Resistance and susceptibility QTL identified in a rice MAGIC population by screening with a minor-effect virulence factor from Xanthomonas oryzae pv. oryzae

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Summary

Effective and durable disease resistance for bacterial blight (BB) of rice is a continuous challenge due to the evolution and adaptation of the pathogen, Xanthomonas oryzae pv. oryzae (Xoo), on cultivated rice varieties. Fundamental to this pathogens’ virulence is transcription activator-like (TAL) effectors that activate transcription of host genes and contribute differently to pathogen virulence, fitness or both. Host plant resistance is predicted to be more durable if directed at strategic virulence factors that impact both pathogen virulence and fitness. We characterized Tal7b, a minor-effect virulence factor that contributes incrementally to pathogen virulence in rice, is a fitness factor to the pathogen and is widely present in geographically diverse strains of Xoo. To identify sources of resistance to this conserved effector, we used a highly virulent strain carrying a plasmid borne copy of Tal7b to screen an indica multi-parent advanced generation inter-cross (MAGIC) population. Of 18 QTL revealed by genome-wide association studies and interval mapping analysis, six were specific to Tal7b (qBB-tal7b). Overall, 150 predicted Tal7b gene targets overlapped with qBB-tal7b QTL. Of these, 21 showed polymorphisms in the predicted effector binding element (EBE) site and 23 lost the EBE sequence altogether. Inoculation and bioinformatics studies suggest that the Tal7b target in one of the Tal7b-specific QTL, qBB-tal7b-8, is a disease susceptibility gene and that the resistance mechanism for this locus may be through loss of susceptibility. Our work demonstrates that minor-effect virulence factors significantly contribute to disease and provide a potential new approach to identify effective disease resistance.

Introduction

The most significant bacterial disease impacting rice is bacterial blight (BB), caused by Xanthomonas oryzae pv. oryzae (Xoo). In the field, BB severity depends on host genotype, pathogen pressure and environmental conditions. An optimal agronomic practice to manage BB is the deployment of resistant varieties. Historically, available resistance to manage the disease has relied on major resistance (R) genes that detect bacterial virulence effectors. Unfortunately, the efficacy of R-genes is frequently lost over time due to the pathogen’s adaptation to a genotype and the evolution of new pathogenic races (Mew et al., 1992; Quibod et al., 2016). Identifying new sources of resistance that are effective and durable for BB is a pressing challenge for rice production, considering the increasing demand for the grain.

Collectively, effectors dictate bacterial virulence and pathogenicity, and some provide a selective advantage to the pathogen, that is, their loss negatively impacts bacterial fitness (Leach et al., 2001; Ponciano et al., 2003). One strain or pathogen population may carry multiple effectors with redundant function, and some populations can house highly variable effector repertoires (Quibod et al., 2016; Vera Cruz et al., 2000). Furthermore, some effectors have a large effect on pathogen virulence (major effects) and can mask the role of minor-effect virulence factors, effector proteins that have moderate yet significant effects on the host (Bai et al., 2000). Knowing the contribution of both major- and minor-effect factors to pathogen fitness and virulence as well as their gene targets can guide breeding strategies towards stabilizing disease resistance (Leach et al., 2001).

Important to Xoo virulence are transcription activator-like (TAL) effectors, which induce expression of plant host genes (Boch et al., 2014; Hutin et al., 2015; Römer et al., 2010; Yang et al., 2000). Gene induction is dependent on the repeat variable diresidues (RVDs) in the central repeat region (CRR) of the protein. TAL effector RVD’s bind to specific sequences, the effector
binding element (EBE), located in the promoters of target plant genes (Boch et al., 2009; Moscou and Bogdanove, 2009). Frequently, TAL effectors target susceptibility (S) genes in the host genome, enhancing disease severity (Perez-Quintero and Szurek, 2019; Streubel et al., 2013). To avoid pathogen infection, plants have evolved TAL effector-associated resistance [for a review, see (Hutin et al., 2015)]. Included in the latter are dominant TAL effector-dependent resistance mechanisms (Zhang et al., 2015), dominant non-transcriptional dependent (Schornack et al., 2004; Triplet et al., 2016) and resistance through loss of susceptibility (Hutin et al., 2015). The last example includes both natural variation in susceptibility alleles, like Xa5/xa5 (Iyer and McCouch, 2004) and those generated through genome editing technologies where the EBEs of S-genes are edited resulting in resistance (Boch et al., 2014; Oliva et al., 2019).

In a recent comparative genomic study, a total of 181 TAL effectors were identified from ten Philippine Xoo isolates that represented the ten different Philippine races (Quibod et al., 2016). Based on RVD compositions, these effectors were classified into 30 TAL effector families (TEF). The TAL effectors within a family have similar RVD sequences, suggesting functional similarities and shared host gene targets. However, function has only been characterized for nine of the 30 TEF, based on one or two representative major-effect TAL effectors for each family. One TEF containing nine homologues that remains to be characterized is TEF7 (Quibod et al., 2016). Within this family, one member, Tal7b, plays an incremental, but significant role in bacterial virulence and fitness (Bai et al., 2000). Each Xoo strain may have nine or more TAL effectors, with only one or two per strain in the major-effect virulence factor category. Thus, the diversity among the families and the conservation of minor-effect TAL effectors, such as Tal7b, among geographically distinct strains raises the questions ‘why are TAL effectors with minor-effects conserved?’ and ‘what are their host targets?’.

Assuming R-gene durability is linked to the relative importance of effectors to pathogen virulence, one potential strategy to select for long-lasting resistance in the field is to screen for resistance sources that target bacterial fitness factors, including highly conserved minor-effect virulence effectors such as Tal7b (Leach et al., 2001; Vera Cruz et al., 2000). To this end, we used a two-pronged approach aimed at identifying new sources of resistance that target this specific TAL effector. We used a multi-parent advanced generation inter-cross (MAGIC) population generated from eight elite founders from the indica subgroup of rice (Bandillo et al., 2013; Raghavan et al., 2017). This powerful resource captures the genetic diversity of multiple founders that have been recombined over several generations to create a population with large phenotypic diversity ideal for high-resolution trait mapping (Bandillo et al., 2013; Raghavan et al., 2017). In addition, we exploited the functionally and bioinformatically deciphered TAL effector code to predict gene targets for Tal7b within the QTL (Boch et al., 2009; Moscou and Bogdanove, 2009) and to gain insights into the mechanisms for Tal7b resistance and susceptibility.

Here, we report the identification of 18 BB disease resistance QTL, 12 of which target BB strain PXO99A, and six of which are unique to PXO99A overexpressing tal7b. We identified 150 putative Tal7b EBEs within these loci and show that several of the resistance QTL harbour polymorphisms in or near the predicted Tal7b EBE. Our results suggest that Tal7b contributes to Xoo virulence by targeting multiple novel S-genes in rice. Altogether, our data broadens the perspective on effector-triggered susceptibility to emphasize that one effector may target multiple S-genes, that is quantitative susceptibility, and suggests that quantitative resistance can result from inactivation of one or more S-genes.

Results and discussion

Tal7b is a conserved TAL effector in Asian X. oryzae pv. oryzae strains

Tal7b is one of nine TAL effectors in the uncharacterized TAL effector family 7 (Quibod et al., 2016). Originally named ab4.5, Tal7b incrementally contributes to pathogen virulence (Bai et al., 2000), and thus, was selected as a minor-effect virulence factor to screen for resistance sources. The effector gene was cloned from Xoo strain PXO86. Sequence analysis confirmed the cloning of tal7b and showed that the CRR of tal7b is composed of 18 repeats and has RVD sequence: NI HG NI NN HD NS NN NS NN NN HD NN NI NN HD NN NS NG (Figure 1 and Booher et al., 2015).

All complete genomes of X. oryzae were queried for TAL effectors, and their RVD sequences were extracted and classified into TAL effector groups using DiSTAL (Hutin et al., 2015; Perez-Quintero et al., 2015). This analysis showed Tal7b homologues are present in all complete genomes of Asian Xoo strains except in strain JLC28 and were absent in all sequenced Xoo strains of African origin (Figure 1). Tal7b was not detected in X. o. pv. oryzicola (Xoc), the causal agent of bacterial leaf streak of rice. At least one copy of the effector was found in representatives of the ten different Xoo races (Figure 1a). A Tal7b variant with two additional repeats in the CRR was also found in several Asian Xoo strains (Figure 1b). These variants were originally grouped into a separate TAL effector family, TEF26 (Quibod et al., 2016), and are 93% identical to Tal7b at the amino acid level (Figure S1). Although they varied at the DNA level, sequence alignment of Tal7b homologues from all strains indicates 100% similarity in RVD composition and sequence, except from those with the two extra repeats (Figure 1b). Both TEF7 and TEF26, which include Tal7b, are now classified into one Xoo TAL effector group (Group 30) according to the DiSTAL tree (Figure 1; Lang

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Figure 1  Tal7b is conserved among Xanthomonas oryzae pv. oryzae (Xoo) strains of Asian origin. (a) TAL effector profiles from 24 Asian strains of Xoo and one representative strain of African Xoo (BA13), and an Asian X. o. pv. oryzicola (Xoc) strain (BL256) were classified into groups according to DiSTAL tree (Perez-Quintero et al., 2015). TAL groups are listed in order from left to right and conserved TAL effector groups are outlined in red; major TAL effector groups containing PthXo1 and AvrXa7 are boxed in blue; grey/white shading indicates the presence or absence of TAL effector in a strain; black represents copies of that TAL effector or a homolog in that strain. The tree at the top shows the phylogenetic distances between TAL effector groups, and the parsimony tree on the right is based on whole genome sequences. (b) Variation and phylogenetic relationships of Tal7b homologues in Asian strains of Xoo are shown for each TAL effector along with their RVD sequence, TAL ID, strain name and specific geographic origin. The heatmap indicates nucleotide identity for each repeat, where blue represents sequence mismatch as indicated in identity key (top left) and white shading represents 100% match among sequences. Red letters indicate the two RVD where TAL effector family (TEF) 7 and 26 members differ and missing repeats are represented as dashed red lines inside grey boxes.
et al., 2019; Pérez-Quintero et al., 2015). TAL effector groups 23, 30 and 32 are among the most conserved TAL effectors in Xoo genomes (Figure 1A). These results show that Tal7b is among the most conserved TAL effectors in Asian Xoo strains and are consistent with the evolutionary selection for the function of this effector in bacterial fitness.

Tal7b is a minor virulence factor, and its phenotype in interactions with rice is masked by more dominant effector activity (Bai et al., 2000), in the case of PXO99A by PthXo1 (Yang and White, 2004). Thus, to amplify resistance and susceptibility phenotypes in our large-scale screening experiments, we created a Xoo PXO99A strain that overexpresses Tal7b. PXO99A also harbours two Tal7b gene homologues, tal7a and tal8a, that encode proteins 99.6% identical to the amino acid sequence of PXO86 Tal7b and show no variation in RVD sequence (Figure 1b).

We cloned tal7b from PXO86 into the low-copy vector pHMI to create pHMI::tal7b and introduced the clone and empty vector control (pHMI::EV) into PXO99A to create PXO99A pHMI::tal7b and PXO99A pHMI::EV, respectively. We confirmed low expression of Tal7b in the PXO99A wild-type strain and demonstrated moderate overexpression in the PXO99A pHMI::tal7b strains (Figure S2). In the following plant screening experiments, the overexpressing strain PXO99A pHMI::tal7b and the empty vector control (PXO99A pHMI::EV) were used for inoculations.

The indica MAGIC population segregates for resistance to PXO99A pHMI::EV and PXO99A pHMI::tal7b

We took advantage of the high allelic variation in the indica MAGIC population to identify resistance loci for Tal7b (Bandillo et al., 2013; Raghavan et al., 2017). We screened all eight founders alongside 330 F6-8 (6th-8th generation of selfing) advanced intercrossed lines, named Advanced Inbred Lines (AILs), for resistance to PXO99A pHMI::EV or PXO99A pHMI::tal7b. On average, more disease (longer lesion lengths, Tukey adjusted P-value < 0.0001) was observed on AILs challenged with PXO99A pHMI::tal7b compared to those inoculated with PXO99A pHMI::EV indicating potential S-gene targets to Tal7b in the MAGIC population (Figure 2a, b; Tables S1 and S2). When pHMI::tal7b was overexpressed in PXO99A, the overall average lesion length (17.1 cm) was 3.3 cm longer than that for PXO99A pHMI::EV (13.8 cm).

Figure 2 Distribution of the phenotypic response (lesion length in cm) of MAGIC AILs to strain PXO99A with tal7b (pHMI::tal7b) or the empty vector control (pHMI::EV). (a) Violin plots represent the distribution of the least square means (LS-Means) of bacterial blight disease lesion lengths on 330 MAGIC AILs and the eight founders inoculated with PXO99A pHMI::EV (left) or pHMI::tal7b (right). Embedded box and whisker plots represent 50% of the values around the median (bold). The asterisk (*) represents Tukey adjusted P-value < 0.001. (b) Each point on the scatter plot represents the LS-Means of lesion length for each of the 330 AIL screened. The black lines connecting each point highlight the 63 AIL with significantly longer lesion lengths in response to PXO99A pHMI::tal7b (right) compared to PXO99A pHMI::EV (left) (Tukey adjusted P-value < 0.05). Histograms represent the frequency distribution of the lesion length (LS-Means) in the MAGIC AILs after inoculation with (c) PXO99A pHMI::EV and (d) PXO99A pHMI::tal7b. Histograms are overlaid on LS-Means of the lesion lengths for the eight founder parents of the indica MAGIC population. PSBRc82 (F) was not included in the PXO99A pHMI::EV screen in panel c because of poor germination.
These data confirm that Tal7b is a minor-effect virulence factor that contributes incrementally to Xoo virulence.

None of the eight MAGIC founders displayed a strong qualitative resistance phenotype (LL < 5 cm) to either strain tested (Figure 2c, d; Table S3). Of the 330 AILs screened, 24 and 32 showed shorter lesion lengths to PXO99A pHMI::EV and PXO99A pHMI::tal7b than the most resistant parent for each treatment, Samba Mahsuri + Sub1 (LL = 9.8 cm) and PSBRC15B (LL = 11.5 cm), respectively (Figure 2c, d; Table S3). At the opposite extreme, one and 96 of the 330 AILs were more susceptible than the most susceptible parent, IR4527-2B-2-2B-1-1, to PXO99A pHMI::EV (LL = 22.8) and PXO99A pHMI::tal7b (LL = 19.4), respectively (Figure 2c, d; Table S3). The normal distribution and continuous variation observed in the phenotypic response of AIL screened to PXO99A pHMI::EV and PXO99A pHMI::tal7b, in conjunction with the transgressive segregation observed for both strains, indicate that the MAGIC population segregates for resistance and susceptibility to these strains (Figure 2c,d). Moreover, the lack of large reductions or increases in resistance or susceptibility suggests that these traits are controlled by multiple small and moderate effect resistance and susceptibility loci.

Marker-assisted analyses identify loci associated with resistance to Xoo PXO99A pHMI::EV and PXO99A pHMI::tal7b

Interval mapping (IM) and genome-wide association studies (GWAS) were used to map resistance loci for PXO99A pHMI::tal7b (Table S1). IM analyses identified a total of 18 QTL associated with disease resistance to both PXO99A pHMI::EV and PXO99A pHMI::tal7b (Figure S3; Tables S4, S5). Nine QTL, named qBB-PXO99A, were detected in response to PXO99A pHMI::EV on chromosomes 1, 5, 7, 8, 10, 11 and 12 (Figure S3a; Table S4). Similarly, nine QTL were identified to PXO99A pHMI::tal7b and named qBB-tal7b on chromosomes 1, 3, 5, 8, 10, 11 and 12 (Figure S3b; Table S5). Of the 18 QTL detected, only the QTL on chromosomes 5 (qBB-PXO99A-5 and qBB-tal7b-5) and 10 (qBB-PXO99A-10-1 and qBB-tal7b-10-1, qBB-PXO99A-10-2 and qBB-tal7b-10-2) were common to both strains and thus were not unique to Tal7b (Figure S3; Tables S4, S5).

The QTL on chromosome 5, qBB-PXO99A-5 and qBB-tal7b-5, explained 15.7% and 12.6% of the phenotypic variance observed, respectively, and were supported by strong SNP associations in GWAS (Tables S4-S7). Of the 43 significantly associated SNP markers genome-wide, 33 clustered on chromosome 5 (Figure 3; Table S6) and four of these were within the 95% confidence interval of qBB-PXO99A-5 (Table S4, S6). This QTL was previously detected in the indica MAGIC population in screenings for bacterial blight QTL (Bossa-Castro et al., 2018). The QTL and its associated SNP markers overlapped with the mapped bacterial blight susceptibility/resistance gene Xa5/Xa5; a broad-spectrum disease resistance gene that functions independently of TAL effector binding specificity through the naturally occurring recessive allele (xas) of the basal transcription factor TFIIA/CDS (Xa5) (Figures 3, 4a; Blair et al., 2003; Huang et al., 2016; Iyer and McCouch, 2004).

The remaining ten significant markers identified by GWAS for PXO99A pHMI::EV were associated with previously identified QTL for BB on chromosome 11 (qBBR11) (Chen et al., 2012), but did not map to qBB-PXO99A-11, the QTL identified by IM in this study (Figures 3, 4a, Figure S2). The QTL identified on chromosomes 5 and 11, both supported by IM and GWAS, confirm that our approach provides robust data to detect previous and novel QTL conferring resistance to one minor-effect virulence factor and also refine previously identified QTL, as in the case for qBB-tal7b-11, qBB-tal7b-12-1, qBB-PXO99A-12-1 and qBB-PXO99A-12-2 (Figure 4a, Figure S8).

Unique resistance QTL to minor-effect virulence factor Tal7b

For PXO99A pHMI::tal7b, a total of 29 significantly associated SNP markers were identified by GWAS (MLM analysis, P-value < 0.001) on chromosomes 5, 8, 11 and 12 (Figure 3b, Table S7). Twenty of the 29 associated SNP markers for PXO99A pHMI::tal7b clustered on chromosome 5, six of which overlapped with the QTL identified through IM on the same chromosome for PXO99A pHMI::tal7b (qBB-tal7b-5) and PXO99A pHMI::EV (qBB-PXO99A-5) (Figure 3, 4a; Tables S5, S7 and S8). All 20 SNP markers associated with PXO99A pHMI::tal7b on chromosome 5 were also associated with PXO99A pHMI::EV (Tables S6, S7). The remaining nine SNP markers were distributed into two smaller clusters and were unique to PXO99A overexpressing Tal7b (pHMI::tal7b). Four SNP markers overlapped with qBB-tal7b-7-8, the 705 kb resistance QTL to PXO99A pHMI::tal7b on chromosome 8 (Figure 3b, 3d). Four different SNP markers converged on qBB-tal7b-12-2, the 236 Kb QTL on chromosome 12, which also overlapped with the previously described QTL for disease resistance to BB, G-BB-AQBT0219 (Figures 3, 4a; Tables S6, S8) (Tello-Ruiz et al., 2017).

The final marker associated with PXO99A pHMI::tal7b on chromosome 11, S11_28697250, did not overlap with any of the QTL identified by IM on this chromosome in this study, qBB-PXO99A-11 and qBB-tal7b-11 (Tables S4-S7). S11_28697250 did overlap with three previously reported QTL including AQAQ017 (Tello-Ruiz et al., 2017) and RL-RB-qRBr-11.2 (Ashkanai et al., 2013), both of which are associated with rice blast disease resistance, and qBBR11, conferring BB disease resistance (Figures 4a, Figure S8) (Chen et al., 2012). Overall, we identified six resistance QTL to PXO99A pHMI::tal7b, demonstrating the genetic resolution housed in the MAGIC population was sufficient to identify novel QTL for a minor-effect TAL effector.

Tal7b gene targets converge on QTL identified for PXO99A pHMI::tal7b

The deciphered TAL effector code allows for the computational prediction of Tal7b gene targets in the rice genome (Boch et al., 2009; Moscou and Bogdanove, 2009). To explore if putative Tal7b gene targets converged on identified QTL in this study, we used two tools, TALE-NT 2.0 (Doyle et al., 2012) and Talvez (Pérez-Quintero et al., 2013), to predict Tal7b targets in the rice genome (Figure 4b, c). A total of 97,219 genome-wide EBEs were predicted for Tal7b (Figure 4e). Next, the predicted EBEs and their positions were aligned with the physical positions of the significantly associated SNP markers and QTL identified in this study. This analysis revealed 153 EBE in 148 different gene candidates that were (i) predicted Tal7b rice gene targets and (ii) in or near one of the QTL identified in this study (Figure 4d; Table S9). Of the 153 EBE, 126 or 80% converged on qBB-tal7b-1, qBB-tal7b-7-8 and qBB-tal7b-12-2 QTL on chromosome 10 (Figure 4b). The remaining 27 EBEs overlapped with Tal7b unique QTL on chromosome 5, 8, 11 and 12 (Figure 4b; Table S9). Among the top 10, Tal7b gene targets in these QTL are a zinc finger (LOC_Os03g05480), a ribosomal protein (LOC_Os01g14830) and several hypothetical proteins.
Two of the top 10 gene targets with Tal7b EBEs, LOC_Os01g14100 and LOC_Os03g83340, were significantly induced (>2 Log2 Fold Change) in a susceptible rice variety when challenged with PXO99A (Perez-Quintero et al., 2018) and one, LOC_Os01g135770, was significantly down-regulated (Perez-Quintero et al., 2018). Additional genome-wide putative targets include homologues of the nodulin MtN3 family proteins that encode SWEET sucrose transporter genes, which have been reported as major TAL effector susceptibility gene targets in rice, citrus and cotton (Cohn et al., 2014; Cox et al., 2017; Yang et al., 2006). However, the putative Tal7b OsSWEET gene targets are not located within the Tal7b unique QTL identified in this study (Figure 4a; Table S8). Other targets near the identified QTL included a putative brassinosteroid insensitive 1-associated receptor kinase 1 precursor (OsSerk1/OsBAK1, LOC_Os01g135770), a putative dehydrin (DHN, LOC_Os11g26790), a calmodulin-binding protein (CaM-binding protein, LOC_Os12g36920) and a spotted leaf 11 gene (SPL11, LOC_Os12g38210) (Figure 4a; Table S8).

Next, using data from the 3K rice genome project (Wang et al., 2018), we aligned and analysed the promoters of the putative Tal7b rice gene targets in each of the eight indica MAGIC founders. We focused on the targets that are unique to PXO99A pHMI::tal7b, and thus, excluded targets in QTL qBB-tal7b-5, qBBtal7b-10-1 and qBBtal7b-10-2 (Figure 4a). Of the remaining 119 Tal7b putative gene targets, we identified in at least one of the eight MAGIC founders: 23 EBEs with polymorphic sites in the EBE sequence (Figure 5a; Table S9); 54 with downstream or upstream shifts surrounding the EBE sequence (Figure 5b; Figure 3 Detection of QTL conferring resistance to PXO99A carrying pHMI::EV or pHMI::tal7b. Manhattan plots and simple interval mapping show the genomic location on the x-axis and the negative logarithm of the P-values on the y-axis. Solid blue line indicates a significance threshold of P ≤ 0.001. (a, b) GWAS results show significant associations to (a) PXO99A pHMI::EV on chromosomes 5 and 11 (cyan; Table S7) and (b) PXO99A pHMI::tal7b on chromosomes 5, 8, 11 and 12 (magenta; Table S8). (c, d) Simple interval mapping featuring unique QTL to Tal7b on chromosome 8, qBB-PXO99A-8 and qBB-tal7b-8. Shaded regions indicate 1-LOD supporting intervals for (c) qBB-PXO99A-8 to PXO99A pHMI::EV, which is distinct from the peak for (d) qBB-tal7b-8 to PXO99A pHMI::tal7b.
Table S9); 18 with indels in the EBE sequence (Figure 5c; Table S9); and 10 which lacked the EBE sequence in the promoter region of the putative target gene (Figure 5d; Table S9). In a survey of published transcriptome data, 21 of the 119 corresponding putative Tal7b targets were differently expressed in rice inoculated with at least one of the three Xoo strains harbouring Tal7b (Table S9). Overall, the polymorphisms in the EBE sequence or in the promoter region and the corresponding changes in target gene expression are reminiscent of resistance through loss of susceptibility and are consistent with a model that allelic variation at these positions may serve as a source of recessive resistance against Xoo strains harbouring Tal7b (Figure 5f-h).

qBB-tal7b-8 harbours a susceptibility allele to PXO99A pHMI::tal7b

The QTL that mapped to chromosome 8 and was unique for PXO99A pHMI::tal7b, named qBB-tal7b-8, was selected to further characterize the resistance mechanism governing this locus. Fourteen MAGIC AILs (AIL: 13, 24, 156, 163, 197, 210, 219, 222, 225, 229, 234, 237, 250 and 267) with resistant and susceptible marker alleles (Table S7) on chromosome 8 were selected from the 330 AILs originally screened. These lines and Nipponbare, which served as a susceptible control, were re-screened for their responses to PXO99A pHMI::EV and PXO99A pHMI::tal7b. A subset of these lines (13, 219, 229) and Nipponbare showed significantly longer lesions (4.7 cm, P-value < 0.001) in response to PXO99A pHMI::tal7b (average LL = 16.9 cm) than to PXO99A pHMI::EV (average LL = 12.2 cm) (Figure 5e; Table S11). The remaining 11 AILs that were re-screened, including lines 24, 163, 197 and 210 (Figure 5e; Table S11), did not differ in the observed lesion lengths to either bacterial strain, regardless of the presence or absence of pHMI::tal7b. Among these 11 non-responsive lines, the average lesion length for PXO99A pHMI::tal7b was 13.1 cm and 12.5 cm for PXO99A pHMI::EV. The absence of a significant lesion length difference in the majority of lines with variant alleles in qBB-tal7b-8 suggests that this QTL houses a minor yet significant susceptibility allele responsive to Tal7b, and that the resistance observed may be through loss of susceptibility, as proposed in Figure 5g-h.

Conclusions

Long-lasting disease resistance is a highly sought-after trait for crop breeding programmes. However, achieving durable resistance is a challenge because pathogen populations continually evolve to overcome deployed host resistance. The pathogen population changes occur through mutation of the pathogen effector genes targeted by specific plant resistance genes, or through invasion or build-up of novel pathogen populations that do not harbour the targeted effector. Many widely studied Xoo effectors are TAL effectors, including PthXo1 and AvrXa7, which are major virulence factors (Bai et al., 2000; Yang and White, 2004), but these are not widely conserved in the global pathogen population (Figure 1; Lang et al., 2019; Quibod et al., 2016)). In contrast, a number of TAL effectors, such as Tal7b and related effectors, are individually minor contributors to pathogen virulence (Figure 2; Bai et al., 2000), and their distribution is widespread in Xoo populations (Figure 1). We and others postulate that incorporating resistances that target these widespread effectors will help stabilize plant disease resistance, increasing its durability (Leach et al., 2001; Mundt, 2014). Towards this end, we employed an innovative rice MAGIC population that contains the genetic diversity of eight indica parents (Bandillo et al., 2013; Bossa-Castro et al., 2018) to identify potential durable disease resistance loci that target the widespread minor virulence effector Tal7b.

Six QTL specific to the Xoo strain overexpressing Tal7b were identified (Figure 4), suggesting that effector Tal7b may target more than one gene in the rice genome. The use of computationally predicted EBEs for Tal7b allowed for identification of gene candidates in or near each qBB-tal7b-unique QTL (Figure 4b, c), and mapping of the EBE to promoter sequences for candidate genes mined for the MAGIC founder lines allowed detection of several types of promoter variations relative to the EBE (Figure 5). The promoter variants detected in at least one or more of the eight founder parents suggested that the resistance we detected in this study against Tal7b may be through a loss of susceptibility. Indeed, expanded phenotyping of AILs harbouring qBB-tal7b-8, a unique Tal7b QTL, revealed that AILs lacking the Tal7b target EBE in one gene target did not show enhanced susceptibility to Tal7b (Figure 5e). In contrast, AILs with the intact allele were more susceptible to Xoo with Tal7b. Thus, we suggest that the resistance mechanism governing qBB-tal7b-8 is through loss of susceptibility (Antony et al., 2010; Huang et al., 2016; Hutin et al., 2015; Iyer and McCouch, 2004).

Overall, our study demonstrates that pathogen virulence factors that contribute quantitatively to disease can facilitate identification of effective minor disease resistance QTL for bacterial blight of rice. The indica MAGIC population, which captures the genetic diversity of eight parents (Bandillo et al., 2013; Raghavan et al., 2017), allowed for identification of novel QTL targeting Tal7b and for refinement of previously identified regions targeting Xoo. Because the parents of the indica MAGIC population are elite varieties, introgression of the identified resistance loci into commercial, farmer accepted varieties can be accelerated due to minimal linkage drag. While this study focuses on resistance to a bacterial disease of rice, the strategy for finding novel resistance based on knowledge of effector-targeted susceptibility and the application of the MAGIC genetic resources is broadly applicable to other crops and diseases.
Methods

Bacterial strains, inoculum preparation, plant material and growth conditions

The bacterial strains and plasmids used in this study are listed in Table S12 along with relevant characteristics. Xoo strains were cultured on Peptone Sucrose Agar (PSA) (Karganilla et al., 1973) at 28 °C and Escherichia coli strains were cultured on Luria–Bertani (LB) medium (Miller, 1992) at 37 °C. When necessary, antibiotics were added to the medium at the following concentrations: 50 µg/mL streptomycin, 50 µg/mL spectinomycin and 2 µg/mL tetracycline. To prepare inoculum, all strains were cultured for 24 h at 28 °C on PSA amended with streptomycin and spectinomycin. Bacteria were suspended in sterile water to 0.2 OD600, approximately 10^8 CFU/mL.

Rice varieties used in this study include 330 AILs from the indica MAGIC population (Bandillo et al., 2013; Bossa-Castro et al., 2018; Raghavan et al., 2017) at the sixth to eighth generation of selfing (S6-S8), as well as the eight founding parents: Fedearroz 50, IR45427-2B-2-2b-1-1, IR4630-22-2-5-1-3, IR77298-14-1-2-10, PSBRC82, Samba Mashuri + Sub1, Sanhuangzhan-2 and PSBRc158. Nipponbare served as a susceptible control, respectively. Plants were grown and cared for as described (Bossa-Castro et al., 2018). Rice lines were grown in triplicate using an incomplete random block (IRB) design. For inoculations, the two youngest, completely expanded leaves of each six-week-old rice plant were leaf clip-inoculated with bacterial suspensions as described (Kauffman et al., 1973). Lesion lengths (LL) were measured in centimetres (cm) at 14 days postinoculation (dpi).

Lesion lengths were analysed using SAS® software version 9.4 (SAS Institute Inc., 2013). Each of the two inoculated leaves per plant was treated as technical replicates and used to calculate simple means for each biological replicate. All biological replicate value averages were then averaged using the least square means (LS-Means) function in PROC MIXED. Bacterial strain, line and bacterial strain by line interaction were treated as fixed effects. Block and replicates were treated as random effects. P-values were calculated using Tukey’s method for multiple comparisons. AILs with missing phenotypic data were removed from all analyses.

Plasmid constructs

The tal7b gene, previously named a84.5 (Bai et al., 2000), was subcloned from a cosmid clone pXO6-33 derived from Xoo strain PXO86 (Hopkins et al., 1992), GenBank accession No. AJQ83546.1 (Booher et al., 2015), to generate pBS45.5b. The CRR of tal7b was cloned as an Spfi fragment into entry vector pCS466 to generate pCS466::tal7b. pCS466 is a derivative of the Gateway entry vector pCR8-GW (Invitrogen) that contains a truncated form of the X. oryzae pv. oryzicola BLS256 tal1c gene, from which the Spfi fragment that comprises the repeat region has been removed (Verdier et al., 2012). Using Gateway LR Clonase (Invitrogen), the tal7b CRR flanked by the tal1c N- and C-termini was transferred to the destination vector pKEB31 (Cermak et al., 2011), Addgene plasmid 31224 (www.addgene.org), to create pKEB31::tal7b for constitutive expression in Xanthomonas. Finally, tal7b was cloned into the low-copy cosmider vector pHM1 (Hopkins et al., 1992) by digesting pKEB31::tal7b with HindIII HF to extract the tal7b CRR flanked by the tal1c domains (Verdier et al., 2012). The resulting plasmid, pHM1::tal7b was transformed into Xoo PXO99A by electroporation (Choi and Leach, 1994). Transformants were selected on nutrient agar containing appropriate antibiotics. Colony PCR with primers F4: CGCAATG- CACTGACGGTGC and R2458: CATGCAAAGACGCCT- GATCCGG confirmed the presence of tal7b CRR. The PCR programme was 96 °C for 4 min, 25 repeats of 15 s at 96 °C, 30 s at 58 °C, 45 s at 70 °C and a final 70 °C for 4 min. Integrity of tal7b was confirmed by Sanger sequencing.

Genotyping

The 330 AILs and the eight founder parents used in this study were genotyped as part of a larger study conducted by the International Rice Research Institute. Sample preparation, genotyping by sequencing (GBS), data collection and analysis were carried out as described (Bandillo et al., 2013; Raghavan et al., 2017). Briefly, GBS libraries were generated using the 96-plex apeki-based protocol according to (Elshire et al., 2011). Raw reads, publicly available at http://snpsseek.irri.org/download.zul, were trimmed and aligned to the Nipponbare reference genome MSUv7 (Kawahara et al., 2013). SNP calling was conducted prior to filtering as described in Raghavan et al. (2017). With a call rate of >70% and a minor allele frequency of >0.05, a total of 14,561 SNP markers remained to cover the rice genome. Approximate marker density was one SNP marker per 25.6 Kb (Figure S4). Overall SNP marker density allowed for genome wide association studies and QTL detection. SNP markers were named to indicate genomic location, that is the number following S is the chromosome number followed by the bp coordinate of the SNP in the MSU7 Nipponbare reference genome (Tables S6, S7).

Interval mapping

Interval mapping was carried out using the mpIIM function from R package mpMap, a platform designed for multi-parent populations (Huang and George, 2011). A linkage map was generated with the SNP data of each AIL and founder, as well as line pedigree. Interval mapping assumed true marker positions, no covariates and a step size of 1 cM. QTL position, founder effect estimate and allele effect were determined as described (Tables S4, S5; Bossa-Castro et al., 2018; Raghavan et al., 2017).

Genome-wide association studies

GWAS was performed as described (Bossa-Castro et al., 2018). Briefly, a kinship matrix was generated for the MAGIC population, excluding parents, to account for population structure using TASSEL version 5.0.2 (Bradbury et al., 2007). Statistics and allele effects were generated in TASSEL with the mixed linear model (MLM) function. Q-values, which assess the false discovery rate from GWAS, were calculated with q-value package in R (Storey and Tibshirani, 2003). The linear model settings were set to default, where compression level was calculated using the optimum level and the variance component estimated using the P3D method. Results are presented in Manhattan plots generated using the qqman R package (Turner, 2014).

Confirmation of resistance phenotype on chromosome 8

MAGIC AILs with the variant allele in the resistance locus on chromosome 8 (13, 24, 156, 163, 197, 210, 219, 222, 225, 229, 234, 237, 250, 267) were selected to confirm the resistance phenotype identified on chromosome 8. Nipponbare was used as a susceptible control. Six plants per AIL served as biological replicates and two leaves per plant served as technical replicates for each bacterial treatment. The two bacterial treatments were
PXD99\(^a\) pHMI:EV or PXD99\(^a\) pHMI:tal7b. At 14 dpi, lesion lengths were measured and data were analysed using one-way ANOVA. Statistical analyses were carried out using JMP Pro 13 (SAS Institute Inc., Cary, NC).

**Physical map of QTL identified in this study, predicted Tal7b gene targets and known resistance genes/QTL**

All reported BB resistance genes and QTL compiled in (Djedatin et al., 2016) were used to generate a physical map of the QTL identified in this study with respect to all known BB resistance loci. The maps were generated with tidyverse, a collection of R packages that include ggplot2 (Wickham, 2017). The physical position of resistance and susceptibility genes, QTL and predicted Tal7b gene targets were then superimposed on each chromosome.

**Tal7b-predicted targets**

Tal7b gene targets were predicted using Talvez v3.1 and TALE-NT 2.0 (Doyle et al., 2012; Pérez-Quintero et al., 2015). Both programmes were run against the Nipponbare v. MSU7 rice reference genome (Kawahara et al., 2013). Predicted EBE were compared between both Tal prediction programmes, and, those that overlapped and were in close proximity to identified QTL were subjected to further analysis.

**Gene expression data retrieval and analysis**

Gene expression data used in this study were retrieved from the daTALbase \(\text{http://bioinfo-web.mpl.ird.fr/cgi-bin2/datalbase/home.cgi}\) (Pérez-Quintero et al., 2018).

**Western blot analysis of Tal7b**

Overnight liquid cultures of bacterial strains were pelleted and washed twice with deionized (DI) water. Bacterial pellets were resuspended in DI water and adjusted to an OD\(_{600}\) of 0.5. Cells were lysed, and protein was denatured with Laemmli buffer and heat. Total protein was separated on a 10% polyacrylamide gel followed by transfer to nitrocellulose. Tal7b was detected by TAL antibody (a gift from Adam Bogdanove, Cornell University), diluted 1:5000 and goat anti-rabbit HRP secondary antibody, diluted 1:2500.

**Allelic variation in putative EBE sites and target gene promoter regions**

Polymorphisms affecting putative Tal7b EBE sites of 113 target genes were examined in the region spanning the start codon to 1kb upstream of the EBE site. Variant Call Format (VCF) data from the rice 3K genomes project (Wang et al., 2018), based on an alignment to IR64 (reference genome for indica subgroup), were extracted from the genomes of the eight MAGIC parents (https://registry.opendata.aws/3kricgenome/). Using a python script, four types of mutations were identified and characterized: nucleotide substitutions within EBE site, insertions/deletions within the EBE site, insertions/deletions downstream or upstream of EBE site or complete deletion of EBE and/or surrounding promoter region (Table S9). Consensus sequences for each region were generated using the VCF data and were aligned using Clustal Omega for visualization (https://www.ebi.ac.uk/).

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**Conflict of interest**

The authors declare no conflict of interest.

**Author contributions**

A.I.H., E.E.D., A.M.B.-C., H.L., V.V. and J.E.L. designed research; A.I.H., E.E.D., A.M.B.-C., B.W.T., R.C., A.P.-Q and V.V. performed research; A.I.H., E.E.D., A.M.B.-C. and J.E.L. analysed data; A.I.H., E.E.D. and J.E.L. wrote the paper.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Figure S1** Amino acid identity matrix for the 15 Tal7b homologues in geographically distinct Asian strains of X. o. pv. oryzae.

**Figure S2** Western blot confirming overexpression of Tal7b in PXO99A (pHMI::tal7b). WT = PXO99A. The product of the tal7b gene, marked with arrow, is 124 kDa.

**Figure S3** Interval mapping shows QTL detected to Xoo (a) PXO99A pHM1::EV and (b) PXO99A pHM1::tal7b in the indica MAGIC population.

**Figure S4** SNP Marker Density.

**Table S1** Indica MAGIC population response to PXO99A pHM1::EV and pHM1::tal7b.

**Table S2** Mixed model ANOVA for the indica MAGIC population screened in response to PXO99A pHMI::EV and PXO99A pHMI::tal7b.

**Table S3** Response of indica MAGIC parents to PXO99A pHM1::EV and PXO99A pHM1::tal7b.

**Table S4** QTL detected in indica MAGIC population to XooPXO99A pHM1::EV.

**Table S5** QTL detected in indica MAGIC population to XooPXO99A pHM1::tal7b.

**Table S6** Significant SNPs associated with disease resistance to XooPXO99A pHM1::EV in indica MAGIC population.

**Table S7** Significant SNPs associated with disease resistance to XooPXO99A pHM1::tal7b in indica MAGIC population.

**Table S8** Resistance QTL identified to PXO99A pHMI::EV and PXO99A pHMI::tal7b and putative Tal7b gene targets in reference to previously described resistance loci for bacterial blight in rice.

**Table S9** Gene induction of putative Tal7b gene targets in Nipponbare and IR24, two susceptible rice verities to X. o. pv. oryzae, and genomic variation in the EBE and coding sequence of the same genes in the eight MAGIC founders.

**Table S10** Phenotypic response of 14 indica MAGIC advanced inbred lines (AIL) with variant allele in qBB-tal7b-8 on chromosome 8 to PXO99A pHMI::tal7b.

**Table S11** Bacterial strains and plasmids used in this study.