MoSec61β, the beta subunit of Sec61, is involved in fungal development and pathogenicity, plant immunity, and ER-phagy in Magnaporthe oryzae

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ABSTRACT
The process of protein translocation into the endoplasmic reticulum (ER) is the initial and decisive step in the biosynthesis of all secretory proteins and many soluble organelle proteins. In this process, the Sec61 complex is the protein-conducting channel for transport. In this study, we identified and characterized the β subunit of the Sec61 complex (MoSec61β). Compared with the wild-type strain Guy11, the ΔMosec61β mutant exhibited highly branched mycelial morphology, reduced conidiation, high sensitivity to cell wall integrity stress, severely reduced virulence to rice and barley, and restricted biotrophic invasion. The turgor pressure of ΔMosec61β was notably reduced, which affected the function of appressorium. Moreover, ΔMosec61β was also sensitive to oxidative stress and exhibited a reduced ability to overcome plant immunity. Further examination demonstrated that MoSec61β affected the normal secretion of the apoplastic effectors Bas4 and Slp1. In addition, ΔMosec61β upregulated the level of ER-phagy. In conclusion, our results demonstrate the importance of the roles played by MoSec61β in the fungal development and pathogenesis of M. oryzae.

Introduction

Magnaporthe oryzae is the most important pathogen infecting rice, wheat, and other grass species[1]. M. oryzae infection begins after the conidia are spread by wind or dew drops to the host surface. Under proper conditions, the conidium germinates and forms a polarized germ tube. Next, the tip of the germ tube differentiates into a dome-shaped infection structure called the appressorium [2]. Appressorium development is accompanied by energy recycling in which the nucleus, glycogen and lipid droplets in the conidia are degraded by autophagy or transported to the appressorium [3]. After the appressorium matures, it forms a penetration peg to rupture the cuticle of the host through glycerol-derived hydrostatic pressure. Subsequently, the filamentous primary invasive hyphae (IH) extend inside the cell and develop into the bulbous secondary invasive hyphae for transcellular infection. Colonization by invasive hyphae results in the appearance of necrotic lesions.

To respond to pathogen stimulation and invasion, plants have evolved an advanced and complex immune system that is sensitive to stimuli from the outside environment [4]. Recognition of pathogen-associated molecular patterns (PAMPs) by pattern recognition receptors (PRRs) leads to PAMP-triggered immunity (PTI) [5]. Furthermore, when PTI is no longer protective of plants, resistance (R) proteins recognize pathogen-derived effectors and initiate effector-triggered immunity (ETI) [6]. The two immune processes share common defense responses, including an outburst of reactive oxygen species (ROS), secretion of antimicrobial compounds, and reinforcement of plant cell walls [7]. In addition to the plant’s immune response, there is also evidence that pathogens interfere with plant immunity. Two bio-structures are formed where pathogens secrete effectors in response to plant immunity. Host-derived extravasive hyphal membranes (EIHM) are formed to enfold biotrophic IH [8], and host mem-

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brane-rich structures called biotrophic interfacial complexes (BIC) are formed focally at the periphery of invasive hyphae during biotrophic invasion [9]. Additionally, apoplastic and cytoplasmic effectors, two distinct effector secretion systems, have been identified in M. oryzae [10]. Apoplastic effectors are associated with host plants before cytoplasmic effectors and partly decide the outcome of the interaction between pathogens and plants [11]. The apoplastic effector Bas4 and Slp1 are dispersed in the extracellular compartment formed by EIHM and the IH membrane to indirectly act on host plant cells through signal transduction [7,12]. The apoplastic effectors are secreted via the conserved ER (endoplasmic reticulum)-to-Golgi secretion pathway. Conventional protein secretion is the transport route of secretory proteins from the endoplasmic reticulum (ER) to the Golgi apparatus (GA) and subsequently through secretory vesicles or secretory particles into the plasma membrane (PM) [13]. Conversely, the function of cytoplasmic effectors is to impair the normal physiological metabolism inside plant cells [14]. Pwl2 is a classical cytoplasmic effector that preferentially accumulates in BIC after being secreted from IH and subsequently enters the plant host cell [15]. The delivery of cytoplasmic effectors is mediated by the exocyt complex, which is essential for establishing epithelial polarity, morphogenesis, and homeostasis [10,16].

The Sec61 complex is a membrane channel on the ER that is involved in the translocation of newly synthesized precursor polypeptides into the ER lumen or on the ER membrane [17]. Sec61 is a highly conserved multisubunit protein complex that consists of three subunits, Sec61α, Sec61β, and Sec61γ, in eukaryotic cells. In budding yeast, the core proteins are Sec61, Sbh1, and Sss1. The α subunit forms the pore channel through which a polypeptide chain passes [18], whereas the γ subunit stabilizes the protein translocation [19]. Unlike Sec61α and Sec61γ, whose functions have been thoroughly investigated, the function of the β-subunit is still under study. At present, it is known that Sec61β (Sbh1 and Sbh2) is not necessary for the functional integrity of Sec61 in yeast, although it does promote the process of protein translocation [20]. In contrast to the results obtained in Saccharomyces cerevisiae, Sec61β has been determined to be essential for the embryogenesis of Drosophila melanogaster [21]. In addition, a study on barley resistance suggested that Sec61β is required for plant susceptibility to powdery mildew [22]. However, although studies have highlighted the importance of Sec61β, the mechanism by which this protein functions has not been elucidated. To the best of our knowledge, the functions of the Sec61 complex in plant pathogenic fungi have not been studied to date.

In this study, we identified the beta subunit of Sec61 in M. oryzae, designated MoSec61β. Deletion of MoSEC61β was associated with abnormal polarized hyphal growth, reduced conidiation and appressorium turgor pressure, defects in cell wall integrity, attenuated utilization of glycogen and lipid droplets, and weak virulence to rice and barley. Furthermore, penetration assays revealed that the ability to form penetration pegs was impaired in ΔMoSec61β. Although a small number of penetration pegs are formed successfully, the IH still fails to colonize adjacent cells because it no longer has the capacity to eliminate the host ROS. Meanwhile, high expression level of pathogenicity-related gene PR1a in leaves of ΔMoSec61β-challenged rice, and weak capacity in degrading callose of ΔMoSec61β suggested the essential role of MoSec61β in overcoming plant defense responses. Additionally, the distribution of the apoplastic effector Bas4 and Slp1 were disrupted in ΔMoSec61β, indicating the important role played by MoSec61β in ER-to-Golgi transport. Also, the deletion of MoSec61β leads to more intense ER-phagy. In conclusion, we demonstrate the biological function of the β subunit of the Sec61 complex in M. oryzae and provide new evidence for ER-mediated ER-phagy and apoplastic effector secretion in plant pathogenic fungi.

Results

MoSec61β is essential for conidiogenesis, polarized hyphal growth, cell wall integrity, and virulence of M. oryzae

Sequence alignment analysis showed that MGG_03644 shared 48.72% and 57.83% identity with S. cerevisiae Sbh1p and Sbh2p, respectively (Fig. S1). We termed MGG_03644 as MoSEC61β. MoSec61β is predicted to encode 144 amino acids (aa) and spans the membrane once (85–103 aa), with the N-terminus being in the cytosol. Then, the biological functions of MoSec61β were explored using target gene replacement. In addition, the full-length genomic copy of MoSEC61β was reintroduced into ΔMoSec61β for genetic complementation analysis, and we named the complementation strain Mosec61βc.

The growth rate of ΔMoSec61β is lower than that of the wild-type Guy11 and the complementation strain Mosec61βc (Figure 1(a,c)). Microscopic observation showed that the pear-shaped conidia of Guy11, ΔMoSec61β, or Mosec61βc were distributed in a concentric axis at the top of the fascicular conidiophore (Figure 1(b)). However, statistical analysis
Figure 1. MoSec61β is involved in hyphal growth, asexual reproduction, cell wall integrity, and pathogenicity. (a) Growth of Guy11, ΔMoSec61β, and MoSec61βc on CM plates. Strains were inoculated on CM plates for 8 days. (b) Conidiophores of wild-type, ΔMoSec61β, and MoSec61βc strains. Bar = 50 μm. (c) Mycelial diameters of the wild-type, ΔMoSec61β, and MoSec61βc strains. (d) Conidiation of Guy11, ΔMoSec61β, and MoSec61βc. (e) Morphology of vegetative hyphae of Guy11, ΔMoSec61β, and MoSec61βc on cover glass. Bar = 50 μm. (f) Vegetative hyphae of wild-type, ΔMoSec61β, and MoSec61βc stained with CFW. White arrows point to the septa. Bar = 20 μm. (g) Strains were incubated on MM plates supplemented with various stress inducers at 25°C for 7 days. Growth of strains in media supplemented with 0.0025% SDS, 100 μg/mL CFW, and 50 μg/mL Congo red (CR). (h) Pathogenicity on barley leaves. Twenty microliters conidial drops (5 × 10⁴ mL⁻¹) were inoculated on barley leaves. Photographs were taken after 4 dpi. (i) Pathogenicity of rice seedlings. Conidia (5 × 10⁴ mL⁻¹) were sprayed on 21-day-old rice seedlings. Photographs were taken after 7 dpi. (j) Disease score assays for Guy11, ΔMoSec61β, and MoSec61βc. The proportion of lesion areas in 5-cm leaves was measured by Photoshop CS6. Error bars represent the standard deviation. Significant differences between the mutant and wild-type strains, as estimated by Duncan’s test: **P < 0.01, *P < 0.05.

showed that the number of conidia produced by ΔMoSec61β was clearly decreased. Compared to the wild-type strain, the sporulation in ΔMoSec61β was decreased to half of that in Guy11 (Figure 1(d)). Additionally, compared with the wild-type Guy11 and the complementation strain MoSec61βc, the ΔMoSec61β mycelia produced more apical and subapical branches, and the wild-type Guy11 and MoSec61βc maintained the vertical growth of mycelium without sub-top branches (Figure 1(e)). To clearly observe the morphology of hyphae, we stained the mycelia of Guy11, ΔMoSec61β and MoSec61βc using calcofluor white (CFW). The mycelia of wild-type Guy11 and MoSec61βc were straight, and the cell intervals were largely equidistant (Figure 1(f)). In contrast, the interval of ΔMoSec61β mycelia had irregular branches, and the spacing interval of each cell was shorter than that of Guy11 and was not evenly spaced (Figure 1(f)). MoSec61β was determined to be involved in the morphogenesis and separation of vegetative mycelia.

To identify the role of MoSec61β in cell wall integrity, we monitored the effects of various cell wall perturbing agents on the ΔMoSec61β mutant. Mycelial growth was measured on MM plates supplemented with CFW, Congo red (CR) and SDS (sodium dodecyl sulfate), compounds known to cause cell wall stress. As
shown in Figure 1(g), the sensitivity of the ΔMosec61β mutant to CFW and Congo red was significantly different from that of both the wild-type Guy11 and the complementation strain Mosec61βc. The growth inhibition rates of ΔMosec61β to CFW and Congo red were significantly different from those of Guy11 and Mosec61βc (Fig. S2). These results indicated that the loss of MoSec61β affects cell wall integrity.

To investigate whether MoSec61β is related to the virulence of *M. oryzae*, two susceptible hosts, rice (CO-39) and barley (ZJ-8), were employed for the assessment of the pathogenicity of strains. After inoculation on barley leaves with conidia suspension (5 × 10⁵ conidia/ml) for 4 days, the wild-type Guy11 and the complementation strain Mosec61βc caused yellow and brown lesions with rotten plant tissues, while inoculation with ΔMosec61β resulted in tiny lesions (Figure 1(h)). Similarly, when inoculated onto barley leaves with mycelium plugs, ΔMosec61β also caused weaker disease lesions compared to the wild-type Guy11 and the complementation strain Mosec61βc (Fig. S3). When sprayed onto 21-day rice seedlings with conidia suspension (5 × 10⁴ conidia/ml), ΔMosec61β caused small necrotic flecks, while the wild-type Guy11 and the complementation strain Mosec61βc caused typical spindle-like, gray centered blast lesions and many merged lesions (Figure 1(i)). The disease lesion areas in 5-cm-long infected leaves caused by ΔMosec61β (10.23 ± 1.23%) were significantly smaller than those caused by the wild-type Guy11 (51.30 ± 4.87%) and the complementation strain Mosec61βc (49.75 ± 1.96%) 7 days post-inoculation (dpi) (Figure 1(j)). Thus, MoSec61β was observed to play an important role in pathogenicity.

**MoSec61β is involved in the development of invasive hyphae in *M. oryzae***

To determine the reasons for the reduced virulence of ΔMosec61β, we first analyzed conidium germination and appressorium formation. However, there were no significant differences in conidia germination, appressorium formation, or appressorium morphology among Guy11, ΔMosec61β, and Mosec61βc (Table S1, Fig. S4). Then, we performed penetration assays on barley leaves. Few appressoria of ΔMosec61β could form penetration pegs (Figure 2(a,b)). Additionally, the appressorial penetration rate of ΔMosec61β on barley leaves was significantly decreased compared to that of the wild-type and complementation strains (Figure 2(a)). At 48 hpi (hours post-inoculation), the appressorial penetration rate of ΔMosec61β was approximately half of that in the wild-type and complementation strains Mosec61βc (Figure 2(c)). At 72 hpi, more than 90% of Guy11 appressoria penetrated barley cells. However, only 30% of the appressoria of ΔMosec61β penetrated barley cells (Figure 2(c)). Further observation showed that ΔMosec61β displayed defects in IH extension. At 48 hpi, nearly 50% of IH in the wild-type and Mosec61βc strains showed transcellular infection, and less than 10% IH of ΔMosec61β invaded neighboring cells (Figure 2(d)). At 72 hpi, more than 80% of IH in the wild-type and Mosec61βc strains colonized other cells, but only 10% of IH in ΔMosec61β expanded into adjacent cells (Figure 2(d)). These data demonstrated that the infection and IH extension of ΔMosec61β are notably impaired.

**MoSec61β affects turgor pressure in appressoria and mobilization of glycerol and lipid droplets from conidia to appressoria***

The process of penetration mediated by the appressorium requires a large amount of inner turgor pressure, which can enable the appressorium to produce adequate mechanical strength and facilitate rupture of the host cuticle with narrow-penetration hyphae [23]. As described previously, most of the appressoria of ΔMosec61β cannot form penetration pegs, and we investigated if the appressorium turgor pressure of ΔMosec61β was impaired. Specifically, we performed an incipient cytorrhysis assay to measure the turgor pressure exerted by mature appressoria. In this assay, a 0.5–1.5 M concentration of glycerol solution was applied to examine the collapse rate of appressoria. Under treatment with 1 M glycerol, approximately 30% of 24 h appressoria of the wild-type strain and the complementation strain Mosec61βc collapsed. However, 60% of ΔMosec61β collapsed with this treatment (Figure 3(a,b)). With the increase of glycerol concentration to 1.5 M, the collapse rate of ΔMosec61β appressoria increased to nearly 90% and was higher than that of the wild-type and Mosec61βc strains (Figure 3(b)). It was concluded that the turgor pressure of the ΔMosec61β appressoria was decreased compared with that of the wild-type strain.

The glycerol in appressoria is primarily produced by the transfer and utilization of glycerol and lipids in conidia. To investigate the glycerogen distribution and lipid turnover of ΔMosec61β, we employed potassium iodide to stain glycerogen and employed Nile red to stain lipids during appressorium development. From 0 to 4 h in the preliminary stage of appressorium development on a hydrophobic surface, abundant lipids were seen in conidia and appressoria of Guy11 and ΔMosec61β (Figure 3(c)). At 8 hpi, more than 60% lipids in conidia of the wild-type Guy11 and
ΔMosec61β strains were transported to appressoria. At 24 hpi, a significantly higher proportion of ΔMosec61β conidia contained lipids (Figure 3(d)). Compared to Guy11, more than 60% of ΔMosec61β appressoria contained lipids at 16 and 24 hpi (Figure 3(e)). Similarly, the distribution of glycogen exhibited the same pattern as lipids in ΔMosec61β, as shown in Figure 3(f–h). The above results indicate that MoSEC61β is required for lipid droplets and glycogen mobilization from conidia to appressoria.

Activity to scavenge host ROS is reduced in ΔMosec61β

When plants are attacked by pathogens, the plant cells generate massive reactive oxygen species (ROS) to inhibit or directly kill invading pathogens [24]. We hypothesize that the reason ΔMosec61β IH failed to colonize adjacent cells was related to its abnormal ability to scavenge ROS. To confirm this possibility, we determined the ability of ΔMosec61β to resist redox stress. The mycelial plugs of Guy11, ΔMosec61β, and Mosec61βc were inoculated onto CM supplemented with the oxidative stress agent H2O2 (5 mM) (Figure 4(a)). The growth inhibition rate of ΔMosec61β was significantly higher than that of Guy11 (Figure 4(b)). This result indicates that the ΔMosec61β mutant is more sensitive to oxidative stress than the wild-type strain.

Then, 3,3′-diaminobenzidine (DAB) staining was used to capture hydrogen peroxide (one type of ROS) produced in the infected barley leaves. At 38 hpi, Guy11 was able to invade the surface cells and was observed to expand to the surrounding cells, and no obvious ROS were dyed. In contrast, ΔMosec61β IH was confined in the first cell and surrounded by obvious reddish-brown dyed ROS (Figure 4(c)). Similar results were obtained when the strains were incubated on a rice sheath (Figure 4(d)). There was an abundant accumulation of dark brown granules among IH of ΔMosec61β (67.88 ± 1.50%) in rice cells. However, few brown granules (19.78 ± 0.19%) were observed in the wild-type Guy11 and the complementation strain (Figure 4(e)). The dark brown granules indicated the immune response of rice, further demonstrating that the capacity of ΔMosec61β to scavenge host ROS was reduced.

Furthermore, an inhibitor of NADPH oxidase diphenyleneiodonium (DPI) was applied to the rice sheath to determine whether ROS accumulation is
Figure 3. (a) Collapse of appressoria at 0.5 M glycerol. Bar = 20 μm. (b) A 0.5–1.5 molar concentration of glycerol solution was applied to examine the collapse rate of appressoria in Guy11, ΔMosec61β, and Mosec61βc. Arrows indicate the collapsed appressoria. (c) Cellular distribution of lipid droplets during appressorium development. Samples were stained with Nile red and observed with UV epifluorescence. The lipid droplets show a red signal fluorescence. Bar = 20 μm. (d) The percentage of conidia containing lipid droplets during appressorial development. (e) The percentage of appressoria containing lipid droplets. (f) Cellular distribution of glycogen during appressorium development. Bar = 20 μm. Samples were stained with KI/1 solution at the indicated time phase. The glycogen appears as dark brown deposits. (g) The percentage of conidia containing glycogen. (h) The percentage of appressoria containing glycogen during appressorium development. Error bars represent the standard deviation. Significant differences compared with the wild-type strain were estimated by Duncan’s test: **P < 0.01, *P < 0.05.
the primary factor causing IH defects. Under normal conditions (without DPI), less than 10% of the appressoria of ΔMos61β penetrated into adjacent cells at 48 h, but more than 50% of the wild-type appressoria successfully developed IH in the neighboring cells. Under the condition of 0.2 μM DPI treatment, the extension of ΔMos61β IH to adjacent cells increased threefold at 48 h. However, there were no obvious changes in the wild-type strain (Figure 4(f)). These data confirmed that the attenuated biotrophic growth of ΔMos61β was due to host ROS accumulation.

**Figure 4.** ΔMos61β is sensitive to oxidative stress and cannot scavenge ROS. (a) *M. oryzae* strains grown on 5.0 mM H₂O₂. (b) Relative growth of mycelial colonies on 5.0 mM H₂O₂. (c) Reactive oxygen species (ROS) capture by 3,30-diaminobenzidine (DAB) staining in infected barley leaves. Bar = 20 μm. (d) ROS capture by DAB staining in infected rice leaf sheaths at 48 hpi. White thick arrows point to the invasive hyphae, white thin arrows point to orange brown granules. Bar = 20 μm. (e) The percentage of *M. oryzae* strains that can induce brown granules. (f) Percentage of appressorium-mediated penetration and infectious hyphae development of Guy11 and ΔMos61β in DPI-treated rice sheaths. Error bars represent the standard deviation. Significant differences compared with the wild-type strain were estimated by Duncan’s test: **P < 0.01, *P < 0.05.
**ΔMosec61β is defective in overcoming plant defense responses**

To further confirm that the defects of ΔMosec61β in infectious growth were caused by the failure to resist the immunity of plants, we used pathogens to infect heat-killed plant cells (Figure 5(a)). The penetration assay was carried out with rice leaf sheaths incubated at 75°C for 25 min before inoculation. After 48 hpi, almost all of the appressoria formed by the wild type (98.24 ± 1.04%) and ΔMosec61β (96.73 ± 0.97%) penetrated the epidermal cells of rice leaf sheaths (Figure 5(c)). In addition, it is obvious that the ΔMosec61β IH are no longer confined in the first cell, and the percentage of IH extending to the neighboring cells is greatly

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**Figure 5.** Penetration assays with heat-killed rice leaf sheaths and onion epidermis. (a) Conidia solution was inoculated into normal or heat-treated leaf sheaths, and infection was observed after 48 hpi. Bar = 20 μm. Black arrows point to IH. White arrows point to IH, which was expanded in adjacent cells. (b) Conidia solution was inoculated into normal or heat-treated onion epidermal cells, and infection was observed after 48 hpi. Bar = 20 μm. Black arrows point to IH. White arrows point to IH, which was expanded in adjacent cells. (c) Statistical analysis of the infection rate of the appressoria of the *M. oryzae* strains in leaf sheaths at 48 hpi. (d) The percentage of invasive hyphae extended to adjacent cells in normal or heat-treated leaf sheaths at 48 hpi. Error bars represent the standard deviation. Significant differences compared with the wild-type strain were estimated by Duncan’s test: **P < 0.01, *P < 0.05.**
increased. There was no significant difference compared with the wild-type Guy11 (90.14 ± 2.30%) and the ΔMosec61β mutant (89.63 ± 2.10%) (Figure 5(d)). We reperformed the penetration assays with heat-killed onion epidermal cells, and similar infectious patterns were observed in the wild-type and ΔMosec61β strains (Figure 5(b)).

Pathogenesis-related (PR) proteins have been used as markers of plant defense responses. In order to further investigate whether the plant defense genes were stimulated by infection with ΔMosec61β, the expression patterns of PBZ1 and PR1a genes were analyzed through quantitative RT-PCR. PBZ1, a probenazole-inducible gene from rice, triggers non-race specific resistance in rice plants against rice blast fungus [25]. PR1a is one of the PR proteins, which accumulates after blast fungus infection in rice [26]. The expression level of PBZ1 was up-regulated in leaves of blast-fungus-challenged rice at 24 hpi, 32 hpi, and 48 hpi, whereas there were no obvious differences in ΔMosec61β or Guy11-challenged rice leaves (Fig. S5a). In contrast, the expression level of PR1a in leaves of ΔMosec61β-challenged rice was higher than that in leaves of Guy11-challenged rice at 24 hpi and 48 hpi (Fig. S5B). The expression level of PR1a in leaves of ΔMosec61β-challenged rice at 24 hpi was 4 folds as high as that in leaves of Guy11-challenged rice (Fig. S5b). The induction of plant defense responses in ΔMosec61β-challenged rice may contribute to the retardation of IH development.

Callose, a β-1,3-glucanase induced by plant defense responses, provides chemical defenses at the cellular sites of attack [27]. To investigate the induction of callose by infection of the ΔMosec61β mutant and the wild-type Guy11 respectively, we observed the accumulation of callose deposits at the cell wall crossing sites around IH using aniline blue staining. As shown in Figure 6(a), the number of the callose deposits around

![Figure 6](image-url)

**Figure 6.** Callose deposition in barley leaves. (a) Deposited callose in leave cells under different treatment. Arrows indicate the callose depositions. app: appressorium. Bar = 10 μm. (b) The number of callose deposition per 50 μm length of side wall at 32 h. Error bars represent the standard deviation. Significant differences compared with the wild-type strain were estimated by Duncan’s test: **P < 0.01, *P < 0.05. (c) The expression level of callose synthase-encoding genes (OsGSL1, OsGSL3, and OsGSL5) in Guy11 and ΔMosec61β-challenged rice leaves.
the ΔMosec61β IH was significantly more than that around the Guy11 IH at 32 dpi. Treatment of DPI decreased the number of the callose deposits around the ΔMosec61β IH (Figure 6(a,b)). Meanwhile, the expression levels of 3 callose synthase-encoding genes (OsGSL1, OsGSL3, OsGSL5) were investigated using quantitative RT-PCR. The expression level of OsGSL1 reached the peak at 32 hpi in the leaves of ΔMosec61β-challenged rice, and it was 20 folds as high as that in the leaves of wild-type Guy11-challenged rice. In addition, the expression of OsGSL3 and OsGSL5 genes was also slightly induced in leaves of the ΔMosec61β-challenged rice, with higher induction levels as compared to those in leaves of the Guy11-challenged rice at 32 hpi (Figure 6(c)). These data indicated that ΔMosec61β was defective in overcoming plant defense responses because of its weak capacity in degrading callose. The above results suggested that MoSec61β is necessary for overcoming plant defense responses.

**ΔMosec61β partially disrupts the localization of the apoplastic effectors Bas4 and Slp1**

To assess whether the MoSec61β protein is involved in the secretion of effectors, we performed live cell imaging of biotrophic invasion by transformants expressing the cytoplasmic effector fusion protein Pwl2-GFP and the apoplastic effector fusion protein Bas4-mCherry, Slp-GFP. During the early infection stage at 48 hpi, Pwl2-GFP localized to a single punctate BIC in the wild-type Guy11 and the ΔMosec61β mutant (Figure 7(a)), indicating that secretion of cytoplasmic effectors was not impaired in ΔMosec61β. In wild-type Guy11, the Bas4-mCherry fusion protein outlines the IH and occupies an inner layer of the BIC. In contrast, the localization of Bas4-mCherry for the ΔMosec61β mutant was partially disrupted, while localization of the Pwl2-GFP fusion appeared to be normal. Localization patterns of Bas4-mCherry in the

![Figure 7](image-url)
ΔMosec61β mutant could be further divided into two types, as shown in Figure 7(b). With type 1, the fluorescence of Bas4-mCherry (32%) was uniformly wrapped the IH and was located in EIHM, which is the same as with the wild-type strain. In type 2, the localization of Bas4-mCherry could not be correctly accumulated in EIHM (68%). In contrast, nearly all randomly imaged infection sites of the wild-type strain showed the type 1 pattern (Figure 7(c), Bas4-mCherry). Similar results were observed in apoplastic effector Slp1 (Figure 7(c), Slp1-GFP). About 98% of Slp1-GFP could trace out IH and locate in EIHM in the wild-type Guy11. In contrast, two types of Slp1-GFP localization patterns were also observed in the ΔMosec61β mutant, type 1 (∼ 40% of Slp1-GFP could be correctly accumulated in EIHM) and type 2 (∼ 60% could not be correctly accumulated in EIHM) (Figure 7, Slp1-GFP). These results confirmed that the localization patterns of Bas4-mCherry and Slp1-GFP were seriously impaired in ΔMosec61β. As previously described, MoSec61β is an ER transmembrane protein, and its deletion may cause abnormal secretion of effector proteins. These results indicate that proper localization of apoplastic effectors in EIHM depends in part on MoSec61β.

**MoSec61β negatively regulates ER-phagy**

In our study, ΔMosec61β exhibited defects similar to those of the atg-deficient mutants, such as reduced turgor pressure, retarded utilization of lipids and glyco- gen, and reduced virulence. However, neither conidia nuclei degradation nor GFP-MoAtg8 degradation under nutritional deprivation conditions showed the dysregulated autophagy process (Fig. S6, S7). Autophagic cell death and macroautophagy were not impaired in ΔMosec61β. Interestingly, we unexpectedly observed that the growth of ΔMosec61β is sensitive to ER stress under the ER stress factors tunicamycin (TUNI) or dithiothreitol (DTT) (Figure 8(a)). Under treatment with 0.5 μg/mL TUNI, the growth inhibition rate of the wild-type strain was significantly lower than that of ΔMosec61β. Similarly, under 5.0 mM DTT treatment, the growth inhibition rate showed the same differences between the wild-type and ΔMosec61β strains (Figure 8(b)). The growth of the ΔMosec61β strain was inhibited by ER stress factors.

In response to ER stress, ER-mediated autophagy (ER-phagy) occurs to maintain the normal function of the ER. Therefore, we hypothesized that ER-phagy of ΔMosec61β is impaired under ER stress. MoSec63-GFP is an integral ER membrane protein. It has been reported that 5 μM DTT can induce MoSec63-GFP cleavage, consistent with the activation of selective autophagy by ER stress. We constructed the MoSec63-GFP fluorescent vector and transformed it into the wild-type Guy11 strain and the ΔMosec61β mutant strain. Under the induction of DTT, the expression of GFP differed significantly between Guy11 and ΔMosec61β (Figure 8(c)). At 0 h, there was little free GFP in Guy11 and more free GFP in ΔMosec61β. At 4 h, the amount of GFP-containing fragments was substantially enhanced in ΔMosec61β compared with

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**Figure 8.** Responses of *M. oryzae* strains to ER stress. (a) Mycelial colonies of Guy11, ΔMosec61β, and Mosec61βc cultured on MM media containing 0.5 μg/mL TUNI and 5.0 mM DTT at 25°C with a 16 h light and 8 h dark cycle for 8 days. (b) Growth inhibition rate of mycelial colonies on 0.5 μg/mL TUNI and 5.0 mM DTT. (c) ER-phagy of Sec63-GFP in Guy11 and ΔMosec61β. Total proteins were extracted from the Sec63-GFP expressed strains exposed to nitrogen starvation conditions with 5 μM DTT for 0 and 4 h. Full-length Sec63-GFP and free GFP were detected using GFP antibodies as described in the Materials and Methods. The extent of ER-phagy was estimated by calculating the amount of free GFP compared with the total amount of intact Sec63-GFP and free GFP. Quantitative analysis of the individual bands was performed using ImageJ software. Error bars represent the standard deviation. Significant differences between the mutant and wild-type strains, as estimated by Duncan's test: **P < 0.01, *P < 0.05.
Guy11 (Figure 8(c)). These results indicate that the absence of MoSec61β stimulates ER-mediated autophagy.

**Discussion**

In eukaryotic cells, protein translocation can occur co- or post-translationally, depending on the hydrophobicity of the precursor protein. These mechanisms require the heterotrimeric Sec61 complex for transmembrane transport [28]. In this study, we reported the biofunctions of Sec61β in *M. oryzae*. By homology alignment, we found only one gene encoding the Sec61β protein in rice blast fungus, MGG_03644, and we named that gene *MoSEC61β*. Loss of *MoSEC61β* had no apparent influence on conidial germination or appressorium formation but showed significant pathogenesis defects. During infection, ΔMoSec61β showed restricted IH and caused a host ROS burst that induced rice innate immunity because of impaired effector secretion.

Although Sec61β has not been studied in other plant pathogenic fungi, the maintenance function of ER under stress has been widely researched. As an essential organelle in cells, the ER keeps continuous renovation to maintain its function and integrity when faced with changes in the external and internal environment [29]. The ER performs quality control through the unfolded protein response (UPR), which leads to the upregulation of chaperone, decrease of protein synthesis and reverse transport of misfolded protein to the cytosol for degradation [30]. In *M. oryzae*, a double knockout strain for ERAD (ER-associated degradation, ERAD) genes, ΔMohrdΔModer1, lost its pathogenicity. Compared with the wild-type strain, ΔMohrdΔModer1 shows an unfolded protein response under normal conditions, and the secretion of pathogenic effector proteins is affected [31]. Another study found that ERQC (ER glycoprotein folding quality control) components, such as Sgl1, Gls2, and Gtb1, were N-glycosylated and were involved in the process of mycelial growth, conidiation, and invasive hyphal growth in host cells [32]. Similarly, the ER chaperone protein Lhs1 is also involved in rice infection in the blast fungus. The UPR target genes, including *SIL1*, *KAR2*, *PDI1*, and *SCF1*, are upregulated, and the function and secretion of AVR-Pita in Δlhs1 are impaired [33]. All these results indicate the importance of ER homeostasis for protein secretion and pathogenicity of rice blast.

During plant-pathogen interactions, pathogens produce effectors to resist and escape the immune response of plants. Our results found that loss of *MoSEC61β* impacts the localization of the apoplastic effectors Bas4 and Srp1 and thus faces a strong plant immune response, including defects in ROS scavenging and callose degradation, and high expression of *PR1a*. Accordingly, in heat-killed plant cells, the growth of the invasive hypha of ΔMoSec61β was not affected. This result is in keeping with the fact that the secretion of apoplastic effectors is transported via the ER-to-Golgi pathway. We thus conclude that unlike in yeast, where Sec61β is not necessary, in *M. oryzae*, the β subunit is necessary for the transposon function of the Sec61 complex. A previous study also suggested other roles played by Sec61β in addition to its function as a component of the translocon. In yeast, Sec61β was found to interact with Sec4p and the exocyst complex component, including Sec8p and Sec15p [34]. It is well-known that the transport of cytoplasmic effectors is mediated by the exocyst complex. Loss of the exocyst components Exo70 and Sec5 causes defects in proper secretion of the cytoplasmic effector Pw12 [10]. We therefore investigated whether the loss of MoSec61β in *M. oryzae* would also influence the ability of exocysts and ultimately affect the function of cytoplasmic effectors. We observed the localization of Pw12 during ΔMoSec61β invasion and found similar localization of Pw12 between the ΔMoSec61β and wild-type strains. These results showed that the β subunit is necessary for the transposon function of the Sec61 complex and that the absence of Sec61β did not affect the normal function of exocysts in the rice blast fungus.

In addition to the defects observed during the infection process, we also found that the mutant exhibited deficiencies in its ability to infect the host. Although ΔMoSec61β forms appressoria with normal morphology, the turgor pressure of the appressoria is generally low and cannot successfully penetrate the host surface. The maturation of the appressorium requires the mediation of core autophagy. Autophagy helps the appressorium to produce glycerol and turgor pressure by transporting and degrading inclusions inside the conidia (mainly glycerol and lipid droplets) [35]. In our experiments, we also found that the normal degradation of glycerol and lipid droplets in the mutant was inhibited, suggesting dysregulated autophagy. However, both the nuclear degradation and GFP-MoAtg8 degradation tests indicated that autophagy can occur normally in ΔMoSec61β.

As the channel connecting the ER to the cytoplasm, Sec61 also plays a role in ERAD, mediating the secretion of misfolding proteins in the ER back into the cytoplasm [36]. It is possible that the dysfunction of the Sec61 complex inhibits the ERAD process, thereby causing the accumulation of misfolding proteins. In fact, except for ERAD, which degrades proteins...
through the ubiquitin-proteasome system, ER also employs the specified autophagy ER-phagy to maintain its inner balance [37]. Therefore, we utilized the ER membrane protein Sec63-GFP to monitor the ER-phagy level inside ΔMosec61β. Our results showed that under normal conditions, Sec63 is expressed at a higher level in ΔMosec61β. After 4 h of treatment with DTT to induce ER stress, the degradation of Sec63-GFP was also more significant, indicating a higher level of ER-phagy in ΔMosec61β than in the wild-type Guy11 strain. We hypothesized that ΔMosec61β requires a stronger level of ER-mediated autophagy to compensate for the restriction of ERAD.

In summary, our results reveal that MoSec61β is necessary for the vegetative growth, asexual development, appressorium penetration, plant immunity evasion and pathogenesis of rice blast fungus. MoSec61β plays pivotal roles in turgor pressure by influencing the mobilization and degradation of glycogen and lipids. Deletion of MoSec61β and M. oryzae is deficient in overcoming plant defense responses. MoSec61β controls pathogenicity via the secretion of the apoplastic effector Bas4 and Slp1. In addition, MoSec61β is involved in ER-phagy in response to ER stress.

Materials and methods

Strains and culture conditions

The Magnaporthe oryzae strain Guy11 and derivative transformants were cultured on complete medium (CM) in a growth chamber at 25 or 28°C with a 16 h light and 8 h dark cycle [38]. Target gene replacement method was used to generate null mutants (Fig. S8, Table S2). For the oxidative stress test, the strains were cultured on CM media with 5.0 mM H2O2 and cultured in dark at 28°C. For cell wall integrity tests, the strains were cultured on minimal medium (MM) medium supplemented with 0.0025% SDS, 100 μg/mL CFW and 50 μg/mL CR in dark at 28°C. For ER stress tests, the strains were cultured on MM with 0.5 μg/ml tunicamycin (TUNI) and 5.0 mM dithiothreitol (DTT) and cultured under a 16 h light and 8 h dark cycle at 25°C.

Phenotypic characterization

For fungal growth and conidiation assays, a 6 x 6 mm mycelium plug of M. oryzae strains was inoculated on complete medium (CM) under the conditions of 25°C, 16 h light/8 h dark for 8 days. The diameter was measured after 8 days. Conidiation was detected by the strains grown for 8 days. The spores were washed with 3 mL water, filtered and diluted to 2 mL. The number of conidia was determined by a counter. To measure conidial germination and appressorium formation, 20 μL of spore suspension (5 x 10^4 conidia/mL) was dropped onto plastic coverslips and incubated at 22°C. Conidial germination and appressorium formation were observed for 4–24 h hpi. To observe conidiophore development of M. oryzae strains, vegetative hyphae inside the medium were sliced into thin pieces, and the pieces were cultured under 16 h light/8 h dark for 24 h at 25°C [39].

Pathogenicity and plant infection assays

Two-week-old seedlings of rice (Oryza sativa cv CO-39) were used in plant infection assays. Conidia harvested from 8-day-old growth on CM plates were resuspended in 0.2% (w/v) gelatin solution for a concentration of 1 x 10^5 conidia/mL. The suspension was sprayed evenly onto rice leaves using an artist’s airbrush (Badger Co., Franklin Park, Illinois). The inoculated plants were placed in a dew chamber at 25°C for 48 h in the dark and then transferred to a growth chamber with a cycle of 16 h of light and 8 h of dark using fluorescent lights. The plants were examined for lesions 7 days after inoculation. Disease severity was rated with the scale developed.

Infection assays were carried out three times for host penetration assays, and leaf segments were excised from 8-day-old seedlings of barley (Hordeum vulgare cv ZJ-8). Drops (20 μL) of conidial suspension (5 x 10^4 conidia/mL) were deposited onto the upper surface of the excised leaves in a dew chamber at 25°C. The leaves were examined for disease lesions at 24, 48, 72, and 96 h after inoculation after decoloration by methanol and preservation in lactophenol, as described previously [40].

Polarity growth and CFW staining

Sterilized coverslips were inserted obliquely on the CM solid medium, a suitable amount of mycelium was picked with a toothpick and inoculated onto the CM plates with coverslips, and strains were cultured in a constant temperature incubator at 25°C under 16 h light and 8 h dark. When the aerial hyphae of the strain grew to the middle of the cover glass, the cover glass was gently extracted from the CM plate and placed under a microscope to observe the polarity growth of the strain. CFW solution was added dropwise to the coverslip with aerial hyphae, and after staining for 5 min, it was placed under a fluorescence microscope for observation and photograph recording. The experiment was repeated three times.
Incipient cytorrhysis assays

Incipient cytorrhysis assays were used to determine the appressorium turgor [41]. In a humid environment, drops (20 μL) of conidial suspension (1 × 10^5 conidia/mL) were incubated on plastic coverslips for 24 h. After that step, the water was carefully removed and replaced with an equal volume of a glycerol solution with a concentration ranging from 0.5 M to 1.5 M. After 1 min of incubation in glycerol solution, the number of appressoria that had collapsed was recorded. The experiments were repeated three times, and more than 200 appressoria were observed each time.

Rice leaf sheath assays

The fresh spore suspension (1 × 10^5 conidia/mL) was injected into the leaf sheath by a disposable syringe (without bubbles). The leaf-sheath tube filled with spore suspension was cultured in an incubator at 25°C with a light cycle of 16 h and darkness of 8 h. Then, the transparent thin layer of the leaf sheath was carefully cut off with a blade, and the surface of the leaf sheath in contact with the spores was turned upward to make a slide of the leaf sheath for fluorescence microscope observation. The experiment was repeated three times.

Lipid droplets and glycogen staining

Tricyclazole (1 μL of 10 μg/mL to inhibit the formation of melanin of the attached cells and reduce the influence on the experimental observation) was added to 1 mL (1 × 10^5 conidia/mL) spore suspension. Drops (20 μL) of the spore suspension were inoculated on a hydrophobic membrane at 25°C in the dark and moisturized for 0 h, 8 h, 16 h, and 24 h, respectively. The water was carefully removed and replaced with an equal volume of a Nile red staining solution. After 30 min of incubation in Nile red staining solution, the transport and degradation of red lipid droplets were observed under a fluorescence microscope. Similarly, glycogen was stained in I2/KI solution for 1 min. The experiment was repeated 3 times, and 200 conidia were counted for each treatment.

Callose deposition staining

For callose deposition staining, leaves of barley (at 32hpi) were fixated and destained in 1:3 acetic acid/ethanol, the saturated destaining solution was replaced until the material was transparent (usually overnight). Fixated and destained leaves or seedlings were washed in 150 mM K2HPO4 for 30 min. Leaves of barley were incubated for at least 2 h in 150 mM K2HPO4 and 0.01% aniline blue (staining solution) in a 2 mL tube wrapped in aluminum foil for light protection.

RNA preparation and RT-PCR analysis

Fresh plant leaves (0.4 g) which inoculated with spore suspension were ground to powder with liquid nitrogen, and RNA was extracted using RNA Iso Plus (TaKaRa, Japan) according to manufacturer instructions. Reverse transcription of total RNA was carried out using PrimeScript™ RT regent Kit with gDNA Eraser (Takara, Japan). The qRT-PCR was then performed using TB Green® Premix EX Taq™ (Tli RNaseH Plus) (TaKaRa, Japan) to analyze the expression level of plant defense-related genes (PR1a, PBZ1) and callose synthase-encoding genes (OsGSL1, OsGSL3, OsGSL5).

Western blot analysis

For Sec63-GFP and cleaved GFP assays, the Sec63-GFP vector was transformed into the Guy11 and mutant strains using ATMT. The Sec63-GFP-expressing strains were grown in CM medium for 7 days and then shifted to nitrogen starvation (SD-N) medium with 5 μM DDT for 4 h to induce ER-phagy. Total proteins were separated on SDS-PAGE gels and transferred to polyvinylidene difluoride membranes for Western blot analysis. Sec63-GFP was detected with a primary anti-GFP antibody (GFP-Tag Rabbit mAb, Huabio, Hangzhou, China) and a secondary antibody (goat anti-rabbit IgG HRP, Beyotime, Shanghai, China). GAPDH antibodies (Huabio, Hangzhou, China) were used to confirm equal protein loading of each strain.

Disclosure statement

No potential conflict of interest was reported by the authors.

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