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Improved bacterial leaf blight disease resistance in the major elite Vietnamese rice cultivar TBR225 via editing of the OsSWEET14 promoter

Phuong Nguyen Duy1*, Dai Tran Lan1,2*, Hang Pham Thu1, Huong Phung Thi Thu1, Ha Nguyen Thanh1, Ngoc Phuong Pham1*, Florence Auguy3, Huong Bui Thi Thu4, Tran Bao Manh5, Sebastien Cunnac3, Xuan Hoi Pham1*

1 Department of Molecular Pathology, Institute of Agricultural Genetics, Vietnam Academy of Agricultural Sciences, Hanoi, Vietnam, 2 Faculty of Natural Sciences, Department of Applied Biology and Agriculture, Quynhon University, Quynhon, Vietnam, 3 PHIM Plant Health Institute, Univ Montpellier, IRD, CIRAD, INRAE, Institut Agro, Montpellier, France, 4 Vietnam National University of Agriculture, Hanoi, Vietnam, 5 ThaiBinh Seed Corporation, Thaibinh, Vietnam

* These authors contributed equally to this work.
Current address: Faculty of Biology, Hanoi University of Sciences, Hanoi, Vietnam
*xuanhoi.pham@gmail.com

Abstract

TBR225 is one of the most popular commercial rice varieties in Northern Vietnam. However, this variety is highly susceptible to bacterial leaf blight (BLB), a disease caused by Xanthomonas oryzae pv. oryzae (Xoo) which can lead to important yield losses. OsSWEET14 belongs to the SWEET gene family that encodes sugar transporters. Together with other Clade III members, it behaves as a susceptibility (S) gene whose induction by Asian Xoo Transcription-Activator-Like Effectors (TALEs) is absolutely necessary for disease. In this study, we sought to introduce BLB resistance in the TBR225 elite variety. First, two Vietnamese Xoo strains were shown to up-regulate OsSWEET14 upon TBR225 infection. To investigate if this induction is connected with disease susceptibility, nine TBR225 mutant lines with mutations in the AvrXa7, PthXo3 or TalF TALEs DNA target sequences of the OsSWEET14 promoter were obtained using the CRISPR/Cas9 editing system. Genotyping analysis of T0 and T1 individuals showed that mutations were stably inherited. None of the examined agronomic traits of three transgene-free T2 edited lines were significantly different from those of wild-type TBR225. Importantly, one of these T2 lines, harboring the largest homozygous 6-bp deletion, displayed decreased OsSWEET14 expression as well as a significantly reduced susceptibility to a Vietnamese Xoo strains and complete resistance to another one. Our findings indicate that CRISPR/Cas9 editing conferred an improved BLB resistance to a Vietnamese commercial elite rice variety.
Introduction

Bacterial leaf blight (BLB) caused by *Xanthomonas oryzae pv. oryzae* (*Xoo*) is a major bacterial disease that causes 10%-20% annual reduction in rice production worldwide [1]. The use of improved rice varieties resistant to *Xoo* is probably the most efficient, economical and environmentally-friendly way to control BLB.

The virulence of *Xoo* depends on the transcriptional activation of specific host disease-susceptibility (*S*) genes by a subgroup of bacterial type III effectors, called transcription activator-like effectors (TALEs) [2]. Upon translocation into the plant cell, TALEs bind to specific host nuclear gene promoter sequences termed Effector-Binding Elements (EBEs) and induce target gene expression to the benefit of the pathogen. The central repetitive domain of TALEs is responsible for DNA target sequence binding. DNA binding involves recognition principles that have been largely deciphered and applied to the computational prediction of TALEs target DNA sequences [3,4]. This and earlier work has fostered the identification of TALEs transcriptional targets in the rice genome and ultimately, of rice BLB *S* genes [2].

All *Xoo* strains recurrently target *S* genes belonging to the *SWEET* gene family and coding for transmembrane sugar exporter proteins [3]. The over accumulation of SWEETs due to TALE induction is presumed to provide an additional ration of apoplastic carbohydrates for full bacterial pathogen multiplication and disease expression [5]. Although all five rice clade III *SWEET* genes can function as *S* genes for bacterial blight, only three, namely *OsSWEET11*, *OsSWEET13* and *OsSWEET14*, are known to be targeted by several unrelated TALEs in nature [6–11]. *OsSWEET11* is activated by PthXo1 [6], *OsSWEET13* is targeted by different variants of PthXo2 [11,12], while *OsSWEET14* is a target of multiple TAL effectors, including AvrXa7, PthXo3, TalC and TalF [7–9].

Previous studies established that rice resistance to *Xoo* resulting from "TALE-unresponsive" alleles can be conferred by natural DNA polymorphisms or targeted editing of EBEs located in *OsSWEET* genes promoters of rice germplasm accessions or engineered rice varieties, respectively [6,13–16]. For example, early resistance engineering work has used TALENs to individually alter the AvrXa7, TalC or TalF EBEs in the *OsSWEET14* promoter and successfully obtained resistance to some Asian *Xoo* strains [13,15]. However, strains collected in Asian countries such as China, Japan, Philippines, Taiwan, Thailand, India, Nepal or South Korea can express combinations of up to three major TALEs redundantly targeting clade III *OsSWEET* genes with either PthXo3 or AvrXa7 being occasionally associated with PthXo2 [11,17]. Broad BLB resistance engineering thus required multiplex *OsSWEET* promoters EBE editing using the CRISPR/Cas9 system [11,12].

The clustered regularly interspaced short palindromic repeats/CRISPR-associated protein-9 nuclease (CRISPR/Cas9) system is a simple and efficient gene-editing tool developed in the past few years [18,19]. Moreover, the targeted mutations generated by CRISPR/Cas9 can be stably transmitted to the next generation. Thus, CRISPR/Cas9 has become a routine tool in plant laboratories around the world to create various mutants for many applications, including the genetic improvement of crops [20].

BLB is a major rice disease which occurs in many rice cultivating areas of Vietnam [21,22]. Most Vietnamese commercial rice varieties, including TBR225, are susceptible to BLB, resulting in annual yield loss of about 15–30% on average [23]. A few studies have identified rice resistance genes effective against Vietnamese *Xoo* lineages [22,23]. However, no information is currently available on the nature of Vietnamese *Xoo* TALEs and their corresponding *S* genes. Despite the large number of mapped rice BLB resistance genes [24,25], there is an need for alternative breeding approaches that enable the rapid introduction of broad BLB resistance in elite varieties in order to cope with swift pathogen populations adaptive shifts in the fields [11,26].
Here, we report on the identification of OsSWEET14 as a transcriptional target of Vietnamese Xoo. CRISPR/Cas9-mediated mutagenesis of the OsSWEET14 promoter in TBR225, a major elite variety in rice production areas of North Vietnam, is shown to confer BLB resistance without detectable yield penalty. The current study found this quintessential S gene to be associated with the virulence of Vietnamese Xoo strains. This is an important step for the future design and implementation of broad-spectrum BLB-resistance in elite rice varieties using genome editing in Vietnam.

Materials and methods

Plant and pathogen materials

Rice cultivar TBR225 (Oryza sativa L. ssp. indica) were obtained from ThaiBinh Seed Cor. [27]. All edited and wild-type (WT) TBR225 plants were grown in a net-house under the following average conditions: 30˚C for 14 h (light) and 25˚C for 10 h (dark) with 80% humidity. The Xoo VXO_11 and VXO_15 strains used in this study were isolated from diseased leaves collected in Hanoi-Vietnam in 2013 and 2016, respectively. Bacteria were cultured as described in Zhou et al. (2015) [28].

Gene expression analysis

Gene expression analyses were carried out as described previously [29] by RT-PCR method. The rice leaves were infiltrated with the indicated bacterial strains and used for total RNA extraction 48 h post inoculation using the TRIZol reagent (Invitrogen, USA). One microgram of RNA was used for each RT-PCR with oligo (dT) primer followed by PCR with OsSWEET14-specific primers (forward 5'-ACTTGCAAGCAAGACAGTGT-3' and reverse 5'-ATGTTGCCTAGGAGACCAAAGG-3'). An Eppendorf Mastercycler ep Gradient S was used for 35 PCR cycles. The OsEF1α gene was used as a constitutive control [15] using specific primers (forward 5'-GAAGTCTCATCCTACCTGAAGAAG-3' and reverse 5'-GTCAAGAGCCTCAAGCAAG-3').

gRNA design

The OsSWEET14 promoter (GenBank, accession number: AP014967.1) was amplified by PCR with forward primer 5'-TTGCGGCTCATCAGTTTCTC-3' and reverse primer 5'-CTAGGAGACCAAAGGCGAAG-3' from genomic DNA of TBR225 rice plants and ligated in pGEM-T Easy vector (Promega) for sequencing. The gRNA target sequence (Fig 1A) for editing the TBR225 OsSWEET14 promoter was designed based on the sequence of the cloned TBR225 OsSWEET14 promoter using a combination of two bioinformatics tools CRISPR-P v2.0 [30] and CCTop [31]. A gRNA sequence with high on-target and low off-target scores in both prediction tools was chosen for vector construction.

Vector construction

The Cas9 rice expression vector (pUbi-Cas9) [32] and the sgRNA expression vector (pENTRsgrRNA) under the control of the OsU6 promoter [33] were used to construct the pCas9/OsSWEET14-gRNA expression vector. The complementary oligonucleotides with appropriate 4-bp overhangs were synthesized by Macrogen (Korea). After heat denaturation, the complementary oligonucleotides (5'-gtgtGGTGCTAAGGGCTCAAGCC-3' and 5'-aadacGGCTTGATGAGCTTAGCACC-3') were first annealed to each other, phosphorylated, and ligated into the Bsal-digested vector pENTR-sgRNA. The integrity of the inserted fragment was verified by sequencing. Subsequently, the sgRNA cassette was cloned into pUbi-Cas9 using the Gateway
LR clonase (Life Technologies) (Fig 1B). The resulting construct was confirmed by Sanger sequencing of the insertion junctions.

**Agrobacterium*-mediated rice transformation**

The pCas9-OsSWEET14-gRNA was electroporated into *Agrobacterium tumefaciens* EHA105 and the resulting strain was used to transform rice using the method described by Hiei et al. (1994) [34]. The presence of the transgene in the genome of T₀ hygromycin-resistant plants or segregating T₁ individuals was evaluated by PCR using 5'-ATGGCCCCAAAGAAGAAG-3' and 5'-GCCTCGGCTGTCTCGCCA-3' primers specific for Cas9. T₁ individuals were analyzed by PCR using Cas9, OsSWEET14-gRNA (5'-GGATCATGAACCAACG-3' and 5'-GAATTCGATATCAAGCTT-3') and HPT (5'-AAACTGTGATGGACGACACCGT-3' and 5'-GTGGCGATCCTGCAGCTCC-3') specific diagnostic primer pairs together with a positive control pair (5'-TTGCGGCTCATCAGTTTCTC-3' and 5'-TGGATCAGATCAAAGGCAAC-3') specific to the OsS-WEET14 promoter.

**Bacterial blight inoculation**

Rice cultivation and disease assays were done according to the methods of Blanvillain-Baufumé et al. (2017) [15]. Bacteria were cultured in PSA media (10 g/liter peptone, 10 g/liter sucrose, 1 g/liter glutamic acid, 15 g/liter Bacto Agar) at 28°C for two days [35] and inoculated at an optical density (OD₆₀₀) of 0.5 (infiltrations) or 0.4 (leaf clipping) in water. For lesion length measurements, at least three inoculated leaves per plant and three plants for each line were measured 14 days after inoculation (DAI), and scored as follows: high resistance (lesion length < 8 cm), moderate resistance (lesion length 8–12 cm) and susceptibility (lesion...
length > 12 cm). For gene expression analyses, 4-cm leaf sections infiltrated with bacterial suspensions were collected at 48 h after inoculation for RNA extraction. Experiments included samples from three pooled biological replicate leaves. The plants inoculated with distilled water only were used as negative controls.

**Analysis of OsSWEET14 edited allele sequences**

To determine the nature of the mutation at the target site, all transgenic T₀ or T₁ plants were analyzed by PCR using genomic DNA (50 ng) as a template and OsSWEET14 specific primers (5’-TTGGCGGTCTCATCAGTTTCTC-3’ and 5’-TGGATCAGATCAAAGCAAC-3’). The PCR products were directly sequenced using the Sanger method. The sequencing chromatograms were decoded using the Degenerate Sequence Decoding method [36] in order to identify the mutations.

**Evaluation of major agronomic traits under net-house conditions**

WT and selected mutant plants were planted under net-house conditions in a randomized pot design experiment. At maturity, five plants of each line were investigated for the following agronomic traits: growth duration, plant height, number of tillers per plant, number of grains per panicle, number of filled grains per panicle and yield (seed mass) per plant. The experiment was repeated three times, so a total of fifteen plants were evaluated for each line.

**Analysis of potential off-target editing**

Off-target sequences were predicted with the CCTop tool [31] against the OsSWEET14 promoter sgRNA and the rice Nipponbare genome with default parameters. A total of 18 potential off-target sequences were identified. Three of them were located in coding regions (S2 Table). These regions were amplified by PCR using the specific primers listed in S2 Table and analyzed by sequencing.

**Results**

**Vietnamese Xoo strains induce OsSWEET14 during infection of the TBR225 rice variety**

OsSWEET14/Os11N3 was previously identified as a susceptibility gene for Xoo strains relying on either of the AvrXa7, PthXo3, TalF (formerly Tal5) or TalC TALEs for infection of the rice cultivars Nipponbare and Kitaake [11]. Because Xoo strains tend to frequently target this gene, we first sequenced a region of the OsSWEET14 promoter from rice cultivar TBR225 to examine if it also carries documented target EBEs. Based on the Nipponbare genome sequence in database (AP014967.1), the region encompassing 1343 bp sequence upstream and 52 bp sequence downstream of the predicted transcription start site of OsSWEET14 gene from TBR225 rice cultivar was PCR amplified and sequenced (S1 Fig). The promoter region including the putative TATA box (TATAAA) and the AvrXa7, PthXo3, TalF/Tal5 and TalC EBEs (Fig 1A), located 319 bp to 216 bp upstream of the ATG initiation codon, showed 100% identity to the Nipponbare sequence. This therefore implied that in principle, the TBR225 OsSWEET14 promoter can be recognized by characterized major Xoo TALEs.

As shown in Fig 2A, we also challenged TBR225 plants with two Vietnamese Xoo strains VXO_11 and VXO_15, both originating from the Hanoi province, using leaf clipping assays. We consistently obtained typical extended disease lesions 14 days after inoculation (25.5 cm and 26.6 cm average lesions length for VXO_11 and VXO_15, respectively in the experiment of S2 Fig), indicating that the TBR225 variety is susceptible to BLB.
To test if OsSWEET14 is a potential direct virulence target of Vietnamese Xoo strains, we infiltrated TBR225 rice leaves with the two Vietnamese Xoo strains. Forty-eight hours post infiltration, TBR225 plants inoculated with VXO strains displayed a strong induction of OsSWEET14 relative to water controls (Fig 2B). These results suggest that OsSWEET14 is a transcriptional target of VXO strains and that it may act as a susceptibility gene in TBR225.

CRISPR/Cas9 design for OsSWEET14 promoter editing

Our main objective was to engineer resistance to BLB caused by Vietnamese Xoo strains. To this end, we subsequently sought to specifically modify the OsSWEET14 promoter in TBR225 rice with CRISPR/Cas9-mediated editing. Previous work revealed that while African Xoo strains rely on TalC and occasionally, TalF, all Asian Xoo strains use either PthXo3- or AvrXa7-like TALEs to activate OsSWEET14 [11]. Because the talC gene is currently exclusively found in African strains, we reasoned that it is unlikely that Vietnamese strains carry a talC copy. Thus, to maximize our chances to perturb all remaining documented EBEs, we selected a 20-bp nucleotide target site overlapping the PthXo3, AvrXa7 and TalF EBEs and having a predicted cut site located near the 3'-end of the AvrXa7 EBE (Fig 1A). The recombinant binary plasmid pCas9/OsSWEET14-gRNA for CRISPR/Cas9 mediated editing of OsSWEET14 was transformed into the rice variety TBR225 via Agrobacterium-mediated transformation (SI Table). A total of nine TBR225 transformants were selected from 10 independent PCR-validated transgenic T0 plants to further investigate CRISPR/Cas9-targeted mutagenesis of the OsSWEET14 promoter. In order to decipher the nature of the editing events in OsSWEET14, the promoter sequencing data of transgenic lines were analyzed using the Degenerate Sequence Decoding software [36]. All 9 T0 transgenic plants harbored at least an editing event (Fig 1C): two were heterozygous mutant/wild type, two had homozygous mutations, and five had bi-allelic mutations. Regarding the type of mutations, 66.7% were nucleotide deletions, 11.1% of the mutations were nucleotide insertions and no substitution was detected (Table 1).

Inheritance of CRISPR/Cas9-induced mutations in the T1 generation

To assess the inheritance of the CRISPR/Cas9-induced OsSWEET14 mutations in the next generation, all T0 mutant transgenic plants (Fig 1C) were allowed to self-pollinate, and T1 transgenic plants were randomly selected in the progeny of T0 plants for sequencing and
analysis of their edited site (Table 2). All T₁ individuals derived from T₀ plants previously genotyped as homozygous possessed the same allele as their parent, suggesting stable inheritance of the mutations to the next generation. Similarly, the T₁ progeny of each of both bi-allelic and heterozygous mutation T₀ lines showed a segregation ratio which is consistent with Mendelian segregation ($\chi^2 < \chi^2_{0.05, 2} = 5.99$), indicating that the CRISPR/Cas9-induced mutations in T₀ plants were transmitted as expected to the next generation. Interestingly, no new mutant allele was detected in the T₁ generation of both heterozygous mutants L-21 and L-27, even though most of them still carried the transgene. Overall, consistent with previous similar studies, our results indicate that the CRISPR/Cas9-mediated mutations generated here are stably transmitted to the next generation in a Medelian fashion.

### Selection of transgene-free mutant TBR225 rice lines

To identify T-DNA free T₁ rice plants containing a mutation in EBEs of the OsSWEET14 promoter, PCR analysis was carried out using primers specific to Cas9, sgRNA and HPT sequences (Table 2). A T₁ individual was considered devoid of the transgene if the control amplification of the OsSWEET14 promoter was successful and if none of the PCR reactions with independent primer pairs designed on the T-DNA produced a detectable diagnostic band. The results of this PCR screen show that the T-DNA could be segregated out in the progeny of most T₀ lines, with 88.9% of the T₀ lines generating T-DNA-free progeny. In total, 44 of 221 analyzed edited T₁ plants did not generate a specific amplicon from the T-DNA construct and 15 of them were homozygous mutant harboring the desired OsSWEET14 modifications. Our results demonstrate that transgene-free, homozygous mutant individuals could be obtained in the segregating progeny of selfed T₀ individuals.

### Table 1. Frequencies of mutant genotypes and target mutation types in T₀ transgenic plants.

<table>
<thead>
<tr>
<th>Mutant genotype ratios$^a$ (%)</th>
<th>Mutation type ratios$^b$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heterozygote</td>
<td>Homozygote</td>
</tr>
<tr>
<td>22.2 (2/9)</td>
<td>22.2 (2/9)</td>
</tr>
</tbody>
</table>

$^a$ (Number of on-target mutant genotype/total number of on-target mutant genotypes) x 100%.

$^b$ (Number of allele mutation type/number of all allele mutation types) x 100%.

https://doi.org/10.1371/journal.pone.0255470.t001

### Table 2. Transmission of CRISPR/Cas9 editing events to the T₁ generation.

<table>
<thead>
<tr>
<th>T₀ plant</th>
<th>Genotype</th>
<th>Allele(s)</th>
<th>No. of T₁ plants tested</th>
<th>Mutation inheritance in the T₁ generation</th>
<th>No. of T-DNA-free plants</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Alleles segregation</td>
<td>$\chi^2 (1:2:1)$</td>
</tr>
<tr>
<td>L-4</td>
<td>Bi-allelic</td>
<td>-5/-3</td>
<td>32</td>
<td>10 (-5), 18 (-5/-3), 4 (-3)</td>
<td>2,750</td>
</tr>
<tr>
<td>L-5</td>
<td>Bi-allelic</td>
<td>-6/+1</td>
<td>44</td>
<td>9 (-6), 22 (-6/+1), 13 (+1)</td>
<td>0,727</td>
</tr>
<tr>
<td>L-7</td>
<td>Bi-allelic</td>
<td>-4/-3</td>
<td>38</td>
<td>14 (-4), 17 (-4/-3), 7 (-3)</td>
<td>3,000</td>
</tr>
<tr>
<td>L-15</td>
<td>Homozygote</td>
<td>+1</td>
<td>5</td>
<td>5 (+1)</td>
<td>-</td>
</tr>
<tr>
<td>L-21</td>
<td>Heterozygote</td>
<td>-3</td>
<td>26</td>
<td>3 (-3), 13 (-3/wt), 10 (wt)</td>
<td>3,769</td>
</tr>
<tr>
<td>L-27</td>
<td>Bi-allelic</td>
<td>-5/-4</td>
<td>7</td>
<td>1 (-5), 3 (-5/-4), 3 (-4)</td>
<td>1,286</td>
</tr>
<tr>
<td>L-29</td>
<td>Heterozygote</td>
<td>-5</td>
<td>33</td>
<td>6 (-5), 19 (-5/wt), 8 (wt)</td>
<td>1,000</td>
</tr>
<tr>
<td>L-31</td>
<td>Homozygote</td>
<td>-3</td>
<td>15</td>
<td>15 (-3)</td>
<td>-</td>
</tr>
<tr>
<td>L-54</td>
<td>Bi-allelic</td>
<td>-3/-2</td>
<td>21</td>
<td>3 (-3), 12 (-3/-2), 6 (-2)</td>
<td>1,286</td>
</tr>
</tbody>
</table>

$^+$ and $-$ indicate respectively, insertion and deletion, of the indicated number of nucleotides.

"w", wild type.

$^*$Number of homozygous mutant plants without T-DNA.

https://doi.org/10.1371/journal.pone.0255470.t002
TBR225 OsSWEET14 promoter editing confers resistance to Vietnamese Xoo

To characterize the BLB-resistance phenotype of the generated rice mutants, three T-DNA-free, homozygous TBR225 edited lines, namely, L-5.7(-6), L-31.12(-3) and L-15.4(+1) with OsSWEET14 promoter alleles corresponding respectively to L-5-a1 (6bp deletion), L-31 (3bp deletion) and L-15 (1bp insertion) in Fig 1C, were established. Selected T1 individuals were propagated to obtain T2 seeds which were used to perform BLB susceptibility assays. Edited T2 and WT TBR225 plants were inoculated by leaf-clipping with the VXO_11 and VXO_15 strains at the eight-week stage. The inoculated leaves of wild type TBR225 plants and of edited lines L-15.4(+1) and L-31.12(-3) developed long water-soaked lesions typical of BLB, ranging from 18.3 cm to 29.0 cm in length. In contrast, the edited line L-5.7(-6), harboring a longer 6-bp deletion at the target site, displayed high (1.2 cm average lesion length) and moderate (7.3 cm average lesion length) resistance to VXO_11 and VXO_15 strains, respectively (Fig 3). Means comparisons with a Tukey’s HSD test further indicated that irrespective of the inoculated strain, the mean lesion lengths measured on the L-15.4 (+1), L-31.12(-3) or wild type lines were not significantly different. In contrast, the mean lesion lengths recorded on the L-5.7(-6) mutant line were significantly different from those obtained on the wild type and the two other edited lines challenged with either of the Vietnamese strains (Fig 3B). Furthermore, our off-target editing analysis on line L-5.7(-6) did not reveal unintended modifications of other annotated rice loci (S2 Table and S5 Fig), indicating that the 6-bp deletion in the OsSWEET14 promoter is probably responsible for this phenotype.

Consistent with disease assays and as shown in Fig 3C, whereas a semiquantitative RT-PCR signal for OsSWEET14 expression was detected on the parental variety and the L-15.4(+1) and L-31.12(-3) edited lines following VXO_11 and VXO_15 infiltration, this amplicon was undetectable in the resistant L-5.7(-6) line.

In conclusion, this data shows that the 6-bp deletion in the AvrXa7/PthXo3 EBE reduces dramatically OsSWEET14 expression following VXO strains inoculation and confers resistance to these strains. In contrast, shorter modifications on the 3’-end of this EBE are insufficient to perturb OsSWEET14 expression after inoculation and do not confer detectable protection against the corresponding strains. Finally, while these results strongly support the view that OsSWEET14 functions as a unique susceptibility gene in the interaction between strain VXO_11 and the TBR225 rice variety, the resistance to strain VXO_15 is not as dramatic and may suggest that other mechanisms partially counteract the effects of the AvrXa7/PthXo3 EBE 6-bp deletion in edited TBR225 plants.

TBR225 OsSWEET14 promoter edited lines agronomic performances are undistinguishable from the parental variety

To determine if mutations in the OsSWEET14 promoter affect agronomic traits of TBR225 rice plants, three independent homozygous mutant lines were analyzed by measuring their growth duration, plant height, number of tillers per plant, number of grains per panicle, number of filled grains per panicle, yield per plant and amylose content under net-house conditions (see picture of S3 Fig). ANOVA tests and Student’s t tests showed that the mutant lines displayed no significant difference to TBR225, in terms of the examined agronomic traits, under our net-house conditions (Table 3). These results suggest that the tested CRISPR/Cas9-induced mutations in the OsSWEET14 promoter did not negatively impact the main agronomic traits of TBR225.
Discussion

Recently, the CRISPR/Cas9 system has emerged as a powerful tool for gene editing in many organisms including plants. Because of its specificity and efficiency, this system has been widely used to improve important agronomic traits of major crops such as rape, tomato, soybean, rice, wheat and maize [37]. Excluding easy-to-transform reference accessions such as Nipponbare and Kitaake that are widely used in the laboratory, the number of reports on the
improvement of agriculturally relevant elite rice cultivars for pertinent traits using the CRISPR/Cas9 technology (see for example [38–42]) is gradually increasing but is still limited. TBR225 [27], a major commercial rice variety cultivated in large areas of Northern Vietnam, has the advantages of early maturity, high and stable yield, as well as cooking quality. However, it is very susceptible to BLB. Here, the CRISPR/Cas9-mediated editing method was applied in order to rapidly improve the BLB resistance of TBR225 by modifying the AvrXa7, PthXo3 and TalF EBEs on the promoter of OsSWEET14. Of the three generated homozygous mutant lines tested for resistance, the one carrying the largest deletion at the target site (6 bp) showed a significantly improved resistance to infection with two Xoo strains VXO_11 and VXO_15. Therefore, using the major commercial rice variety TBR225 as an example, we illustrate the advantages of CRISPR/Cas9 tool for rice breeding.

In the present study, the frequency of individuals with CRISPR/Cas9-induced mutations in T0 transgenic plants was 90%, which is similar to previous observation [33]. We obtained only two heterozygous mutant/wild type lines versus seven homozygous or bi-allelic mutant lines. This high frequency of mutated alleles is another proof that the CRISPR/Cas9 system is indeed an efficient tool for gene editing in plant. We also observed the stable transmission of edited alleles to subsequent generations. This is a common phenomenon that has been repeatedly documented for rice plants carrying CRISPR/Cas9-induced mutations [38,40,43,44]. In this study, we obtained only two types of induced mutations in T0 plants: insertion (11.1%) and deletion (66.7%), but no substitution were observed. In some earlier studies, new mutations were continuously obtained in the T1 offspring of heterozygous T0 mutants because the Cas9 complex remains active on edited targets until the seed or PAM regions cease to be functional [35,37,43]. In contrast, here, all the T1 plants generated from both heterozygous lines L-21 and L-29, regardless of whether they had a CRISPR/Cas9 T-DNA transgene integrated in their genome, did not show any new mutation. We could also readily obtain transgene-free plants from most of the T1 segregation populations without any laborious crossing or backcrossing steps, which illustrates an advantage of the CRISPR/Cas9 technology compared to conventional breeding.

Clade III SWEET family proteins are involved in a number of biological processes such as seed and pollen development or pathogen susceptibility [45]. Their inactivation has previously been shown to cause pleiotropic and/or detrimental effects. For example, both ossweet11 single and ossweet11-ossweet15 double Kitaake rice mutants showed defects in endosperm development and filling [46]. In addition, RNA-mediated silencing of either Os11N3/OsSWEET14 [7] or Os8N3/OsSWEET11 [6] in BLB resistant Kitaake lines causes negative effects on seed production. In contrast, here, we show that T-DNA-free TBR225 plants harboring homoygous mutations generated with the CRISPR/Cas9 system in the AvrXa7/PthXo3 EBE of the OsSWEET14 promoter exhibited enhanced Xoo resistance but did not show any significant

<table>
<thead>
<tr>
<th>Lines</th>
<th>Growth duration (day)</th>
<th>Plant height (cm)</th>
<th>No. of tillers per plant</th>
<th>No. of grains per panicle</th>
<th>No. of filled grains per panicle</th>
<th>Amylose content (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>108.4 ± 1.1</td>
<td>86.6 ± 3.2</td>
<td>5 ± 0.7</td>
<td>144.4 ± 4.9</td>
<td>125 ± 4.5</td>
<td>13.2 ± 0.38</td>
</tr>
<tr>
<td>L-5.7(-6)</td>
<td>108 ± 1.2</td>
<td>86.4 ± 4.3</td>
<td>5.2 ± 0.4</td>
<td>144.2 ± 4.4</td>
<td>123.4 ± 5.5</td>
<td>13.7 ± 0.35</td>
</tr>
<tr>
<td>L-15.4(+1)</td>
<td>107.8 ± 0.8</td>
<td>86.4 ± 5.0</td>
<td>4.8 ± 0.4</td>
<td>147.8 ± 5.1</td>
<td>121.8 ± 3.0</td>
<td>13.5 ± 0.41</td>
</tr>
<tr>
<td>L-31.12(-3)</td>
<td>108 ± 1.2</td>
<td>88.4 ± 4.3</td>
<td>5.4 ± 0.5</td>
<td>144.6 ± 5.3</td>
<td>124.2 ± 7.4</td>
<td>13.8 ± 0.21</td>
</tr>
</tbody>
</table>

Five plants per line were measured. Experiments were repeated three time.
Means followed by the same letter do not differ significantly (P < 0.05).

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difference in all examined agronomic traits compared to wild-type plants under net-house growth conditions. It is conceivable that limited modifications in promoter regions do not affect the normal expression of SWEET genes in contrast to KO or silenced lines. Our findings are consistent with the previous work of Oliva et al. [11] who studied 30 combinations of EBE mutations in the OsSWEET11, OsSWEET13 and OsSWEET14 promoters of the IR64 or Ciherrang-Sub1 varieties and detected only a single line with abnormal agronomic traits.

Some individual Xoo strains have evolved a set of distinct TALE effectors that collectively target several members of the clade III SWEET family. The presence of these redundant TALEs thereby trumps single “loss-of-tale-responsiveness” resistance alleles [11,12,17,47]. For example, Kitaake lines carrying TALEN-induced mutation in the SWEET14 promoter [13,15] exhibit resistance to strains which depend exclusively on matching AvrXa7/PthXo3 for clade III SWEET family induction. Likewise, the natural xa13 allele [48] or CRISPR/Cas9-induced mutation in the SWEET11 promoter [11] exhibit resistance to strains such as PXO99 which depend exclusively on PthXo1, for virulence. However, the BLB resistance of the Kitaake lines harboring mutations in both AvrXa7/PthXo3 (OsSWEET14) and PthXo1 (OsSWEET11) EBEs was defeated by Xoo strains expressing simultaneously the AvrXa7/PthXo3 and PthXo2B TALEs [11]. Recently, the stacking of EBE-edited alleles in several OsSWEET promoters have overcome this limitation and was shown to achieve a broad spectrum of resistance to strains from most BLB-prone countries in Asia [11,12].

All of the three T2 lines tested for BLB resistance were affected for the AvrXa7/PthXo3 EBE and conserved an otherwise wild type TalF EBE (Fig 1C). The homozygous mutant TBR225 line L-5.7(-6) carrying a 6-bp deletion in the AvrXa7/PthXo3 EBE exhibited a significantly enhanced resistance to two Vietnamese Xoo strains compared to WT TBR225. The L-15.4(+1) and L-31.12(-3) lines that harbored more subtle alterations in the 3’-end of this EBE (a 1-bp insertion and a 3-bp deletion, respectively) in contrast remained susceptible to VXO strains. Our OsSWEET14 expression analysis after Vietnamese Xoo strains inoculation (Fig 1C) suggests that these editing events did not alter the EBE sequence sufficiently to compromise promoter recognition by an AvrXa7/PthXo3-like Vietnamese TALE. With less than 2 cm average lesion length, the resistance of line L-5.7(-6) (6-bp deletion) to the VXO_11 strain is rather extreme (versus average lesion length of 20.1 cm on wild type plants). Moreover, in this line, OsSWEET14 expression following bacterial inoculation is strongly reduced relative the parental line and the two other edited lines, which suggest that in this case, recognition by an AvrXa7/PthXo3-like Vietnamese TALE is abrogated. Consistent with OsSWEET14 expression analysis and as shown in S4 Fig, the Talvez [49] target prediction scores for AvrXa7 and PthXo3 on the OsSWEET14 promoter L-5-a1 allele sequence of line L-5.7(-6) are markedly lower than on the wild type promoter sequence. This is not the case however for the edited alleles carried by lines L-15.4(+1) and L-31.12(-3) (respectively L-15 and L-31 in S4 Fig) whose Talvez scores are identical or slightly lower than those of the wild type promoter sequence.

The magnitude of the effect of the 6-bp deletion allele on susceptibility to VXO_11 is comparable to the dramatic effect of previously characterized alterations of the same EBEs in the Kitaake background against the PXO86 strain that possesses a single TALE, AvrXa7, targeting OsSWEET14 for clade III OsSWEET gene induction [15]. By analogy, this suggests that OsSWEET14 is also the only clade III OsSWEETs target of VXO_11 in the TBR225 background but, in order to confirm this hypothesis an examination of other clade III OsSWEET genes expression patterns in response to this strain would be required. The situation with the VXO_15 strain is not as straightforward to interpret and will require further investigations. Although the 6-bp deletion in the AvrXa7/PthXo3 EBE did provide an increased resistance to the edited plants, the VXO_15 strain caused intermediate disease severity (7.3 cm average lesion length on Fig 3). This incomplete resistance is unlikely to result from the partial but still...
productive recognition of subsequences of the altered EBE by a VXO_15 AvrXa7/PthXo3-like TALE because OsSWEET14 expression is similarly decreased in response to either this strain or VXO_11 (Fig 3C). Alternatively, contrary to all Asian Xoo examined so far, but similar to African Xoo [11,15], VXO_15 may have the intrinsic potential to cause disease in the absence of clade III OsSWEET gene induction, a phenomenon that seems to be dependent on the edited rice variety genetic background [42]. More likely, analogous to other Asian strains, VXO_15 may encode alternative TALEs, such as PthXo2B or PthXo1 that compensate the loss of OsSWEET14 induction by targeting other clade III OsSWEET genes. In this regard, deciphering clade III OsSWEET genes expression patterns in combination with long read genome sequencing will ultimately help describe TALEs variability in Vietnamese Xoo strains and its functional impact on OsSWEET genes induction.

In conclusion, we showed that editing specific EBEs of Xoo TALEs via CRISPR/Cas9 tool is an efficient method for improving BLB resistance of elite rice varieties such as TBR225 without detectable yield penalties. This also uncovered the potential diversity of TALEs in Vietnamese Xoo population, which will thus require future investigations to address the TALE repertoires of Vietnamese Xoo strains in order to generate broad-spectrum BLB-resistant rice varieties in Vietnam.

Supporting information
S1 Fig. Nucleotide sequence of the OsSWEET14 promoter in TBR225. (TIF)
S2 Fig. Virulence of Vietnamese Xoo strains VXO_11 and VXO_15 on TBR225 rice. Grey points correspond to individual lesion length measurements while the black points indicate the calculated average value. The line range represents standard deviation. (TIF)
S3 Fig. Picture of an individual plant from the homozygous mutant rice lines L-5.7(-6). (TIF)
S4 Fig. Talvez scoring of AvrXa7, PthXo3 and TaLF target EBES in the edited OsSWEET14 promoter allele sequences. Score values are represented both by the length of the horizontal bar and a fill color scale. Higher Talvez prediction scores reflect a better match between a predicted EBE and the sequence of RVD of the query TALE. (TIF)
S5 Fig. Amplicon sequencing of predicted off-target sites for the OsSWEET14 promoter-sgRNA in annotated exons of the TBR225 edited line L-5.7(-6). Potential unintended target sequences including the PAM are highlighted in boxes. They are all identical to the expected wild type Nipponbare sequences. (TIF)
S1 Table. Key figures on the TBR225 transformation procedure for OsSWEET14 promoter editing. (DOCX)
S2 Table. Output of the CCTop tool used with the OsSWEET14 promoter sgRNA for off-target prediction on the rice Nipponbare genome. (DOCX)
S1 Raw images. Original photograph used in Fig 2 for the RT-PCR gels panel. (DOCX)
S2 Raw images. Original photograph used in Fig 3C for the RT-PCR gels panel.

(DOCX)

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Author Contributions

Conceptualization: Phuong Nguyen Duy, Sebastien Cunnac, Xuan Hoi Pham.

Data curation: Phuong Nguyen Duy.

Formal analysis: Phuong Nguyen Duy, Dai Tran Lan, Sebastien Cunnac, Xuan Hoi Pham.

Funding acquisition: Phuong Nguyen Duy, Tran Bao Manh, Xuan Hoi Pham.

Investigation: Phuong Nguyen Duy, Dai Tran Lan, Hang Pham Thu, Huong Phung Thi Thu, Ha Nguyen Thanh, Ngoc Phuong Pham, Florence Auguy, Sebastien Cunnac.

Methodology: Phuong Nguyen Duy, Dai Tran Lan, Hang Pham Thu, Ha Nguyen Thanh, Sebastien Cunnac, Xuan Hoi Pham.

Project administration: Xuan Hoi Pham.

Resources: Hang Pham Thu, Ha Nguyen Thanh, Florence Auguy.

Supervision: Sebastien Cunnac, Xuan Hoi Pham.

Validation: Sebastien Cunnac, Xuan Hoi Pham.

Writing – original draft: Phuong Nguyen Duy, Dai Tran Lan.

Writing – review & editing: Huong Bui Thi Thu, Tran Bao Manh, Sebastien Cunnac, Xuan Hoi Pham.

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