G-Protein Subunit Gα<sub>i</sub> in Mitochondria, MrGPA1, Affects Conidiation, Stress Resistance, and Virulence of Entomopathogenic Fungus *Metarhizium robertsii*

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G proteins are critical modulators or transducers in various transmembrane signaling systems. They play key roles in numerous biological processes in fungi, including vegetative growth, development of infection-related structures, asexual conidiation, and virulence. However, functions of G proteins in entomopathogenic fungi remain unclear. Here, we characterized the roles of MrGPA1, a G-protein subunit Gα<sub>i</sub>, in conidiation, stress resistance, and virulence in *Metarhizium robertsii*. MrGPA1 was localized in the mitochondria. MrGpa1 deletion resulted in a significant reduction (47%) in the conidiation capacity, and reduced expression of several key conidiation-related genes, including *fluG*, *flbD*, *briA*, *wetA*, *phiA*, and *stuA*. Further, MrGpa1 disruption resulted in decreased fungal sensitivity to UV irradiation and thermal stress, as determined based on conidial germination of ΔMrGpa1 and wild-type (WT) strains. Chemical stress analysis indicated that MrGpa1 contributes to fungal antioxidant capacity and cell wall integrity, but is not involved in tolerance to antifungal drug and osmotic stress. Importantly, insect bioassays involving (topical inoculation and injection) of *Galleria mellonella* larvae revealed decreased virulence of ΔMrGpa1 strain after cuticle infection. This was accompanied by decreased rates of appressorium formation and reduced expression of several cuticle penetration-related genes. Further assays showed that MrGpa1 regulated intracellular cyclic AMP (cAMP) levels, but feeding with cAMP could not recover the appressorium formation rate of ΔMrGpa1. These observations suggest that MrGpa1 contributes to the regulation of conidiation, UV irradiation, thermal stress response, antioxidant capacity, and cell wall integrity in *M. robertsii*. This gene is also involved in insect cuticle penetration during infection. These findings raise the possibility of designing powerful strategies for genetic improvement of *M. robertsii* conidiation capacity and virulence for killing pests.

**Keywords:** *Metarhizium robertsii*, Gα<sub>i</sub>, protein, appressorium, conidiation, virulence
INTRODUCTION

G protein with GTP-hydrolase activity is a type of signaling protein that binds to guanine nucleotides (Robishaw and Berlot, 2004). It participates in signal transduction pathways linking activated cell-surface receptors with intracellular effectors, including adenylyl cyclase and phospholipase through a series of signaling cascades involved in the regulation of physiological and biochemical processes (Ortiz-Urquiza and Keyhani, 2015; Chakravorty and Assmann, 2018). In fungi, G protein is associated with sexual and asexual reproduction, virulence, and response to external signal stimuli (Ivey et al., 1996; Liu and Dean, 1997; Guo et al., 2016b).

The heterotrimeric G protein is composed of three subunits α, β, and γ, wherein α subunit binds to GDP, and β and γ subunits form a heterodimer (Birnbaumer, 2007; Lambert, 2008; McIntire, 2009). When the heterotrimeric G protein is stimulated by a G-protein-coupled receptor (GPCR) that senses external signals, GDP is exchanged for GTP, and Gα subunits dissociate (Wedegaertner, 2012). Then, Gα-GTP and Gβγ complexes dissociate (Barren and Artemyev, 2007). The cycle is reset by the hydrolysis of GTP to GDP, and Gα recombining with Gβγ and GPCR (Gilchrist et al., 1999; Slessareva and Dohlman, 2006).

In mammals, G-protein α (GPA) subunits are divided into four classes, Gαi, Gαq, Gαz, and Gα12, based on the amino acid sequence identity (Neer, 1995). Further, the Gαi family is composed of four subfamilies, Gαia, Gαiq, Gαiz, and Gα12z. The functions of Gα family proteins are diverse and include regulation of adenylyl cyclase, K+ and Ca2+ channels, and cGMP phosphodiesterase activities (Simon et al., 1991). The conserved functional motif of Gαi protein is characterized by possession of N-myristoylation and ADP-ribosylation (Buss et al., 1987).

The functions of the Gαi subunit have been characterized in some fungi. In Saccharomyces cerevisiae, two kinds of Gα proteins have been identified (Gpa1 and Gpa2). Gpa1 is involved in pheromone regulation (Jahng et al., 1988), while Gpa2 regulates pseudohyphal development via cyclic AMP (cAMP)-dependent pathways and heat resistance (Kubler et al., 1997). However, these proteins have different functions in other filamentous fungi. For example, Fusarium oxysporum f. sp. cubense possesses three Gα proteins (Ga-fga1, -fga2, and -fga3), and the deletion of encoding genes leads to phenotypic defects in colony morphology, reduced conidiation, increased heat tolerance, reduced virulence and intracellular cAMP levels (Jain et al., 2002; Guo et al., 2016a,b). Further, three different Gα proteins control unique signal transduction pathways in Magnaporthe grisea, influencing fungal vegetative growth, conidiation, conidium attachment, appressorium formation, mating, and pathogenicity (Liu and Dean, 1997; Zhang et al., 2012). GNA-1 protein is also required for the extension of basal hypha, growth, conidiation, and formation of female reproductive structures in Neurospora crassa (Ivey et al., 1996; Yang and Borkovich, 1999). In entomopathogenic fungus, some upstream and downstream genes for Gpa, such as GPCRs gene BbGpcr3, and regulators of the G protein signaling (RGS) genes BbRgs1 and Mrcag8, have been only characterized (Fang et al., 2007, 2008; Ying et al., 2013), but the function of Gpa in insect pathogenic fungi is poorly understood. Thus, it is necessary to characterize the Gα proteins, and figure out whether the Gα proteins are involving in vegetative growth, conidiation, stress resistance, and virulence in entomopathogenic fungi, such as Metarhizium robertsii.

Metarhizium robertsii, an important entomopathogenic fungus, has been developed as an environmentally friendly alternative to chemical insecticides (Frazzon et al., 2000; Lord, 2005; Wang and Wang, 2017). Unfortunately, commercialized broad application of M. robertsii formulations is limited by the low conidiation rate, failure of conidia germination under high-temperature and UV stress, slow killing speed, and inconsistent field performance (Faria and Wraight, 2007; Fang et al., 2012; Muniz-Paredes et al., 2017). Genetic improvements of this mycoinsecticide require extensive understanding of the molecular mechanisms and M. robertsii genes involved in stress tolerance and virulence (Zhang and Feng, 2018).

In the current study, we aimed to investigate the role of Gα proteins in M. robertsii. BLASTP search against the assembled draft genome sequence of M. robertsii identified four putative Gα proteins. Among these Gα proteins, MrGPA1 (EFZ00892) shared the highest identity (96.32%) with GNA-1. We then characterized the biological function of MrGPA1 by constructing and analyzing MrGPA1 gene deletion mutant. We show that MrGPA1 influences conidiation, stress resistance, virulence, and intracellular cAMP levels in M. robertsii.

MATERIALS AND METHODS

Fungal Strains and Culture

In the present study, M. robertsii strain ARSEF 23 was the wild-type (WT) strain. All M. robertsii strains were inoculated onto potato dextrose agar (PDA, 20% potato, 2% glucose, and 2% agar, w/v), and cultured at 25°C for 10 days. Conidial suspensions were obtained by vortex-mixing in 0.05% (v/v) Tween-80, and filtered through sterile non-woven fabric to remove mycelial debris.

Sequence Analysis

To construct the phylogenetic tree of Gα proteins and analyze the structural domains of guanine nucleotide-binding site, amino acid sequences of the Gα subunits were downloaded from the National Center for Biotechnology Information1 and phylogenetic analysis was performed using MEGAX software2.

Construction of MrGPA1-GFP Fusion Vector and Analysis of Subcellular Localization of MrGPA1

To monitor subcellular localization of MrGpa1, gfp and MrGpa1 gene fragments were amplified by polymerase chain reaction (PCR), using gfp-F/gfp-R, and gfpMrGpa1-F/gfpMrGpa1-R primers (Supplementary Table S1), high-fidelity Taq DNA polymerase (KOD Plus Neo, Toyobo, Osaka, Japan), and

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1https://www.ncbi.nlm.nih.gov/
2https://www.megasoftware.net/
M. robertsii genomic DNA as a template. The amplification products were inserted into the EcoRI restriction site in pDHt-SK-bar vector (kindly provided by Dr. Chengshu Wang; the vector conferred resistance against glufosinate-ammonium) (Fang et al., 2006) containing a strong promoter and terminator to generate vector pDHt-MrGpa1-gfp for Agrobacterium tumefaciens transformation. The corresponding transformants resistant to glufosinate ammonium were obtained, and verified by PCR using the primers gfp-F and gfp-R (Supplementary Table S1).

The MrGPA1-GFP strain was cultured on sabouraud dextrose agar medium containing yeast extract (SDAY, 4% glucose, 1% peptone, 2% agar, and 1% yeast extract powder, w/v) at 25°C for 2 days. The hyphae were then washed off the plate with sterile water and mixed with 500 nM MitoTracker Red CMXRos (Invitrogen, Shanghai, China), a dye specific to mitochondria. Subcellular localization of MrGPA1 was evaluated using a laser scanning confocal microscopy (LSCM, Zeiss LSM880). Before using laser scanning confocal microscopy, Wolf PSORT software was used for prediction subcellular localization by analyzing protein sequence of MrGPA1.

**Gene Deletion and Complementation**

To disrupt MrGpa1 gene, the 5′- and 3′-flanking regions of MrGpa1 were obtained by using MrGpa1-5F/MrGpa1-5R and MrGpa1-3F/MrGpa1-3R primers, genomic DNA (using the Plant Genomic DNA Kit; Tiangen, Beijing, China) extracted as a PCR template and high-fidelity Taq DNA polymerase (KOD Plus Neo, Toyobo, Osaka, Japan). The amplification products were then inserted into the pDHt-SK-bar vector (containing glufoisinate resistance gene) (Fang et al., 2006) digested with the SmaI, BamHI, and XbaI restriction enzymes to generate vector pDHt-MrGpa1-bar for A. tumefaciens transformation (Fang et al., 2006). ΔMrGpa1 strains were obtained by selection for glufoisinate resistance, and two fragments containing the MrGpa1 upstream-bar-MrGpa1-downstream cassette subsequently verified by PCR and southern blotting (RT)-PCR using primer pairs MrGpa1-F/MrGpa1-R, upMrGpa1-F/upMrGpa1-R, dnMrGpa1-F/dnMrGpa1-R, gpd-F/gpd-R, and bar-F/bar-R (Supplementary Table S1).

For gene complementation, the entire MrGpa1 gene and the 1000-bp upstream sequence and 600-bp downstream sequence were inserted into vector pDHt-SK-ben (containing benomyl resistance gene) digested with the SpeI restriction enzyme for fungal transformation. The 3050-bp fragment was ectopically integrated into ΔMrGpa1 strain by the same method as that used for gene deletion. Complemented strains (cpΔMrGpa1) were obtained by selection for benomyl resistance, and verified by PCR using primer pairs MrGpa1-F/MrGpa1-R and ben-F/ben-R (Supplementary Table S1).

**Phenotype Assays**

For phenotypic assays, these experiments were performed with three technical and biological replicates per strain (WT, ΔMrGpa1, and cpΔMrGpa1). Fungal conidiation ability was evaluated as previously described (Meng et al., 2017). Briefly, 30 µl of conidial suspension (1 × 10^6 conidia/ml) was spread on PDA plate (35-mm diameter). After culturing at 25°C for 14 days, the conidia on each plate were collected into 30 mL of 0.05% Tween-80 by vortex-mixing, and conidial density was determined using a hemocytometer and converted to the number of conidia per square centimeter of colony.

To evaluate the fungal vegetative growth, 1 µl of WT, ΔMrGpa1, and cpMrGpa1 conidial suspensions (1 × 10^6 conidia/ml) was spotted on PDA and 1/4 SDAY (1/4 dilution of SDAY) media, and incubated in the dark at 25°C for 10 days. Colony diameters were then measured.

For conidial germination assay, 10 µl of conidial suspension (5 × 10^6 conidia/ml) were spread on PDA medium. The conidial germination was observed by microscope (Olympus BX 51, Tokyo, Japan) at 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24 h after incubated at 25°C. Conidia were considered to be germinated when the length of the germ tube reaches or longer than the length of the conidia (Wang et al., 2014). Three hundred conidia were counted at least per plate and the germination rates were calculated by comparing the number of germinated conidia with the 300 counted conidia, and the median germinate time (GT\textsubscript{50}) was calculated using the SPSS software.

For heat stress tolerance assays, 1 ml of conidial suspensions (5 × 10^6 conidia/ml) of WT, ΔMrGpa1, and cpΔMrGpa1 strains were placed in 1.5-ml Eppendorf tubes, and then incubated in a water bath at 42 or 28°C (as control) for 1 h. Then 10 µl of the suspension were spread on PDA medium, incubated at 25°C. Conidial germination was observed under a microscope (Olympus BX 51, Tokyo, Japan) after 16 and 24 h. Three hundred conidia were counted at least by per plate and the relative germination rates were calculated by comparing the number of germinated conidia with had not been heat stressed (Wang et al., 2019).

To determine fungal tolerance to ultraviolet B (UV-B) light, 10 µl of conidial suspensions (5 × 10^6 conidia/ml) of WT, ΔMrGpa1, and cpΔMrGpa1 was taken to PDA medium. The plates were then exposed to UV-B irradiation (312-nm wavelength at 100 µJ cm\textsuperscript{-2}) using HL-2000 Hybrilinker (UVP, CA, United States) (Yao et al., 2010) or exposed to sunlight (as control). Relative UV-B tolerance was assessed and calculated by aforementioned methods to assess relative germination rate of tolerance to UV-B.

To examine the fungal tolerance to chemical stress, 1 µl conidial suspensions (1 × 10^6 conidia/ml) of WT, ΔMrGpa1, and cpΔMrGpa1 strains were spotted onto PDA medium containing carbendazim (2 µg/ml), NaCl (0.5 M), H\textsubscript{2}O\textsubscript{2} (2 mM), or Congo red (2 µg/ml), and incubated in the dark for 10 days at 25°C. Colony diameter was then measured and the relative inhibition rate was calculated (Ying and Feng, 2011; Wang et al., 2017).

To assess the effects of MrGpa1 disruption on virulence, bioassays with Galleria mellonella larvae (RuiQing Bait, Shanghai, China) were performed as described previously (Zhou et al., 2018). The larvae were immersed in conidial suspension (1 × 10^6 conidia/ml) for 90 s or injected (into the hemocoel) with 10 µl of conidial suspensions (1 × 10^5 conidia/ml) and

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\[https://wolfsort.hgc.jp\]
incubated at 25°C, each treatment was performed in triplicate, with 18 larvae in each group. The experiment was repeated three times. Larva mortality was evaluated every 24 h, and the median lethal time (LT<sub>50</sub>) was calculated using the SPSS software.

The appressorium formation assay was performed as described previously (Gao et al., 2013). Briefly, to test the appressorium formation on a hydrophobic surface, 1 ml of conidial suspension (1 × 10<sup>6</sup> conidia/ml) in MMGly (minimal medium amended with 1% glycerol) was spread on a sterile plastic Petri dishes (3.5-cm diameter), followed by 24 and 48 h incubation at 25°C. At least 300 conidia of each strain were evaluated microscopically, and the induction rates of appressorium formation were quantified by observing different microscopic fields (inverted microscope, Olympus IX 71, Tokyo, Japan).

Quantitative RT-PCR (RT-qPCR)
To analyze the expression of conidiation-related genes, 200 µl of conidial suspensions (1 × 10<sup>6</sup> conidia/ml) of WT, ΔMrGpa1, and cpΔMrGpa1 strains were plated on PDA medium, and cultured in the dark at 25°C for 2.5 days. The samples were collected and milled in liquid nitrogen to extract total RNA. To analyze the expression of virulence genes related to cuticle infection, G. mellonella larvae were dipped in conidial suspensions (5 × 10<sup>7</sup> conidia/ml) of WT, ΔMrGpa1, and cpΔMrGpa1 strains for 1.5 min, transferred to 25°C for 48 h, and then placed in liquid nitrogen for total RNA extraction. Total RNA was extracted by using Trizol reagent (Invitrogen, Foster City, CA, United States). cDNA was obtained by using the PrimeScript<sup>™</sup> RT reagent kit with gDNA Eraser (TaKaRa, Dalian, China), and used as a template for RT-qPCR. The gene expression analysis was performed by using the CFBR96<sup>™</sup> Real-Time PCR System (Bio-Rad, Hercules, CA, United States) and SYBR<sup>®</sup> PremixEx Taq<sup>™</sup> II (TaKaRa). Three biological repeats of each treatment were analyzed. The qPCR primers are listed in Supplementary Table S2. The expression of the gpd gene (MAA_07675, encoding glyceraldehyde 3-phosphate dehydrogenase) was used as an internal control (Fang and Bidochka, 2006). The relative gene expression was calculated by using the 2<sup>−ΔΔCt</sup> method (Livak and Schmittgen, 2001).

cAMP Assay
To measure the intracellular cAMP levels of WT, ΔMrGpa1, and cpΔMrGpa1 strains, the method was performed previously (Liu et al., 2012). The cAMP levels were measured by high-performance liquid chromatography (HPLC) analysis as described previously (Liu et al., 2016).

To test whether exogenous cAMP could recover appressorium formation rate of ΔMrGpa1, the conidial suspensions (1 × 10<sup>6</sup> conidia/ml) were amended with the final concentration of 5 mM cAMP and left for appressorium formation on a hydrophobic surface as mentioned above.

Statistical Analysis
All data are presented using GraphPad Prism version 6.0. Data are expressed as the mean ± standard error (SE) of the mean, from three biological replicates. Statistical analysis was performed by one-way analysis of variance (ANOVA). For multiple comparisons, Tukey’s multiple comparison test was used to analyze statistical the significance. p < 0.05 was considered to be significant, and p < 0.01 was considered to be extremely significant.

RESULTS
Sequence Characteristics of MrGPA1
From M. robertsii
We chose the sequences of GNA-1 proteins of the model fungus N. crassa as reference sequences to retrieve their M. robertsii orthologs. We thus identified four genes encoding GPA subunit [GenBank accession numbers EFZ00892.1, EFY98464.1, EFY99066.2, and EFZ00060.2, named MrGpa1, MrGpa2, MrGpa3, and MrGpa4 (i.e., the M. robertsii Gpa1, Gpa2, Gpa3, and Gpa4 genes), respectively] in the genome of M. robertsii ARSEF 23.
Further bioinformatics analysis indicated that MrGpa1 (MAA-03488) is a single copy gene encoding GPA subunit (353-aa protein) in M. robertsii. A BLASTP search of MrGPA1 homologs in NCBI revealed that the protein shares 100% amino acid similarity with GPA from Metarhizium acridum (XP_007811324), Metarhizium anisopliae (KFG82129), Metarhizium brunneum (XP_014542423), and Metarhizium rileyi (OAA44756). Phylogenetic tree of GPA proteins from
**FIGURE 3** | MrGpa1 is required for fungal stress response. (A) Germination rate and $GT_{50}$ values for three fungal strains after growth on PDA. (B) Relative germination rate of three fungal strains after exposure to heat stress and UV-B treatment. (C) Inhibition ratio and colony phenotype values for three fungal strains, on PDA medium containing Carbendazim (2 $\mu$g/ml), NaCl (0.5 M), $H_2O_2$ (10 mM), or Congo red (2 mg/ml). *$p < 0.05$, **$p < 0.01$. 

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**FIGURE 4** | MrGpa1 deletion impacts fungal virulence. (A) Survival of *G. mellonella* after topical application of conidial suspensions from three fungal strains. The control insects were treated with sterile water. (B) *LT* 50 (days) of three fungal strains after topical inoculation of larvae. (C) Survival of *G. mellonella* after injection of conidial suspension of three fungal strains. The control insects were treated with sterile water. (D) *LT* 50 (days) of three fungal strains after injection. **p < 0.01.

*Metarhizium* spp., and related fungal species was constructed with *S. cerevisiae* as an outgroup (Supplementary Figure S1A). All GPA proteins from the genus *Metarhizium* formed an independent branch (100% support value). These *Metarhizium* GPA proteins are closely related to GPA proteins from *Pochonia chlamydosporia* (99.72%), *Moelleriella libera* (99.43%), and *Purpureocillium lilacinum* (99.15%).

A conserved domain database search demonstrated that MrGPA1 contains a highly conserved guanine nucleotide-binding site (34-347-aa protein), which is the key identification domain of the GPA subunit (Supplementary Figure S1B). Furthermore, homologous alignment revealed the presence of *N*-myristoylation and ADP-ribosylation sites (two conserved positions in the GPAi subunit) in the conserved functional motif of MrGPA1. Hence, MrGPA1 is a member of the Gαi family.

### MrGPA1 Is a Mitochondria Protein

To investigate the subcellular localization of MrGPA1, the Wolf PSORT software was first used. The analysis predicted that the protein is localized in the mitochondria.

To verify this prediction, we generated *MrGpa1-gfp* strain (Supplementary Figures S2A,B). As shown in LSCM images in Figure 1, the mitochondria in hypha is stained with a mitochondrial dye (red), in a punctate patterns, while punctate green fluorescence was also observed in vegetative hyphae. Red and green fluorescence was detected and overlapped (Figure 1), suggesting that MrGPA1 is a mitochondria protein.

### Construction of MrGpa1 Knockout and Complementation

The result indicated the presence of a 1,222-bp fragment corresponding to the partial *MrGpa1* gene sequence in the WT and cpΔMrGpa1 stains, but not in the ΔMrGpa1 strain. In addition, a partial 806-bp *bar* gene fragment was detected in ΔMrGpa1 and cpΔMrGpa1 strains, and a partial 785-bp *ben* gene fragment was detected in the cpΔMrGpa1 strain. Furthermore, PCR analysis indicated the presence of a fragment containing upstream sequence of *MrGpa1* and a partial *bar* gene (2689 bp) and a fragment containing downstream sequence of *MrGpa1* and a partial *bar* gene (1975 bp), and detected by using the primer sets up
MrGpa1-F/upMrGpa1-R and dnMrGpa1-F/dnMrGpa1-R, respectively, in the ΔMrGpa1 strain. Finally, RT-PCR analysis verified the loss or regain of the MrGpa1 gene expression in ΔMrGpa1 and cpΔMrGpa1 strains, accordingly. These observations indicated a successful construction of the MrGpa1 knockout and complementation strains (Supplementary Figure S2).

**MrGpa1** Contributes to Fungal Conidiation but Is Not Involved in Vegetative Growth

To examine the effect of MrGpa1 on the growth and development of *M. robertsi*, we evaluated mycelial growth and conidial yield of WT and mutant strains on PDA and 1/4 SDAY medium, respectively. The 14-day-old colonies of WT, ΔMrGpa1, and cpΔMrGpa1 strains formed 6.18 × 10^7, 3.28 × 10^7, and 6.55 × 10^7 conidia/cm^2, respectively (Figure 2A). The loss of MrGpa1 caused a significant, 47% reduction in conidiation, but little difference in the growth rate of WT, ΔMrGpa1, and cpΔMrGpa1 strains on PDA and 1/4 SDAY media was apparent (Figures 2A,B). We also examined the expression of genes involved in conidiation in *M. robertsi* by RT-qPCR. The expression of fluG, flbD, brlA, wetA, phiA, and stuA genes in the MrGpa1 strain was significantly reduced compared with that in the WT and cpΔMrGpa1 strains (Figure 2C). Collectively, these observations indicate that while MrGPA1 plays an important role in the conidiation of *M. robertsi*, it is not involved in vegetative growth.

**MrGpa1** Is Important for Heat and UV Stresses Tolerance, and Is Involved in Antioxidant Capacity and Cell Wall Integrity of *M. robertsi*

The GT50 values for WT, ΔMrGpa1, and cpΔMrGpa1 were 11.99 h, 6.16 h (*p* < 0.01, compared with WT strain), and 12.16 h, respectively (Figure 3A). Conidial germination speed of ΔMrGpa1 strain on PDA medium was significantly faster than that of WT and cpΔMrGpa1 strains.

To investigate the effects of MrGpa1 deletion on UV irradiation and thermal stress, the relative germination rate of conidia exposed to these stresses was determined 16 or 24 h after stress exposure. We found that the sensitivity of ΔMrGpa1 strain to 42°C heat stress was reduced. For example, compared with the WT, ΔMrGpa1 germination rates at 16 and 24 h increased by 51% (*p* < 0.01) and 24% (*p* < 0.05), respectively (Figure 3B). Similar results were obtained for conidial tolerance of UV irradiation; compared with the WT, ΔMrGpa1 germination rates at 16 h increased by 83% (*p* < 0.01), but only by 8% (*p* < 0.05) at 24 h (Figure 3B). Hence, it appears that MrGPA1 plays an important role in conidial tolerance of both UV irradiation and thermal stress.

To evaluate the role of MrGpa1 in fungal growth under different chemical stress conditions, we investigated the mycelial growth of the WT and mutant strains on PDA containing carbendazim, NaCl, H_2O_2_, or Congo red. The antioxidant capacity and cell wall integrity of ΔMrGpa1 strain were significantly different than those of the WT and cpΔMrGpa1 strains. For instance, compared with the WT, the relative inhibition of ΔMrGpa1 growth was decreased by 68.6% (*p* < 0.01) on PDA containing H_2O_2_, while the sensitivity to Congo red was increased by 47.4% (*p* < 0.01). However, the relative inhibition of ΔMrGpa1 growth in the presence of carbendazim and NaCl was not markedly different from that of the control strains (Figure 3C). These observations indicate that MrGpa1 contributes to fungal antioxidant capacity and cell wall integrity, but is not involved in antifungal ability and osmotic stress.
**MrGpa1** Plays an Important Role in Insect Cuticle Penetration via Appressorium Formation

We next used *G. mellonella* bioassays to assess the consequences of MrGpa1 deletion on fungal virulence. In topical infection bioassays, the mean lethal times to death (LT₅₀) in insects infected with ΔMrGpa1, WT, and cpΔMrGpa1 strains were 7.2 ± 0.45, 11.8 ± 0.54, and 8.3 ± 0.71 days, respectively, with a significant (p < 0.05) attenuation of virulence in *G. mellonella* (Figures 4A,B). The treatment with ΔMrGpa1 also resulted in an increased survival rate of the larvae compared with the WT and cpΔMrGpa1 strains treatment. By contrast, in the injection bioassays, we did not observe any differences in LT₅₀ values between larvae infected with ΔMrGpa1 (3.69 ± 0.15 days), and WT (3.53 ± 0.14 days) or cpΔMrGpa1 strains (3.57 ± 0.12 days) (Figures 4C,D).

We then determined the expression of insect virulence-related genes during cuticle penetration by RT-qPCR. Indeed, the expression of several genes involved in the adhesion (*mad1*, 45% expression in ΔMrGpa1 strain compared with the WT strain), appressorium formation (*mpl1*, 72%, and *gpa*, 52%), and cuticle penetration (*pr1A*, 93%, and *pr1C*, 96%) was significantly decreased in the ΔMrGpa1 strain compared with their expression in the WT strains (Figure 5A).

To determine the mechanism of the virulence defect of ΔMrGpa1 strain, we then assayed appressorium formation on a hydrophobic surface. We observed that the loss of MrGpa1 impaired appressorium differentiation, compared with the control strains. Specifically, 24 h after induction, ΔMrGpa1 strain did not form appressoria, while the appressorium formation rate of the WT strain was 80% (Figure 5B). Further, 48 h after induction, the appressorium formation rate of ΔMrGpa1 was only approximately 20% and were significantly reduced (by 76.5%) compared with WT strains (Figure 5C). Therefore, MrGpa1 plays an important role in cuticle penetration by impacting appressorium formation.

**MrGpa1** Regulates Intracellular cAMP Levels, but Further Feeding With cAMP Cannot Recover the Appressorium Formation Rate of ΔMrGpa1

To determine whether MrGpa1 regulates cAMP levels in *M. robertsii*, the intracellular cAMP levels were measured in the hyphal stage. We found that the intracellular cAMP levels of ΔMrGpa1 were significant decreased by 52.8% (p < 0.01), compared with WT strains (Figure 6A). The result indicated that MrGpa1 is a positive regulator in intracellular cAMP levels of *M. robertsii*. However, the appressorium formation rate of ΔMrGpa1 cannot recover with the addition of 5 mM exogenous cAMP (Figure 6B).

**DISCUSSION**

G-proteins are key components of various signal transduction pathways and play show important biological roles in the control of cell proliferation, behavior, and development in higher organisms (Ivey et al., 1996; Regenfelder et al., 1997; Harashima and Heitman, 2005). In the current study, we described the identification of MrGpa1 gene encoding the Gaᵢ subunit in *M. robertsii*, and showed that this Gaᵢ protein plays an important role in conidiation, stress resistance, and virulence in the host fungus.

In filamentous fungi, genes encoding three GPA subunits were reported in *N. crassa*, *F. oxysporum* f. sp. cubense, *Aspergillus nidulans*, and *M. grisea* (Ivey et al., 1996; Hicks et al., 1997; Liu and Dean, 1997; Guo et al., 2016b). However, while genes of four G protein α subunits were identified in *M. robertsii*, two of the open reading frames (ORFs) are highly similar (MrGpa2 and MrGpa4). Further, the
existence of Gαi has been demonstrated in filamentous fungi including the *M. robertsii* in this study, but not in the yeasts *S. cerevisiae* or *Schizosaccharomyces pombe* (Kubler et al., 1997; Kallal and Fishel, 2000).

Previously, to be activated by GPCR that sense external signals in plasma membrane (PM), G protein was considered to locate at the PM (Marrari et al., 2007). However, recent studies revealed G-protein localization beyond the PM, e.g., in the mitochondrion, endoplasmic reticulum, and Golgi apparatus (Michaelson et al., 2002; Nair et al., 2017). Wolf PSORT prediction in the current study indicated a possible location of MrGPA1 in the mitochondrion. Indeed, by using a fluorescent protein fusion, we showed here that MrGPA1 is located in the mitochondria, which is consistent with location of Gαi in HEK293T cells (Lysand and Bajjalieh, 2007). It has been reported that the roles of mitochondria in fungi is involved in the aging, conidiation, tolerance to adverse stresses and pathogenesis (Lorin et al., 2006; Kretschmer et al., 2012; Kitagaki and Takagi, 2014; Khan et al., 2015). For example, previous report in *Magnaporthe oryzae* shows that mitochondrial fusion protein MoFis1 mediates conidiation and is required for full virulence (Khan et al., 2015). Moreover, Kretschmer et al. (2012) found that defects in mitochondrial influence virulence from *Usilago maydis*. Therefore, the biological roles of MrGPA1 localized to mitochondria are further investigated in the present study.

According to multiple studies, GPA subunits play a role in the conidial yield in some fungi. For example, the conidiation of magB deletion mutant of *M. grisea* and Δfga1 mutant of *F. oxysporum* f. sp. *cubense* is impaired (Liu and Dean, 1997; Guo et al., 2016b). A similar phenomenon was also observed in the current study. Because a conserved conidiation regulatory pathway in *M. robertsii* contains Mr-BrlA, Mr-AbaA, and Mr-WetA (Zeng et al., 2017), these key genes were downregulated in the MrGpa1 deletion mutant too. All these indicated that the MrGpa1 gene is involved in the conidiation of *M. robertsii* by regulating the expression of conidiation-related genes.

We also observed that ΔMrGpa1 strain is more tolerant to UV irradiation and thermal stress than the WT and cpΔMrGpa1 strains. This observation was consistent with findings for *F. oxysporum* f. sp. *cubense* and *N. crassa* (Yang and Borkovich, 1999; Guo et al., 2016b). According to the two cited studies, intracellular cAMP levels are reduced in the Gαi deletion mutants, suggesting that the cAMP pathway might be involved in the response to thermal stress and tolerance of UV irradiation in some fungi.

In the *M. robertsii*, MrGpa1 deletion resulted in a marked attenuation of virulence in the *G. mellonella* model, with an increased LT50 value (by 63.9%) compared with that of the WT strain. Further analysis indicated that the reduced virulence is associated with an impaired appressorium differentiation in the mutant. This was similar to the effect of magB deletion in *M. grisea*, namely, blocked appressorium formation in a deletion strain (Liu and Dean, 1997). The virulence of plant pathogenic fungi is regulated by multiple pathways, such as the mitogen activated protein kinase (MAPK) cascades and the cAMP-PKA pathway (Dean, 1997). Further, the Gαi family contributes to pathogenicity in many plant pathogenic fungi. For example, the pathogenicity of the deletion strains *M. grisea* ΔmagB, *Botrytis cinerea* Δbcg1, and *F. oxysporum* f. sp. *cubense* Δfga1 is markedly reduced (Liu and Dean, 1997; Gronover et al., 2001; Guo et al., 2016b). Moreover, MAPK cascades and cAMP-PKA pathway are also involved in virulence of entomopathogenic fungus (Fang et al., 2009; Jin et al., 2012; Chen et al., 2016). In this study, the decrease of the intracellular cAMP levels in ΔMrGpa1 strains demonstrates that MrGpa1 involved in cAMP-PKA pathway. However, feeding with cAMP cannot recover the appressorium formation rate of ΔMrGpa1. In contrast to *magB* deletion in *M. grisea*, the appressorium differentiation is dependent on cAMP (Liu and Dean, 1997). We speculated that MrGpa1 located in mitochondria and *magB* are different G proteins, and MrGpa1 is involved in not only cAMP-PKA pathway, but also other regulatory pathways affecting on appressorium differentiation, such as MAKP cascades or RGS pathway.

**CONCLUSION**

MrGPA1 is located in the mitochondria of *M. robertsii* cell and is a member of Gαi family. MrGPA1 controls unique signal transduction pathways, and thus plays important role in conidiation, stress resistance, virulence, and intracellular cAMP levels in that fungus. These findings raise the possibility of designing powerful strategies for genetic improvement of *M. robertsii* conidiation capacity and virulence for killing pests.

**DATA AVAILABILITY STATEMENT**

All datasets generated for this study are included in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

BH and ZW conceived and designed the study. YT and HW wrote the manuscript, conducted the experiments, and analyzed the data. ZL did a part of the experiments. BH edited the manuscript and supervised the project. All authors read and approved the manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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