การตรวจสอบเครื่องหมายโมเลกุลสำหรับคัดเลือกความต้านทานเพลี้ยกระโดด สีน้ำตาลและโรคไหม้ในพันธุ์ข้าวเหนียวปรับปรุง Validation of Brown Planthopper and Blast Resistance Markers in Improved Aromatic Glutinous Rice

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ABSTRACT

Bph32 brown planthopper (BPH) resistance gene was incorporated into Hom Xebangfai 4 (HXBF4) together with maintaining two of QTLs BL resistance qBL1 and qBL11. This breeding was processed by a single cross with the BPH and BL rsistant glutinous line, RGD13117-115-52-B. Marker-assisted selection was done in F_2 population. In F_3 , the 11 selected lines were evaluated for two BPH resistance from Singburi (SBR) and Ayutthaya (AYY) by a modified standard seedbox screening method. The results showed that 10 lines that detected *Bph32 Bph3* and *TPS* had high resistance against both BPH populations. However, the other 1 line carrying *Bph3* and *TPS* had high resistance to AYY but moderately resistant to SBR. After that, the 16 F_4 lines derived from the 1 line of F_3 were evaluated with 7 mixed Thai BL isolates in a greenhouse condition. The results showed that all of the 16 F_4 lines carrying both of qBL1 and qBL11 had resistance against 7 mixed BL isolates and they were more effective rather than that one BL QTL. Therefore, the phenotyping in this study also strongly suggested that the genotyping with high-throughput markers of BPH and BL were very accurate and trustable.

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ยืนที่ด้านทานเพลี้ยกระโดดสีน้ำตาล (*Bph32*) ถูกถ่ายทอดสู่พันธุ์ข้าวเหนียว Hom Xebangfai 4 (HXBF4) พร้อมกับคงดำแหน่ง QTL ที่ด้านทานต่อโรคไหม้ qBL1 และ qBL11 โดยการผสมข้ามกับสายพันธุ์ RGD13117-115-52-B ซึ่งมียืนดังกล่าว จากนั้นจึงใช้เครื่องหมายโมเลกุลช่วยในการคัดเลือกในประชากรชั่วที่ 2 โดยในชั่วที่ 3 นำสายพันธุ์ที่ถูกคัดเลือกจำนวน 11 สายพันธุ์ มาประเมินความด้านทานต่อเพลี้ยกระโดดสี น้ำตาลประชากรสิงห์บุรี (SBR) และอยุธยา (AYY) ด้วยวิธี modified standard seedbox screening method ผลการทดลองพบว่า สายพันธุ์ที่ถูกคัดเลือก 10 สายพันธุ์ ที่มียืน *Bph32, Bph3* และ *TPS* มีความด้านทานต่อ เพลี้ยกระโดดสีน้ำตาลทั้งสองกลุ่มประชากรในระดับที่สูง อย่างไรก็ตาม อีก 1 สายพันธุ์ ที่มีเฉพาะ *Bph3* และ *TPS* มีความต้านทานต่อ AYY ในระดับสูง แต่ต้านทานต่อ SBR ในระดับปานกลาง จากนั้นนำสายพันธุ์ชั่วที่ 4 จำนวน 16 สายพันธุ์ ที่คัดเลือกจาก 1 สายพันธุ์ของชั่วที่ 3 ที่มี qBL1 และ qBL11 มาประเมินความต้านทาน ต่อโรคไหม้จากเชื้อผสม 7 กลุ่ม พบว่า สายพันธุ์ของชั่วที่ 3 ที่มี qBL1 และ qBL11 มาประเมินความต้านทาน ต่อโรคไหม้จากเชื้อผสม 7 กลุ่ม พบว่า สายพันธุ์ของชั่วที่ 3 ที่มี qBL1 และ qBL11 มาประเมินความต้านทาน เชื่อถือไก้ในระดับสูง จึงสรุปได้ว่า ยืน *Bph32* และ 2 BL QTLs, qBL1 และ qBL11 สามารถใช้เป็นเครื่องหมาย ในการคัดเลือกในโปรแกรมการปรับปรุงพันธุ์พืชให้ด้านทานเพลี้ยกระโดดสีน้ำตาลและโรคไหม้ในเครื่องหมาย และลาวได้.

คำสำคัญ: ข้าวเหนียว ความต้านทานเพลี้ยกระโดดสีน้ำตาล ความต้านทานโรคใบไหม้ เครื่องหมายช่วยในการคัดเลือก

Introduction

Brown planthopper (Nilaparvata lugens Stål) is the most destructive insect pest for rice production in Southeast Asia (Wu *et al.*, 2018) including Lao PDR (Inthapanya *et al.*, 2011). Planting resistant cultivars is the most ecological friendly strategy to reduce production loss from the insect. Over 30 BPH resistance genes have been identified in cultivated and wild species of Oryza (Prahalada *et al.*, 2017; Brar *et al.*, 2009). Only five Bph genes, Bph14, Bph26, Bph3, Bph29, and *Bph32* have been successfully cloned (Ren *et al.*, 2016). The Sri Lankan rice cultivar Rathu Heenati was found strong and broad-spectrum resistance against BPH (Lakshminarayana & Khush, 1977).

Besides, Ikeda & Kaneda (1981) reported that Rathu Heenati was resistant to all four known biotypes of brown planthopper (BPH). The *Bph32*, also previously known as Bph3 on chromosome 6 was identified by Jairin *et al.* (2007). It was incorporated into KDML105 background and the new breeding lines showed broad resistance to BPH populations in Thailand (Jairin *et al.*, 2009). Bph3 on chromosome 4 was identified and cloned by Liu *et al.* (2015). This locus contains a cluster of three genes encoding plasma membrane-localized lectin receptor kinases which are OsLecRK1, OsLecRK2 and OsLecRK3. Lectin receptor kinase genes function together to confer broad-spectrum and durable insect resistance. Terpene synthase gene (TPS) on chromosome 4 was identified and found to be induced by BPH feeding (Kamolsukyunyong *et al.*, 2013). TPS may involve in antixenosis BPH resistance mechanism.

Blast disease caused by Magnaporthe oryzae (anamorph: Pyricularia oryzae) is one of the most devastating diseases for rice-growing countries worldwide (Nalley et al., 2016; Asibi et al., 2019). The disease caused about 10 to 20% yield loss in regular seasons and as high as 100% yield loss in years with BL epidemics (Dean et al., 2005). It is also the most serious disease reducing yield substantially in the rain-fed lowland in Laos PDR (Teng & Revilla, 1996; Gnanamanickam, 2009). Over 100 resistance genes and 350 quantitative trait loci (QTLs) have been identified in Oryza sativa L. Only 25 BL resistance genes or Pi genes have been successfully cloned and applied in breeding programs (Ashkani et al., 2015). Many allelic genes have been reported such as Pish/Pi35 on chromosome 1 and Pikh/Pikm/Pik/Pikp/Pi1 on conferring chromosome 11 broad-spectrum resistance to rice BL (Hua et al., 2012; Takahashi et al., 2010). Two QTLs for broad-spectrum resistance, qBL1 and qBL11 were also identified in Thai cultivar Jao Hom Nin (JHN) (Noenplab et al., 2006). These QTLs confer high resistance against BL isolates from Thailand and Lao PDR (Wongsaprom

et al., 2010; Korinsak *et al.*, 2011). The tightly linked markers RM212- RM319 and RM224 - RM144 were developed for the selection of the QTLs (Noenplab

et al., 2006). These linked markers were successfully applied in rice breeding programs in Thailand and Lao PDR (Manivong *et al.*, 2014; Khanthong *et al.*, 2018; Srichant. *et al.*, 2019).

Hom Xebangfai 4 is an aromatic glutinous rice. It was officially released in Lao PDR in 2017

for blast resistance (qBL1, qBL11), submergence tolerance (Sub1C), aroma (badh2) and brown planthopper resistance (Bph3, TPS). It performs well in many aspects but the level of brown planthopper resistance was moderate due to the lacking of *Bph32*. A breeding program for the incorporation of *Bph32* into HXBF4 was initiated in 2017. Desirable traits in this project include blast resistance, submergence tolerance and brown planthopper resistance. A total of seven highthroughput markers were used in the genotyping of F2 populations. The objective of this study was to validate the BPH resistance in F3 families and the BL resistance in F4 families.

Materials and Methods

Plant materials

HXBF4 carrying Sub1C, badh2, qBL11 and gBL1 was used as a female parent. RGD13117-115-52-B line (RGD13117), an aromatic glutinous introgression line carrying Bph32, Bph3, TPS, Sub1C, badh2, and qBL11 developed by Rice Gene Discovery Unit (RGDU), National Center for Genetic Engineering and Biotechnology (BIOTEC), Kasetsart University Kamphaeng Saen Campus, Thailand, was used as a male parent. Bph32 was introgressed from the male parent to the female parent (HXBF4) using marker-assisted selection (MAS). The MAS was done in F2 generation at the segregation loci of Bph32 and qBL1 including other loci of Bph3, TPS, Sub1C, badh2, qBL11. F3 families derived from the selected F2 individual plants carrying positive homozygous alleles of Bph32 were evaluated for BPH resistance. F4 families derived from the selected F3 individual plant carrying positive homozygous alleles of qBL1 and gBL11 were evaluated for BL resistance (Fig 1).

Rice growth conditions

This experiment was conducted from September 2017 to December 2020 at RGDU, Kasetsart University, Kamphaeng Saen Campus, Nakhon Pathom, Thailand. The rice plants were seeded in a field nursery. After 30 days, the rice seedlings were transplanted into the paddy field. The F2, F3, and F4 selected plants were grown in 1 row/line of 2.5×2.5 m with a spacing of 25×25 cm. The management practices were performed following conventional high-yield cultivation approaches.



Figure 1 Development of aromatic glutinous rice lines with BPH and BL by using marker-assisted selection

Genotyping

High-throughput genotyping was conducted in F2 population by using SNP markers developed by RGDU, BIOTEC, Thailand. DNA from the leaf sample was extracted by using DNA trapping method (DNA technology laboratory) (Nubankoh *et al.*, 2020). Polymerase Chain Reaction (PCR) was conducted following the KASP genotyping protocol (LGC Ltd). Allele discrimination was read by using the QuantStudio[™] 12K Flex machine (Applied Biosystems[™]).

Table 1	I The SNP	markers	used	for o	detect	ted	target	i trai	ts
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Traits	Gene/QTL	Chro	Markers	Reference
	Bph3	4	sLecRK3Bph32	Liu <i>et al.,</i> (2015)
Brown planthopper resistance	Terpene synthase	4	OsSTPS2	Kamolsukyunyong <i>et al.</i> , (2013)
	Bph32	6	Bph32	Jairin <i>et al</i> ., (2007): Ren <i>et al</i> ., (2016)
Blast resistance	qBL1	1	TBGI055578	Wongsaprom <i>et al.</i> , (2010)
Didet recipitance	qBL11	11	TBGI454717	Wongsaprom <i>et al.</i> , (2010)
Submergence tolerance	Submergence tolerance Sub1C 9 Sub1A_SNP1 Siangl		Siangliw <i>et al.</i> , (2003)	
Aromatic	badh2	8	Aroma	Wanchana <i>et al.</i> , (2005)

Validation for brown planthopper resistance

Insect materials: BPH used in this experiment were collected from a rice field in Singburi (SBR) and Ayutthaya (AYY) in 2013. The two populations were reared in 20x30x30 cm3 40-mesh nylon cages in a controlled room at 26-28 °C with 15 hour-light/9 hour-dark in 50-60% relative humidity. Taichung Native 1 (TN1) seedlings were used as feed and habitat for the insect. New seedlings were replenished every four days. Mass rearing was done for insect multiplication in a greenhouse.

BPH infestation and scoring: Eleven F3 lines were evaluated for BPH resistance with modified standard seedbox screening described by Saxena (1989) and this method has been used in many experiments (Jairin et al., 2005; 2006; 2009). Germinating seeds of each line were planted in a row (10 seeds/row). Rathu Heenati (resistant check), TN1 (susceptible check), HXBF4 (female parent) and RGD13117-115-52-B (male parent) were also included as checks. At 25 days old, a mix of 2nd- and 3rd-instar BPH nymphs were released at the rate of 25-30 insects/plants. The infestation was recorded as damage scores (DS) according to the Standard Evaluation System (SES) for Rice (IRRI, 2013). The DS was recorded when all susceptible check plants (TN1) died at nine days after infestation (DAI). The DS was recorded again at 17 DAI for durability of the resistance. This experiment was conducted in a completely randomized design with three replications. A score under 3.0 is considered resistance (R). Scores from 3.1 to 5.0 are considered moderately resistant (MR). Scores from 5.1 to 7.0 are considered moderately susceptible (MS). Score from 7.1 to 9.0 are considered susceptible (S).

Validation for blast resistance

Disease materials: Forty-nine blast isolates used in this study were grouped into 7 mixed BL isolates. These blast isolates were collected from different regions in Thailand by RGDU. The collected isolates were classified by genetic diversity cluster analysis by using Amplified Fragment Length Polymorphism (Rice Gene Discovery, Thailand, unpublished).

BL inoculation and scoring: The 16 F4 lines were evaluated for BL resistance under greenhouse conditions. Sariceltik (susceptible check), KDML105, and RD6 (moderate resistance check), Jao Hom Nin (resistant check), HXBF4 (female parent) and RGD13117-115-52-B (male parent) were also included in the experiment. Four germinating seeds of each line were planted in plastic trays with 6 x 12 holes. The experiment was conducted in a completely randomized design with three replications. The seedlings were maintained in a greenhouse under high humidity for 21 days The preparation of rice plants and the old. inoculation were followed the protocol described by Korinsak et al. (2011), and the lesion score (LS) was recorded at 7 days after inoculation (DAI) on a 0 to 6 scale. Plants exhibiting reactions that

scored 0-2 were considered resistant (R), 3-4 as moderately resistant (MR) and 5-6 as susceptible (S).

Results

Genotyping in F₂ population

Marker-assisted selection was done in 240 F2 individual plants. Step-wise genotyping 60 lines for BPH (*Bph32*) and 10 lines for BL resistance (qBL1) were selected respectively to reduce the sample size. Genotyping at other loci including Bph3, TPS, qBL11, Sub1C and badh2 was also confirmed of existence in progeny lines. Ten F2 individual plants with three homozygous positive

alleles for BPH resistance loci and two BL resistance loci were selected (Table 2). The RGD17020-MS373 with a negative allele for *Bph32* was also maintained to compare between progeny

lines. Eleven F3 families were obtained from selffertilization of the selected F2. All 11 families were evaluated for brown planthopper resistance.

 Table 2 Genotypes of 11 F₂ selected lines compared with their parents, resistant and susceptible checks for

 BPH and BL

	SNP on E	3PH resista	Blast resistance genes		
Line	Bph3	TPS	Bph32	qBL1	qBL11
Rathu Heenati (R check) ^{1/1}	G/G	T/T	G/G	C/C	C/C
TN1 (S check) ^{1/2}	C/C	C/C	C/C	C/C	T/T
Jao Hom Nin (R check) ^{2/1}	C/C	C/C	C/C	T/T	T/T
Sariceltik (S check) ^{2/2}	C/C	C/C	C/C	C/C	C/C
HXBF4 (female parent)	G/G	T/T	C/C	T/T	T/T
RGD13117-115-52-B (male parent)	G/G	T/T	G/G	C/C	T/T
RGD17020-MS30	G/G	T/T	G/G	T/T	T/T
RGD17020-MS36	G/G	T/T	G/G	T/T	T/T
RGD17020-MS42	G/G	T/T	G/G	T/T	T/T
RGD17020-MS45	G/G	T/T	G/G	T/T	T/T
RGD17020-MS87	G/G	T/T	G/G	T/T	T/T
RGD17020-MS94	G/G	T/T	G/G	T/T	T/T
RGD17020-MS143	G/G	T/T	G/G	T/T	T/T
RGD17020-MS166	G/G	T/T	G/G	T/T	T/T
RGD17020-MS194	G/G	T/T	G/G	T/T	T/T
RGD17020-MS338	G/G	T/T	G/G	T/T	T/T
RGD17020-MS373	G/G	T/T	C/C	T/T	T/T

Note: ^{1/1} = Positive for BPH validation

^{1/2} = Negative for BPH validation

^{2/1} = Positive for BL validation

2/2 = Negative for BL validation

BPH resistance in F₃ generation

The experiment on BPH evaluation in F3 generation was well controlled which the resistant check showed the lowest DS and the susceptible check showed the highest DS from the infestation by both BPH populations. Different BPH populations were similar to the pattern of infestation. However, SBR population made more damage than that of the AYY in all selected plants and checks. The levels of damage were also obviously observed and shown in Fig. 2 and Fig. 3.

According to the SES, ten of F3 lines containing all three homozygous Bph resistance loci were scored under 3.0 at 9 DAI. They performed as well as the resistant check and the male parent. This suggests that the three Bph loci conferred high levels of resistance to both BPH populations. On the other hand, RGD17020-MS373 with a negative allele of *Bph32* was resistant to AYY, but it had moderate resistance against SBR at 9 DAI.

For durability, DS from AYY population was still lower than 3.0 even at 17 DAI for ten F3 lines.

Besides, their damage from SBR population was still lower than 5.0 which was moderately resistant at 17 DAI. RGD17020-MS373 with a negative allele of *Bph32* was susceptible to SBR population at 17

DAI. However, it was resistant to AYY population with a DS of 3.7 at 17 DAI. The damage trend of RGD17020-MS373 was the same as the female parent.



BPH Damage score

🖾 SBR 9 DAI 🛛 SBR 17 DAI 🖳 AYY 9 DAI 🖄 AYY 17 DAI

Figure 2 Average damage scores of 11 F₃ lines after infested with Singburi and Ayutthaya BPH populations at 9 DAI and 17 DAI. Scoring 0-3 = resistant (R), 3-4 = moderately resistant (MR), 5-7 = moderately susceptible (MS), and 7-9 =susceptible (S)



Figure 3 Symptoms of 25 days old F₃ seedlings infested with Singburi BPH population 9 DAI compared with resistant check (RH), susceptible check (TN1), HXBF4 (female parent) and RGD13117 (male parent)

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BL resistance in F_4 generation

The 16 F4 lines derived from a single F3 plant were subjected to BL screening with their parents by using 7 mixed BL isolates groups. The three susceptible checks, (Sariceltik, RD6 and KDML105) and a resistant check (JHN) were included as controlled varieties. The result showed that Sariceltik was susceptible to all of BL isolates, while RD6 and KDML105 were susceptible (Fig 4 and 5). JHN was resistant to all of the mixed isolates. The male parent carrying only qBL11 was resistant to the mixed 2, 3, 4, 5, and 6 and moderately resistant to the mixed 1 and 7. While female parent carrying both qBL1 and qBL11 showed resistance to all of the 7 mixed BL isolates. In the same way, the 16 F4 tested lines carrying these two BL QTLs were resistant to all of the 7 mixed BL isolates as well as the HXBF4.



Figure 4 Average damage scores of 16 F₄ lines carrying qBL1, qBL11 inoculated with 7 mixed BL isolates compared with Sariceltik and RD6 (susceptible checks), HXBF4 (female parent) and RGD13117 (male parent). 0-2 = resistance (R), 3-4 = moderately resistant (MR), and 5-6 = susceptible (S)



Figure 5 Symptoms of rice leaves on 21 day-old F₄ seedlings inoculated MIX 1 compared with Sariceltik and RD6 (susceptible checks), HXBF4 (female parent) and RGD13117 (male parent)

Discussion

DS scores of BPH in the F3 lines with three BPH resistance genes of Bph32, Bph3 and TPS were lower than those of the line without Bph32 and the susceptible check-in both SES and durability tests. This suggested that more resistance genes of Bph gave better BPH resistance than a single gene. The results solidified the gene pyramiding concept that had been proved by other researchers in insect and disease resistance. Hu et al. (2010) reported that Shanyou 63 improved hybrid rice carrying Bph14 and Bph15 had higher resistance than that of Bph14 or Bph15 alone. Besides, the pyramiding of Bph6 and Bph9 in LuoYang 69 improved hybrid rice was found to show stronger resistance than that of a single gene (Wang et al., 2017). Moreover, two Bph genes Bph3 and Bph32 were effective for BPH resistance. The Bph3 that was identified in Rathu Heenati was found that it had resistance to four BPH biotypes

(Pathak & Heinrichs, 1982). The Bph3 has also been used and widely recommended in rice breeding programs for over 30 years (Jena *et al.*, 2015). *Bph32* still an unknown domain-containing protein, but the results in the experiment of Ren *et al.* (2016) were found that *Bph32* was highly expressed of resistance in leaf sheath, and might inhibit feeding in BPH. Besides, it was confirmed that *Bph32* was very valuable for rice defense against BPH (Ren *et al.*, 2016; Jairin *et al.*, 2007).

In this study, all 10 F3 lines carrying *Bph32*, Bph3 and TPS were resistant against two SBR and AYY populations at 9 DAI and 17 DAI. RGD17020-MS373 carrying Bph3 and TPS was resistant against AYY but moderately resistant against SBR at 9 DAI and susceptible at 17 DAI. This suggests that the F3 plants that carrying all three Bph genes especially, the lines carrying *Bph32* in a combination were expressed more resistance levels than those of the progenies carrying only two genes.

The 16 F4 lines derived from one single F3 plant were screened for BL resistance. The results showed that the 16 F4 lines carrying two BL QTLs, qBL1 and qBL11 exhibited a high level of resistance to 7 mixed BL isolates as well as their parent 'HXBF4'. The results were resisted to BL similar to the reported of Manivong et al. (2014) by reporting that, the improved lines in the F5 population from three-way crossed by F5 lines carrying two BL QTLs, qBL1 and qBL11 were resisted against all 15 Lao PDR BL isolates and all 42 Thai BL isolates. This indicated that the two BL QTL used in this study were board-spectrum resistance to BL and numerous studies have also reported the success of BL resistant gene introgression by using these two BL QTLs. Sreewongchai et al. (2010) reported that, the improved lines in F4 population between IR64 and Jao Hom Nin carrying four BL QTLs on chromosomes 2, 12, 1, and 11 showed resistance against 11 Thai blast isolates while the report of Wongsaprom et al. (2010) to qBL1 and qBL11 in BC4F2 of improved lines between IR64 and Jao Hom Nin showed resistance against 8 Thai BL isolates. Besides, the development of aromatic glutinous rice of Khanthong et al. (2018) was reported that the F5 new improved lines carrying qBL1 and qBL11 gave a high level of blast resistance against 4 blast isolates from rice production areas in Thailand, and not only that, the reported of Ruengphayak et al. (2015), Khanthong et al. (2018) and Srichant et al. (2019) were give a similar resistance by using two of these BL QTLs "qBL1 and qBL11".

Conclusion

The validation in the F_3 lines with three *Bph* genes of *Bph3, TPS* and *Bph32* by screening both BPH populations was confirmed that these *Bph*

genes were very effective against BPH. The pyramiding of all three Bph genes in a combination was shown more effectiveness than two genes in a combination (Bph3 and TPS). It also give durability resistance to rice seedlings at 17 DAI. The result of two BL QTLs, qBL1 and qBL11 were shown broad-spectrum resistance to all of 7 mixed BL isolates more than one BL QTL. Thus, it is also strongly suggested that these BPH and BL QTL can be used as the donor parents in other breeding programs. Besides, the genotyping with highthroughput markers for BPH and BL is very accurate and trustworthy. Therefore, Bph32, qBL1 and gBL11 are recommended to use in the breeding program against BPH and BL in Thailand and Lao PDR. However, more validation should be conducted by using other BPH populations and BL isolates to ensure broad-spectrum resistance of these loci.

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