INTRODUCTION

Rice (Oryza sativa) is one of the most important cereal crops cultivated around the world. Rice cultivation often suffers from several types of diseases, including rice sheath blight (RSB), caused by the basidiomycete fungus Rhizoctonia solani. RSB is one of the most devastating rice fungal diseases globally and causes significant yield losses and reductions in rice grain quality (Moni et al., 2016; Rao et al., 2019; Wang et al., 2018). RSB often occurs under warm and humid conditions, particularly in tropical areas. In China, RSB affects over 17 million hectares per year and results in a yield reduction as high as 50% in severely affected areas (https://www.natesc.org.cn).

Abstract

Rice sheath blight, caused by the soilborne fungus Rhizoctonia solani, causes severe yield losses worldwide. Elucidation of the pathogenic mechanism of R. solani is highly desired. However, the lack of a stable genetic transformation system has made it challenging to examine genes' functions in this fungus. Here, we present functional validation of pathogenicity genes in the rice sheath blight pathogen R. solani by a newly established tobacco rattle virus (TRV)-host-induced gene silencing (HIGS) system using the virulent R. solani AG-1 IA strain GD-118. RNA interference constructs of 33 candidate pathogenicity genes were infiltrated into Nicotiana benthamiana leaves with the TRV-HIGS system. Of these constructs, 29 resulted in a significant reduction in necrosis caused by GD-118 infection. For further validation of one of the positive genes, trehalose-6-phosphate phosphatase (Rstps2), stable rice transformants harbouring the double-stranded RNA (dsRNA) construct for Rstps2 were created. The transformants exhibited reduced gene expression of Rstps2, virulence, and trehalose accumulation in GD-118. We showed that the dsRNA for Rstps2 was taken up by GD-118 mycelia and sclerotial differentiation of GD-118 was inhibited. These findings offer gene identification opportunities for the rice sheath blight pathogen and a theoretical basis for controlling this disease by spray-induced gene silencing.

KEYWORDS

disease control strategy, host-induced gene silencing, pathogenicity gene, Rhizoctonia solani, rice sheath blight
R. solani exhibits a broad host range and causes catastrophic diseases in numerous plant species, including Asteraceae, Brassicaceae, Fabaceae, Poaceae, and Solanaceae, as well as some woody ornamental plants and forest trees (Ghosh et al., 2014; Pan et al., 1999; Zou et al., 2000). In nature, R. solani produces no asexual spores and exists mainly on crop residues and in the soil in the form of mycelia and sclerotia (Shu et al., 2019). Unfortunately, the molecular basis for RSB pathogenesis and rice–Rhizoctonia interaction is largely unknown (Molla et al., 2020), mainly due to the lack of rice cultivars with stable and perfect resistance to RSB and of efficient techniques for functional analysis of R. solani genes (Singh et al., 2019). Due to the broad host range, high genetic variability, and survival as sclerotia (Taheri & Tarighi, 2011; Tiwari et al., 2017; Yellareddyari et al., 2014; Zhu et al., 2019), farmers heavily rely on the use of chemical fungicides such as ARMURE and jinggangmycin (validamycin) to control RSB. However, excessive use of chemical fungicides may cause environmental pollution and increase the risk of fungicide resistance in the RSB pathogen (Singh et al., 2019).

In the last decades, elucidation of the pathogenic mechanism of the RSB pathogen has been accelerated thanks to de novo genome sequencing and transcriptome sequencing efforts (Singh et al., 2019; Zheng et al., 2013). In the earliest stage of RSB pathogenesis, sclerotia in the soil germinate to produce pyxhaya and attach to plants’ surfaces. With the continuous branching, the hyphae gather to form infection cushions and penetrate rice cells or tissues (Taheri & Tarighi, 2011). Finally, water-soaked lesions develop and expand on rice tissues. During the infection process, R. solani produces cell wall-degrading enzymes (e.g., pectinase, laccase, and xylanase) and some metabolites, including phytotoxins, facilitating its penetration into and killing of host cells (Rao et al., 2019; Singh & Subramanian, 2017; Yang et al., 2011). In recent years, three polygalacturonase (PG) genes, Rspg1, RsPG3, and RsPG4, have been reported to induce tissue necrosis in rice sheath as essential virulence genes in the host–pathogen interaction (Chen et al., 2018; Yang et al., 2012). Recent studies reported effector-like proteins based on necrosis induction or cell death signalling in plant cells (Li et al., 2019; Yamamoto et al., 2019; Zheng et al., 2013). However, few pathogenicity genes have been identified in R. solani.

RNA interference (RNAi), or gene silencing, which is characterized by targeted mRNA degradation by the introduction of sequence-specific double-stranded RNA (dsRNA) into cells, has become an essential tool for studying gene functions (Kuo & Falk, 2020). Small interfering RNA duplexes (siRNAs) are derived from the processing of the dsRNA by an RNAse III-like enzyme. Then siRNAs are recruited into a multiprotein complex known as RNA-induced silencing complex, which interacts with the target RNA to mediate cleavage in a catalytic fashion (Weigel et al., 2015). Virus-induced gene silencing (VIGS) is a particular specific gene-silencing phenomenon triggered by dsRNA. It has been applied in examining gene functions in plants and adapted for high-throughput functional genomics (Su et al., 2020). Tobacco rattle virus (TRV), a bipartite virus, has been chiefly used as a vector in VIGS. In this case, pTRV1 encodes the replication and viral movement proteins, and pTRV2 encodes the coat protein and harbours the sequence used for VIGS. Inoculation of Nicotiana benthamiana with two different Agrobacterium tumefaciens strains, carrying pTRV1 and pTRV2, results in gene silencing (Robertson, 2004).

The exploration of mechanisms of cross-kingdom RNAi in the virulence and pathogenesis of pathogens, as well as host resistance, could significantly enhance our understanding of the R. solani–rice pathosystem. In 2010, Nowara et al. (2010) silenced GTF1 and GTF2 in the wheat and barley powdery mildew pathogen Blumeria graminis using the virus-induced gene silencing (VIGS) system with barley stripe mosaic virus (BSMV), which ultimately enhanced resistance in barley. From then on, a new procedure to determine the gene function of microbes by VIGS was proposed, and a new concept of host-induced gene silencing (HIGS) was put forward (Nowara et al., 2010). HIGS is an RNAi-based strategy that includes expressing a suitable RNAi structure in the host plant, targeting pathogen pathogenicity-related genes, and transferring dsRNA or siRNA into the pathogen during the interaction, followed by silencing of the target gene and suppression of virulence of the pathogen (Song & Thomma, 2016). The HIGS technique has been widely used in the study of biotrophic and necrotrophic fungi, including Puccinia striiformis (Zhu et al., 2017), Verticillium dahliae (Su et al., 2018; Su et al., 2017), and Colletotrichum gloeosporioides (Mahto et al., 2020).

RSB resistance is believed to be a quantitative trait that is under the control of many genes. To our knowledge, there are only two examples that showed increased RSB resistance by silencing pathogenicity genes of R. solani via stable rice transformation (Rao et al., 2019; Tiwari et al., 2017). To improve our understanding of the pathogenic mechanism of the RSB pathogen, in the present study we established a rapid and stable HIGS screening system for the identification of pathogenicity genes in the RSB fungus using the virulent R. solani AG-1 IA strain GD-118, which is also virulent to N. benthamiana. We showed reduced virulence of GD-118 to tobacco subjected to HIGS of 29 candidate pathogenicity genes. Stable rice transformation of siRNA targeting a trehalose 6-phosphate phosphatase gene of R. solani (Rstps2) increased resistance to RSB. This observation seemed to be attributable to the uptake of dsRNA of Rstps2 by R. solani cells, resulting in a reduction of trehalose biosynthesis. Thus, we demonstrated that the HIGS system can be used to identify virulence factors in R. solani.

2 | RESULTS

2.1 | Screening of candidate pathogenic genes in R. solani AG-1 IA

We conducted a knowledge-based screening of candidate pathogenicity genes from the R. solani AG-1 IA reference genome (Zheng et al., 2013). All the predicted protein sequences were searched against the Pathogen-Host Interaction Database (PHI-base) (Urban et al., 2017) and the carbohydrate-active enzyme (CAZyme) database CAZy (Cantarel et al., 2009) to screen candidate pathogenicity genes. After compilation, 33 genes involved in diverse biological
processes (energy metabolism, development, protein secretion, and others) were selected for a functional test (Table S1). Gene expression patterns of these genes in *R. solani* AG-1 IA were clustered into four groups based on transcriptome data (Zheng et al., 2013) (Figure S1). Group A comprised genes showing the highest expression levels at 32 to 72 hr after inoculation (hai) of the pathogen. This group contained three genes encoding enzymes involved in trehalose metabolism. Group B contained genes showing the highest gene expression levels at 18 or 24 hai. Two CAZyme genes (AG1IA_00742 and AG1IA_01403) were included in this group. Group C comprised genes showing the highest gene expression levels at 24 or 32 hai. This group contained two genes encoding capsular-associated proteins (AG1IA_03049 and AG1IA_03050). Group D comprised genes showing the highest gene expression levels at 10 hai. It had two CAZyme genes (AG1IA_01330 and AG1IA_00669). These results supported the reliability of our gene annotation-based selection of candidate pathogenicity genes. No significant sequence similarity was found between these candidate genes and internal genes in *N. benthamiana*. No possible off-target effect for the quantitative PCR (qPCR) primers was validated by SI-FI software (http://labtools.ipk-gatersleben.de/).

### 2.2 Functional test of the candidate genes by TRV-HIGS

In this study, 35 *Agrobacterium* constructs, including a positive control construct pTRV2::PDS, a negative control construct pTRV2::GFP (RNAi-0), and 33 candidate gene constructs pTRV2::[RNAi-(1–33)], were agro-infiltrated into *N. benthamiana* leaves. Infiltration of pTRV2::PDS caused photobleaching at 10 days postinfiltration (dpi) (Figure S1a). Bleaching leaves appeared on the top of *N. benthamiana* initially and gradually expanded onto the whole plant until up to 20 dpi (Figure S2a). By contrast, infiltration of RNAi-0 produced no photobleaching (data not shown). These results indicated the appropriateness of our gene silencing system. The infiltrated tobacco with RNAi-0 allowed the growth of *R. solani* AG-1 IA strain GD-118 mycelia at 1 dpi and exhibited necrotic spots at 2 dpi (Figure S2b). After 5 days, almost all leaves withered, wilted, and tended to die (Figure S2b). By contrast, infiltrated tobacco plants with all candidate gene constructs exhibited lower disease index (DI) values than RNAi-0 tobacco plants but varied among each other (Figure 1). The DI value of RNAi-0 plants at 5 dpi was 86.11, while those of RNAi-(1–33) plants ranged from 22.22 to 69.44. Four RNAi constructs did not significantly reduce pathogenicity of GD-118 (Figure 1). These results suggested that 29 of the 33 candidate genes are involved in the pathogenic fitness of *R. solani*.

### 2.3 Validation of the transient silencing effect of Rstps2 on pathogenicity of *R. solani*

To validate the effect of silencing AG1A-04214 (RNAi-25), which targets a gene that encodes a trehalose 6-phosphate phosphatase (Rstps2), the tobacco plants transiently expressing pTRV:Rstps2...

**Figure 1** Disease index values in host-induced gene silencing (HIGS) assays in *Nicotiana benthamiana* for 33 candidate pathogenicity genes in *Rhizoctonia solani*. Highly resistant lines, moderately resistant lines, and slightly resistant lines are highlighted in red, blue, and yellow, respectively. Error bars represent standard errors of biological triplicates. *p < .05, one-way analysis of variance with Tukey’s post hoc test.
were further characterized. The tobacco plants silenced for Rstps2 exhibited no noticeable phenotypic difference to those of the negative control (Figure 2a). Reverse transcription (RT)-PCR assays were used to detect transcript levels of the targeted genes Rstps2, GFP, and PDS (Figure 2b). The agro-infiltrated tobacco silenced for Rstps2 exhibited more resistance to GD-118 than the green fluorescent protein (GFP) control (Figure 2c). The increased resistance in Rstps2-infiltrated tobacco was supported by the decrease of visibly infected leaf area (Figure 2d), lower amount of biomass of the pathogen (Figure 2e), and smaller leaf area with reactive oxygen species (ROS) accumulation (Figure 2f). The relative infection area of GFP plants was 47.0% but that of Rstps2 plants was 6.02% (p < .01; Figure 2d). Compared with GFP plants the fungal biomass of Rstps2 plants decreased by 86% (p < .01; Figure 2e). The leaf area stained with 3,3′-diaminobenzidine (DAB) in Rstps2 plants was less than half of that of GFP plants (p < .001; Figure 2g). As expected, the expression level of Rstps2 plants was significantly lower (24.8%) than that of GFP plants (Figure 2h). These results indicated that silencing of Rstps2 in GD-118 by the TRV-HIGS technique in N. benthamiana decreased virulence and inhibited mycelial growth.

Noteworthily, ROS accumulation was observed throughout the leaves of Rstps2 plants. In GFP plants, severe disease symptoms appeared on the leaves' edges, and ROS was detected throughout the whole leaf (Figure 2f). By contrast, the Rstps2 plant leaves had no prominent disease spots. The accumulation of ROS was observed as uniformly distributed spots over the whole leaf, indicating that GD-118 colonized the leaf successfully. These results indicated that the virulence of strain GD-118 decreased by transient silencing of Rstps2 in N. benthamiana and that N. benthamiana generated moderate amounts of ROS to prevent further invasion of the pathogen.
2.4 | HIGS of Rstps2 by stable transformation in rice

Validation of putative transgenic rice lines (T₀ generation) was conducted by detecting Rstps2 and the selection marker hygromycin resistance gene (hyg) using genomic PCR with gene-specific primers (Table S2). Among 14 lines with hygromycin resistance, 11 lines were positive for both genes (Figure S4a). Three randomly selected T₀ generation rice plants exhibited more resistance against GD-118 than nontransformants, as determined by detached leaf assays (Figures S4b–d and S5a–d). In three of these T₀ lines with confirmed transgene integration by genomic Southern blot (Figure S4e), the fungal biomass decreased with statistical significance (Figure S4d). We used this line (named TPS-3) for further verification.

To check the inheritance of the increased resistance to GD-118, we analysed two T₂ transgenic lines, TPS-3-2 and TPS-3-6, which were derived from TPS-3. These T₂ lines harboured hyg and Rstps2 genes and exhibited no differences in rice plant height and the number of effective tillers between the RNAi lines and nontransformants (Figure 3a–d). Genomic DNA Southern blot indicated stable inheritance of the transfer DNA (T-DNA) into the T₂ generation (Figure 3e). No obvious morphological abnormalities were observed in the RNAi transgenic rice lines (data not shown). We observed increased resistance to GD-118 in these T₂ rice lines as well (Figure 4a). Only a few mycelia and disease spots were observed in homozygous T₂ transgenic detached rice leaves at 60 and 120 hpi, respectively, whereas dense disease spots were observed in the wild-type (WT) rice leaves. The mycelial growth and fungal biomass significantly decreased. The relative infection area in T₂ transgenic rice lines was 9.34%, which was significantly reduced when compared with the WT (59.45%) at 5 dpi (Figure 4b,c). Intact rice plants grown in pots were used to confirm the increased resistance to GD-118 in the T₂ transgenic plants. Disease symptom development was less severe in the transgenic rice lines, and the height of exhibited disease spots in nontransformants was higher than in homozygous T₂ transgenic plants at 10 dpi (Figure 4d,e). Inheritance of increased resistance to GD-118 was observed in detached leaves of T₂ rice lines derived from T₂ Tps-3-2 and Tps-3-6 (Figure S6). These results indicated stable inheritance of silencing of Rstps2 and increased RSB resistance, although the results are based only on one transgenic line.

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**FIGURE 3** Molecular analysis and morphological characteristics of homozygous T₂ transgenic rice. (a,b) PCR analysis of T₂ transgenic plants using Rstps2 gene-specific primers. (c) Plant height. (d) Number of tillers. (e) Southern blot analysis showing the integration of the transgene in the T₂-Tps2 lines. Data are shown as the mean ± SE of triplicates. ns: no significant changes observed, one-way analysis of variance and Tukey's test.
2.5 Concomitancy among silencing of Rstps2, reduced trehalose content, and increased levels of siRNA for Rstps2 in GD-118 mycelia grown on HIGS rice

Trehalose 6-phosphate phosphatase (EC: 3.1.3.12) catalyses the reaction from trehalose 6-phosphate to trehalose, the last biochemical step in trehalose biosynthesis. We assumed that transformation of the RNAi construct for Rstps2 led to an accumulation of siRNA for Tps2, silencing of Tps2, and inhibition of trehalose biosynthesis in GD-118 inoculated on the transgenic rice. As expected, the trehalose content in R. solani-(TPS-3-2) and R. solani-(TPS-3-6) was decreased by 31.17% and 34.61% when compared with the R. solani-(WT), respectively (Figure 5a). To test the persistence of siRNA, we subcultured the R. solani-(Tps-3-2) controls at 5 and 10 days postinoculation (dpi). **p < .01, ***p < .001, two-way analysis of variance Tukey’s test [Correction added on 02 September 2021, after first online publication: Figure 4 has been updated in this version.]
to G4, but there was a significant increase in G5 (p < .001). This result indicated that the effect of RNAi can continuously affect gene expression, leading to gene down-regulation. However, the effect of gene silencing decreased with multiple subgenerations (Figure 5b). For siRNA, we observed a significant increase of Rstps2-associated reads in small RNA sequencing (Figure 5c). The length distributions of siRNAs looked shifted to increased length (from 22 nucleotides [nt] in a nontransformant to 23 nt in the transgenic line). These results support the mechanistic view that siRNAs derived from Rstps2 can enter pathogenic fungi from plants and silence the target gene to improve host resistance.

2.6 | Supplementation of exogenous dsRNA for Rstps2 affects the expression of Rstps2

To further verify the mechanism of our HIGS technique in R. solani, we examined the effects of externally supplemented dsRNA for Rstps2 on GD-118 cells. Coincubation of GD-118 mycelia with the dsRNAs reduced the gene expression levels of Rstps2. We observed 56.70%, 67.19%, 78.75%, and 83.69% reductions at 24, 48, 72, and 96 hr, respectively. There was no significant difference between 24, 48, 72, and 96 hr. The gene expression changes could be due to the result of the trafficking of the dsRNAs (Figure 6a). The mycelial growth rate and colony morphology of GD-118 showed no significant difference when grown on potato dextrose agar (PDA) plates (control) and PDA plates with dsRNA (500 ng/ml) (data not shown), but our study found that sclerotial differentiation was inhibited on PDA plates with dsRNA by 17.73% compared to controls (Figure 6b). We validated this by fluorescence microscopy (Figure 6c). A stronger fluorescence signal was observed in the hyphae at 24 hr, indicating GD-118 mycelia absorbed and/or adsorbed the labelled dsRNA from the medium. Thus, these results show a potential application of spray-induced gene silencing (SIGS) for bio-control of R. solani.

3 | DISCUSSION

To date, HIGS has been reported in many important plant diseases caused by fungi and oomycetes, including Puccinia striiformis, Puccinia triticina, Verticillium dahliae, Bremia lactucae, Phytophthora parasitica, and Phytophthora infestans (Govindaraju et al., 2015; Panwar et al., 2013; Sanju et al., 2015; Yin et al., 2011; Zhang et al., 2011, 2016). In this paper, we report an application of HIGS in R. solani, for which no genetic transformation techniques are available. Our HIGS system using N. benthamiana and GD-118 offers a relatively high-throughput experimental scheme with potential usefulness in other transformable plant species. At least, we demonstrated that the HIGS system is applicable to rice, which cannot undergo agro-infiltration-based VIGS. The TRV-HIGS system takes only 5–6 weeks to analyse the effect of silencing of pathogenicity gene candidates. Thus, the establishment of this HIGS system could accelerate functional validation of candidate pathogenicity genes and elucidation of the pathogenic mechanisms of R. solani diseases such as sheath blight and foliar blight.

In recent years, cross-pathogenicity of R. solani in N. benthamiana has been recognized, although tobacco plants were thought to be nonhost plants in nature (Anderson et al., 2016; Gonzalez et al., 2011). In our previous trials, we inoculated mycelial plugs or sclerotia to the stem base of N. benthamiana. This method resulted in propagation of mycelia and stem rot disease, but was not suitable for quantitative phenotyping of disease status (data not shown). Hence, in the present study, we sprayed a mycelial fragment suspension of GD-118 onto N. benthamiana using a spray bottle. This method allowed the observation of many disease spots with ROS accumulation on N. benthamiana leaves. Finally, we observed dead leaf tissues of N.
Trehalose-6-phosphate phosphatase (Tps2), and (b) synthesis from UDP-glucose by trehalose-6-phosphate synthase (Tps1) and (c) synthesis from glucose-1-phosphate by trehalose phosphorylase (Thammahong et al., 2017). However, the former pathway seems to be the primary one in *R. solani* because our HIGS reduced trehalose content, and these data are based only on one transgenic line and are thus not conclusive. Nevertheless, a Tps2 mutant in *U. maydis* resulted in a lack of trehalose production with extremely reduced virulence to maize (Cervantes-Chavez et al., 2016). Deficiency of Tps2 in *F. graminearum* reduced virulence by 99%, while abnormal septum development and nuclear distribution in its conidia and lower mycelial growth were observed (Song et al., 2014). The results of our Tps2 silencing experiments in GD-118 with dsRNA showed lower sclerotal dry weight. The role of trehalose in *R. solani* diseases might be the detoxification of ROS species from the host, as reported in a study on *Candida albicans* (Lu et al., 2011). Yamamoto et al. (2019) hypothesized that *R. solani* secretes a series of ROS detoxification enzymes for their pathogenicity.

We report a potential application of gene silencing of virulence factors in *R. solani* by genetic modification of crop plants. Transfer of dsRNA and siRNAs from plants to various pathogens is a common phenomenon. With no surprise, we found that *R. solani* could directly take up dsRNA from the growth environment with concomitant gene silencing over time (Figure 6). Due to the absence of completely resistant germplasm of rice against RSF (Lavale et al., 2018; Mulualem & Bekeko, 2016; Richa et al., 2017), the application of SIGS may offer an alternative crop protection approach for rice.

4 | EXPERIMENTAL PROCEDURES

4.1 | Fungal strain and plant materials

The virulent *R. solani* AG-1 IA strain GD-118 (Yang et al., 2012) was grown on PDA at 28 °C in the dark for a few days. The mycelia were
collected and used for inoculation onto tobacco/rice plants or stored at −80 °C until RNA extraction.

*N. benthamiana* seeds were sown on soil in pots covered with plastic films to increase humidity. Then 10-day-old *N. benthamiana* seedlings were transplanted into individual pots with soil and grown for 2 weeks at 25 °C under 55% relative humidity (RH) and a 16/8-hr light/dark period.

Seeds of a japonica rice cultivar (Nipponbare) were distributed by Wuhan Biorun Biological Technology Co., Ltd. The seeds were used for genetic transformation with RNAi constructs by the same company.

### 4.2 | DNA extraction, total RNA extraction, and cDNA synthesis

Genomic DNA from rice tissues was isolated using the cetyltrimethylammonium bromide method (Hormaza, 2002). Total RNA was extracted using the MiniBEST Universal RNA Extraction Kit (Takara) according to the manufacturer’s instructions. Approximately 2 μg of DNase-treated total RNA was used for first-strand cDNA synthesis using the Goldenstar RT6 cDNA Synthesis Kit (Beijing Tsingke Biotech Co. Ltd.) according to the manufacturer’s instructions. Synthesized first-strand cDNA was stored at −20 °C for later use.

### 4.3 | Construction of TRV-HIGS vectors and agro-infiltration in *N. benthamiana*

Short cDNA fragments of 33 candidate genes in *R. solani* were amplified by RT-PCR using gene-specific primers with EcoRI and BamHI linkers (Table S1). These primers were designed based on the reference genome of AG-1 IA (GenBank accession GCA_000334115.1). The resultant PCR product of each gene was digested with EcoRI and BamHI and cloned into a linearized pTRV2 vector. A TRV2-based construct harbouring GFP was used as a negative control of VIGS and HIGS (Senthil-Kumar & Mysore, 2014; Su et al., 2014). A TRV2-based construct harbouring phytoene desaturase (PDS) from *N. benthamiana* (pTRV2::PDS) was used as a positive control for VIGS efficiency (Senthil-Kumar & Mysore, 2014). Finally, pTRV1 and these PTRV2-based constructs were separately transformed into *Agrobacterium tumefaciens* GV3101 by electroporation.

Subsequently, *Agrobacterium* cell culture of each construct was diluted to $OD_{600} = 0.4$ with infiltration buffer (10 mM MgCl$_2$, 10 mM MES, pH 5.6, 200 μM acetosyringone). pTRV1-transformed Agrobacterium culture cell was mixed in a 1:1 ratio with each pTRV2-based construct and incubated at room temperature for 4 hr. Then the mixed Agrobacterium solution was infiltrated into the first and second leaves of 2-week-old *N. benthamiana* using a needleless syringe. The *N. benthamiana* plants were grown at 25 °C in a growth chamber under a 16 hr light/8 hr dark photoperiod and 50% RH for at least 14 days.

### 4.4 | RNAi construct preparation and rice transformation

A partial cDNA fragment for Rstps2 was amplified from the Rstps2 cDNA by PCR with gene-specific primers that include attB1 and attB2 sequences in the termini and then subcloned into pDONR-221 via BP recombination with Gateway BP Clonase II (Invitrogen). The cloned Rstps2 fragment was subcloned into two sites of a binary vector pBDL03 by LR recombination reactions with Gateway LR Clonase II (Invitrogen) (Figure S3). The T-DNA sequence of the resultant plasmid pDONR-221::Rstps2 was verified, and the plasmid was transformed into rice via an *Agrobacterium*-mediated method by a commercial company (Wuhan Biorun Biological Technology Co., Ltd.). Hygromycin-resistant transgenic rice lines were subjected to PCR analysis using Rstps2- and hyg-specific primers (Table S2). The positive T$_0$ rice lines were phenotyped by inoculation of GD-118 onto detached rice leaves. One T$_0$ rice line and two T$_2$ rice lines originating from the T$_0$ rice line were tested by genomic Southern blot analysis as described elsewhere (Southern, 1975). Namely, 20 μg of genomic DNA from the transgenic rice line (with increased resistance) and nontransformants was digested with EcoRV, and a 983-bp partial fragment of the T-DNA, which was amplified by PCR (for primers, see Table S2), was labelled with the digoxigenin DNA Labeling Kit (Roche) and used as a probe for Southern blot analysis.

### 4.5 | Disease phenotyping of VIGS-treated *N. benthamiana* and transgenic rice

VIGS-treated *N. benthamiana* plants were sprayed with 5 ml of *R. solani* AG-1 IA strain GD-118 mycelial suspension (OD$_{600} = 1.0$) and grown in an illumination incubator (90% RH, 16 hr light, 8 hr dark). After 5 days, the DI (Su et al., 2020) was measured using the following equation: $DI = (no.\ of\ plants \times DS score)/\text{total no. of plants} \times \text{highest DS score} \times 100$, where “DS score” is defined as follows: 0, no spot and no wilt; 1, less than two leaves have disease spots or wilting; 2, three or four leaves have disease spots or wilting; 3, five to seven leaves are wilting or chlorotic; and 4, plant death or near-death.

To further determine the resistance level, the third, fourth, and fifth leaves of *N. benthamiana* were selected to measure the relative lesion area using the “Lasso” tool in Adobe Photoshop CC2018 (Adobe Inc.).

To conduct disease phenotyping of transgenic rice lines at the individual plant level, one fresh sclerotium of GD-118 was inoculated on the bottom part of each tiller at the late tilling stage. The inoculated plants were then kept in a humidity chamber (RH > 90%, c.30 °C) for 5–6 days and then shifted to a normal glasshouse. At 10 days after GD-118 inoculation, the disease severity was scored by highest relative lesion height (HRLH), which was calculated as follows: $\text{HRLH} = \text{length of the highest lesion (cm)}/\text{plant height (cm)} \times 100%$ (Tiwari et al., 2017).
For disease phenotyping of detached rice leaf blades, one fresh sclerotium of GD-118 was inoculated on the centre of each detached leaf blade. Then the leaves were incubated in a humidity chamber (RH > 90%, c.30 °C) for 60 and 120 hr. The Photoshop “Lasso” tool was used to measure the percentage of the leaf area occupied by the disease area.

4.6  |  RT-qPCR

qPCR was performed using a CFX96 real-time PCR system (Bio-Rad) and TransStart Top Green qPCR SuperMix (TransGen) with three biological replicates with three technical replicates. Genomic qPCR assays for the 5.8S rDNA fragment in R. solani AG-1 IA (GenBank accession number KX674524.1) were conducted for the estimation of fungal biomass by the method of (Tzima et al., 2012). EF1-α of N. benthamiana (GenBank accession number AF120093) and the 18S rRNA gene of rice (GenBank accession number X00755) were used as internal controls for quantitative gene expression assays (Schmidt & Delaney, 2010; Bo-Ra Kim et al., 2003). GAPDH (GenBank accession number ELU44274.1) was selected as the internal control of gene expression in GD-118. The relative amounts of DNA or transcripts were analysed using the 2−ΔΔCt method (Livak & Schmittgen, 2001). The sequences of gene-specific primers used for qPCR are listed in Table S2.

4.7  |  Measurement of trehalose content

Trehalose contents of GD-118 mycelia were measured using a kit (HT-2-Y) (Suzhou Comin Biotechnology Co., Ltd.) according to the manufacturer’s protocol. GD-118 was reisolated from the infected rice leaves, named Rs-(sample). After growing on PDA for 2 days, the mycelia were collected and the trehalose content was determined. Briefly, about 0.1 g of mycelia was homogenized for the extraction of trehalose. Extracted solution (0.25 ml) was mixed with 1 ml of reaction solution and the reaction proceeded at 95 °C for 10 min. After cooling to room temperature, the absorbance (A) was recorded at 620 nm. Trehalose content was calculated as follows: trehalose content (mg/g of fresh weight) = 0.112 × (A620−0.0729)/0.1.

4.8  |  Small RNA sequencing

GD-118 mycelia were prepared using the abovementioned method for trehalose content measurements and frozen in liquid nitrogen. The mycelia were used for small RNA sequencing by Novogene Co., Ltd. (Beijing, China). Briefly, total RNA of mycelia was extracted and used for the construction of small RNA libraries by the NEBNext Multiplex Small RNA Library Prep Set for Illumina (New England Biolabs). Short DNA fragments with a length of 140–160 bp were size-fractionated by polyacrylamide gel electrophoresis. Recovered short DNA fragments were sequenced on an Illumina HiSeq 2500 platform to generate 50-bp single-end reads.

The raw fastq files were processed by a data analysis pipeline for quality control by Novogene Co., Ltd. In this process, ambiguous sequence repeats “poly-N,” the adaptor sequences, reads containing poly(A), poly(G), poly(C), and poly(T), and low-quality sequence reads were removed. Resultant high-quality short reads were mapped on the R. solani genome sequence by Bowtie (Langmead et al., 2009) for estimation of gene expression levels of micro-RNAs using transcript per million (TPM) values as described elsewhere (Zhou et al., 2010).

4.9  |  In vitro synthesis and supplementation of fluorescently labelled dsRNA

dsRNA molecules for Rstps2 were prepared by in vitro transcription. The partial cDNA fragment of Rstps2 was amplified by PCR using a primer set, of which 5’ termini were linked with the T7 promoter sequence (5’TATAACGACTCACTATAGGG-3’) (Table S2). The resultant PCR products were purified using an AxyPrep PCR Clean-Up Kit (Axygen). Complementary fluorescently labelled dsRNAs were transcribed in vitro by T7 RNA polymerase (Thermo Fisher Scientific) according to the manufacturer’s protocol as follows. Each 20-μl reaction solution contained 500 ng of purified PCR product, 2 μl of fluorescein RNA-labelling mix, 4 μl of transcription buffer, and 2 μl of T7 RNA polymerase. The synthesized dsRNAs were collected by ethanol precipitation.

External supplementation of dsRNA was carried out as described by McLoughlin et al. (2018). The dsRNA fraction was supplemented to GD-118 mycelia grown on potato dextrose broth (PDB) for 2 days. This incubation was conducted in 6 ml PDB with dsRNA at a final concentration of 500 ng/ml with shaking at 200 rpm at 28 °C. The mycelia were collected at 0, 24, 48, 72, and 96 hpi for gene expression analysis by RT-qPCR.

To visualize dsRNA trafficking into the hyphae of GD-118, the fungal mycelia were grown on PDA plates containing 500 ng/ml dsRNA. Cover slides were obliquely inserted into PDA, and the hyphae were visualized under an Eclipse 90i fluorescence microscope (Nikon) with elicitation of the fluorescent signals.

To determine the effect of Rstps2 silencing on sclerotial differentiation (SDIF), GD-118 was cultured in PDA with dsRNA (500 ng/ml) or without dsRNA (control) for 10 days. Then the dry weight (DW) of mature sclerotia was determined by placing fresh sclerotia in a drying oven at 70 °C until a stable weight was reached. SDIF was calculated as follows: SDIF = (DW)dsRNA/(DW)control (Wang et al., 2018).

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DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available in the supplementary material of this article.

ORCID
Naoki Yamamoto https://orcid.org/0000-0002-2162-7203
Erxun Zhou https://orcid.org/0000-0003-1193-4661
Canwei Shu https://orcid.org/0000-0003-2463-4924

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