β-Glucosidase VmGlu2 Contributes to the Virulence of Valsa mali in Apple Tree

Yan Huang1†, Chunlei Yu1†, Cuicui Sun1, Muhammad Saleem2, Pingliang Li1, Baohua Li1 and Caixia Wang**

1 Key Laboratory of Integrated Crop Pest Management of Shandong Province, Shandong Province Key Laboratory of Applied Mycology, College of Plant Health and Medicine, Qingdao Agricultural University, Qingdao, China, 2 Department of Biological Sciences, Alabama State University, Montgomery, AL, United States

The apple tree canker is caused by Valsa mali, which produces major pathogenic factors involving multiple cell wall-degrading enzymes (CWDEs) and toxins. The β-glucosidases are among the main CWDEs, and thus, they play important roles in the virulence of necrotrophic pathogens. However, the specific roles of β-glucosidases in the virulence of V. mali remain largely unknown. In this study, we identified a β-glucosidase gene, VmGlu2, which was upregulated during the V. mali infection. We found that VmGlu2 protein had high enzyme activity of β-glucosidase using p-nitrophenyl-β-D-glucopyranoside (pNPG) as a substrate, while the VmGlu2 could convert phloridzin to phloretin with the release of glucose. The deletion and overexpression of VmGlu2 showed no effect on vegetative growth, but gene deletion mutants of V. mali showed significantly reduced pycnidia formation. The gene deletion mutants had lower β-glucosidase activities and toxin levels as compared to the wild-type strain. Therefore, these mutants showed a reduced virulence. Moreover, the overexpression of VmGlu2 did not affect toxin levels, but it significantly enhanced β-glucosidase activities, which resulted in an increased pathogenicity. Thus, we conclude that VmGlu2 is required for the full virulence of V. mali. These results provide valuable evidence to the complex role of CWDEs in the fungal pathogenicity.

**Keywords: Valsa mali, β-glucosidase activity, toxin level, pathogenicity, apple tree

INTRODUCTION

The hydrolytic enzymes of pathogens are important factors in pathogenesis because they determine the accumulation of secondary metabolites and degradation of host plant tissues (Morrissey and Osbourn, 1999; Schirawski et al., 2006; Kubicek et al., 2014). The carbohydrate active enzymes degrade cell wall polymers cell wall-degrading enzymes (CWDEs), and facilitate pathogens to invade plant tissues for nutrient acquisition (Hématy et al., 2009; Zhao et al., 2013). These enzymes that regulate the breakdown of cellulose, xylan, and pectin, are particularly important for phytopathogenic fungi, because they lack specialized penetration structures (Kikot et al., 2009; Gibson et al., 2011). There are evidences that the disruption or modification of genes encoding CWDEs may reflect their direct involvement in the infection and disease (Zhang et al., 2009; Yu et al., 2018). For example, the deletion of xylanase genes Xyn11A and BcXyl1 had a marked effect on the ability of Botrytis cinerea to infect host plants (Brito et al., 2006; Yang et al., 2018), while the gene deletion mutant of VdCUT11, encoding a cutinase, impaired the virulence of Verticillium
In *Phytophthora sojae*, the deletion and overexpression of xylglucanase gene *PsXEG1* severely reduced its virulence (Ma et al., 2015). However, the specific roles of the majority of CWDEs in the virulence remain largely unknown.

The β-glucosidases (EC 3.2.1.21) play an important role in the degradation of the cellulose that catalyze the hydrolysis of glucosides and oligosaccharides by releasing glucose (Kubicek et al., 2014). β-glucosidases are predominantly found in the glycoside hydrolases 1 (GH1) and GH3 families. Both GH families hydrolyze their respective target substrates with a net retention of the configuration of the anomeric carbon (Collins et al., 2007; Tiwari et al., 2013). These enzymes have distinct biological roles, and thus have been well characterized for their wide applications in the biomedical industry (Tokpohozin et al., 2016; Salgado et al., 2018). In addition, these enzymes also play a vital role in the production of different energy sources (biofuel) during biomass conversion (Bayer et al., 2007).

Nevertheless, the specific roles of β-glucosidases in virulence are reported in several pathogens. In *B. cinerea*, the positive correlation was found between the β-glucosidase activities and the pathogenicity (Sasaki and Nagayama, 1994, 1996), while the deletion of *BcSUN1*, encoding a β-glucosidase, had a marked effect on the production of reproductive structures as well as the ability of the pathogen to infect bean, tomato and tobacco plants (Pérez-Hernández et al., 2017). Jourdan et al. (2018) demonstrated that β-glucosidase *BglC* plays an important role in the virulence of *Streptomyces scabiei*, by affecting the intracellular accumulation of signals that trigger the thaxtomin A biosynthesis. Thaxtomin is the main phytotoxin produced by *S. scabiei*. However, the underlying mechanism of *BglC* in the virulence of *S. scabiei* is more complex than commonly perceived (Jourdan et al., 2018).

*Valsa mali*, a typical necrotrophic fungus, is the causative agent of *Valsa* canker on apple tree via the production of toxic compounds (phytotoxins) and CWDEs (Chen et al., 2012; Wang et al., 2014). A whole-genome analysis also revealed that *V. mali* contained a number of genes associated with the hydrolytic enzymes and secondary metabolite biosynthesis (Yin et al., 2015). Transcriptome profiling had also suggested that cell wall degradation is important for the infection of apple tree by the *V. mali* (Ke et al., 2014). Some studies investigated the role of CWDEs such as xylanase, polygalacturonases and ferulic acid esterases in the virulence of *V. mali* (Xu et al., 2016, 2018; Yu et al., 2018). Previous studies have found that *V. mali* secretes β-glucosidases with high activity, and its encoding genes are up-regulated during the pathogen infection (Chen et al., 2012; Li et al., 2017). This indicates that β-glucosidases play important roles in the virulence of this organism. In addition, the toxic compounds produced by *V. mali* are the degradation products of phlorizin, and the first step in the degradation process involves the conversion of phlorizin into phloretin with the release of glucose (Natsume et al., 1982). Gaucher et al. (2013) demonstrated that high β-glucosidase activity in the apple tree infected by *Erwinia amylovora* were responsible for the conversion of phlorizin into phloretin accompanied by the release of large amounts of glucose that stimulated pathogen growth in the diseased tissues. However, we know little about the comprehensive role of β-glucosidase in the virulence of *V. mali*, and thus it requires immediate research efforts.

Here, we identified a gene, *VmGlu2*, from *V. mali*, which is up-regulated in apple tree during the pathogen infection. The *VmGlu2* is a member of GH1 with β-glucosidase activity when p-nitrophenyl-β-D-glucopyranoside (pNPG) or phlorizin as a substrate. Moreover, *VmGlu2* is required for full pathogenicity of *V. mali* in apple tree, and involved in pycnidia formation. Our results indicate that *VmGlu2* has a major role in the virulence of *V. mali* and provides important information for us to understand the pathogenicity of necrotrophic fungus.

### MATERIALS AND METHODS

#### Strains and Culture Conditions

The *V. mali* wild-type strain LXS080601 was grown on the potato dextrose agar (PDA, 200 g of potato, 20 g of dextrose, and 15 g of agar per liter) at 25°C in the dark. The gene deletion and complemented strains were cultured on PDA supplemented with 100 mg/mL of hygromycin B or geneticin G418 (Sigma, St. Louis, MO, United States). The *Escherichia coli* strains were grown in the lysogeny broth (LB) with appropriate antibiotics at 37°C. The measured colony diameter was used to calculate the growth rate of different strains on PDA medium.

#### Identification of *VmGlu2* in *V. mali* and Sequence Analysis

The genomic DNA and total RNA were extracted from the *V. mali* mycelium. The cDNA was synthesized using the Prime Script RT reagent Kit with gDNA Eraser (TaKaRa, Dalian, China) using an oligo (dT)12–18 primer. The gene *VmGlu2*, with putative β-glucosidase activity and high transcript levels during *V. mali* infection was cloned. Primer pairs for the cloning of *VmGlu2* were synthesized by TsingKe (Beijing, China) (Supplementary Table 1). The PCRs were performed with PrimerSTAR Max DNA polymerase and cloned to T-Vector pMD 19 Simple (TaKaRa, Dalian, China), according to the manufacturer’s instructions.

The amino acid sequences of β-glucosidase from other strains in this study were obtained from the NCBI GenBank. All the homology searches were carried out on the NCBI BLAST server. The obtained sequences were compared with the sequences from *V. mali* (KX013493). Maximum likelihood (ML) method, as implemented in MEGA7, was used to infer the phylogenetic tree with 1000 bootstrapping replicates. Multiple sequence alignments of *VmGlu2*, and other well characterized β-glucosidase genes were performed using the DNAMAN (version 6.0) with all the parameters set at the default values.

#### Expression and Purification of Recombinant Protein

The cDNA fragment of *VmGlu2* was subcloned into the pET-32a vector by homologous recombination using the ClonExpress II One Step Cloning Kit (Vazyme, Nanjing, China). The vector construct was transformed into *E. coli* strain Rosetta while the
soluble recombinant protein of VmGlu2 was obtained after induction with 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) for 16 h at 15°C (Shi et al., 2015). The purification of recombinant VmGlu2 protein from the culture was performed using Ni-NTA Spin Column (Qiagen, Beijing, China). The expression and purification of recombinant protein were detected by the sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and Western blot.

Assays for β-Glucosidase Activities

The β-glucosidase activities were determined using pNPG as a substrate according to a previously established method (Alconada and Martínez, 1996). The enzyme activity was measured at 50°C in 50 mmol/L sodium-citrate buffer with a pH of 5.5. One unit of enzyme activity was defined as the amount of product (μmol) formed per min under the assay conditions.

The activities of β-glucosidase were also measured using the phlorizidin as a substrate by high performance liquid chromatography (HPLC) according to Gaucher et al. (2013) with some modifications. Activity was assayed in 1 ml of reaction mixture containing 0.5 mmol/L phlorizidin and about 20 μg purified recombinant protein in 50 mmol/L phosphate buffer (pH 7.0) at 25°C under continuous stirring conditions. The samples were analyzed by HPLC on an Agilent 1200 (Agilent Technologies, Santa Clara, United States), and separated on a C18 column (250 mm × 4.6 mm, 5 μm) at 30°C, with a detection wavelength at 285 nm.

Detection of Gene Expression by qRT-PCR

The mycelia plugs of V. mali were used to inoculate apple twigs described by Yu et al. (2018). The lesion border of apple bark tissues was sampled at different time points (0, 6, 12, 24, 48, 72, and 120 hpi). For samples at 0 hpi, bark tissues around inoculation sites containing mycelium plugs were collected. The RNA was extracted from bark tissues using the RNAiso Plus Kit (TaKaRa, Dalian, China), and then the cDNA was synthesized with a Reverse Transcription Kit. All of the qRT-PCR experiments were conducted in a LightCycler 480II PCR Detection System (Roche, Mannheim, Germany) with SYBR Master Mix (TaKaRa, Dalian, China) following the manufacturer’s protocol. The EF1-a of V. mali was used as an internal control, while primers are given (Supplementary Table 1). The relative expression level of VmGlu2 was calculated using the 2−ΔΔCT method (Livak and Schmittgen, 2001). Data from three biological replicates were used to calculate the means and standard deviation. The whole experiment were repeated twice.

Generation of VmGlu2 Deletion and Complementary Mutants

To obtain the VmGlu2 deletion mutants, the polyethylene glycol (PEG)-mediated homologous recombination was performed as described previously (Yu et al., 2018). VmGlu2 and about 1200 bp flanking sequences of the gene were amplified from the genomic DNA of the wild-type strain LXS080601. The hygromycin B phosphotransferase gene (HPH) gene was amplified with primers HPH-F and HPH-R from the vector pBS (Supplementary Table 1). Two flanking fragments and HPH resistance cassette were constructed into a fusion fragment (3,383 bp) using a nested PCR reaction. The fusion fragment was later transformed into the protoplasts of V. mali strain LXS080601, and the transformants were screened by culturing on medium with 100 μg/mL of hygromycin (Yu et al., 2018). More than 600 transformants were detected by PCR using the detection primer pairs VmGlu2-F/R. The putative gene deletion mutants were then validated by PCR using four primer pairs (Supplementary Table 1). The gene deletion mutants were finally verified by Southern hybridization with the DIG DNA Labeling and Detection Kit II (Roche, Mannheim, Germany) following the instruction manual.

To generate a complemented strain, the entire VmGlu2 gene with upstream fragment was amplified from genomic DNA using the primer pair 1491-CM-F/R (Supplementary Table 1). The PCR products were cloned into the plasmid pYF11 using yeast gap repair and then verified by sequencing analysis. The recombinant vector pYF11-Glu2 was then transformed into the gene deletion mutant via PEG-mediated method. The transformants were selected using G418, and confirmed by PCR with the corresponding primers (Supplementary Table 1).

Generation of VmGlu2 Overexpression Transformants

The full-length fragment of VmGlu2 was amplified from the genomic DNA of V. mali and then cloned into plasmid pYF11 using the ClonExpress II One Step Cloning Kit (Vazyme, Nanjing, China). The construct was verified by sequencing and then transformed into the wild-type strain LXS080601 as described above. Transformants were screened by PCR with corresponding primers outside the cloning sites of the pYF11. Relative transcript levels of VmGlu2 in mycelia grown on PDA and inoculated apple twigs were determined as described above. Transformants were further verified by Western blot with Anti-GFP antibody (Abcam, United Kingdom).

Vegetative Growth, Pycnidia Formation, and Pathogenicity Assays

The vegetative growth of VmGlu2 deletion mutants and overexpression transformants was determined as described by Yu et al. (2018). Colony shape and color were observed, and then the colony diameters were measured. For determination of the dry weight of mycelia, plugs were inoculated into PDB media (potato dextrose broth) at 150 rpm and 25°C for 7 days. For pycnidia formation, all the strains were grown on PDA under UV-light (365 nm) at 25°C for 4 weeks. The assays were carried out three times and each experiment with four replicates.

For pathogenicity assays, we selected healthy apple leaves and 1-year old twigs (Malus domestica Borkh. cv. “Fuji”), which were sterilized with 75% ethanol, and spaced wounds were made as described by Yu et al. (2018). The mycelia plugs of all strains were used to inoculate the wounds. The inoculated leaves and twigs were placed in trays to maintain humidity at 25°C in the dark. Then, the lesion length was measured and the photography was...
performed at different time intervals. The assays were repeated three times, with at least 10 leaves and twigs per treatment.

**Determination of Toxins Production**

For toxins production, the mycelium plugs of different strains were inoculated into apple branch extract media. After culturing for 7 days at 150 rpm/min and 25°C with a 12-h photoperiod, the supernatant was collected. The liquid media inoculated with PDA was used as a control. Toxins were extracted and detected by HPLC according to the method described by Wang et al. (2014). Each experiment was done in triplicate.

**Statistical Analysis**

All statistical analysis was conducted using SPSS software (Version 19.0, SPSS Inc., Shanghai, China). All the data collected were subjected to analysis of variance (ANOVA) followed by Duncan's multiple range tests. The asterisks indicate a statistically significant difference with the wild-type strain ($P < 0.05$).

**RESULTS**

**Sequence Identification and Analysis of VmGlu2**

The VmGlu2 gene was amplified by PCR using cDNA of V. mali as template, and then confirmed by the sequencing (Supplementary Figure 1A). The full-length cDNA of VmGlu2 is 1,677 bp containing 5’- and 3’-non-coding regions, as well an open reading frame of 1,491 bp that encodes a protein with a calculated molecular mass of 54.6 kDa. The DNA fragment was also 1,491 bp, with no intron. Using the Signal P5.0 server, VmGlu2 was predicted that no signal peptide is present. NCBI's conserved domain database showed that VmGlu2 had a β-glucosidase (Bg1B) conserved domain, which belonged to a GH1 family (Supplementary Figure 1B). The sequence comparison demonstrated that VmGlu2 exhibited high homology with the well-characterized β-glucosidase. A phylogenetic tree was constructed with β-glucosidase from other strains (Figure 1A). The sequence alignments of VmGlu2 with four characterized GH1 β-glucosidases showed that VmGlu2 had the catalytic residues among these enzymes that are related to their activity, in which Glu395 acts as a nucleophile and the Glu181 acts as an acid/base (de Giuseppe et al., 2014). The sequence alignment is shown (Figure 1B).

**Confirming the β-Glucosidase Activity of VmGlu2 Protein**

In order to determine the enzymatic activity of Bg1B, the VmGlu2 was cloned into pET-32a for heterologous expression in E. coli with a six histidine tag fused to the N-terminus part. The VmGlu2 recombinant protein with the expected molecular weight (MW) was obtained and verified by Western blot with Anti-His antibody (Figure 2A). We got the purified recombinant protein through Ni-nitrilotriacetic acid (Ni-NTA) Spin Column, and checked the protein by SDS-PAGE (Figure 2B).

The β-glucosidase activity of VmGlu2 protein was measured using pNPG as substrate at different temperatures (from 30 to 70°C) and pH values (pH 2.5–7.5). The enzyme activity remained constant from 30 to 40°C, dramatically increased to the highest value of 1.74 U/mL at 50°C, then declined abruptly to 36% of the maximal activity at 70°C. The optimal pH of VmGlu2 is around pH 5.5 as the enzyme maintained high activity between pH 3.5 and pH 6.5, while its activity declined rapidly to 30 and 37% of its optimum at pH 2.5 and pH 8.5, respectively (Figure 2C). In addition, we determined its activity when phloridzin as substrate by high performance liquid chromatography (HPLC). The Figure 2D showed that VmGlu2 had high activity to convert phloridzin into phloretin, with 78 and 100% conversion rate at 24 and 72 h, respectively.

**Expression Profile of VmGlu2**

To examine the expression level of VmGlu2 in different apple tissues, we sampled infected apple phloem and xylem tissues at 72 h post inoculation (hpi) and compared the expression level with that of mycelia grown on PDA. qRT-PCR analysis showed that the expression level of VmGlu2 was significantly up-regulated in the apple phloem (9.7-fold change) as compared to its expression in the mycelia grown on PDA. During the infection stage, the up-regulation trend of VmGlu2 in apple xylem (1.7-fold change) had no significant difference relative to its expression levels in PDA (Figure 3A).

We also determined the expression level of VmGlu2 during the interaction of V. mali with apple twigs (Figure 3B). To do this, we sampled infected apple phloem tissues at seven different time points (0, 6, 12, 24, 48, 72, and 120 hpi), and then compared gene expression levels to those of mycelia grown on PDA. The transcript level of VmGlu2 was significantly up-regulated from 12 hpi, gradually increased and reached the highest with a 13.5-fold change at 48 hpi, and then declined sharply from 72 to 120 hpi (4.8-fold change). Overall, the high induction of VmGlu2 during infection suggests its potential role in the pathogenicity of V. mali.

**VmGlu2 Is Not Necessary for the Vegetative Growth but Pycnidia Formation of V. mali**

To verify the functional role of VmGlu2 in V. mali, we successfully knocked out the gene by PEG-mediated protoplasts transformation method. The mutants were examined via PCR assays and confirmed by Southern blot (Supplementary Figure 2). When probed with HPH fragment, the wild-type (WT) sample showed no hybridization signal. However, a clear hybridization signal of the predicted size was appeared on analysis of the gene deletion mutants. For the complementation of VmGlu2 deletion mutant, a complementary construct was generated and transformed into the gene deletion mutant. All the complementation transformants were confirmed by PCR assays (Supplementary Figure 3A). In addition, RT-PCR analysis were also performed to confirm that the target VmGlu2 was knocked out in the gene deletion mutants and that the complementation strains contained the target gene (Supplementary Figure 3B).
To determine whether VmHbhs plays a role in V. mali growth and development, we investigated the colony, growth rate, and pycnidia formation of the VmGlu2 deletion mutants and the wild-type strain. However, no obvious differences either in colony morphology or growth rate were observed between the VmGlu2 deletion mutants and the wild-type strain (Figures 4A,B). Assessment of the pycnidia formation by ΔVmGlu2 was performed on PDA under UV-light (365 nm) at 25°. The VmGlu2 deletion mutants produced less than 13 pycnidia per plate, while the wild-type strain produced more than 126 pycnidia per plate. Furthermore, the complementation strain ΔVmGlu2-C by reintroducing VmGlu2 restored the wild-type strain pycnidia formation (Figure 4C).

**Deletion of VmGlu2 Reduced the Virulence of V. mali**
To investigate whether four VmGlu2 play role in the virulence, we performed pathogenicity assays of the wild-type strain and gene deletion mutants on detached apple leaves and twigs. Our
FIGURE 2 | The determination of VmGlu2 recombinant protein expression, purification and enzyme activities. (A) Detection of the expression of VmHbh2 recombinant protein by Western blot. Lane M, protein molecular weight marker (Low); Lane 1 and 3, the empty vector as control (about 20 µg of total protein); Lane 2 and 4, the supernatant of the induced cells at 15 °C (about 30 µg of total protein). (B) Detection of the purified VmHbh2 recombinant protein by SDS-PAGE. Lane M, protein molecular weight marker (High); Lane 1, the empty vector as control; Lane 2, the supernatant of the induced cells at 15 °C; Lane 3, the purified recombinant protein. (C) Enzymatic activity of VmGlu2 recombinant protein when using pNPG as a substrate. (D) Enzymatic activity of VmGlu2 recombinant protein when using phloridzin as a substrate, and the data show the conversion of phloridzin into phloretin as determined by HPLC. (1), the standard of phloridzin and phloretin; (2), VmGlu2 recombinant protein and phloridzin (1 mg/mL) incubation for 0 h; (3), VmGlu2 recombinant protein and phloridzin (0.91 mg/mL) incubation for 24 h; (4), VmGlu2 recombinant protein and phloridzin (not detected) incubation for 72 h. The data next to the peak are the retention times for each compound.
results showed that VmGlu2 deletion strains were significantly less virulent toward the apple leaves and twigs at 3 or 5 dpi, as compared to the wild-type strain, which typically displayed diseased symptoms of necrosis and canker (Figure 5). However, smaller lesions were found on the ΔVmGlu2 inoculated apple leaves and twigs than those caused by the wild-type strain (Figures 5A, B). The ΔVmGlu2 strains demonstrated a more than 60 and 76% reduction in the average lesion size on apple leaves at 3pi and twigs at 5 dpi. The complementation strain ΔVmGlu2-C recovered the high virulent phenotype that showed same symptoms on the apple leaves and twigs after reintroduction of VmGlu2 into the ΔVmGlu2 mutant (Figure 5C).
Overexpression of VmGlu2 Increased the Virulence of V. mali

To further determine the function of VmGlu2 on the pathogenicity of V. mali, the gene overexpression transformants were generated and detected by PCR (Supplementary Figure 4). The overexpression transformants were confirmed by Western blot analysis. When probed with Anti-GFP antibody, the wild-type strain showed no GFP protein signal was found, however, two transformants of VmGlu2 overexpression (OE) displayed specific GFP detected bands with the expected size. Moreover, when probed with Anti-GAPDH antibody, all the strains exhibited specific immune signals (Figure 6A). Two VmGlu2 overexpression transformants (VmGlu2-OE-78 and VmGlu2-OE-92) were selected for further analysis. Compared with the wild-type strain, the transcript levels of VmGlu2 in VmGlu2-OE-78 and VmGlu2-OE-92 increased 8.6- and 7.6-fold, respectively, during the infection apple twigs (Figure 6B). Further analysis showed that overexpression of VmGlu2 did not affect vegetative growth and pycnidia formation (Figures 7A–C). However, VmGlu2 overexpression transformants significantly enhanced the virulence of V. mali. Larger lesions were observed on the VmGlu2 overexpression transformants inoculated apple leaves (17% increase), especially apple twigs (25% increase) than those caused by the wild-type strain (Figures 7D–F). The phenotype and pathogenicity of the empty vector transformants and the wild-type strain were comparable (data not shown). These results verify that VmGlu2 is a major virulence factor involved in the V. mali pathogenicity.

Deletion and Overexpression of VmGlu2 Affects β-Glucosidase Activity in V. mali

To detect VmGlu2 activity, the wild-type strain, gene deletion mutants, and overexpression transformants were cultured in the apple branch extract media. We determined β-glucosidase activity in the culture filtrates of all strains and uses pNPG as a substrate. The gene deletion strain ΔVmGlu2 showed 50 and 39% reduction in the β-glucosidase activity at 1 and 3 dpi, respectively. The complementation strain ΔVmGlu2-C with the native gene restored β-glucosidase activity to the wild-type levels (Figure 8A). In contrast, the VmGlu2 overexpression transformants exhibited an enhanced β-glucosidase activity, with 3.0- and 3.3-fold increases compared with the wild-type strain (Figure 8B). Thus, the results in enzyme activity assays suggest the deletion of VmGlu2 significantly reduced the β-glucosidase activity.
activity, however, the gene overexpression enhanced the β-glucosidase activity.

**VmGlu2 Deletion Affects Toxins Production by *V. mali***

To further reveal the mechanism underlying the virulence of *VmGlu2* deletion and overexpression, we examined the toxin levels in the apple branch extract media for different strains. All tested strains produced five kinds of toxins, however, the Δ*VmGlu2* mutants exhibited different degrees of reduction in the levels of five toxic compounds (Figure 8C). The *VmGlu2* deletion strains exhibited 20 (phloroglucinol) to 71% (protocatechuic acid) reduction in each toxic compounds than the wild-type strain. Moreover, the total toxins produced by the Δ*VmGlu2* mutants showed 48% decrease than the wild-type strain, whereas the complementation strain with the native gene restored the phenotype. In contrast, the toxin levels produced by the *VmGlu2* overexpression transformants were not significantly different than those produced by the wild-type strain (Figure 8D).

**DISCUSSION**

The role of CWDEs in the virulence of plant pathogens has been described by previous studies, while these enzymes are considered vital for the generation of disease symptoms and pathogenesis (Brito et al., 2006; Kubicek et al., 2014; Pérez-Hernández et al., 2017; Gui et al., 2018). The β-glucosidases are among the main CWDEs, and thus play important roles especially in the virulence of necrotrophic pathogens infection and disease development (Sasaki and Nagayama, 1996; Pérez-Hernández et al., 2017; Jourdan et al., 2018). *V. mali*, the causal agent of apple tree Valsa canker, infects host plants and causes typical symptoms through secreted CWDEs and toxins. Although previous studies have demonstrated that several CWDEs play crucial function in the pathogenicity of *V. mali*, the role of β-glucosidases in the pathogen virulence is still largely unknown. In the present study, we identified a β-glucosidase gene *VmGlu2* from *V. mali* that contains GH1 glycosyl hydrolase motif with β-glucosidase activities, while it is required for the full virulence of *V. mali*.

The bioinformatics analyses showed that *VmGlu2* protein shared the characteristics of β-glucosidases and multiple conserved motifs (Brunner et al., 2002; de Giuseppe et al., 2014). The recombinant protein was successfully obtained by the prokaryotic expression system, while the purified *VmGlu2* protein showed high activity (1.74 U/mL) of β-glucosidase using pNPG as a substrate at pH 5.5 and 50 °C, thus suggesting that the *VmGlu2* encoded a β-glucosidase and participated in the degradation of cellulose during the infection of *V. mali*. This result was in line with a previous study, which demonstrated that a 66 KDa protein of β-glucosidase from *F. oxysporum*, a phytopathogenic fungus, had high enzyme activities when using pNPG as a substrate at the similar reaction conditions (Alconada and Martinez, 1996).

Interestingly, in this study, the *VmGlu2* protein could also catalyze the hydrolysis of phloridzin glucosides, and produce phloretin, which is the first step during toxin production in *V. mali* (Natsume et al., 1982; Gosch et al., 2010). Although the toxic compounds, leading to necrosis on apple tree, had long been identified, however, the specific functional genes involved in the degradation process of toxins in *V. mali* remain largely unknown. Previous study demonstrated that the candidate phloridzin-degrading enzyme gene *Vmlph1* had relationship with the vegetative growth. Moreover, it also participated in the virulence, conidiation and melanin biosynthesis in *V. mali*. Nevertheless,
Huang et al. VmGlu2 Contributes to the Virulence

Overexpression of VmGlu2 does not affect the growth and development of V. mali, but increases the virulence of V. mali in the leaves and twigs of M. domestica Borkh. cv. Fuji. (A) Colony morphology of WT and VmGlu2 overexpression (OE) transformants for 4 days incubation on PDA. (B) Growth rate of WT and VmGlu2 overexpression transformants on PDA for 3 days. (C) Number of pycnidia produced in the 9.0 cm per petri plate. Bars indicate standard deviation of means of four replicates. (D) Pathogenicity assay of the WT and VmGlu2 overexpression (OE) transformants on apple leaves at 3 dpi. (E) Pathogenicity assay of WT and VmGlu2 overexpression transformants on the apple twigs at 5 dpi. (F) Lesion sizes produced by different strains on the apple leaves at 3 dpi and apple twigs at 5 dpi. The mean lesion length was calculated from 6 apple leaves and 10 apple twigs. Bars represent the standard deviation. Asterisks indicate significant differences in OE transformants as compared to that in the WT ($P < 0.05$).

FIGURE 7 | Overexpression of VmGlu2 does not affect the growth and development of V. mali, but increases the virulence of V. mali in the leaves and twigs of M. domestica Borkh. cv. Fuji. (A) Colony morphology of WT and VmGlu2 overexpression (OE) transformants for 4 days incubation on PDA. (B) Growth rate of WT and VmGlu2 overexpression transformants on PDA for 3 days. (C) Number of pycnidia produced in the 9.0 cm per petri plate. Bars indicate standard deviation of means of four replicates. (D) Pathogenicity assay of the WT and VmGlu2 overexpression (OE) transformants on apple leaves at 3 dpi. (E) Pathogenicity assay of WT and VmGlu2 overexpression transformants on the apple twigs at 5 dpi. (F) Lesion sizes produced by different strains on the apple leaves at 3 dpi and apple twigs at 5 dpi. The mean lesion length was calculated from 6 apple leaves and 10 apple twigs. Bars represent the standard deviation. Asterisks indicate significant differences in OE transformants as compared to that in the WT ($P < 0.05$).
the deletion mutant of Vmlph1 did not affect the ability of the pathogen to degrade phloridzin (Zhu et al., 2018). In addition, the function of Vmlph1 protein to degrade phloridzin still has not been verified. In this study, our results indicated that VmGlu2 had high enzymatic activities to hydrolyze phloridzin, thus suggesting that VmGlu2 also plays crucial roles in toxins production by V. mali.

The β-glucosidases have been reported as pathogenicity factors in several plant pathogens. For example, the β-glucosidases involved in the virulence of B. cinerea and
S. scabies (Sasaki and Nagayama, 1996; Jourdan et al., 2018), and disruption of the BcSUN1 gene resulted in different cell surface alterations affecting the infection of B. cinerea, therefore decreasing its virulence potential (Pérez-Hernández et al., 2017). As in these examples from fungi and bacteria, VmGlu2 is a critical pathogenicity factor in V. mali, as evidenced by its high expression during plant infection. Furthermore, we also found that VmGlu2 exhibited higher transcript levels in the apple phloem than in the apple xylem. Based on the finding from previous studies that V. mali could grow rapidly and survived for a long time in the apple xylem, but it did not develop diseased symptoms, until the pathogen reached the phloem (Chen et al., 2016; Wang et al., 2018), we speculate that the expression pattern of VmGlu2 is in line with their roles in the pathogenicity of V. mali.

Subsequently, we generated VmGlu2 deletion and complementation stains to investigate the role of VmGlu2 in the growth, development, and pathogenicity of V. mali. The ΔVmGlu2 mutants exhibited similar morphology and growth rate of mycelia as compared to the wild-type strain. Intriguingly, our results revealed that VmGlu2 was necessary for pycnidia formation and pathogenicity of V. mali. Moreover, the ability of producing pycnidia and pathogenicity was restored in the VmGlu2 complementation strains. Similar results were demonstrated in other CWDEs in phytopathogens, for example, the deletion of cutinase VdCUT11 did not affect mycelia growth and colony morphology, but it contributed to the virulence of V. dahliae. Therefore, in this study, the VmGlu2 deletion and overexpression strains, respectively, were demonstrated in other CWDEs in phytopathogens, for example, the deletion of cutinase VdCUT11 did not affect mycelia growth and colony morphology, but it contributed to the virulence of V. dahliae, and the deletion mutants of feruloyl esterases FAEs exhibited a significant reduction in the pathogenicity. But it had no effect on the vegetative growth and development of V. mali (Gui et al., 2018; Xu et al., 2018). In addition, we obtained VmGlu2 overexpression transformants, and found that the gene overexpression did not affect the vegetative growth and development, but it increased the pathogenicity of V. mali. Based on the deletion and overexpression of VmGlu2 as well as pathogenicity assays, VmGlu2 was confirmed to be involved in the virulence of V. mali in apple trees.

The conidia production is a vital phase in the life cycle of pathogens, therefore, it is generally believed that inhibition of pycnidia formation by V. mali could alleviate or effectively control the occurrence of apple tree Valsa canker. However, the relationship between conidiation and pathogenicity of V. mali is not yet clear. In a previous study, the deletion of VmE02, encoding the pathogen-associated molecular pattern (PAMP), demonstrated attenuated conidiation but not the virulence (Nie et al., 2019). Furthermore, the deletion of mitogen-activated protein kinase gene, VmPmk1, could lead to a reduced growth rate, and a decreased pathogen virulence. Meanwhile, it also leads to absence of pycnidia, thus implying no pycnidia production in mutants (Wu et al., 2017). The deletion mutant of VmXyl1, encoding the xylanase, produced normal growth rate and exhibited decreased pycnidia formation and virulence (Yu et al., 2018). Moreover, the deletion mutants of VmVelA and VmVelB, Velvet protein family members, showed increased conidiation and melanin production, but they exhibited reduced virulence (Wu et al., 2017, 2018; Yu et al., 2018). In the present study, the ΔVmGlu2 mutants exhibited normal growth rate and decreased pycnidia formation and pathogenicity. These results suggest that conidiation and virulence of V. mali are not necessarily correlated.

The major pathogenic factors of V. mali involve multiple CWDEs and toxins (Natsume et al., 1982; Chen et al., 2012; Wang et al., 2014). Therefore, in this study, the VmGlu2 activities of β-glucosidase and production of toxins were investigated. The β-glucosidase activities of the ΔVmGlu2 mutants were significantly lower, while the enzyme activities of the VmGlu2 overexpression transformants were notably higher, when compared with those of the wild-type strain, which was consistent with the reduced and enhanced virulence of the gene deletion and overexpression strains, respectively. Similarly, the deletion mutants of VmXyl1 and most of the FAEs that exhibited significant decreases in virulence had significantly lower enzyme activities than the wild-type strains (Xu et al., 2018; Yu et al., 2018). Moreover, in B. cinerea, the deletion of Xyn11A resulted in a reduction in the xylanase activity and virulence, however, further study demonstrated that Xyn11A mainly contributed to the fungal virulence with its inducing necrosis of the infected plant tissue and not with its enzyme catalytic activity (Brito et al., 2006; Noda et al., 2010). In contrast, interruption of a xylanase gene, FGSG_03624, from F. graminearum demonstrated a significant reduction in the xylanase activity, but it did not impair its virulence (Sella et al., 2013). These complexities might be attributed to the functional redundancies of the CWDEs genes. By analysis the genome sequence of V. mali, we found three genes of GH1 family encode β-glucosidase with 40.7% identity. The three genes all have typical catalytic residues of GH1 enzymes in which two Glu act as the nucleophile and acid/base, respectively (de Giuseppe et al., 2014). In our previous study, we cloned VmGlu1 (KY646110) and found the gene significantly up-regulated during V. mali infection (Li et al., 2017). However, the biological function of VmGlu1 needs to be further studied.

In addition, our data also showed that toxin production by the ΔVmGlu2 mutants were significantly lower than the wild-type strain, which also corresponds to the results of the reduced β-glucosidase activities and virulence. However, the overexpression of VmGlu2 did not increase the toxin production. The degrading rates of phloridzin were both almost up to 100% for the wild-type and overexpression strains at 7 days after incubation in the apple branch extract media (data not shown). Therefore, we speculated that the β-glucosidases produced by V. mali were excessive to involve in the degradation of phloridzin. In apple and E. amylovora interaction, β-glucosidase activity was significantly higher in the susceptible genotype MM106 than that in the resistant genotype Evereste, which contributes to the fast multiplication of the bacteria in the leaves of MM106 (Gaucher et al., 2013). In the interaction of apple and V. mali, the phloridzin is converted into phloretin with the release of glucose by the action of VmGlu2. The phloretin is the precursor of five toxic compounds produced by V. mali, which determine the severity of necrosis on the apple tissues.
On the other hand, the release of glucose could favor the growth of *V. mali* in apple tissues. Taken together, these data suggest that the role of β-glucosidases is complex in the virulence of *V. mali*. However, the functions of β-glucosidase genes and their cooperative roles deserve to be further investigated.

**CONCLUSION**

We have demonstrated that the protein encoded by *VmGlu2* has high β-glucosidase activity when using the pNPG as a substrate, and it can convert phloridzin into phloretin while releasing the glucose. In addition, we reveal that *VmGlu2* is required for full pathogenicity of *V. mali* in apple tree, and it significantly influences its pycnidia formation. These results provide a vital evidence to explore further the complex functions of CWDEs in fungal pathogenicity.

**DATA AVAILABILITY STATEMENT**

Bioinformatics analysis data was downloaded from the National Center for Biotechnology Information (NCBI) database (URL: https://www.ncbi.nlm.nih.gov/) under the accession numbers BAA74958, AAF74209, BAA74959, QGZ11144, and AAM94393.

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AUTHOR CONTRIBUTIONS

YH, CY, and CS carried out the experiments and analyzed the data with the help of MS. PL determined the toxins levels by HPLC. YH and CW wrote the manuscript with help from all authors. BL performed the manuscript revision and provided part of the financial support. MS reviewed and edited the manuscript. All authors approved its final version.

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**SUPPLEMENTARY MATERIAL**

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