation of avirulence to virulence in the progeny derived from crossing Y90-71 and 3514-R-2 on Hattan 3 was in a ratio of 95:90, and consistent with a 1:1 ratio<sup>24</sup>, but that of the parasexual recombinants (BI-BS-resistant parasexual recombinants) on Hattan 3 was not consistent with a 1:1 ratio. These results suggested that *Avr-Hattan 3* was in a linkage group with introduced antibiotic-resistance genes. While the segregation of *avr/vir* in the progeny derived from crossing Y90-71 and 3514-R-2, in BI-BS-resistant parasexual recombinants and in the BI-BS-resistant sexual progeny was consistent with a 1:1 ratio on line K59-1, which suggested that *Avr-Pit* was not in a linkage group with introduced antibiotic-resistance genes. Parasexuality in imperfect fungi is important in maintaining genetic diversity. It appears to be an adaptation that occurs predominantly in asexual fungi, such as the rice blast fungus. In plant pathogens, a parasexual cycle might lead to variable pathogenicity, which is a critical problem in controlling plant disease.

Although many comprehensive studies on parasexuality in the rice blast fungus have been conducted<sup>3,5,6,8,13,14,20,22,25</sup>, little direct evidence exists of the parasexual cycle in nature. Nevertheless, parasexuality must be able to make a considerable contribution to genetic diversity in field populations of the rice blast fungus based on some observations<sup>2,25</sup>. Using DNA fingerprints, field populations of the rice blast fungus were not strictly clonal, and studies suggested parasexual and sexual recombination<sup>2,25</sup>. Further studies on the population structure and genetics of *M. oryzae* are needed to elucidate the role of parasexuality in nature.

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