Mapping of a major QTL, qBK1Z, for bakanae disease resistance in rice

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Abstract

Background: Bakanae disease is a fungal disease of rice (Oryza sativa L.) caused by the pathogen Gibberella fujikuroi (also known as Fusarium fujikuroi). Recently the disease incidence has increased in several Asian countries and continues to spread throughout the world. No rice varieties have been developed yet to be completely resistant to this disease. With increasing need to identify various genetic resources to impart resistance to local elite varieties, this study was carried out to identify novel quantitative trait loci (QTLs) from an indica variety Zenith.

Results: We performed a QTL mapping using 180 F_{2:9} recombinant inbred lines (RILs) derived from a cross between the resistant variety, Zenith, and the susceptible variety, Ilpum. A primary QTL study using the genotypes and phenotypes of the RILs indicated that the locus qBK1\textsuperscript{z} conferring bakanae disease resistance from the Zenith was located in a 2.8 Mb region bordered by the two SSR markers, RM1331 and RM3530 on chromosome 1. The log of odds (LOD) score of qBK1\textsuperscript{z} was 13.43, accounting for 30.9% of the total phenotypic variation. A finer localization of qBK1\textsuperscript{z} was delimited at an approximate 730 kb interval in the physical map between Chr01_1435908 (1.43 Mbp) and RM10116 (2.16 Mbp).

Conclusion: The development of a rice variety with a higher level of resistance against bakanae disease is a major challenge in many rice growing countries. Introducing qBK1\textsuperscript{z} or pyramiding with other previously identified QTLs could provide effective genetic control of bakanae disease in rice.

Background

Bakanae disease, which means foolish seedling in Japanese, was firstly identified in 1828 in Japan (Ito and Kimura 1931), and has widely distributed in temperate zone as well as tropical environment and occurring throughout rice growing regions of the world (Gupta et al. 2015).

Four Fusarium species including F. andiyazi, F. fujikuroi, F. proliferatum and F. verticillioides in the G. fujikuroi species complex have been associated with Bakanae disease in rice (Wulff et al. 2010). This disease typically seed-borne fungus, but may occur when the pathogen is present in plant material or soil. Infected seeds/plants result in secondary infections (Ora et al. 2011), which spread through wind or water that carries the fungal spores from one plant to another. Bakanae disease has different symptoms starting from pre-emergence seedling death to grain infection at maturity such as tall, lanky tillers with pale green flag leaves. Infected plants have fewer tillers, and plants surviving till maturity bear only empty panicles (Singh and Sunder 2012), resulting in yield loss (Mew and Gonzales 2002; Ou 1985). Low plant survival and high spikelet sterility (Singh and Sunder 2012) may account for yield losses up to 50% in Japan (Ou 1985), 3.0-95% in India (Sunder and Satyavir 1997; Fiyaz et al. 2014; Gupta et al. 2015), 3.7-14.7% in Thailand, 5-23% in Spain, 40% in Nepal (Singh and Sunder 1997), 6.7-58.0% in Pakistan (Yasin et al. 2003), 75% in Iran (Saremi et al. 2008), and to 28.8% in Korea (Park et al. 2009). Hot water immersion and fungicide treatment are most common ways for seed disinfection (Hayasaka et al. 2001; Gupta et al. 2015; Lee et al. 2018). However, both the hot water treatment and application of fungicide are insufficient to control bakanae disease. Thermal effect does not reach the pericarp of the severely infected rice seeds. The application of
fungicides is not functioning well for destroying the spores of this fungal pathogen, and some pathogen showed resistance to the fungicides (Ogawa 1988; Park et al. 2009; Kim et al. 2010; Lee et al. 2011). Therefore, the genetic improvement of rice using the QTLs/genes providing the bakanae disease resistance would be a more effective way to control bakanae disease.

Several QTLs associated with bakanae disease resistance have been identified and those can be used for marker-assisted selection in rice breeding as well as for understanding the mechanisms of resistance. Yang et al. (2006) identified two QTLs located on chromosome 1 and chromosome 10 by in vitro evaluation of the Chunjiang 06/TN1 doubled haploid population. Hur et al. (2015) identified a major QTL, qBK1, on chromosome 1 from 168 BC2F4 near isogenic lines (NILS) generated by crossing the resistant indica variety Shingwang with susceptible japonica variety Ilpum. Lee et al. (2019) delimited the location of qBK1 to 35 kb interval between two InDel markers, InDel 18 (23.637 Mbp) and InDel 19-14 (23.672 Mbp). Fiyaz et al. (2016) identified three QTLs (qBK1.1, qBK1.2, and qBK1.3) on chromosome 1 and one QTL (qBK3.1) on chromosome 3 from the highly resistant variety Pusa 1342. Physical location of qBK1.1 is close to qBK1 (Fiyaz et al. 2016; Lee et al 2019). Ji et al. (2017) mapped a major QTL (qFfR1) in 22.56–24.10 Mbp region on chromosome 1 using 180 F2:3 lines derived from a cross between a resistant Korean japonica variety Nampyeong and a susceptible Korean japonica line, DongjinAD. Volante et al. (2017) identified qBK1_628091 (0.6 Mbp to 1.0 Mbp on chromosome 1) and qBK4_31750955 (31.1 Mbp to 31.7 Mbp on chromosome 4) by GWAS (Genome Wide Association Study) approach using 138 japonica rice germplasm. Kang et al (2019) discovered the QTL qFFR9 at 30.1 centimorgan (cM) on chromosome 9 from a japonica variety Samgwang. Lee et al. (2018) found the QTL qBK1WD located between markers chr01_13542347 (13.54 Mb) and chr01_15132528 (15.13 Mb) from the japonica variety Wonseadaesoo. They also found that resistance of gene pyramided lines harboring two QTLs, qBK1WD and qBK1 was significantly higher than those with only qBK1WD or qBK1. This finding relates to the problem of varieties with single resistance gene losing its effect against new populations of fungal isolates (Wang and Valent 2009). Hence, identifying new resistance genes from diverse sources is important for rice breeding programs to acquire durable resistance against bakanae disease by enhancing the resistance level and/or help to overcome the breakdown of resistance genes.

In this study, we aimed to provide a new genetic source, qBK1^2 with detailed gene locus information for developing resistant rice lines which contains single or multiple major QTLs to enhance bakanae disease resistance.

Results

Bakanae disease bioassay in parents and F2:3 RILs

The proportion of healthy Zenith (resistant) and Ilpum (susceptible) plants was evaluated with 10 biological replicates after inoculation of the virulent F. fujikuroi isolate CF283 (Kim et al. 2014). Most of Zenith plants did not exhibit a thin and yellowish-green phenotype, which is a typical symptom of bakanae disease as unlike Ilpum (Fig. 1A). The proportion of healthy Zenith
plants was 63.2± 11.8% (ranging from 42.7% to 79.3%), which was significantly different from that of Ilpum (14.3± 11.4%, ranging from 3.3% to 37.0%) (Fig. 1B). Zenith and Ilpum were further inoculated with green fluorescent protein (GFP)-tagged *F. fujikuroi* isolate CF283. Ten days after inoculation, plants with typical disease symptom of each variety were subjected to a confocal microscopy analysis. Confocal imaging of radial and longitudinal sections of the basal stem showed that the fungus penetrated and was localized easily and abundantly at vascular bundle, mesophyll tissue and hypodermis in the susceptible Ilpum variety while only background level of GPF signal was detected in the resistant Zenith (Fig. 2).

QTL analysis and mapping of *qBK1z* using 180 F_{2:9} RILs

Based on the bakanae disease bioassay (proportion of healthy plants), the 180 F_{2:9} RIL population exhibited continuous distribution (ranged from 0% to 98.0%; Fig. 3), which quantitatively confirmed the inheritance of bakanae disease resistance.

We selected 164 SSR markers among 1,150 SSR markers from the polymorphism survey between Ilpum and Zenith from the Gramene database (http://www.gramene.org) covering the whole rice chromosome (S1 Fig). The genetic linkage map of Ilpum and Zenith for primary mapping was constructed with 164 polymorphic markers covering a total length of 3,140 cM with average interval of 19.14 cM as described by Lee et al. (2017). Primary QTL mapping using the 180 F_{2:9} populations showed that a significant QTL associated with bakanae disease resistance at the seedling stage was located between the SSR markers, RM1331 and RM3530 on chromosome 1, and it was designated *qBK1z*. The LOD score of *qBK1z* was 13.43, which accounted for 30.9% of the total phenotypic variation (Table 1).

A finer localization of *qBK1z* was determined by analyzing the chromosome segment introgression lines in the region detected from primary mapping. The *qBK1z* region between RM1331 and RM3530 from primary mapping was narrow downed with additional 55 SSR markers and designed 12 InDel markers for the insertion/deletion sites based on the differences between the *japonica* (http://www.gramene.org) and *indica* (http://rice.genomics.org.cn) sequences. Four SSR markers and six InDel markers were selected as polymorphic markers between the parents to narrow down the position of the *qBK1z* region (S2 Table). Finally, seven homozygous recombinants were selected from the F_{2:9} lines using 14 markers in the 2.8 Mb region around the SSR markers RM1331 and RM3530 (Fig. 4 and Fig. 5).

The proportion of healthy plants of the seven homozygous recombinants was evaluated with three biological replicates according to Duncan’s new multiple range test. Based on this bioassay, lines classified to Group a were regarded as resistant, and Group b as susceptible (Fig. 4). Considering the genotype and the phenotype of the recombinants, it is clear that *qSTV11Z* conferring resistance to bakanae was an approximate 730 kb interval delimited by the physical map between Chr01_1435908 (1.43 Mbp) and RM10116 (2.16 Mbp).

Discussion
Rice varieties with a single resistance gene are at an increased risk of being overcome by new pathological races (Lee et al. 2017; Lee et al. 2018). The development of a rice variety with a higher level of resistance against bakanae disease is a major challenge in many countries (Cumagun et al. 2011; Bashyal et al. 2014; Fiyaz et al. 2016; Hur et al. 2015; Hur et al. 2016). In this study, we identified \( qBK1^z \) locus related to bakanae disease resistance based on genotype and phenotype analyses of homozygous recombinants on the recombinant progeny of Ilpum and Zenith, using SSR and newly developed InDel markers.

It was reported that successful infection of \( \text{Fusarium} \) species is a complex process that includes adhesion, penetration (through wounds, seeds, stomatal pores) and subsequent colonization within and between cells (Rana et al, 2017; Jansen et al 2005). Lee et al. (2018) revealed that the fungus \( F. fujikuroi \) was more abundant in the stem of the susceptible variety than it was in the resistant one. Elshafey et al. (2018) indicated that \( F. fujikuroi \) prefer to grow in aerenchyma, pith, cortex, vascular bundle of both sheath and stem of rice. In this study, we examined both the localization and abundance of \( F. fujikuroi \) isolate CF283 in the basal stem of rice using GFP-tagged \( F. fujikuroi \) isolate CF283. Consistent with previous reports (Elshafey et al. 2018; Lee et al. 2018), \( F. fujikuroi \) isolate CF283 was extensively observed on the vascular bundle, mesophyll tissue and hypodermis of infected stem in susceptible Ilpum variety, whereas this was rarely observed in resistant Zenith.

Many QTLs on bakanae disease resistance have been identified on chromosome 1. Three QTLs, \( qBK1^z, qBK1.2 \) and \( qBK1.3 \), were found in a similar region in spite of the different source of resistant varieties (Fig 6). Fiyaz et al. (2016) mapped \( qBK1.2 \) to a 260 kb region between markers RM10153 and RM5336, and \( qBK1.3 \) to a 3.49 Mb region between markers RM10271 and RM578 from the Pusa 1121/Pusa1342 cross. Location of \( qBK1^z \) was found upstream to \( qBK1.2 \). Fiyaz et al. (2016) mapped \( qBK1.1 \) to a 20 kb region between markers RM9 and RM11232 from the Pusa 1121/Pusa1342 cross. These authors hypothesized that \( qBK1.1 \) and \( qBK1 \) (Hur et al. 2015) might be the same QTL as they had overlapping positions. Ji et al. (2017) found that QTL \( qFFR1 \) was located in a 230 kb region of rice chromosome 1 in Korean \( japonica \) variety Nampyeong, and suggested that the three QTLs \( qBK1, qBK1.1, \) and \( qFFR1 \) might indicate the same gene. Lee et al. (2019) narrowed down the position of the \( qBK1 \) locus to a 35 kb region between InDel 18 and InDel 19-14, and revealed that location of \( qBK1 \) is close to those of \( qBK1.1, \) and \( qFFR1, \) does not overlapping each other. Two additional QTLs including \( qB1 \) from Chunjiang 06 (Yang et al. 2006) and \( qBK1^{WD} \) from Wonseadaesoo (Lee et al. 2018) also were also found on chromosome 1. Gene pyramiding via phenotypic screening assays for crop breeding is considered to be difficult and often impossible due to dominance and epistatic effects of genes governing disease resistance, and the limitation of screenings being all year-round (Sundaram et al. 2009). Pyramiding of multiple resistant QTLs/gene by using marker-assisted breeding (MAB) in a single plant might confer higher and/or durable resistance against bakanae disease. The effects of pyramiding resistance genes have been observed for several plant-microbe interactions. Pyramiding three bacterial blight resistance genes resulted in a high level of resistance and were expected to provide a durable pathogen resistance (Singh et al. 2001; Pradhan et al. 2015). On the other hand, pyramiding of resistant genes resulted in a level of resistance that was comparable to or even lower to than that of the line with a single gene. For example, Yasuda et al. (2015) reported rice lines with pairs of blast resistance genes to be only comparable to lines with a single gene which may have a stronger
suppressive effect. Similarly, pyramiding of brown planthopper (BPH) resistance genes in a susceptible variety IR24 markedly improved BPH resistance against four BPH populations in the Philippines than the single BPH introgressed lines (Jena et al. 2017). In our previous study of bakanae disease resistance (Lee et al. 2018) revealed that the gene pyramided lines harboring $qBK1^{WD} + qBK1$ had a much higher level of resistance than those possessing either $qBK1^{WD}$ or $qBK1$. The novel QTL, $qBK1^Z$, identified in this study can be utilize in MAB and gene pyramiding to achieve higher resistance in many bakanae disease prone rice growing areas.

Conclusions

In this study, we identified a new major QTL $qBK1^z$ conferring bakanae disease resistance from a new genetic source of indica variety, Zenith. Through QTL analysis and fine mapping, we narrowed down the $qBK1^z$ locus into 730 kb on the short arm of chromosome 1 where is a novel locus compared with all the previously identified bakane disease resistance QTLs. Together with the previously identified QTLs of the bakane disease resistance, the new $qBK1^z$ can be introduce to the elite favorable background varieties by a marker-assisted backcrossed breeding. Furthermore, the information of the localization and different abundance on the vascular bundle, mesophyll tissue and hypodermis of infected stem of infected rice between resistant and susceptible varieties will be useful for further studying an interaction between the pathogen ($F. fujikuroi$) and rice host plants.

Methods

Plant materials

Zenith, a medium grain type indica variety of USA released in 1936, was identified as resistant to bakanae disease in a preliminary screening of rice germplasm (data not shown) using the large-scale screening method developed by Kim et al. (2014). We generated 180 $F_{2:9}$ RILs from a cross between susceptible variety, Ilpum, and resistant variety, Zenith for QTL analysis. The population was developed in the experimental fields of the National Institute of Crop Science of the Rural Developmental Administration in Miryang, Korea.

Evaluation of bakanae resistance and statistical analysis

The inoculation and evaluation of bakanae disease were conducted using a method that described by Lee et al. (Lee et al. 2019). The isolate CF283 of $F. fujikuroi$ (imperfect stage of $G. fujikuroi$) were obtained from National Academy Agricultural Science in Korea. Isolate was inoculated in potato dextrose broth (PDB) and cultured at 26 °C under continuous light for one week. The $F. fujikuroi$ culture was washed by centrifugation with distilled water. Regarding the standardization of the density of the homogenized inoculum, the fungal spore concentration was adjusted to $1 \times 10^6$ spores/mL with a hemocytometer. Forty seeds per each line were placed in a tissue-embedding cassette (M512, Simport, Beloeil, QC, Canada). Before inoculation, the seeds in the in the tissue-embedding cassette were surface sterilized with hot water (57 °C) for 13 min, then allowed to drain and cool. Subsequently, the seeds were soaked in the spore suspensions ($1 \times 10^6$
spores/mL) for 3 days for inoculation with gentle shaking four times a day for equilibration. After inoculation, thirty seeds per line were sown in commercial seedling tray, and seedlings were grown in a greenhouse (28 ± 3 °C day/23 ± 3 °C night, 12 h light). Bakanae disease symptom on each line was evaluated by calculating the proportion of healthy plants at 1 month after sowing. The healthy and non-healthy plants are classified as described by Kim et al. (2014). The plants exhibit elongation with thin and yellowish-green, elongation then stunted growth, stunted growth, dead seedling were classified as non-healthy plants. The plants showing same phenotype with untreated plants, slight elongation then normal growth without thin and yellowish-green was regarded as healthy plants.

Statistical differences between means were analyzed using Duncan's multiple range test after one-way analysis of variance (ANOVA). The level of significance was designated as P < 0.05 and was determined using the SAS Enterprise Guide 4.3 program (SAS Institute Inc., Cary, NC, USA).

**Localization of** *F. fujikuroi* **in Zenith and Ilpum plants**

Shoot base of 10 day-old seedlings derived from Zenith and Ilpum seeds inoculated by GFP-tagged *F. fujikuroi* isolate CF283 (Lee et al. 2018) was observed under confocal laser-scanning microscopy (LSM-800, Zeiss, Germany) at GFP channel, and images were obtained using Zeiss LSM Image Browser. All experiments were conducted twice with at least three replicates.

**DNA extraction and polymerase chain reaction**

Genomic DNA from young leaf tissue was prepared according to the CTAB method (Murray and Thompson 1980) with minor modifications. Polymerase chain reaction (PCR) was performed in 25-µL reaction mixture containing 25 ng template DNA, 10 pmol of each primer, 10× e-Taq reaction buffer, 25 mM MgCl2, 10 mM dNTP mix and 0.02 U of SolGent™ e-Taq DNA polymerase (SolGent, Daejeon, South Korea). The reaction conditions were set as follows: initial denaturation at 94°C for 2 min; 35 cycles of denaturation at 94°C for 20 s, annealing at 57°C for 40 s and extension at 72°C for 40 s; and a final extension at 72°C for 7 min. The amplification products were electrophoresed on a 3% (w/v) agarose gel and visualized by ethidium bromide staining.

**QTL analysis of the F_{2:9} population and development of InDel markers for fine mapping**

Polymorphic SSR markers (n = 164) that were evenly distributed on rice chromosomes were selected from the Gramene database (http://www.gramene.org). These markers were used to construct a linkage map and for QTL analysis of the F_{2:9} populations. The linkage map was constructed using Mapmaker/Exp v.3.0, and the genetic distance was obtained using the Kosambi map function (Lander et al. 1987). Putative QTLs were detected using the composite interval mapping (CIM) function in WinQTLcart v.2.5 (WinQTL cartographer software; Wang et al. 2007), with the threshold being set based on the results of 1000 permutation tests. A logarithm of the odds (LOD) ratio threshold of 3.0 was used to confirm the significance of a putative QTL. InDel markers were developed based on the fragment size differences in the sequence (in the range of 20 bp) between *japonica* (Gramene database http://www.gramene.org) and *indica* (BGI-RIS;
http://rice.genomics.org.cn) in the target region on chromosome 1. The primers were designed using Primer3 software (http://web.-bioneer.co.kr/cgi-bin/primer/primer3.cgi).

List Of Abbreviations

QTLs: quantitative trait loci
LOD: log of odds
RILs: recombinant inbred lines
SSR: simple sequence repeat
InDel: insertion/deletion
MAS: marker-assisted selection
PDB: potato dextrose broth
GFP: green fluorescent protein

Declarations

Ethics approval and consent to participate
Not applicable

Consent for publication
Not applicable. The Author transfers to Springer the publication rights.

Availability of supporting data
The datasets supporting the conclusions of this article are provided within the article and its additional files.

Competing interests
We declare that the funding agencies and organizations are acknowledged and there are no financial or non-financial competing interests.

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Author's contributions
SB, YJ, and DS carried out the molecular genetic studies and drafted the manuscript. SM, JY, JHC, JHL and YC participated in the QTL analysis. M, SR and JM contributed to the phenotyping and result interpretation. NK, YS and JK carried out microscopic analysis using GFP-tagged \textit{F. fujikuroi} isolate CF283GFP.

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**Tables**

**Table 1.** Putative quantitative trait locus (QTL) associated with bakanae disease resistance detected at the seedling stage by composite interval mapping of the 180 F_{2:9} populations derived from a cross between Ilpum and Zenith

| QTL  | Chromosome | Position (cM) | Marker interval | LOD^a | PVE^b (%) | Additive effect | Dominant effect |
|------|------------|---------------|----------------|-------|-----------|----------------|----------------|
| qBK1^Z | 1 | 4.0 | RM133 \_1 ~ RM353 \_0 | 13.43 | 30.93 | -22.10 | -1.09 |

^a LOD: log of odds score; ^b PVE: percentage of phenotypic variation explained.

**Figures**
Figure 1

Bioassay of parents (Ilpum and Zenith) against F. fujikuroi isolate CF283. (A) Plant phenotype from the pathogen-inoculated seeds of Ilpum and Zenith. (B) Proportion of healthy plants from the pathogen-treated seeds.
Figure 2

Confocal imaging of shoot base of 10 days-old-seedling derived from Zenith and Ilpum rice plants infected with *F. fujikuroi* (CF283 isolates) containing GFP expression cassette. (A) Radial and (B) longitudinal sections of the basal stem (Scale bar = 50 μm).
Figure 3

Frequency distribution of the proportion of healthy F2:9 NILs derived from a cross between Ilpum and Zenith after bakanae disease inoculation. The mean proportion of healthy plants of Ilpum and Zenith is indicated by black arrow heads.
Figure 4

QTL analysis of qBK1Z on chromosome 1 using 180 RILs derived from a cross between Ilpum and Zenith. (a) In primary mapping, qBK1Z was located in 2.8 Mb regions between the RM1331 and RM3530 markers on chromosome 1. (b) Location of qBK1Z was narrowed down to 730 kb region between the markers Chr01_1435908 and RM10116 in secondary mapping using selected seven homozygous recombinants. Black bars show the homozygous regions for zenith alleles; white bars indicate the homozygous regions for Ilpum alleles. R; resistant to bakanae disease, S; susceptible to bakanae disease. The proportion of healthy plant was calculated from three biological replications. Values (%) of the proportion of healthy plant with different letters are significantly different by Duncan's multiple range test at $P = 0.05$. 
Figure 5

Phenotypic responses to bakanae disease infection of seven homozygous recombinants used in secondary mapping.
Figure 6

Physical locations of bakanae disease resistance QTLs identified on chromosome 1.

Supplementary Files

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- TableS1.docx
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