

Fine mapping of a strong QTL of field resistance against rice blast, *Pikahei-1(t)*, from upland rice Kahei, utilizing a novel resistance evaluation system in the greenhouse

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Abstract Field resistances (FR) against rice blast are highly evaluated by breeders for their durability, in contrast to the conspicuous but often less durable true resistances. However, lack of efficient systems for evaluation of resistance has delayed their practical application. Kahei, an upland domestic cv., is known for its very high FR against rice blast. We fine-mapped its highest quantitative trait loci (QTL), *qBFR4-1*, using residual heterozygosity of recombinant inbred lines (RILs) and our semi-natural rice blast inoculation/evaluation system in the greenhouse, with comparable accuracy to the true resistance genes. This system enabled reproducible high-density infection, and consequently allowed quantification of the resistance level in individual plants. The target region was first narrowed down to about 1 Mb around at 32 Mb from the top of chromosome 4 in the Nipponbare genome, with the upland evaluation system assessing the F₇ generation of Koshihikari (lowland, FR: very weak) × Kahei (upland, FR: very strong)

RILs. Then, F₉ plants (4,404)—siblings of hetero F₈ plants at the region—were inoculated with rice blast in a greenhouse using the novel inoculation system, and individual resistance levels were diagnosed for fine QTL analysis and graphical genotyping. Thus, the resistance gene was fine-mapped within 300 kb at 31.2–31.5 Mb on chromosome 4, and designated *Pikahei-1(t)*. By annotation analysis, seven resistance gene analog (RGA) ORFs of nucleotide-binding-site and leucine-rich-repeat (NBS-LRR)-type were found in the center of the region as the most likely candidate counterparts of the resistance gene. This is similar in structure to the recently reported *Pik* cluster region, suggesting that most of the other dominant QTLs of the FRs may have similar RGA structures.

Introduction

As disease control of crops is now being required to reduce both environmental and labor costs, as well as to secure food safety, breeding control with disease resistance genes seems to have better prospects than chemical control, although even the latter is also shifting toward the use of inducers of plant resistance, such as probenazole (Iwata et al. 2004). Resistance against plant diseases can be classified roughly into two types: complete (qualitative) or true resistance and partial (quantitative) or field resistance (FR) (Ezuka 1972), although these respective groups of definition are not completely overlapping. These resistances were studied extensively in rice against its most serious disease, rice blast disease caused by *Magnaporthe oryzae*. True resistance is effective in a very clear-cut way against blast isolates with the corresponding avirulence gene, but often very vulnerable to mutations of that avirulence gene, especially in modern large scale monoculture systems. On the

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other hand, in most cases although FR allows slight blast infection, it shows effective suppression of outbreaks and such varieties are highly valued for their durability (Bonman 1992).

To attain durability, breeding with true resistances genes requires rather complicated strategies, such as multi-line or gene pyramiding. Although the Koshihikari-BL multi-(NIL-)line project, which began in 2005 (Ishizaki et al. 2005) and covers the whole of Niigata prefecture, is having good success at present, it requires a well-organized breeding and race monitoring system and complicated line management, and it is still too early to draw definitive conclusions regarding its success. Most FR consists of multiple quantitative trait loci (QTLs), and reconstruction of FR with their combinations is difficult even with the aid of molecular markers because of the absence of an efficient and accurate assay system. Although there have been some reports of greenhouse assays of FR (Hayashi 2005), the main system used for assaying FR is field diagnostic testing, which requires a large area for replication and can be performed only once a year. Therefore, fine mapping or cloning is difficult, and to date only one recessive blast FR gene has been cloned (Fukuoka et al. 2007).

“Kahei” is a domestic upland rice cv. found in southern Kyushu, which has very strong FR. Analysis of F_2 plants with F_3 progeny of Kahei \times Koshihikari identified *qBFR4-1* as the most highly contributing QTL of rice blast resistance located on the long arm of chromosome 4 (Miyamoto et al. 2001).

As a first step in its fine mapping, we analyzed a sibling population from those with residual heterozygosity in the QTL region in recombinant inbred lines (RILs), using the upland late-sowing field assay and localized it within about 2.2 Mb. This gene was tentatively named *Pikahei-1(t)*, because of confusion regarding the still-formal numbering system of *Pi* genes. To increase the efficiency and accuracy of the mapping data, we utilized our newly developed blast resistance assay system in the greenhouse (Kawasaki et al. 2008), mimicking the natural infection, by placing the testing plants for 4–7 days in an inoculation tent in a greenhouse, with diseased plants as the source of spores. With this system, we could diagnose the symptoms or phenotypes of the individual plants in ten grades, and QTL analysis was performed. Further refining of the conditions allowed the graphical genotyping of close recombinant individuals. Using this novel resistance evaluation system in 4,404 F_8 heterozygous siblings of RI lines in the greenhouse, we could narrow down the *Pikahei-1(t)* gene region to within \sim 300 kb. The cluster of NBS-LRR type RGAs found in the center of the region was consistent with our hypothesis that identity of the most of FR components may be the weak type of these resistance (R-) genes.

Materials and methods

Plant and fungal materials

A set of F_4 RILs of Koshihikari (lowland, FR very weak) \times Kahei (upland, FR very strong) with heterozygosity only in the strongest QTL *qBFR4-1*, and homozygous in other QTL, *qBFR4-2*, and *qBFR4-3* (Miyamoto et al. 2001), along with the parental strains Kahei and Koshihikari were kind gifts from Dr Miyamoto and Dr Hirasawa of the Agricultural Research Institute, Ibaraki Agricultural Center (Mito, Ibaraki, Japan). From the F_6 generation, siblings of the above RILs, which were confirmed to be heterogeneous at *qBFR4-1*, a total of 400 F_7 lines were obtained, and these lines were planted in a segregated upland field in June 2003 at the National Institute of Agrobiological Sciences (NIAS) Tsukuba, Japan, for field diagnosis as described in the following section.

For greenhouse diagnosis, the F_7 lines segregating resistant/susceptible on upland field, set on heterozygous F_6 at *qBFR4-1*, were grown in a lowland field to propagate F_8 seeds. For genotyping of *qBFR4-1*, about 60 F_8 lines (34 seeds each; F_9 generation) were sown in the grain-seeding mats of a 17×34 hole matrix with resistant (Kahei) and susceptible (Koshihikari) reference lines, and checked for the line's heterozygosity in the greenhouse, as described in the following section. The seeds (F_9) of the hetero-confirmed F_8 lines were later utilized for large-scale screening of close recombinants around the target *qBFR4-1* locus. Standard cv. Nipponbare was obtained from the NIAS Gene Bank.

The rice blast isolate Kyu9439013 (047.0) was a kind gift from Dr N. Hayashi of NIAS.

Diagnosis of blast resistance in the field

Field resistance diagnosis was first made in the segregated upland field by late sowing of the seeds around mid-June. Field diagnosis of the RI lines of F_6 to F_7 generations was done as follows. For each F_6 individual thought to be heterozygous at *qBFR4-1*, around 20 lines F_7 were selected and their seeds (about 20 grains of F_8 for each F_7 line) were sown in a line 40 cm in length in the segregated upland field as shown in Fig. 1. The line interval was 20 cm, and each of 20 lines was accompanied by reference lines of Kahei (R resistant) and Koshihikari (S susceptible). This set of 22 lines constituted a unit of testing lines, roughly corresponding to an F_6 parent. Four such sets comprised one row and the total field accommodated about ten rows. The test was performed with two repeats (in two population), i.e. 20 sets of F_6 corresponding to 400 lines of F_7 were tested in duplicate; in total, 800 of F_8 lines were tested. As spore spreaders, a line of Koshihikari was placed

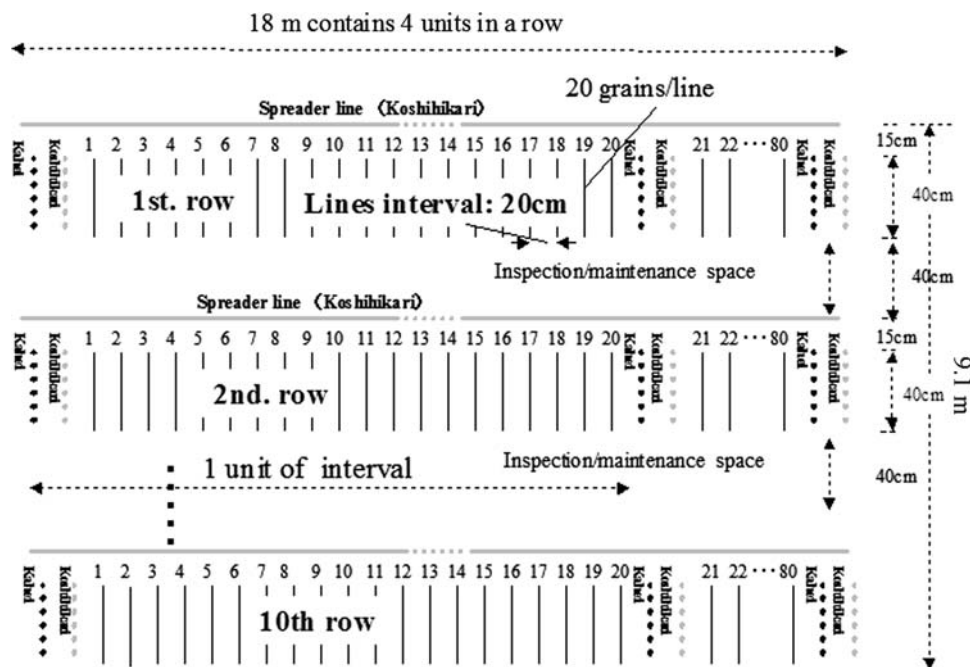


Fig. 1 Field rice sowing scheme for diagnosis of field resistance in a segregated upland field. North is to the top of the figure. In mid-June 2003, samples of about 20 grains of each F_8 generation were sown in a line 40 cm in length. Each line represents the sibling population of an F_7 plant. Diagnoses were made for these F_7 lines and not for the F_8 individuals, because the susceptibility was discernible only as lines. Each row was from four units, consisting of 20 test lines and R (Kahei) and

S (Koshihikari) reference lines at the ends. Each row was accompanied with a spore spreader line of Koshihikari of about the same density as the test lines. Ten rows were surrounded by three lines of Koshihikari spreaders. No artificial spreading of blast was done. The diagnosis was performed three times around the seed maturation time at half-month intervals

at the north rim of each testing row, apart 15 cm from the rim, and the total field was circled with three lines of Koshihikari spreaders. By diagnosing the symptoms of the unit lines of F_7 by over-viewing three times from September to October, the heterozygous F_6 sets were identified as a population segregating R and S in F_7 lines. Only the obviously susceptible F_7 lines in the segregating F_6 sets, and all-susceptible F_6 sets were selected for DNA analysis for fine mapping. In this protocol, the plant density was lower than in conventional field diagnosis methods. Therefore, the seeds of the important lines were harvestable in autumn even from the susceptible lines.

The leaves of apparently susceptible lines were harvested and their DNA was extracted using a Kurabo NA2000 DNA isolation machine (Kurabo, Osaka, Japan), and then the genotypes of the neighboring DNA markers were analyzed with the High Efficiency Genome Scanning (HEGS) system (Nihon Eido, Tokyo, Japan) as described previously (Xu et al. 2005).

Diagnosis of blast resistance in the greenhouse

Greenhouse diagnosis of blast resistance was performed according to the newly developed semi-natural inoculation system for screening of blast-resistant mutants (Kawasaki

et al. 2008), in a greenhouse with a controlled temperature of 17–27°C with a day length of 14 h supplemented with fluorescent lamps in winter. For inoculation, 2- to 3-week-old seedlings, prepared in grain-seeding mats, were covered in a tent for 4–6 days with a blast-infected seedling mat as a source of inoculating spores at the center, in a pool accommodating up to nine such mats. Daily temperature fluctuation of more than 5–7°C was important to secure sufficient dew formation at night. After inoculation, the tent was removed and seedlings were grown under the same conditions for a further 2–3 weeks to allow symptoms to develop fully. The inoculated plants were diagnosed on a scale of 0–10 by evaluating symptoms as described in the manual by Hayashi (2005) or as shown in Table 1 around 20 days after inoculation. Conventional spray inoculation as a control was performed according to the standard protocol (Hayashi 2005) for seedlings of the same age as above at a conidia concentration of 3×10^5 /ml with incubation for 24 h in a humid chamber at 25°C at a relative humidity of 100%, but diagnosis was made at 3 weeks after inoculation.

The F_9 generation seedlings obtained from heterozygous F_8 lines, which were confirmed by checking the genotypes of flanking markers and segregation of the resistant and susceptible individuals, were inoculated with the rice blast race Kyu9439013 (047.0).

Table 1 Diagnosis table of the symptoms of blast-infected individual rice seedlings 3 weeks after inoculation with semi-natural spores spread for 5 days in a humidity-saturated tent

Scores	Symptoms of the plants	Diagnosis
0	No blast lesions	
1	Small brown specks the size of pinpoints were observed	
2	Small brown specks were observed	Resistant
3	Significant numbers of small brown specks were observed	
4	Larger brown specks were observed	
5	Significant numbers of larger brown specks were observed	
6	Large progressive blast lesions were observed	
7	Leaf tips dying with blast	
8	Single leaf of a plant died with blast	Susceptible
9	A few leaves of a plant died with blast	
10	Almost the whole plant died	

The 3-week-old rice seedling mats in a water-pool were covered with a transparent vinyl or polyethylene tent with a mat of diseased plants as the inoculation seeds. The level of infection was diagnosed 3 weeks after inoculation. Symptom grades 1–6 were determined by the infected leaves that had developed during the period of inoculation, and grades 7–10 were analyzed by the symptoms of whole diseased plants at a glance. The threshold for diagnosis as R or S, here between 5 and 6, was selected to be consistent with the results of graphical genotyping

Genotyping

DNA was extracted from leaves 2 cm in length from seedlings at the age of approximately 1 week after sowing by a high-throughput CTAB method (Xu et al. 2005). From the selected close recombinants, samples of approximately 5 µg of DNA were re-extracted (1 g fresh weight) using a Kurabo NA2000 DNA isolation machine. Thirteen SSR markers from IRGSP (The International Rice Genome Sequencing Project 2005) and Gramene (<http://www.gramene.org/>) were analyzed with the HEGS system (Nihon Eido, Yushima, Tokyo, Japan) as described previously (Xu et al. 2005), and six single nucleotide polymorphism (SNP) markers were subjected to single strand conformation polymorphism analysis in an HEGS system (HEGS-SSCP; Xu et al., in preparation) setting the gel running temperature at 6–8°C with cooling jackets (Nihon Eido) connected in tandem to a water cooler/circulator.

Fine QTL analysis

Quantitative trait loci analysis was performed with Windows QTL Cartographer V2.5 (Wang et al. 2006) via single marker diagnosis of the leaf symptoms as grade 0–10 described in Table 1. To clarify the fine structure of QTL, data of the close recombinants between markers flanking the target gene were used. The map addresses in megabases were taken from the marker location on chromosome 4 from IRGSP and Gramene. A minimum LOD (logarithm of the odds) threshold of 2.5 was used to indicate significance.

Physical map construction

The physical locations of SSR markers and ORFs were determined in the Gramene marker database (<http://www.gramene.org/>) and RiceGAAS analysis in RGP (<http://ricegaas.dna.affrc.go.jp/>), respectively, in correlation with the close recombinant map constructed as described above and graphical genotyping of the markers. The physical map was constructed on the basis of that of Nipponbare from IRGSP (<http://rgp.dna.affrc.go.jp/IRGSP/>), again in correlation with the results of close recombinants analyses. The ORFs were also localized on the physical map if SNPs were detected between the parents, to correlate the results with the graphical genotypes.

Construction of fine SNP markers

Within the physical map, novel markers were constructed by checking the target bacterial artificial chromosome (BAC) clones or ORF sequences in IRGSP (*ssp. japonica* cv. Nipponbare, <http://rgp.dna.affrc.go.jp/IRGSP/>) by BLAST analysis (<http://www.ncbi.nlm.nih.gov>) against the complete rice genome sequence. Sequences found to be unique in the genome were used to design primers with Primer3 (http://fokker.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi). Rather longer PCR product size, around 1.5 kb, was preferred to detect more SNPs. The uniqueness of each primer in the genome was again confirmed by BLAST analysis.

PCR fragments amplified from the parent cvs. and confirmed to be single-banded on 1% agarose gels were

directly sequenced using an ABI BigDye[®] Terminator v3.1 Cycle Sequencing Kit and ABI 3100 DNA sequencer (Applied Biosystems, Foster City, CA, USA) to detect SNPs between parents. The confirmed SNPs were selected to redesign HEGS-SSCP primers with Primer3 to obtain a suitable PCR product size (100–300 bp). As far as possible, the SNP was positioned in the center although we did not observe a significant “position effect” during HEGS-SSCP electrophoresis.

Results

Mapping of *qBFR4-1* by upland field evaluation of the heterozygosity-remaining population at the locus

The most contributing locus to blast resistance in Kahei, *qBFR4-1*, was fine-mapped with an upland late sowing field test. Seeds of the F₈ generation from 400 F₇ lines obtained from 20 parents (F₆ generation) with residual heterozygosity at *qBFR4-1* but homozygous (mostly susceptible) at other QTLs, *qBFR4-2* and *qBFR4-3*, were sown in duplicate as shown in Fig. 1 in a segregated upland field for blast resistance evaluation. The sowing protocol was sparser than in the conventional system (Ezuka 1972) to secure harvesting of seeds even from the susceptible lines. The phenotypes of the F₇ lines were diagnosed by overviewing the corresponding lines in the field, and 96 clearly susceptible lines were checked for their DNA markers around the *qBFR4-1*. From the results of graphical genotyping of the close recombinants around the target region, the target locus was narrowed down to a region of 32–33 Mb of Chromosome 4 (<http://www.gramene.org>), close to the end of the long arm, in the Nipponbare physical map (refer to Fig. 3). However, further narrowing down of the locus was difficult because of the unstable field condition in the subsequent years, probably due to repeated upland cultivation of rice. As these mapping data indicated that this blast resistance was due to a single resistance gene, it was named *Pikahei-1(t)*—the *Pi-numbering* system was avoided due to present confusion due to several uncontrolled gene reports and lack of information of the gene origin and their clustering.

Diagnosis with a semi-natural blast spore inoculating system in a greenhouse

To increase the accuracy of mapping for gene cloning, a novel greenhouse blast semi-natural inoculation and diagnosis system, developed for recessive mutant screening (Kawasaki et al. 2008), was applied for fine mapping of *Pikahei-1(t)*. The main difficulty in diagnosing field or quantitative resistance against rice blast in the greenhouse by conventional spray inoculation with conidia suspension

is that the reproducing high density of lesion development is often difficult even with a concentrated conidia suspension. The novel system described here mimics natural infection by covering the test seedlings with a lesion-developing rice mat as a spore source in a tent for 5–7 days in a greenhouse. With this system, a far great number of lesions developed per leaf within a few weeks from the start of covering (Fig. 2), thus enabling fine quantification of the blast resistance of the rice strains. Only covering in a tent without diseased mat after spray inoculation resulted in far less lesion numbers especially on the main target of 3rd–4th leaves, less than even the conventional humid chamber inoculation, indicating that natural conidia spreading is the key source of infection (Fig. 2).

For fine mapping of the *Pikahei-1(t)* gene, siblings of the residual heterozygotes at this locus were tent-inoculated as described above with the diseased seedling mat infected with the race Kyu9439013. Twenty-one days after commencement of infection, the symptoms of the leaves were diagnosed on the original scale shown in Table 1, which was constructed from observation of these inoculated populations. The phenotype was clear and consistent with the results of genotyping of markers close to the target region. This system was both consistent and efficient, and valid even in the middle of summer or winter, seasons that are not optimal for conventional spray inoculation, even with the presence of air conditioner.

From the lowland field-propagated plants (F₈), lines heterozygous for *Pikahei-1(t)* were selected in this system as lines with segregation of susceptible individuals at a ratio of 1/4. The siblings from these confirmed heterozygous F₈ lines, a total of 2,965 seedlings (F₉), were used for diagnosis of the phenotypes and genotyping of the neighboring markers. These lines were mostly homozygous in the genome except the target region, enabling stable phenotype assessment.

Fine QTL map and physical map around *Pikahei-1(t)*

To increase the accuracy of the target region, QTL analysis was performed for the 616 individuals among the above 2,965 F₉ lines with recombination between the distant flanking markers RM6748 and RM1153, identified from field diagnosis as described earlier. QTL analysis with the data obtained as described above revealed a sharp single peak around 31.2–31.5 Mb on chromosome 4 between the markers RM 17496 and RM6629, and RM5473 at the crest; peak LOD score of 33 and half height width of about 1 Mb (Fig. 3). This was also consistent with the expectation that a single gene is responsible for the resistance. The asymmetry of the peak was due to the occurrence of a recombination event on the left side some generations prior to the diagnosed population. Despite selecting the recombined

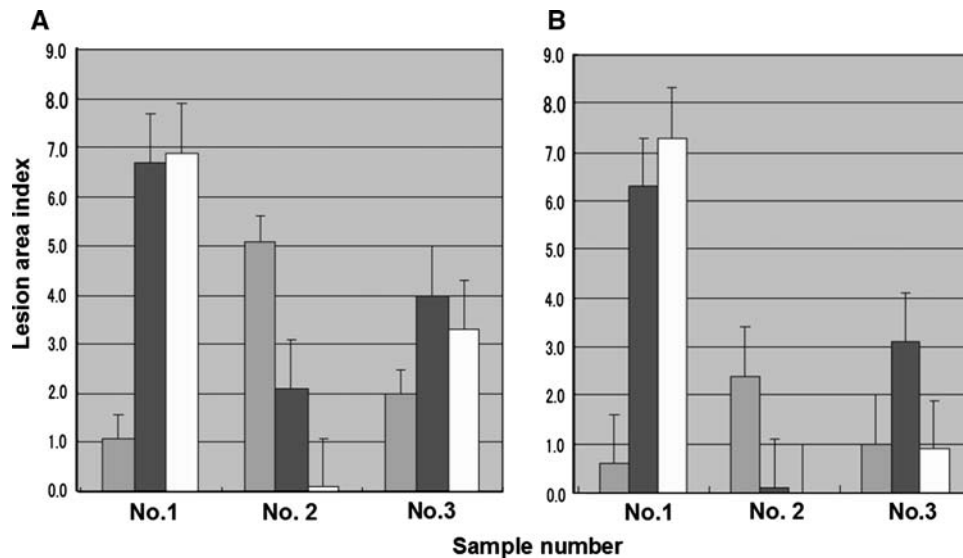


Fig. 2 Comparison of leaf blast development in rice seedlings inoculated with the novel semi-natural infection system and conventional spray inoculation. Rice seedlings in the sowing mats were inoculated with the different systems at 2 weeks of growth in a greenhouse after sowing, and the symptoms were checked on the 3rd week. The lesion area index was determined with a field resistance diagnosis system (0 no lesion to 10 died out) as described in the Hayashi manual (2005) for the 2nd, 3rd, and 4th leaves, and the results are shown in histograms with standard deviation bars of left (*gray*), center (*black*), and right

(*white*), respectively, in each group. Blast fungal race was Kyu9439013 (047.0). A: cv. Koshihikari. B: cv. Nipponbare. Sample No. 1: novel semi-natural inoculation system, 3 weeks from starting of 5 days of covering in the inoculation tent with the lesion-developed plants as an inoculation seed. No. 2: spray inoculated as in no. 3, but kept in the inoculation tent as no. 1 without diseased plants. No. 3: infected by the conventional spray inoculation system with a suspension of 3×10^5 conidia/ml, and kept for 24 h in the humid chamber

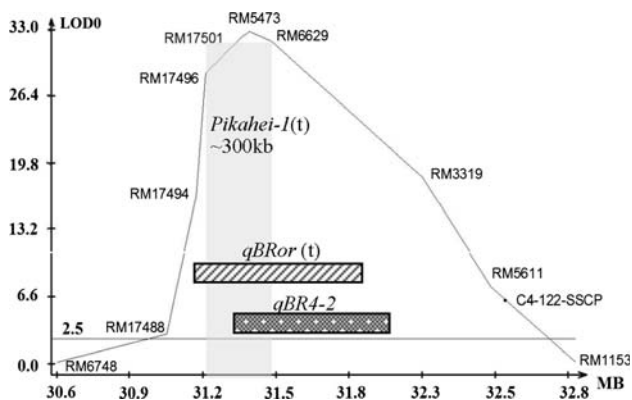


Fig. 3 Fine QTL map around *Pikahei-1(t)* from analysis of 616 recombinants between the flanking markers RM6748 and RM1153, selected from a population of 2,965 F_9 siblings of heterozygous F_8 plants in this region, using the single marker analysis method with WinQTL-Cart 2.5. A single strong QTL peak of *Pikahei-1(t)* with LOD score 33 and half height width of about 1 Mb was observed. The peak region indicated by the *gray rectangle* is close to the blast resistance QTL from *O. rufipogon* (adapted from Hirabayashi et al. 2005 and named as *qBRor(t)* although referred by them as *Pi37(t)*) and *qBR4-2* from Owarihatamochi (adapted from Fukuoka et al. 2003) on the long arm of chromosome 4. The asymmetry of the peak is due to the occurrence of a close recombination event on the *left side* some generations before

population, the LOD score was comparable to that of the original data of *qBFR4-1* of Kahei reported by Miyamoto et al. (2001). This region seems to be a blast R-gene-rich cluster, including *qBFR4-2* of Owarihatamochi (Fukuoka

et al. 2003) and a QTL from the *Oryza rufipogon* referred as *Pi37* by Hirabayashi et al. (2005), indicated as *qBRor(t)* in Fig. 3.

Graphical genotyping

Although graphical genotyping with the above close recombinants was mostly consistent with the results of QTL analysis, there were slight contradictions in phenotypes (data not shown). Therefore, an additional experiment was performed with more refined conditions of elongated infection period and temperature control with 1,495 hetero-siblings of the F_9 generation. A very consistent graphical genotype was constructed based on the results of this experiment, as shown in Fig. 4. From these data, the *Pikahei-1(t)* gene was delineated again between the markers RM17496 and RM6629, separated by a distance of about 300 kb, in the Nipponbare genome. Considering the genotypes of the close marker genotypes, the threshold value for diagnosis as S or R was set between 5 and 6. The results of diagnosis were highly consistent with the neighboring DNA markers around the *Pikahei-1(t)*, even at the individual level. Only a very slight inconsistency in the phenotype was observed for a single close recombinant, 13–28, but other three recombinants at the same site showed a consistent phenotype (Fig. 4). Although from Fig. 4, this diagnosis seems to differentiate the R-homo individuals from the hetero

Markers	RM 17488	970-60K2	RM 17494	RM 17496	970-149K	N-SNP	RM 17501	Symptom	645-80K	RM 5473	RM 3843	RM 6629	669-40K	RM 3319	RM 5611
1--2	R	H	H	H	H	H	H	5	H	H	H	H	H	H	H
1--3	R	H	H	H	H	H	H	5	H	H	H	H	H	H	H
7--22	H	R	R	R	R	R	R	2	R	R	R	R	R	R	R
1--50	S	H	H	H	H	H	H	4	H	H	H	H	H	H	H
4--12	S	H	H	H	H	H	H	4	H	H	H	H	H	H	H
14--55	S	H	H	H	H	H	H	2	H	H	H	H	H	H	H
3--42	S	S	S	H	H	H	H	5	H	H	H	H	H	H	H
5--37	S	S	S	H	H	H	H	3	H	H	H	H	H	H	H
6--6	S	S	S	H	H	H	H	5	H	H	H	H	H	H	H
4--42	R	R	R	H	H	H	H	5	H	H	H	H	H	H	H
12--29	R	R	R	H	H	H	H	3	H	H	H	H	H	H	H
2--94	H	H	H	S	S	S	S	7	S	S	S	S	S	S	S
3--79	H	H	H	S	S	S	S	8	S	S	S	S	S	S	S
4--44	H	H	H	S	S	S	S	6	S	S	S	S	S	S	S
4--88	H	H	H	S	S	S	S	8	S	S	S	S	S	S	S
7--77	H	H	H	S	S	S	S	8	S	S	S	S	S	S	S
8--40	H	H	H	S	S	S	S	6	S	S	S	S	S	S	S
12--39	H	H	H	S	S	S	S	8	S	S	S	S	S	S	S
3--56	H	H	H	H	S	S	S	8	S	S	S	S	S	S	S
6--47	H	H	H	H	S	S	S	7	S	S	S	S	S	S	S
5--64	H	H	H	H	R	R	R	1	R	R	R	R	R	R	R
6--58	R	R	R	R	H	H	H	5	H	H	H	H	H	H	H
1--76	S	S	S	S	H	H	H	5	H	H	H	H	H	H	H
7--93	S	S	S	S	H	H	H	4	H	H	H	H	H	H	H
13--28	S	S	S	S	H	H	H	6	H	H	H	H	H	H	H
9--70	S	S	S	S	H	H	H	4	H	H	H	H	H	H	H
9--87	H	H	H	H	H	R	R	2	R	R	R	R	R	R	R
7--13	R	H	H	H	H	H	H	5	H	H	H	S	S	S	S

Fig. 4 Graphical genotype of the close markers around the *Pikahei-1(t)* region, and the blast diagnosis scores of the close recombinants, selected from 1,439 siblings of the neighboring locus heterozygous plants, obtained by the tent inoculation and greenhouse evaluation system. This enabled diagnosis at the level of single plants, and even graphical genotyping of the close markers in close recombinants for this field resistance component. Symptom scores of six or more and of five or less were considered S and R, respectively, and were mostly consistent with the results of genotyping of the close markers. The

symptoms also seemed to be dose-related, and a score of two or less seemed to correspond approximately to R homozygotes. The margins of recombination are indicated with a *bold line*. The region of *Pikahei-1(t)* is also marked with a *bold rectangle* covering the central markers and their genotypes. The heterozygotes and susceptible homozygotes are shown in light and dense hatching, respectively, including the corresponding symptom scores. The validity of diagnosis of the *critical line* 9–87 as R-homo requires confirmation in the next generation

individuals, with threshold between 2 and 3, verification of the critical recombination of 9–87 may require additional experiments.

Physical mapping of *Pikahei-1(t)* in Nipponbare genome

To physically map the *Pikahei-1(t)* locus in the Nipponbare genome, the above identified close SSR markers were “landed” on three BAC clones, OSJNa0058K23, OSJNb0085C12, and OSJNa0053K19, in the Gramene and IRGSP database (Fig. 5). With annotation by GAAS analysis, 52 possible ORFs were identified in the region (Table 2). These included seven resistance gene analogs (RGAs) with an NBS-LRR domain. By PCR amplification of their counterparts from the Kahei genome, only RGAs 2 and 3 were properly amplified as full ORFs, and were located in the central region of graphical genotyping. Counterparts of RGAs 1, 4, and 6 were partially amplified, but their graphical genotypes were not related to the *Pikahei-1(t)* region (data not shown). RGAs 5 and 7 were not amplified from

the Kahei genome. The functions of other ORFs were mostly unknown, and only three were probably transposon-related and were excluded from consideration. Although the RGAs are the most likely candidates, all the possibilities should still be considered, as the dominant FR gene has yet to be cloned.

Discussion

Field resistance, also known as quantitative resistance, especially against rice blast, has been studied and attempts have been made to introduce FR into practical cvs., especially by rice breeders, to achieve durability and wide-spectrum race specificity (Bonman 1992). However, the molecular identification of FRs as genes and introduction into elite cvs. have been hindered by their multigenicity and lack of methods to fine map the respective constitutive components. Even after dissecting their components genetically by QTL analysis of RI lines, fine mapping was difficult

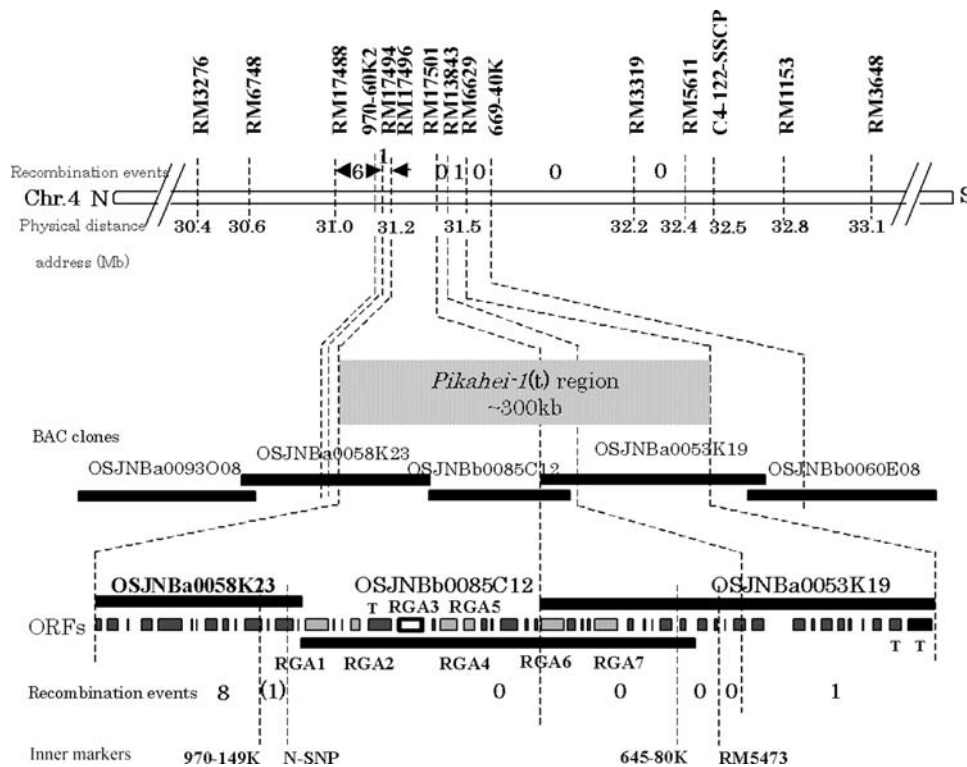


Fig. 5 Physical maps around the *Pikahei-1(t)* region on the long arm of chromosome 4 based on the Nipponbare RGP database. Marker names and recombination event numbers in the data, as presented in Fig. 4, are shown above the chromosome bar. Addresses on chromosome 4 in physical distances from the *top* are indicated below the chromosome bar in megabases (Mb). The details of the *Pikahei-1(t)* region in the Nipponbare physical map are shown at the *bottom* with 52 predicted ORFs, among which the transposon-related three ORFs are indicated by T on the *right* end. The new markers in the *Pikahei-1(t)* region at the bottom were developed from SNPs between Nipponbare, as

shown in Table 3, and the possible recombination event estimated from 9-87 diagnosed as R-homo in Fig. 4 is shown in *parentheses*. All annotations of the ORFs and corresponding RGAs are shown in Table 2. There were seven candidate RGAs (RGA 1–7) as possible counterparts of the actual *Pikahei-1(t)* in the resistant cv. genome. By graphical genotyping of the RGAs with other markers, only parts of RGAs 2 and 3 were shown to be located in the *Pikahei-1(t)* region and are shown by *blank boxes*, and RGAs 1, 4, and 6 were assigned outside the region. RGAs 5 and 7 were not properly amplified in the Kahei genome

with the conventional upland late sowing method, because it requires large areas of special isolated fields for diagnosis, and the number of lines that can be diagnosed in several replicates is still limited; especially, field diagnosis can only be performed once a year.

We attempted to fine map the most highly contributing FR QTL of domestic upland rice cv. Kahei in Kyushu, *qRBF4-1* (Miyamoto et al. 2001) first by the near conventional upland late sowing method using the RI lines with residual heterozygosity at *qRBF4-1*, but homozygous, mostly susceptible, with the other QTLs. By selecting 96 susceptible F_7 lines from this field analysis, the tentative locus was assigned to an area of 32–33 Mb from the top on chromosome 4, near the end of the long arm, but a yearly repeat to confirm the result was hindered by poor field conditions, seemingly due to long repeated upland cultivation of rice. Therefore, we applied a new greenhouse diagnosis system to evaluate FR against rice blast. This is a half-natural blast inoculation system in the greenhouse developed for screening recessive blast resistant mutants (Kawasaki

et al. 2008) from culture-mutated M_2 rice plant populations (Miyao et al. 2003). Seedlings 2–3 weeks old were semi-naturally infected from diseased plants within a tent with sufficient dew formation caused by daily fluctuation of temperature of >5 – 7°C . Spores showed sufficiently uniform dispersal without displacing the seedling mats during the covering time. This system worked very consistently and reproducibly, even in the winter or summer both of which are normally avoided as non-optimal seasons for blast inoculation, if the greenhouse temperature control can provide the prescribed conditions. In these seasons, it is often difficult to achieve reproducible inoculation using a conventional spray inoculation system even in well-equipped greenhouses and with a good humid chamber.

As shown in Fig. 2, this inoculation system can provide very high-density lesions by increasing the inoculation period (5–7 days) with a high degree of reproducibility, often difficult in the conventional spray inoculation system. It is possible to achieve any level of infection of the test plants by changing the inoculation (tent-covering) period.

Table 2 Annotation data of the putative ORFs in the *Pikahei-1(t)* region in the Nipponbare genome database

ORF no	Proteins coded by predicted genes	Predicted length (bp)
1	Unknown protein	208
2	Putative F-box family protein/WD-40 repeat family protein	684
3	Hypothetical protein	28
4	Putative ATBAG1 (ARABIDOPSIS THALIANA BCL-2-ASSOCIATED ATHANOGENE 1); protein binding	504
5	Putative Canalicular multispecific organic anion transporter 1 (ATP-binding cassette sub-family C member 2) (multidrug resistance-associated protein 2) (canalicular multidrug resistance protein) (epithelial basolateral chloride conductance regulator)	1,636
6	Hypothetical protein	79
7	Hypothetical protein	35
8	Putative remorin family protein	659
9	Hypothetical protein	187
10	Hypothetical protein	27
11	Putative SIT4 phosphatase-associated family protein	714
12	Unknown protein similar to Os04g0620600	93
13	Putative tobacco nucleolin	729
14	Hypothetical protein similar to Os04g0620800	117
15	Putative NBS-LRR disease resistance protein homolog (RGA1)	1,435
16	Hypothetical protein	51
17	Unknown protein	162
18	Putative NB-ARC domain-containing protein, expressed (RGA2)	376
19	<i>Putative MuDR family transposase domain containing protein</i>	1,357
20	Putative NBS-LRR disease resistance protein homolog (RGA3)	1,716
21	Putative polyprotein	125
22	Putative XA1 (RGA4)	1,084
23	Putative XA1 (RGA5)	648
24	Unknown protein	339
25	Hypothetical protein	120
26	Putative SWIM zinc finger family protein	1,152
27	Hypothetical protein	137
28	Hypothetical protein	121
29	Putative XA1 (RGA6)	1,886
30	Unknown protein similar to PREDICTED OSJNBa0056006.23 gene product	303
31	Hypothetical protein	148
32	Unknown protein similar to CDR1	193
33	Putative XA1 (RGA7)	1,822
34	Unknown protein similar to PREDICTED OSJNBa0056006.23 gene product	457
35	Unknown protein	147
36	Hypothetical protein	12
37	Putative amine oxidase family protein	502
38	Hypothetical protein	103
39	Putative exonuclease family protein	261
40	Hypothetical protein	41
41	Hypothetical protein	73
42	Putative (S)-2-hydroxy-acid oxidase (EC 1.1.3.15)	319
43	Putative (S)-2-hydroxy-acid oxidase (EC 1.1.3.15)	429

Table 2 continued

ORF no	Proteins coded by predicted genes	Predicted length (bp)
44	Putative Aminomethyltransferase, mitochondrial precursor (Glycine cleavage system T protein) (GCVT)	355
45	Hypothetical protein	54
46	Unknown protein	396
47	Hypothetical protein	177
48	Putative microtubule-associated protein 1 light chain 3	120
49	Hypothetical protein	74
50	Hypothetical protein	434
51	<i>Putative gypsy-type retrotransposon RIRE2 protein</i>	833
52	<i>Putative gag-pol precursor</i>	1,998

Although Nipponbare does not have a functional *Pikahei-1(t)* gene, there are seven probable counterpart RGAs, indicated by bold letters, and numbered in parentheses corresponding to Fig. 5. The transposon-related proteins in italics were excluded from further consideration

Table 3 PCR primer sequences of newly developed SNP and HEGS-SSCP markers near *Pikahei-1(t)* for the Kahei/Nipponbare map

Primer	Forward	Reverse
970-60K2	5'CCACGAAAAACGGAATAAGG3'	5'GCAAATGAAGCACAAAAGCA3'
970-149K	5'TTTGTTGAAGCTGCTGGATG3'	5'TATGCACTTTGTGGCTCGAC3'
N-SNP	5'TTCAGCGGTGGAAGAGACTT3'	5'ATCAAGTCCAAGCAGCGAGT3'
645-80K	5'TTGGTTTCATGCAGAAGCAG3'	5'GTAGCGGCTAGCCATGTTGT3'
669-40K	5'AAGGGACCGTGTGCTTAGTG3'	5'CCAGATCTCCCTATGCTTCG3'
C4-122-SSCP	5'GACGACAAGTCCAGTTGCATT3'	5'TCACCAGTTCCAAACTCG3'

Their locations are indicated in Fig. 5. They were developed from sequences of Nipponbare

This allows quantification of the disease resistance of the strains or even of individual plants, as shown in Fig. 4, which is difficult with the standard spray inoculation method. Thus, application of this system for *Pikahei-1(t)* enabled diagnosis at the level of single plants, and the mapping resolution could be increased in proportion to the sample numbers. Therefore, QTL analysis (Fig. 3) and graphical genotyping (Fig. 4) were possible in the greenhouse system. Thus, although recombination around the target gene was suppressed, with 1439 samples the *Pikahei-1(t)* region was narrowed down to within a region of 300 kb (Fig. 5), around 31.2–31.5 Mb on chromosome 4 from the top end. This region was shifted slightly from the above assigned region based on limited field assay data, suggesting insufficient accuracy in the limited sampling diagnosis. This region is close to other rice blast FR QTLs: one from *O. rufipogon* and *qBR4-2* from Owarihatamochi. These genes probably form a cluster as in the case of other blast resistance genes, such as *Pik* and *Pita* groups.

Previously, the number of plants in study populations for rice FR QTL analyses has not exceeded about 300 (Wang et al. 1994; Chen et al. 2000; Bagali et al. 2000; Miyamoto et al. 2001; Fukuoka and Okuno 2001; Sirithunya et al. 2002; Zenbayashi et al. 2002; Fukuta et al. 2004; Tabien et al. 2002; Liu et al. 2004; Talukder et al. 2004, 2005; Wu

et al. 2005; Nguyen et al. 2006; Sato et al. 2006). Here, we analyzed *Pikahei-1(t)* with more than ten times the number of samples than conventional methods in a very short time (~5 weeks), although this was partly due to the strong resistance conferred by *Pikahei-1(t)*. This system will be applicable even for weaker FR QTL analyses and for breeding, using line evaluations in the seedling mats in combination with QTL analysis.

A physical map of the region around *Pikahei-1(t)* was constructed based on Nipponbare BAC clones (Fig. 5). The GAAS data of the region revealed the presence of seven candidates of RGAs, probably including the counterpart of the resistance gene in Kahei. The clustering of these several RGAs indicates that this region constitutes a rice blast R-gene cluster comparable to those of *Pik* or *Pita* in rice, as suggested by the co-localization of other blast FR QTLs (Fig. 3). Such clustering of RGAs in the center of the region is surprisingly similar to the *Pik* cluster region in Nipponbare (Xu et al. 2008). By graphical genotyping of these seven RGAs with their SNPs, RGAs 1, 4, and 6 were shown to be located outside of the region (data not shown), probably excluding them from the candidate counterpart list. This also indicates a highly mutable character of this region of the genome, often observed in various R-gene clusters (Michelmore and Meyers 1998).

From these results, as a hypothesis, we are speculating that the most of the QTLs of the conventional multi-component FRs are NBS-LRR type R-genes with weak defense responses than the true resistance genes: i.e., FRs are natural gene pyramiding of WEAK R-genes. That will secure natural suppression of *Avr*-mutated races by the competition with wild-type races because *Avr*-mutations are expected to lower the fitness value of the pathogen. On the contrary, the true resistances may be providing sterile condition without competition, for the *Avr*-mutated races, by completely suppressing wild-type races, thus making the true resistance less durable.

However, as the possibility of other ORFs as candidate counterparts could not be excluded based on the present results, we are currently increasing the population of hetero-siblings to further narrow down the region. In parallel, a new Kahei genome BAC library is being constructed, from which we hope to be able to clone and identify the *Pikahei-1(t)* gene by constructing a physical map of Kahei genome around the gene in the near future.

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References

- Bagali PG, Hittalmani S, Shashidhar SY, Shashidhar HE (2000) Identification of DNA markers linked to partial resistance for blast disease in rice across four locations. In: Tharreau D, Lebrun MH, Talbot NJ, Nottoghem JL (eds) Advances in rice blast research. Kluwer Academic Publishers, Dordrecht, pp 34–44
- Bonman JM (1992) Durable resistance to rice blast disease—environmental influences. *Euphytica* 63:115–123
- Chen DH, Nelson RJ, Wang GL, Inukai T, Mackill DJ, Ronald PC (2000) Characterization of blast resistance in the durable resistant rice cultivar Moroberekan. In: Tharreau D, Lebrun MH, Talbot NJ, Nottoghem JL (eds) Advances in rice blast research. Kluwer Academic Publishers, Dordrecht, pp 17–27
- Ezuka A (1972) Field resistance of rice varieties to rice blast disease. *Rev Plant Prot Res* 5:1–21
- Fukuoka S, Okuno K (2001) QTL analysis and mapping of *Pi21*, a recessive gene for field resistance to rice blast in Japanese upland rice. *Theor Appl Genet* 103:185–190
- Fukuoka S, Kudo S, Okuno K (2003) Genetic analysis of qBR4–2, a QTL for field resistance to rice blast using backcrossed progeny lines. *Breed Res* 5(Suppl. 2):332
- Fukuoka S, Shimizu T, Saka N, Koga H, Ebana K, Hirochika H, Yano M, Okuno K. (2007) Molecular cloning of a QTL, *pi21*, controlling field resistance to blast in rice. *Plant & Animal Genomes XV Conference*, San Diego, CA, W369
- Fukuta Y, Yanoria MJT, Mercado-Escueta D, Ebron LA, Fujita A, Araki E, Khush GS (2004) Quantitative traits loci (QTL) reactions to rice blast isolates from Japan and Philippines. In: Kawasaki S (ed) Rice blast: interaction with rice and control. Kluwer Academic Publishers, Dordrecht, pp 113–121
- Hayashi N (2005) MAFF microorganism genetic resources manual, vol vol. 18. Rice blast fungus, National Institute of Agrobiological Sciences, Tsukuba
- Hirabayashi H, Sato H, Kato H, Ando I, Imbe T, Ogawa T (2005) Mapping of a partial resistance gene for rice blast in *Oryza rufipogon* (IRGC-Acc104814). *Breed Res* 7(Suppl. 1–2):244
- International Rice Genome Sequencing Project (2005) The map-based sequence of the rice genome. *Nature* 436:793–800
- Ishizaki K, Hoshi T, Abe S, Sasaki Y, Kobayashi K, Kasaneyama H, Matsui T, Azuma S (2005) Breeding of blast resistant isogenic lines in rice variety Koshihikari and evaluation of their characters. *Breeding Sci* 55:371–377
- Iwata M, Umemura K, Midoh N (2004) Probenazole (*Oryzemat*[®])—a plant defense activator. In: Kawasaki S (ed) Rice blast: interaction with rice and control. Kluwer Academic Publishers, Dordrecht, pp 163–171
- Kawasaki S, Awato H, Takahashi A, Xu X, Miyao A, Kato H, Hirochika H (2008) Development of an efficient screening system of recessive rice blast resistance mutants and its application. *Breed Res* 10(Suppl. 1):162
- Liu B, Zhang SH, Zhu XY, Yang QY, Wu SZ, Mei MT, Mauleon R, Leach J, Mew T, Leung H (2004) Candidate defense genes as predictors of quantitative blast resistance in rice. *Mol Plant Microbe Interact* 17:1146–1152
- Michelmore RW, Meyers BC (1998) Clusters of resistance genes in plants evolve by divergent selection and a birth-and-death process. *Genome Res* 8:1113–1130
- Miyamoto M, Yano M, Hirasawa H (2001) Mapping of quantitative trait loci conferring blast field resistance in the Japanese upland rice variety Kahei. *Breed Sci* 51:257–261
- Miyao A, Tanaka K, Murata K, Sawaki H, Takeda S, Abe K, Shinozuka Y, Onosato K, Hirochika H (2003) Target site specificity of the *Tos17* retrotransposon shows a preference for insertion within genes and against insertion in retrotransposon-rich regions of the genome. *Plant Cell* 15:1771–1780
- Nguyen TT, Koizumi S, La TN, Zenbayashi KS, Ashizawa T, Yasuda N, Imazaki I, Miyasaka A (2006) *Pi35(t)*, a new gene conferring partial resistance to leaf blast in the rice cultivar Hokkai 188. *Theor Appl Genet* 113:697–704
- Sato H, Takeuchi Y, Hirabayashi H, Nemoto H, Hirayama M, Kato H, Imbe T, Ando I (2006) Mapping QTLs for field resistance to rice blast in the Japanese upland rice Variety Norin 12. *Breed Sci* 56:415–418
- Sirithunya P, Tragoonrun S, Vanavichit A, Pa-In N, Vongsaprom C, Toojinda T (2002) Quantitative trait loci associated with leaf and neck blast resistance in recombinant inbred line population of rice (*Oryza sativa*). *DNA Res* 9:79–88
- Tabien RE, Li Z, Paterson AH, Marchetti MA, Stansel JW, Pinson SRM (2002) Mapping QTLs for field resistance to the rice blast pathogen and evaluating their individual and combined utility in improved varieties. *Theor Appl Genet* 105:313–324
- Talukder ZI, Tharreau D, Price AH (2004) Quantitative trait loci analysis suggests that partial resistance to rice blast is mostly determined by race-specific interactions. *New Phytol* 162:197–209
- Talukder ZI, McDonald AJS, Price AH (2005) Loci controlling partial resistance to rice blast do not show marked QTL × environment interaction when plant nitrogen status alters disease severity. *New Phytol* 168:455–464
- Wang GL, Mackill DJ, Bonman JM, McCouch SR, Champoux MC, Nelson RJ (1994) RFLP mapping of genes conferring complete and partial resistance to blast in a durably resistance rice cultivar. *Genetics* 136:1421–1434

- Wang S, Basten CJ, Zeng ZB (2006) Windows QTL Cartographer 2.5. Department of Statistics, North Carolina State University, Raleigh, NC (<http://statgen.ncsu.edu/qtlcart/WQTLCart.htm>)
- Wu JL, Fan YL, Li DB, Zheng KL, Leung H, Zhuang JY (2005) Genetic control of rice blast resistance in the durably resistant cultivar Gumei 2 against multiple isolates. *Theor Appl Genet* 111:50–56
- Xu X, Kawasaki S, Fujimura T, Wang CT (2005) A protocol for high-throughput extraction of DNA from rice leaves. *Plant Mol Biol Rep* 27:291–295
- Xu X, Hayashi N, Wang CT, Kato H, Fujimura T, Kawasaki S (2008) Efficient authentic fine mapping of the rice blast resistance gene *Pik-h* in the *Pik* cluster, using new *Pik-h*-differentiating isolates. *Mol Breed* DOI 10.1007/s11032-008-9175-5.
- Zenbayashi K, Ashizawa T, Tani T, Koizumi S (2002) Mapping of the QTL (quantitative trait locus) conferring partial resistance to leaf blast in rice cultivar Chubu 32. *Theor Appl Genet* 104:547–552