Development of a CRISPR/Cas9 gene editing system for *Fusarium oxysporum* and characterization of an extracellular superoxide dismutase and its contribution to pathogenicity on cotton

by

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Abstract

The *Fusarium oxysporum* species complex (FOSC) is an important group of filamentous fungi that are able to infect both animals and plants. Based on its economic importance, an effort has been made to better understand the genetics and pathogenicity of members of the FOSC; however, efficient reverse genetic techniques including gene disruption/deletion methods have been limited. One aim of this research was to establish an efficient, stable CRISPR/Cas9 mediated genome editing system for investigating fungal genetics and pathogenicity. In Chapter 2, a transformation method based on Cas9 ribonucleoproteins was developed and used to obtain *URA3* and *URA5* mutants, proving that the Cas9 RNPs could be transferred into fungal protoplasts and function. In addition, the *FoBIK1* gene from a secondary metabolite biosynthetic cluster was mutated using this system and the maximum efficiency of this gene disruption was ~50%. In Chapter 3, the CRISPR/Cas9 system was used to carry out endogenous gene tagging (EGT) for the study of protein subcellular localization using two different integration strategies, homology-independent targeted integration (HITI) and homology-dependent recombination integration (HDRI). The HITI strategy was able to facilitate integration of a large DNA fragment, ~8 kb in length, into the genome of *F. oxysporum* at the sgRNA cleavage site, and was used to insert a C-terminal 3×sGFP tag to the *FoCHS5* gene and a N-terminal *mCherry* tag to the *FoSSO2* gene. The HDRI strategy was used to tag the paralogous gene, *FoSSO1*, with a C-terminal mCherry marker. In Chapter 4, an extracellular SOD protein (*FoSOD5*) from *F. oxysporum* was characterized. Expression of
FoSOD5 was dependent on external environmental stimuli and the protein was localized to various cellular structures in different environments, and a lacZ expression reporter strain indicated FoSOD5 was induced in alkaline conditions. Importantly, a ΔFoSOD5 strain was more sensitive to reactive oxygen species and has attenuated virulence on cotton. Above all, a stable and efficient CRISPR/Cas9 system has been developed for F. oxysporum genome editing and was used for investigating gene regulation and protein subcellular localization.
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Chapter 1.

Introduction and review of the literature

1.1 *Fusarium* spp.

*Fusarium* is a diverse genus with at least 20 species complexes of filamentous fungi that are widely spread in our environment, especially in the soil (Geiser et al., 2013). Although most species are saprobes and rarely cause infections, several *Fusarium* species are important pathogens of humans, animals, and plants. One of the best studied *Fusarium* spp. is *F. graminearum* the predominant agent causing Fusarium head blight (FHB), a devastating disease of wheat and barley. The fungus rapidly causes premature bleaching symptoms on wheat spikelets after infection and produces the well-known mycotoxin deoxynivalenol which can lead to strong emetic effects after consumption (Goswami and Kistler, 2004).

The *F. oxysporum* species complex contains more than 100 formae speciales with a wide range of hosts including plants and humans (Michielse and Rep, 2009). Unlike *F. graminearum*, *F. oxysporum* has no known sexual life cycle. Several well-known formae speciales are responsible for vascular wilt diseases (Table 1.1); for example, *F. oxysporum* f. sp. *cubense* causes Fusarium wilt of banana (Panama disease) and is widely distributed in Africa, Asia, and South America while *F. oxysporum* f. sp. *lycopersici* causes wilt disease on tomato (Ploetz, 2006). The symptoms of Fusarium wilt disease includes browning of the xylem tissue, leaf
yellowing, chlorosis, necrosis, stunting, and wilting. Besides infecting plants, some species can infect humans and cause invasive fusariosis, in particular infecting immunocompromised patients (Nucci and Anaissie, 2007). Although some forma speciales are considered agricultural or human pathogens, several are used as biological agents to control other plant diseases. Nonpathogenic *F. oxysporum* strains could effectively inhibit Fusarium or Verticillium wilts in suppressive soils (Fravel et al., 2003; Veloso and Diaz, 2012). For instance, the nonpathogenic isolate *F. oxysporum* 47 (Fo47) induces expression of plant resistance genes and significantly reduces disease symptoms (Veloso and Diaz, 2012). Increasing evidence has demonstrated that strains of *F. oxysporum* and *F. solani*, and not other Fusarium species, are able to more efficiently establish the suppressive property in soils (Fravel et al., 2003).

### 1.2 Fusarium wilt of cotton caused by *F. oxysporum* f. sp. *vasinfectum*

*F. oxysporum* f. sp. *vasinfectum* (FOV) is a soil-borne pathogen that causes Fusarium wilt of cotton and threatens cotton production throughout the world. The disease was first identified in 1892 in cotton growing areas of Alabama in the United States, and subsequently appeared in Egypt, India, Brazil, China, and Australia (Chakrabarti et al. 2011, Holmes et al. 2009). Eight races of FOV have been identified based on their pathogenicity on different cotton cultivars or alternative host plants (Skovgaard et al., 2001). However, this classification system is time consuming, unstable, and influenced by environmental conditions. More recent research has focused on identifying genetic markers associated with each “race” of FOV, which may provide a simple and more reproducible method for identification (Assigbetse et al., 1994; Skovgaard et al., 2001).
Multiple factors can lead to cotton yield losses within a region. When planting susceptible cultivars, farmers may experience high yield losses, and in California there have been reports of total cotton crop loss in some regions. The variability in virulence of FOV populations has an influence on the development of cotton in the field. In addition, factors such as climate conditions, soil fertility, soil type, and interactions with nematodes also contribute to high yield losses (Davis et al., 2006).

FOV initially infects the plant through the root system, enters the vascular tissue via the cortex, and spreads rapidly within the host. It is able to infect cotton at any stage of development (Davis et al., 2006). In the cotton seedling stage, infection results in uneven stands, causing problems for cotton production throughout the season. After infection, older cotton plants show various symptoms such as wilting, chlorosis, leaf necrosis, and dieback (Davis et al. 2006, Chakrabarti et al. 2011). However, it is difficult to distinguish this disease from only these symptoms since the symptoms between Fusarium wilt and Verticillium wilt, another vascular disease of cotton caused by a different pathogen, are similar.

### 1.3 *F. oxysporum* pathogenicity

Several *F. oxysporum* genomes have been sequenced in recent years (Ma et al., 2010; Schmidt et al., 2016), and revealed these fungi carry mobile chromosomes that contain a high proportion of transposable elements and are involved in pathogenicity. These chromosomes are capable of being transferred horizontally among isolates of *F. oxysporum*, allowing a non-pathogenic isolate to become pathogenic on a particular host plant (Ma et al., 2010).

Plant pathogens may secrete small proteins or chemicals into the host cells to facilitate
infection. *F. oxysporum* is able to secrete numerous proteins into the host xylem tissue, many of which have been identified and referred to as SIX (secreted in xylem) proteins and some have been shown to function as effectors. Fourteen Six proteins have been identified in xylem sap after infection with *F. oxysporum* (Houterman et al., 2007), and among them, Six1, Six3, Six5, and Six6 showed involvement in fungal pathogenicity (Gawehns et al., 2014; Rep et al., 2004). Many of the SIX genes including SIX5, SIX6, and SIX7 reside on the mobile chromosomes, suggesting these effector encoding genes can be transferred horizontally to other isolates of *F. oxysporum*, potentially leading to the observed host range expansion (Ma et al., 2010). Besides the Six proteins, *F. oxysporum* secretes a plant alkalinizing peptide to alter the extracellular pH, ultimately promoting fungal growth during infection and contributing to fungal pathogenesis (Masachis et al., 2016).

*F. oxysporum* can synthesize some key secondary metabolites involved in fungal virulence. Fusaric acid (FA) was shown to be involved in virulence on both plant and animal hosts (Lopez-Diaz et al., 2018). FA is able to interfere with the water balance in banana leaves, accelerating development of Fusarium wilt symptoms (Ding et al., 2018; Dong et al., 2012). In addition, *F. oxysporum* is able to secrete siderophores, a small, membrane-transmissive, high-affinity iron-chelating compound, which can regulate fungal growth and enhance fungal virulence in low iron conditions (Leal et al., 2013). Some strains of *F. oxysporum* are also able to produce isoleucine- and leucine-conjugated jasmonate (JA-Ile/Leu) and use these compounds to promote infection of the root (Cole et al., 2014).
1.4 ROS role for fungal developments and virulence

Reactive oxygen species (ROS) such as peroxides, superoxide, and hydroxyl radical have long been known as a natural byproduct of normal metabolism and play an important role in cell signaling. The importance of ROS was first recognized as part of the innate immune system of animals where large quantities of ROS, referred to as the ‘oxidative burst’, can be rapidly released by phagocytes and used for killing invasive microbes (Rada and Leto, 2008). Increasing evidence has shown fungi can regulate ROS levels intercellularly and intracellularly for growth development and virulence (Heller and Tudzynski, 2011).

NADPH-dependent oxidases (Nox) and superoxide dismutases (SOD) are the main enzymes that contribute to the balance of ROS levels. Nox, the most important enzymatic ROS generating class of enzymes, are found in a wide range of organisms. For example, the mammalian Nox2 gene is responsible for the neutrophil oxidative burst defense response. In fungi, the Nox gene family is composed mainly of three distinct clusters, NoxA, NoxB, and NoxC (Tudzynski et al., 2012). Nox genes are involved in regulating sexual development in filamentous fungi, as in Aspergillus nidulans NoxA was required for the development of cleistothecia while the homologous gene was necessary for perithecia development in Neurospora crassa (Lara-Ortiz et al., 2003). Additionally, Nox proteins participate in the regulation and development of specific fungal structures including sclerotia and appressoria. Loss of these fungal structures lead to a reduction in fungal survival or virulence (Tudzynski et al., 2012). In Botrytis cinerea, two copies of Nox genes, NoxA and NoxB, were necessary for the development of sclerotia, a compact mass of hardened fungal mycelium that enable fungi to survive in extreme environments (Segmuller et al., 2008).
Since large amounts of ROS can cause various types of cellular damage, superoxide dismutase (SOD) plays a key antioxidant role and is an important enzyme responsible for partitioning superoxide into either oxygen or hydrogen peroxide which can be catalyzed by cellular catalases. Based on the protein folding of the SOD and the required metal cofactors, these proteins can be categorized into three types, Cu/Zn-SOD, Fe/Mn-SOD, and Ni-SOD.

Fungal SOD proteins, primarily the Cu/Zn-SOD and Fe/Mn-SOD types, have been shown to contribute to pathogenicity during fungal infections. In the medically important basidiomycete yeast *Cryptococcus neoformans*, disruption of the Cu/Zn-SOD-encoding gene (*SOD1*) lead to decreased cellular SOD activity and high sensitivity to ROS in vitro. The sod1 mutant was significantly less virulent than the wide-type strain (Cox et al., 2003). Similarly, the homologous *SOD1* genes serve as virulence factors in the phytopathogenic fungi, *F. graminearum* and *S. sclerotiorum* (Veluchamy et al., 2012; Yao et al., 2016). Fe/Mn SODs mainly localize to the mitochondria where endogenous superoxide accumulation occurs (Miller, 2012). The sod2 mutants (Fe/Mn Sod) in *A. fumigatus* have a reduced growth rate when cultured at high temperatures, while in *C. neoformans var. gattii SOD2*, rather than *SOD1*, was shown to be essential for survival during the stationary phase of growth (Lambou et al., 2010; Narasipura et al., 2005).

Recently, a class of extracellular Cu/Zn-SODs have been described that are involved in pathogenesis during fungal infections. These SOD enzymes, containing GPI anchors that allow them to be attached to the fungal cell wall, are regulated by the external environment. Interestingly, some extracellular SODs, like Sod5 from *Candida albicans*, only binds Cu instead of Zn but maintains strong SOD activity. In addition, they can reduce host-derived
superoxide and promote virulence in animal models (Broxton and Culotta, 2016; Frohner et al., 2009).

1.5 The working mechanism of the CRISPR/Cas9 system for genome editing

CRISPR (clustered regularly interspaced short palindromic repeats) is a family of DNA repeats found in ~50% of the sequenced bacterial genomes and ~90% of the archaeal genomes (Hille et al., 2018), and these repeats are homologous to some regions of viruses or other invasive genetic elements. These sequences (pre-crRNAs) are transcribed and later cleaved into short CRISPR RNAs (crRNAs) by one of the Cas proteins. The tracrRNAs, trans-activating crRNAs, direct the maturation of crRNAs by the activities of the widely conserved endogenous RNase III and the CRISPR-associated Csn1 protein (Deltcheva et al., 2011). Cas9, a DNA endonuclease enzyme, can form a simple bacterial immune system with the guide of a crRNA/tracrRNA hybrid to recognize, bind, and ultimately cleave the foreign DNA (Jinek et al., 2012). The crRNA/tracrRNA hybrid was later engineered into a single guide RNA which facilitated experimental design (Jinek et al., 2012).

The recognition of a DNA target region depends on a PAM (protospacer adjacent motif) which is located at the 3’-terminus of a 20 bp target sequence. Once the CRISPR/Cas9 complex recognizes the target DNA sequence, it will generate a double strand break (DSB) at the DNA target region between the final third and fourth nucleotides which will activate the host DNA repair process. Two cellular DNA repair mechanisms, non-homologous end joining (NHEJ) and homologous recombination (HR), play essential roles in precise genome editing and gene manipulation. NHEJ was usually viewed as an error-prone repair mechanism that generates
short insertions or deletions of nucleotides in close proximity to the DSB site(s) (Rodgers and McVey, 2016). If these short insertions or deletions exist in a gene coding region or within a portion of the promoter involved in recruiting proteins involved in transcription, the function of the endogenous gene will be disrupted. Consequently, this procedure is widely used for generating gene knock-outs. However, increasing evidence indicated that NHEJ was not always error-prone (Betermier et al., 2014). Recently, a homology independent targeted integration (HITI) strategy was devised and allowed fragments to be integrated into the genome by NHEJ repair mechanisms. Interestingly, the integration can occur in both dividing and non-dividing cells with a high probability of perfect DNA repair at the gRNA cleavage site (Guo et al., 2018; Suzuki et al., 2016). The second well-known DSB repair pathway is based on homologous recombination (HR). In HR where a template exists, DSBs will be repaired based on the templates, allowing the foreign DNA fragments to be precisely integrated into the host genome. However, HR-mediated DSB repair only occurs in dividing cells, limiting the application of this method to a broad range of uses (Fig 1).

1.6 Various strategies for developing CRISPR/Cas9 tools in filamentous fungi

Unlike animal cell lines, fungi have thick cell walls and can have a complicated genetic background as it is not uncommon for fungi to have many duplicated loci, have undergone whole genome duplications, or harbor supernumerary chromosomes. Transformation of filamentous fungi usually requires the remove all of the cell wall generating protoplasts or utilizes electrical shock. Consequently, various versions of CRISPR/Cas9 tools have been
developed for fungi and each of these methods may be the preferred methods for a particular fungus. The yeast *Saccharomyces cerevisiae* was the first fungus that CRISPR/Cas9 was used to genetically modify the genome (DiCarlo et al., 2013), and the ability of this fungus to stably maintain plasmids facilitated CRISPR/Cas9 tool development. Usually these plasmids contain the Cas9 gene under constitutive expression, and when coupled with transient gRNA cassette transformation, generates DSBs thereby increasing homologous recombination rates (DiCarlo et al., 2013). Previously, gene manipulation in the clinically relevant yeast *C. albicans* was tedious as the fungus is a diploid; however development of a CRISPR/Cas9 method provided an efficient way to generate homozygous mutations and facilitated genome engineering in this organism (Vyas et al., 2015).

Based on these initial CRISPR/Cas9 developments with yeasts, this gene editing tool was adapted to filamentous fungi, in which several methods currently exist for both model and non-model fungi. A CRISPR/Cas9 system was first established in the filamentous fungus *Trichoderma reesei* using a codon-optimized Cas9 gene that was introduced into the genome using *Agrobacterium tumefaciens*-mediated transformation (Liu et al., 2015). When the expression of Cas9 was under the control of either a constitutive or inducible promoter, there was obvious alteration in the function of other proteins as observed by fungal morphology and various other phenotypes. The successful integration of Cas9 into the fungal genome generated a strain (a ‘Cas9’ strain) that requires only the sgRNA for genome editing through protoplast transformation, providing a method to rapidly target multiple genes simultaneously (Liu et al., 2015).

The random integration of the Cas9 gene into the host genome may induce undesired
mutations, and obtaining a ‘Cas9’ fungal strain may require screening several transformants or optimizing the codon usage of the Cas9 gene due to different codon preference for each species. Another caveat is the constitutive expression of Cas9 and/or the long-time retention in host cells may generate off-target effects (Kuscu et al., 2014). To overcome these limitations, transient expression systems of the Cas9 gene and gRNA in filamentous fungi have been utilized. A transient expression CRISPR/Cas9 system was developed in the model filamentous fungus, N. crassa (Matsu-Ura et al., 2015), where the Cas9 gene was engineered with the trpC constitutive promoter from A. nidulans, and the Small Nucleolar RNA 52 (SNR52) promoter from S. cerevisiae was used to drive gRNA expression. The editing efficiency gradually increased with increasing number of copies of Cas9 and quantity of gRNA plasmids. The replacement of the endogenous promoter of the clr-2 gene with the β-tubulin promoter and the insertion of a codon optimized firefly luciferase cassette suggested that this strategy was feasible and efficient in filamentous fungi (Matsu-Ura et al., 2015).

A CRISPR/Cas9 gene editing tool for the multinucleate filamentous phytopathogenic fungus Sclerotinia sclerotiorum has been reported (Li et al., 2018). Cas9 and gRNA are able to be transiently expressed in the protoplasts and function properly. Interestingly, plasmid fragments of various lengths are integrated into the Cas9/sgRNA cleavage site by the NHEJ repair pathway, generating mutants with large insertions of the plasmid (Li et al., 2018). Using this tool, the Ssoah1 gene which is involved in the accumulation of oxalic acid was disrupted, and the ΔSsoah1 strain was unable to acidify the growth medium and change the pH indicator (Li et al., 2018).

Although filamentous fungi cannot stably maintain most plasmids, several chromosomal
replicators, including AMA1 and MATE elements, have provided a possible mechanism for extrachromosomal maintenance of plasmids in filamentous fungi (Aleksenko and Clutterbuck, 1997). AMA1 plasmids can significantly enhance transformation efficiency, and are able to be easily lost after transformation without selection, allowing the dominant selectable marker(s) to be reused for additional transformation procedures. In addition, the Cas9-sgRNA gene cassette on the AMA1 plasmids is less likely to be inserted into the fungal genome during transformation. Based on these advantages, several CRISPR/Cas9 systems utilize AMA1 plasmids for filamentous fungi (Nielsen et al., 2017; Nodvig et al., 2015; Pohl et al., 2016). In one of the first studies using this method, Nodvig et al. successfully induced mutagenesis of the yA gene in A. nidulans, which was responsible for producing the yellow color of conidia (Nodvig et al., 2015). In this study, the expressions of the gRNA depended on two key components, an intrinsic hammerhead (HH) ribozyme and a hepatitis delta virus (HDV) ribozyme. Both can be self-cleaved and allowed the precise initiation of gRNA (Nodvig et al., 2015). In addition, Pohl et al. knocked out a polyketide synthetase (pks17) from Penicillium chrysogenum, which was involved in formation of a green pigment in spores, resulting in a mutant where the colony color changed from green to white (Pohl et al., 2016).

All of these CRISPR/Cas9 systems in filamentous fungi require a functional host transcription or translation system for the expression of the foreign Cas9 gene and gRNA transcription. Appropriate promoters must be chosen for both the Cas9 and gRNA expression, requiring various promoters to be attempted and perhaps optimized for Cas9 gene expression in different filamentous fungi. In addition, the synthesis of gRNA is controlled by the RNA polymerase III (Pol III) promoters like the U6 promoter and requires precise transcription
initiation. A considerable amount of time and effort is necessary for optimizing a plasmid-based Cas9/gRNA delivery method. Consequently, more recent attempts have used a Cas9 RNP-based delivery method for genome editing in different species, including Arabidopsis, fruit fly, zebrafish, and human cell lines. Cas9 RNPs consist of the purified Cas9 protein and in vitro transcribed gRNA, and both components are assembled in vitro into a complex and directly transfected to the host cells for gene editing. Compared to plasmid-based delivery, the Cas9 RNPs-based delivery has several advantages including: 1) the expression of the Cas9 gene and transcription of the gRNA does not rely on host machinery, and therefore the precise transcription initiation of the gRNA is not a concern; 2) the assembled Cas9 RNPs have immediate cleavage activity and can be directly delivered into the host cells to target the specified DNA locus without having to express Cas9 or transcribe the gRNA; 3) the cleavage efficiency of the different gRNAs can be evaluated in vitro, allowing a more optimal gRNA to be chosen for further experiments; 4) Cas9 RNPs will be removed over a short timeframe from host cells by protein degradation pathways, reducing the possibility of off-target activity of Cas9.

Cas9 RNPs-based delivery methods have been used in filamentous fungi. The well-established polyethylene glycol (PEG)-mediated protoplast transformation technique has been used for Cas9 RNP delivery, in which protoplasts are generated and the Cas9 RNPs added to the recipient fungal protoplasts for genome manipulations. Cas9 RNP is a large ribonucleoprotein, but despite their size Cas9 RNPs can be transferred into fungal protoplasts by PEG-mediated transformation. A simple and universal CRISPR/Cas9 system based on in vitro-assembled Cas9 RNPs has been reported in A. fumigatus (Al Abdallah et al., 2017), where
two strains, the WT and a ΔakuB (disrupting the NHEJ pathway), were used in a study on the CRISPR/Cas9-mediated HR repair mechanism to disrupt pksP, a gene responsible for the green color of the conidia. The hygromycin resistance cassette flanked with two microhomology sequences (35~50 bp) at both ends of the cassette would serve as both the selective marker and the donor template for insertion into the Cas9 RNP cleavage sites. With the ΔakuB laboratory strain, up to 97% of the colonies showed the desired phenotypes regardless of using a 35 bp or 50 bp homology sequence while in the WT strain, the efficiency was 40~74%, and the length of microhomology directly influenced the integration efficiency (Al Abdallah et al., 2017).
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With the gRNA, Cas9 can cleave double stranded DNA (inducing dsDNA breaks, DSBs) at specific sites, activating the host DNA repair mechanism. Two well-known DNA repair pathways, NHEJ and HDR, were utilized for host genome modifications. NHEJ could disrupt the desired genes due to an error in the DNA repair or integration of fragments into the genome by HDR and precise DNA repair.

**Figure 1** CRISPR/Cas9 gene editing system and DNA repair mechanisms.
Chapter 2.

Efficient genome editing in *Fusarium oxysporum* based on CRISPR/Cas9 ribonucleoprotein complexes

This chapter has been published (Wang et al., 2018).

2.1 Introduction

Fungi in the genus *Fusarium* originated during the Cretaceous period and are currently represented by at least 20 species complexes containing isolates that are able to cause diseases in plants, animals, and humans (Geiser et al., 2013; O'Donnell et al., 2013). They are well known to produce a diverse array of secondary metabolites including mycotoxins that contaminate food and poison livestock leading to reduced production (Geiser et al., 2013; Hansen et al., 2012; O'Donnell et al., 2013). One of these groups, the *Fusarium oxysporum* species complex (FOSC), are soil-borne filamentous fungi that have an extensive host range from plants to animals. These isolates carry small supernumerary chromosomes that harbor host-specific virulence factors that are capable of being horizontally transferred to other FOSC isolates and increase host range (Ma et al., 2010; van Dam et al., 2017). In agriculture, *F. oxysporum* is an important plant pathogen that is able to cause significant economic damage on various plant species including tomato, banana, legumes, and cotton (Michielse and Rep, 2009). This fungus infects the root system of a susceptible host and eventually colonizes the
xylem system, where it blocks the translocation of water and nutrients, resulting in chlorosis, necrosis, stunting, and wilting. Clinically, members of the FOSC are responsible for ~20% of the disseminated *Fusarium* infections termed fusariosis (Muhammed et al., 2013; Nucci and Anaissie, 2007) and have been implicated in several disease outbreaks such as nosocomial infections due to contaminated water supplies and contact lens associated keratitis (Chang et al., 2006; Edel-Hermann et al., 2016; O’Donnell et al., 2004).

Reverse genetic approaches utilizing gene deletion/disruption technologies have played an essential role in studies identifying the function of a gene. Recently, the CRISPR (clustered regularly interspaced short palindromic repeats)-Cas9 system, derived from the bacterial and archaeal immune system, has been developed into a powerful gene editing tool (Doudna and Charpentier, 2014; Hsu et al., 2014; Ran et al., 2013). Based on the highly efficient gene targeting editing and low off-target rates, this tool has become popular and has been widely used in model or non-model organisms such as fruit fly, zebrafish, *Arabidopsis*, yeast, mouse, and human cell cultures (Bassett et al., 2014; Jacobs et al., 2014; Jiang et al., 2013; Li et al., 2016; Platt et al., 2014; Shalem et al., 2014). The simple gene editing system includes two components, a single chimeric guide RNA (sgRNA) and the Cas9 endonuclease enzyme. The sgRNA consists of a protospacer sequence which targets the expected DNA region by base pairing, and is able to interact with Cas9 to become a stable complex. Once the base pairing has occurred, Cas9 cuts the double-stranded DNA in vivo, subsequently activating DNA repair mechanisms. One method of DNA repair involves random insertion or deletion of nucleotides at the DNA cleavage site (Non-Homologous End Joining, NHEJ). The NHEJ mechanism tends to repair the DNA incorrectly, and if this repair occurs in a gene coding region, the associated
gene is likely to be inactivated. In contrast, when a template with homologous sequence is present, the cleaved DNA region could be repaired using the homologous sequence as a template. The main benefits of this gene disruption/deletion system are accurate gene modification and highly successful mutation rates (Doudna and Charpentier, 2014; Ran et al., 2013).

The development of CRISPR/Cas9 genome editing tools for fungal research has facilitated functional studies. Several gene deletion systems based on CRISPR/Cas9 have been developed in *Saccharomyces cerevisiae*, *Candida albicans*, *Trichoderma*, *Aspergillus*, *Alternaria* and *Neurospora* (DiCarlo et al., 2013; Liu et al., 2015; Matsu-Ura et al., 2015; Nodvig et al., 2015; Vyas et al., 2015; Wenderoth et al., 2017). While these cases showed successful gene disruptions, some limitations still exist. *Trichoderma reesei* was the first filamentous fungus used for CRISPR/Cas9 research, where *Agrobacterium*-mediated fungal transformation was used to insert a codon-optimized Cas9 gene randomly into the genome leading to a potential unintended mutation. Other CRISPR tools have been developed for filamentous fungi that are highly dependent on AMA1-derived plasmids (Nodvig et al., 2015; Pohl et al., 2016; Wenderoth et al., 2017). The AMA1 element is an inverted duplication of a sequence derived from *Aspergillus nidulans*, and it is able to enhance fungal transformation efficiency (Aleksenko and Clutterbuck, 1997). However, experimental evidence has shown that AMA1 plasmid integration occurred in some filamentous fungi and could lead to long-time retention in fungal transformants (Fierro et al., 1996). Additionally, after successful transformation positive colonies may have to be treated with single spore isolation for several rounds to remove the residual AMA1 plasmids (Nodvig et al., 2015; Wenderoth et al., 2017).
Using the Cas9 protein/sgRNA ribonucleoproteins (RNPs) to perform genome editing has several advantages compared with plasmid transformation and Cas9 mRNA/sgRNA transformation. A major advantage is transformation of Cas9 RNPs alleviates the possibility of integration of genetic material to a non-targeted region of the genome. Additionally, Cas9 and sgRNA are able to form a stable ribonucleoprotein in vitro, so there is less likelihood of RNA degradation compared with Cas9 mRNA/sgRNA transformation. Furthermore, this method is able to evaluate the target cleavage efficiency of different sgRNA in vitro, so a highly efficient sgRNA can be selected for research. This method has been widely used in different organisms such as mouse, human, grape, apple, tobacco, and Arabidopsis (Kim et al., 2014; Malnoy et al., 2016; Woo et al., 2015). In plants, an efficient protoplast transformation method for Cas9 RNPs has been developed and mutant individuals can be generated from the Cas9 RNP-modified protoplasts (Woo et al., 2015); however, few studies involving Cas9 RNP transformation for filamentous fungi have been published.

In this study, we developed and optimized a Cas9 RNP transformation procedure for gene editing in the ascomycete fungus, Fusarium oxysporum. A PEG-mediated transformation method was used to transfer Cas9 RNPs, which efficiently target the locus of interest, into fungal protoplasts. Several genes that have been well characterized in other fungi were disrupted in F. oxysporum with a hygromycin gene cassette using this system, generating mutants displaying strong expected phenotypes based on characterization of orthologous genes. As a proof of concept, the gene encoding the polyketide synthase PKS4 was disrupted using this system and demonstrated this enzyme is responsible for production of the red pigment bikaverin, and in keeping with the established nomenclature of other Fusarium spp., refer to it
as FoBIK1 (Wiemann et al., 2009).

2.2 Materials and methods

2.2.1 Fungal isolate and culture media.

All experiments were conducted using *F. oxysporum* f. sp. *vasinfectum* obtained from the Fungal Genetics Stock Center (FGSC #10442) (McCluskey et al., 2010). This fungus was cultured in Potato Dextrose Broth (PDB), PDB with agar, and minimal medium (M-100) agar plates (Stevens, 1974). All mutants were strictly determined on M-100 agar plates with a hygromycin concentration of 75 μg/mL or a 5-FOA concentration of 1.5 g/L. The wild type and mutant strains were cultured at room temperature (25 °C).

2.2.2 Sequence analysis and fluorescence microscopy.

Amino acid multiple sequence alignment analysis was conducted using Clustal Omega (Sievers et al., 2011), and the nuclear localization signal prediction accomplished using NLStradamus (Nguyen Ba et al., 2009). The subcellular localization of NLS<sub>H2B</sub>-eGFP was carried out using a Zeiss Axiovert 200 fluorescence microscope. The transformant containing NLS<sub>H2B</sub>-eGFP was cultured in PDB medium on a rotary shaker (160 rpm) at room temperature for three days. Before microscopic examination, the hyphae were stained with 4′,6-diamidino-2-phenylindole (DAPI) at a concentration of 1 μg/mL for 20 min in the dark. The excitation for the DAPI dye was 405 nm while the value for eGFP was 488 nm.

2.2.3 Plasmid construction.

The donor template DNA loci containing the hygromycin cassette were constructed using a NEBuilder® HiFi DNA Assembly Master Mix kit. All template plasmids were constructed
in the background of the pUC19 plasmid with HindIII and BamHI restriction sites. For the construction of the pExFO-NLSH2BeGFP plasmid, the basic plasmid pExFO was constructed in the background of pGEM-T where three fragments, the hygromycin cassette including pgpdA promoter amplified from the plasmid pCWHyg1, the trpC promoter amplified from the plasmid pII99, and the tef1 terminator amplified from the plasmid pFFC332, were simultaneously inserted into the SphI and SacI enzyme sites in pGEM-T. Between the trpC promoter and the tef1 terminator, a multiple cloning site (MCS, HindIII-KpnI-NotI-PacI-SalI-SpeI-XbaI) were inserted. The NLS sequence from histone H2B and eGFP sequence were inserted into this MCS region for the construction of pExFO-NLSH2BeGFP. The assembled plasmids were sequenced to confirm they were correct and plasmids were extracted with the E.Z.N.A® plasmid DNA maxi kit (OMEGA, bio-tek) for PEG-mediated transformation.

The plasmid construction for the *E.coli* protein expression and purification was performed with the pHis-parallel1 plasmid (Sheffield et al., 1999). First, the nuclear localization sequence from the H2B gene of *F. oxysporum* was cloned from genomic DNA using primers NLSH2B_BamHIF and NLSH2B_SalIR and inserted into the BamHI and Sall enzyme sites of the pHis-parallel1 plasmid (pHis-parallel1-NLSH2B). The Cas9 gene was cloned from the plasmid pFFC332, and subsequently inserted between the Sall and HindIII sites of the pHis-parallel1-NLSH2B plasmid with the NEBuilder® HiFi DNA Assembly Master Mix kit. The plasmids were sequenced to confirm no mutations for inserted fragments. All the primers used for plasmid constructions and sequencing in this study are listed in Table 2.2.

### 2.2.4 Cas9 protein purification.

The plasmid pHis-parallel1-NLSH2BCas9 was transformed into Rosetta™(DE3) strain
which is able to highly express genes containing rare codons. The correctly transformed *E. coli* isolate was determined by PCR using with primers DetCas9F and DetCas9R (Table 2.2). High expression of this protein in the *E. coli* containing the pHis-parallel1-NLS\textsubscript{H2B}Cas9 plasmid was obtained by first culturing in 50 mL LB liquid medium for 14 h at 37 °C with 100 μg/mL ampicillin. The cultured *E. coli* was used to inoculate 1.5 L of fresh LB liquid medium at a dilution of 1:100. When the OD\textsubscript{600} value of the *E. coli* cell culture concentration reached 0.6, IPTG was added to a final concentration of 0.3 mM and the culture incubated for an additional 5 h at room temperature (25 °C) at 160 rpm on a rotary shaker. After determining the proper protein expression using a SDS-PAGE gel, the culture was centrifuged at 13000 rpm for 15 min and the pellet suspended in 30 mL lysis buffer (20 mM NaH\textsubscript{2}PO\textsubscript{4}, 300 mM NaCl, 1 mM imidazole, 1% TritonX-100, 1 mM PMSF, pH 7.5) and lysed in an ultrasonic bath (15 s + 15 s, 10 min). The mixture was centrifuged at 13000 rpm, 4 °C for 15 min, and the supernatants were filtered with a 0.22 μm filter, and subsequent protein purification conducted using HisPur™ Ni-NTA Chromatography Cartridges (Thermo Fisher Scientific). The eluted Cas9 protein was further dialyzed against the Cas9 buffer (20 mM NaH\textsubscript{2}PO\textsubscript{4}, 300 mM NaCl, 10% glycerol, pH 7.5) for 6-8 h. The dialyzed Cas9 protein was centrifuged and the associated undissolved protein was removed. The Cas9 protein was further concentrated with a 100 K centrifugal filter to a final concentration of 4 mg/mL. The Cas9 nuclease cleavage assay *in vitro* should be conducted immediately. The final purified Cas9 protein was aliquoted into PCR tubes and stored at -30°C for future use. There was no significant difference in cleavage activity of the purified Cas9 protein after a year at -30°C.

2.2.5 sgRNA design and in vitro Cas9 nuclease assay.
The sgRNA for gene mutation was designed with the following parameters: 1) the sgRNA cleavage site was located close to the 5’ end of the gene coding region; 2) the protospacer and PAM sequence was (N)_{20}NGG, and if the beginning nucleotide of the target sequence lacked a G, at least one G was added after the T7 promoter; 3) similarity to the protospacer and PAM sequence was searched by BLASTn to the *F. oxysporum* genome to avoid cleavage at other loci; and 4) the sgRNA forward primer (T7 promoter-(N)_{20}-GTTTTAGAGCTAGAAATAGCAAG) and the sgRNA universal primer (Table 2.2) were used to generate the sgRNA templates by PCR using the EnGen® sgRNA Synthesis kit (New England Biolabs).

For the *in vitro* cleavage assay mediated by Cas9 with different sgRNAs, a PCR fragment ranging from 1000 bp to 3000 bp was first amplified and purified from the agarose gel with E.Z.N.A® Gel Extraction Kit (OMEGA, bio-tek). The purified Cas9 protein was diluted to a concentration of 200 ng/μL. The cleavage assay was conducted in a system with 20 μL total volume (1 μL diluted purified Cas9 protein, 40 ng sgRNA, 1× Cas9 Nuclease Reaction Buffer, 3 μL 20 nM PCR fragment, DEPC-treated water). All items were added together and incubated at 37 °C for 1 h. The final products were analyzed on a 0.8% agarose gel and the associated cleavage efficiency was evaluated with the software, ImageJ (https://imagej.nih.gov/ij/). All sgRNAs and the associated targeted genes used in this study are listed in Table 2.3.

2.2.6 PEG-mediated fungal transformation.

PEG-mediated fungal transformation was conducted according as previously described with slight modifications (Coleman et al., 2011b). About 10^7 conidia were inoculated into 100 mL fresh PDB liquid medium and allowed to germinate at 28 °C at 160 rpm overnight. The
fungal germlings were collected and treated with a mixture of 10 mg/mL Driselase (Sigma) and 15 mg/mL β-Glucanase (Sigma) suspended in 0.7M NaCl for 2-3 h. Fungal protoplasts were collected and washed twice with SuTC buffer. The fungal protoplasts were diluted into a concentration of \(2 \times 10^7\) per mL with SuTC buffer, and 200 μL of the protoplast solution was used for each transformation.

The Cas9 RNPs or template donor DNA was prepared during generation of the protoplasts where the Cas9 RNPs were made as follows: a 1:1 mole ratio of Cas9: sgRNA was added into a 50 μL total volume with 5 μL 10× Cas9 Nuclease Reaction Buffer and DEPC-treated water. This mixture was incubated in a 37 °C water bath for 20 min. 200 μL of fungal protoplasts were mixed with 50 μL Cas9 RNPs or donor DNA (template) at room temperature for 20 min. An equal volume of 60 % PEG was added into the above system and incubated at room temperature for 20 min. The total system was transferred into a 15 mL tube containing 5 mL TB3 and placed on a rotary shaker (160 rpm at room temperature) for 18-24 h. The regenerated fungal protoplasts were mixed with the selective medium and poured into sterile plates. For screening ura3 and ura5 mutants with only Cas9 RNPs transformation, the selective plates with uracil, uridine, and 2 g/L 5-FOA were used. For the ura5 mutant generation with Cas9 RNPs and donor DNA transformation, the selective plates contained uracil, uridine, and a final concentration of 150 μg/mL hygromycin. The Fobik1 mutant selection was accomplished using selective plates with 150 μg/mL hygromycin. All transformants were visible in 3 ~ 5 days after plating on selective media. The transformants were then transferred onto the minimum nutrient agar medium (M100) with the associated antibiotics for further determination.

Three pairs of primers were used to detect the hygromycin cassette location for HR-
directed mutant determination. The first pair (H_inF + H_inR) was used to detect the presence of the hygromycin phosphotransferase gene. For HR-directed *ura5* mutants, three pairs of primers (H_outF+ FoURA5TR, FoURA5TF + H_outR and FoURA5TF + FoURA5TR) were used to determine the insertion location of the hygromycin phosphotransferase gene. Similarly, Det_bik1g1F + Det_bik1g1R and H_outF+ Det_bik1g2R were used to determine HR-directed *Fobik1* gene mutants.

2.7 HPLC and MS analysis.

The wild type and *Fobik1* mutants were inoculated into the fresh PDB liquid medium and grown on a rotary shaker (160 rpm) at room temperature for 7 days. The culture was centrifuged at 13000 rpm for 10 min to remove conidia and hyphae. The supernatant was treated with a 0.22 μm filter. The supernatant medium from wild type and the *Fobik1* mutant was fractionated on a Sonoma phenyl-hexyl 10-μm 100-Å (25 cm x 4.6 mm) HPLC column (ES Industries, West Berlin, NJ, USA) which was equilibrated in 1% formic acid and acetonitrile (ACN). The HPLC programs were set as follows: a linear gradient from 20% to 50% ACN was applied over 15 minutes, and a linear gradient from 50% to 75% ACN was used over 5 minutes. The column was washed with 100% ACN for 5 minutes and then was re-equilibrated as above. Fractions were collected and submitted to MS analysis which was performed on a Bruker Micro-TOF MS with flow injection.
2.3 Results and Discussion

2.3.1 Identification of an efficient endogenous nuclear localization sequence (NLS) from \textit{F. oxysporum}.

The Cas9 RNPs must enter into the nucleus in order to bind to the targeted DNA regions. The classical nuclear localization sequence (NLS) SV40 has been shown to be functional in many organisms (Liu et al., 2015), however this NLS was not suitable for \textit{F. oxysporum} and was unable to efficiently translocate GFP into the nucleus (data not shown), similar to observed results with other organisms such as \textit{Phytophthora sojae} (Fang and Tyler, 2016). In order to facilitate Cas9 entering the \textit{F. oxysporum} nucleus, the endogenous nuclear localization sequence from histone H2B was fused to the N-terminus of Cas9. Histone H2B is conserved in eukaryotes and involved in the assembly of the nucleosomes (Mosammaparast et al., 2001). The histone H2B NLS amino acid sequence was highly conserved among \textit{Fusarium} spp., which indicates that this system may be applicable for use in other \textit{Fusarium} species (Fig. 2.7).

Bioinformatic analysis revealed the \textit{F. oxysporum} histone H2B contained a NLS at its N-terminus (NLS\textsubscript{H2B}, aa: 6-45). The middle and C-terminus amino acids of histone H2B were highly conserved among a diverse group of organisms including other fungi, plants, and metazoans, but the NLS sequences of these organisms has evolved divergently (Fig. 2.1A). In order to confirm the predicted NLS of histone H2B was sufficient to localize a protein to the nucleus of \textit{F. oxysporum}, the first 54 amino acids including the entire NLS sequence were fused to the N-terminus of eGFP under the constitutive promoter of \textit{trpC} (pExFO-NLS\textsubscript{H2B}eGFP). Transformants of \textit{F. oxysporium} expressing the NLS\textsubscript{H2B}-eGFP construct contained eGFP
localized to the nuclei of the cells (Fig. 2.1B), indicating the NLS_{H2B} was suitable to engineer a protein to be translocated into the nucleus.

2.3.2 Design and protein purification of NLS_{H2B}Cas9 and \textit{in vitro} cleavage assay.

The endogenous NLS from the \textit{F. oxysporum} histone H2B gene was fused to the N-terminus of the Cas9 gene and cloned into an \textit{E. coli} protein expression plasmid (pHis parallel1-NLS_{H2B}Cas9) which was subsequently transformed into the \textit{E. coli} RosettaTM (DE3) strain for protein expression and purification. The resulting recombinant protein with a molecular weight of approximately 165 kDa was soluble in water and reached a yield at least 4 mg per liter of bacterial culture (Fig. 2.2A). The ability of the purified Cas9 protein to efficiently cleave at a specified target site was evaluated \textit{in vitro}. A \~{}2.1 kb PCR fragment containing the coding region of the \textit{F. oxysporum} URA5 gene was mixed with a single guide RNA targeting this locus and the Cas9 protein. The NLS_{H2B}Cas9-FoURA5sgRNA complex was able to cleave almost all of the template \textit{in vitro} within an hour and the lengths of cleaved fragments were of the expected size, supporting that the NLS_{H2B} sequence fusion had no significant influence on the Cas9 endonuclease cleavage activity (Fig. 2.2B).

2.3.3 A PEG-mediated transformation method can transfer NLS_{H2B}Cas9 RNPs into fungal protoplasts.

Recently, Cas9 RNPs have been transferred into plant protoplasts for gene-editing (Malnoy et al., 2016; Woo et al., 2015). This system has at least two advantages in filamentous fungi 1) fungal protoplasts are easier to be regenerated into individual transformants (compared to
plants) and 2) Cas9 RNPs will be degraded in the protoplasts, which further minimizes the potential influence from the Cas9 RNPs being maintained in the cell. A RNP mediated transformation method has been developed for two fungi, *Aspergillus fumigatus* and *Penicillium chrysogenum* (Al Abdallah et al., 2017; Pohl et al., 2016). A PEG-mediated transformation method was developed for the *F. oxysporum* optimized NLS<sub>H2B</sub>Cas9. The *URA5* gene encoding orotate phosphoribosyltransferase involved in pyrimidine biosynthesis was initially targeted, as mutants of the ortholog in *S. cerevisiae* and other fungi generates uracil auxotrophs and allows for direct selection on media containing 5-fluoro-orotic acid (5-FOA) (de Montigny et al., 1989). Protoplasts transformed with the NLS<sub>H2B</sub>Cas9-FoURA5sgRNA RNPs should result in cleavage at the target site in the *URA5* coding region and subsequent inactivation of the gene due to NHEJ repair. Seven transformants were generated that were able to grow on 5-FOA selective media containing uracil and uridine. As Cas9 RNPs cleave the dsDNA between the last third and fourth nucleotide at the region targeted by the protospacer element, this region was sequenced to characterize the resulting mutations in the transformants. All these *ura5* mutants contained short nucleotide deletions ranging in size from a single nucleotide to -123 bp in the target region of the guide RNA (Fig. 2.3). The *ura5* mutants were uracil auxotrophs and are able to grow on media containing 5-FOA (Fig. 2.4).

In order to further confirm that PEG-mediated transformation of Cas9 RNPs can be used in *F. oxysporum* protoplasts another sgRNA was generated that targets the homologous gene of *URA3* encoding orotidine 5'-phosphate decarboxylase. This gene is involved in de novo UMP biosynthesis and, similar to *ura5* mutants, *ura3* mutants are uracil auxotrophs and can grow on minimum media containing 5-FOA (Fig. 2.4) (Boeke et al., 1984). The targeted locus of two
randomly selected *ura3* transformants were sequenced and contained short deletions at the Cas9 cleavage site (Fig. 2.3C). Overall, the PEG-mediated transformation of Cas9 RNPs into *F. oxysporum* protoplasts was successful and the transformed particles can cleave the genomic DNA at the target site generating a mutation via an error-prone DNA repair mechanism. Interestingly, among the sequenced transformants, several of the mutants generated using this gene editing tool contained a single nucleotide deletion instead of double or four nucleotide deletions, which suggests the NHEJ repair system of *F. oxysporum* prefers the removal of a single nucleotide at the cleavage site causing a frame-shift mutation in the coding sequence.

2.3.4 Homology directed repair (HDR) can efficiently disrupt targeted genes by insertion of a dominant selectable marker.

To further extend the protocol, experiments were conducted to assess whether donor DNA containing a selective marker and the NLS<sub>H2B</sub>Cas9 RNPs are able to promote CRISPR–Cas9 induced homologous recombination. The NLS<sub>H2B</sub>Cas9-FoURA5sgRNA transformation system used for NHEJ mutation above (Fig. 2.3) was selected as it was previously confirmed to have activity at the specified target site in *URA5*. The donor DNA contained the gene encoding hygromycin phosphotransferase (*hph*), conferring resistance to hygromycin, and had ~ 600 bp of homologous sequence flanking both sides of the cleavage site in the *URA5* gene (Fig. 2.3D). Mutants should grow on the selective medium including uracil, uridine, and hygromycin (Fig. 2.4). Of the 14 transformants generated, three contained the *hph* cassette integrated at the gRNA target site (21.4%; Table 2.1). The *ura5* mutants are unable to grow on minimal medium containing hygromycin without uracil and uridine and were resistant to 5-FOA, supporting the
integration of the hygromycin resistance cassette at the targeted cleavage site in the *F. oxysporum* *URA5* gene (Fig. 2.4).

2.3.5 Cas9 RNP-mediated disruption of a gene encoding a polyketide synthase in a secondary metabolite biosynthetic gene cluster.

Some filamentous fungi including *Aspergillus*, *Penicillium*, and *Alternaria*, produce a pigment on certain agar medium during the early growth phase (Nodvig et al., 2015; Pohl et al., 2016; Wenderoth et al., 2017), and detection of these pigments have been used as a marker for the rapid development of CRISPR methodologies in these fungi. *F. oxysporum* is able to produce red pigmented compounds during the later stages of growth. While the gene cluster responsible for the pigment production has not been functionally characterized in *F. oxysporum*, the biosynthetic genes responsible for a red pigment (bikaverin) biosynthesis in the closely related fungus *Fusarium fujikuroi* are encoded in a large secondary metabolite gene cluster (Wiemann et al., 2009). Bioinformatic analysis showed that there is a similar secondary metabolite cluster that may be involved in the synthesis of bikaverin in *F. oxysporum*. This secondary metabolite cluster is composed of six genes, where the ortholog to the *BIK1* gene encodes a putative polyketide synthase that could be essential for the biosynthesis of bikaverin. As this method can mediate donor DNA and NLS<sub>H2B</sub>Cas9 RNPs to enter *F. oxysporum* protoplasts and trigger the CRISPR–Cas9 induced homology directed repair, the *bik1* gene was selected for optimizing our CRISPR system.

In order to confirm whether the putative *BIK1* gene from *F. oxysporum* (FOTG_08225) was involved in bikaverin synthesis, a sgRNA site was chosen with a location close to the 5′-
terminus of the coding region in the first exon (Fig. 2.5A). The PCR template cleavage efficiency was close to 100% in vitro, demonstrating the chosen sgRNA had high cleavage activity with NLS_{H2B}Cas9 (Fig. 2.5B). In order to optimize this CRISPR system various concentrations of the assembled NLS_{H2B}Cas9-Fobik1sgRNA RNPs were used for transformation. In the absence of the Cas9 RNPs in the transformation system, no bik1 F. oxysporum mutants were generated; however, using concentrations of Cas9/sgRNA between 10/2 (μg/μg) and 15/3 (μg/μg) resulted in generation of Δbik1 transformants of approximately 50% (Table 2.1). Using a Cas9/sgRNA concentration higher than 15/3 (μg/μg) was unable to further increase the frequency of desired mutants. Six randomly selected Fobik1 mutants were unable to produce the red pigment when cultured in conducive conditions such as PDB medium for 7 days (Fig. 2.5C), supporting this secondary metabolite gene cluster is responsible for the synthesis of the red pigment. The identity of the red pigment of F. oxysporum was confirmed to be bikaverin by fractionation on HPLC using a phenyl hexyl column and mass spectrometry (MS). The HPLC profile monitoring absorbance at 400 nm indicated the wide-type isolate contained multiple peaks with one major peak at a retention time (R_T) of 22 minutes, that appeared different to two large peaks seen under highly similar conditions in F. fujikuroi (Fig. 2.5D, Fig. 8). Previous results showed F. fujikuroi synthesized two similarly structured compounds, nor-bikaverin and bikaverin (Wiemann et al., 2009). The peak at R_T~22 min from the medium of the wild-type isolate and the associated region from the medium of Fobik1 mutant were collected for MS analysis. A large peak with a molecular weight of 383 was present in the extract of the wild-type F. oxysporum isolate, which was absent in the fraction from the Fobik1 mutant (Fig. 8). The exact mass determination and isotope ratio suggest a best
fit to $C_{26}H_{14}O_8$ corresponded to bikaverin and initial MS/MS fragmentation of the molecule confirmed the identity as bikaverin (ESI-MS $m/z$ 383: MS/MS (40 eV) $m/z$ (%): 270 (100), 340 (70), 256 (60), 297 (25)). This identification combined with the absence of this secondary metabolite in the mutant demonstrates the disrupted gene ($FoBIK1$) is responsible for the synthesis of bikaverin in $F. oxysporum$.

This CRISPR/Cas9 RNP transformation system is an additional molecular tool to study members of the FOSC. Unlike currently used transformation procedures such as the $Agrobacterium tumefaciens$-mediated and protoplast transformation methods (Coleman et al., 2011a; Mullins et al., 2001), when utilizing the CRISPR/Cas9 RNP system only short flanking sequences are required for homologous recombination, thereby bypassing the need to construct vectors with >1 kb of homologous sequence flanking the selectable marker. Additionally, this system has the potential to be used to target multiple copies of a single gene if the entire PAM sequence and a significant proportion of the protospacer region between the copies are conserved. While the NLS$_{H2B}$Cas9 protein needs to be expressed and purified in vitro prior to transformation, the advantage of using the purified Cas9 protein is it does not rely on the integration and expression of the Cas9 gene within the fungal nucleus, and therefore there is less of a chance of unforeseen effects due to the random integration of the Cas9 gene within the fungal genome.

In summary, we have developed an approach to transfer Cas9 RNPs and/or donor DNA into fungal protoplasts for gene editing as demonstrated by generating several mutants having the expected phenotypes (Fig. 2.6). In addition, we showed experimental evidence that the red-pigmented compound produced by $F. oxysporum$ is bikaverin. This is the first report of a
CRISPR system that could be efficiently used to generate mutations in genes of interest in a member of the FOSC and may facilitate functional studies concerning the broad host range, secondary metabolite production, and supernumerary chromosomes of these fungi. Importantly, the high degree of similarity of the NLSH2B sequence between all the fusaria and the NLSH2B sequence that was fused to the Cas9 protein suggests this transformation system may be applicable to the other members within the genus *Fusarium*.

**Reference**


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Woo, J. W., et al., 2015. DNA-free genome editing in plants with preassembled CRISPR-Cas9 ribonucleoproteins.
Figure 2.1  Determination of an endogenous NLS from the *F. oxysporum* histone H2B gene. (A) H2B amino acid multiple sequence alignment among *F. oxysporum* (Fo), yeast, *Arabidopsis*, human and *Drosophila*. (B) Subcellular localization of *F. oxysporum* hyphae expressing NLS<sub>H2BeGFP</sub> showing it localizes to the nucleus. Scale bars, 10 μm.
Figure 2.2  Protein purification and *in vitro* cleavage activity assay.

(A) SDS-PAGE gel depicting protein expression and purification conditions. Lane 1 represents total protein before inducing; Lane 2 total protein after addition of 0.3 mM IPTG for 5 h; Lane 3 represents about 5 μg purified NLS$_{H2B}$Cas9 protein. (B) Top: diagram depicting the template length, the sgRNA cleavage site, and the lengths of expected fragments after cleavage with Cas9; Bottom: assessment of the cleavage efficiency by gel electrophoresis after incubation at 37 °C for one hour.
Figure 2.3  Disruption of two *F. oxysporum* genes, *URA5* and *URA3*, using the optimized Cas9 RNP transformation system.

(A-C) characterization of individual NHEJ-mediated mutants (A) The multiple sequence alignment of target regions of seven randomly chosen *ura5* mutants is shown. (B) DNA sequences of the wild-type isolate and three *ura5* mutants. (C) The multiple sequence alignment of target regions of two randomly chosen *ura3* mutants is shown. (D) *URA5* homologous-directed repair disruption chart depicting insertion site of hygromycin phosphotransferase.
Figure 2.4  The phenotypes of *ura5* and *ura3* mutants obtained by NHEJ and HDR methods.

About 200 conidia were positioned onto the center of the plates, which were cultured at room temperature. The phenotypes were evaluated after six days. MUU represents M100 medium supplemented with uracil and uridine. The concentrations of 5-FOA and hygromycin are 1.5 g/L and 100 μg/mL, respectively.
Figure 2.5  *FoBIK1* gene encoding a polyketide synthase in a secondary metabolite biosynthetic cluster is the core enzyme responsible for the biosynthesis of a red pigment. (A) The *BIK1* gene structure, the location of the sgRNA in exon 1, and the construction of the associated HDR template. (B) The cleavage efficiency of the sgRNA for *bik1* *in vitro* after incubation at 37 °C for one hour. (C) The phenotypes of three replicative wild type and six randomly chosen *bik1* mutants cultured for seven days. (D) HPLC-DAD analysis of red pigment production between wild type and *bik1* mutants.
Figure 2.6  Schematic diagram of Cas9 RNPs-directed delivery method to *F. oxysporum* protoplasts to efficiently generate gene disruptions/deletions.
**Figure 2.7** Histone H2B amino acid sequence alignment of different *Fusarium* species. Abbreviations for specific species are as follows: Fo, *F. oxysporum*; Fg, *F. graminearum*; Fv, *F. verticillioides*; Ff, *F. fujikuroi*; Fa, *F. acuminatum*; Fs, *F. solani*; Fpse, *F. pseudograminearum*; Flan, *F. langsethiae*; Fman, *F. mangiferae*; Fpro, *F. proliferatum*; and Fpoa, *F. poae*. 
Figure 2.8  Analysis of the compounds in the supernatant of wild type (WT) and a ΔFobik1 transformant by mass spectrometry.

The major peak found in the WT at 383.0722 corresponds to bikaverin and is absent in the Fobik1 mutant profile.
Table 2.1 Mutation frequencies of homolog directed repair using Cas9 RNPs

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<th>Cas9/sgRNA&lt;sup&gt;a&lt;/sup&gt;</th>
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<th>Positive No.</th>
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<sup>a</sup> represent that the unit is µg
### Table 2.2 The sequences of oligonucleotides used in Chapter 2

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Chapter 3.

CRISPR/Cas9-mediated endogenous gene tagging in *Fusarium oxysporum*

This chapter has been published (Wang and Coleman, 2019).

3.1 Introduction

Members of the *Fusarium oxysporum* species complex (FOSC) are soil-borne pathogens which pose a serious threat to agricultural production and human health (Gordon, 2017). In agriculture, *F. oxysporum* is capable of causing several diseases, most notably Fusarium wilt resulting from colonized plant xylem tissue. Infected plants show a range of symptoms including chlorosis, necrosis, stunting, and wilting (Gordon, 2017). In addition, these fungi can be opportunistic human pathogens infecting immunocompromised patients with high mortality rates (Muhammed et al., 2013; Nucci and Anaissie, 2007).

Rapid improvement of high-throughput DNA and RNA sequencing techniques has facilitated researchers to investigate fungal pathogenicity based on a large scale of whole genome and/or transcriptome analysis. Currently, several high-quality, well-annotated *F. oxysporum* genomes are publicly available, and enable further exploration of fungal pathogenesis and disease control (Ma et al., 2010; van Dam et al., 2017). Experimental evidence has shown that horizontal transfer could have contributed to the genetic diversity
among FOSC isolates, as mobile chromosomes harboring virulence factor genes were able to expand the host range of the recipient and increase their environmental nich(es) (Ma et al., 2010; Vlaardingerbroek et al., 2016).

Molecular manipulation of genes and proteins are essential to fully understand the role they have in an organism, and some of the more common techniques to elucidate cellular functions include gene deletion, RNAi knockdown, ectopic gene expression, and protein subcellular localization. The subcellular localization of a protein influence on the function, as improperly localized proteins, including those that have lost their subcellular localization peptides, may result in decreased or abolished protein activity (Scott et al., 2005). Several studies have indicated that cellular location of a fungal protein is critical for proper function inside the cell. For example, the well characterized calcineurin-regulated transcription factor Crz1 which ultimately controls transcription of several stress response genes and cell wall integrity in most fungi (Lev et al., 2012; Schumacher et al., 2008; Spielvogel et al., 2008). Upon an increase in intracellular calcium levels, Crz1 is dephosphorylated resulting in rapid localization into the nucleus activating the expression of downstream genes.

Studies of fungal protein subcellular localization may rely on several methods, where the most commonly employed method is the use of fluorescent markers including GFP, eGFP, RFP, YFP, and mCherry. These fluorescent proteins are usually fused in frame with the coding sequence of the gene of interest and randomly inserted into the fungal genome under the control of a constitutive or native promoter (Gupta et al., 2015; Schuster et al., 2016). Based on different excited marker fluorescence, protein subcellular localization can be determined and protein translocation can be dynamically tracked (Schuster et al., 2016). While this method
enables the protein subcellular localization to be detected or translation of the protein monitored, some limitations still exist including: 1) native promoters may not have been accurately predicted upstream of the coding regions, resulting in no or altered expression of the protein fused with tags; 2) protein expression that is driven by constitutive promoters might cause inaccurate protein subcellular localization, as constitutive expression may disrupt post-translational regulation; 3) ectopic insertion may disrupt other genes or important regions in the fungal genome, resulting in inaccurate protein subcellular localization or dynamic gene expression, necessitating multiple transformants to be assessed. While antibody-based immunofluorescence approaches to determine the subcellular localization of proteins exist (Rasmussen and Glass, 2005), the difficulty in obtaining an antibody with high specificity, compounded by the time required to generate the antibody, have limited the application of this approach.

As molecular methods to generate gene fusions are an important component for functional studies, a rapid and efficient endogenous gene tagging system (EGT) in *F. oxysporum* was developed based on a CRISPR/Cas9 ribonucleoprotein complex transformation method (Wang et al., 2018). With this system, we could not only localize the protein encoded by the endogenous genes of interest, but also detect the relative protein abundance at various fungal structures. Different fluorescent markers were integrated prior to the start/stop codon of the gene, allowing gene fusions to be generated with the gene under its native promoter, at the usual location within the genome, and containing the gene introns. Two strategies, a homology-independent targeted integration (HITI) and homology-dependent recombination integration (HDRI), were used in the development of the EGT system with
fluorescent markers including \textit{sGFP} (a variant of \textit{eGFP}) and \textit{mCherry}. The HITI strategy was initially developed based on the non-homologous end joining (NHEJ) DNA repair approach (Suzuki et al., 2016). We used this strategy to tag endogenous genes at either the N-terminus or the C-terminus, and generated FoChs5-3\times sGFP (C-terminus tagging) and mCherry-FoSso2 (N-terminus) transformants. The protein subcellular localization showed FoChs5 in \textit{F. oxysporum} was present in both conidia and hyphae where a majority accumulated at the hyphal tips although some existed in the septum. FoChs5 was largely found on the surface of newly formed germling tubes, suggesting Chs5 plays an important role in cell wall modelling. The HDRI strategy was used to tag the plasma membrane syntaxin 1 SNARE protein FoSso1 with a mCherry marker at the C-terminus. FoSso1-mCherry was localized to the cell plasma membrane (PM) of hyphae and the septum in conidia while mCherry-FoSso2 showed strong PM localizations in conidia, germlings, fungal tips, hyphae, and phialides. Overall, this CRISPR/Cas9 mediated EGT system was highly efficient and stable for use in protein subcellular localization studies and could provide a genetic engineering reference for gene tagging in other filamentous fungi.

3.2 Materials and methods

3.2.1 Strains and culture conditions.

The \textit{Fusarium oxysporum} f.sp. \textit{vasinfectum} strain (FGSC #10442) was used in this study (McCluskey et al., 2010), and was cultured in PDB (Potato Dextrose Broth), YG (1% yeast extract, 3% glucose) or M-100 (minimum medium). All transformants in this study were isolated on M-100 agar plates containing 75 \(\mu\)g/mL hygromycin B before observation by
confocal microscopy. All transformed strains generated in this study are listed in Table 3.1.

3.2.2 Homologous gene identification.

Two well-annotated CHS5 genes from Neurospora crassa (NCU04352) and Ustilago maydis (UM03204.1) were used as query sequences to identify the Chs5 homologous sequence in F. oxysporum with blastp software in the EnsembleFungi database (https://fungi.ensembl.org/index.html). Several previously identified syntaxin 1 SNARE proteins from Saccharomyces cerevisiae (SSO1: YPL232W and SSO2: YMR183C), N. crassa (SSO1: XP_961235.1) and Zymoseptoria tritici (SSO1: XP_003857391.1) were used as the blastp queries against the homologous sequences in the above database. Since these genes perform similar functions during fungal development, their amino acid sequences were hypothesized to be conserved. To determine the sequence identity or splice variants, the potential homologous genes were compared with the associated genes in closely-related filamentous fungal pathogens including Fusarium oxysporum f. sp. lycopersici 4287, Magnaporthe oryzae 70-15, Fusarium graminearum PH1, and Fusarium solani (Nectria haematococca MPVI 77-13-4), which had well-annotated genomes. The protein domains were predicted using PFAM (Finn et al., 2016). Multiple sequence alignments were conducted using the online bioinformatic program TCOFFEE (Notredame et al., 2000). The phylogenetic tree was generated using online PhyML 3.1 with the maximum likelihood algorithm and 500 bootstraps (http://www.phylogeny.fr/one_task.cgi?task_type=phylm) (Dereeper et al., 2008). All accession numbers of predicted genes in this study are listed in Table 3.1.

3.2.3 The evaluation of fusion protein topology and plasmid construction.

Since the integrated location of the different fluorescent protein markers may alter the
original protein topology, the TMHMM v. 2.0 analysis software (http://www.cbs.dtu.dk/services/TMHMM/) was used to compare the differences of protein topology between the WT proteins and the fusion proteins. All plasmids were maintained, propagated, and stored in DH5α E. coli. All tagging plasmid constructs were based on a pUC19 plasmid, and cloned fragments were inserted between two enzyme sites, BamHI and HindIII, with the NEBuilder® HiFi DNA Assembly Master Mix kit (NEB). For the CHS5 gene tagging vector construction (HITI), the initial vector was first constructed by amplifying the following three fragments by PCR: 1) 1× sGFP from the plasmid pCT74 (Lorang et al., 2001), 2) the Tef-1 terminator from the plasmid pFFC332, and 3) the hygromycin cassette from the plasmid pCWHyg1. These three fragments were simultaneously inserted between the BamHI and HindIII sites in the pUC19 vector in the following order 1×sGFP-Tef-1-HYGB generating the primary vector pUC19-1×sGFP-Tef-1-HYGB. The final CHS5 HITI vector was constructed as follows: 1) a ~700 bp coding sequence prior to the stop codon of FoCHS5 was cloned as a bait sequence which contained a sgRNA cleavage site; 2) to construct a triple sGFP in the vector, the first copy of sGFP (1st sGFP) was cloned with the addition of a 3′-terminus marker sequence (nt: GACTACAAGGACGAC, aa: DYKDD) while the second copy of sGFP (2nd sGFP) was cloned with the addition of the 3′-terminus marker sequence (nt: AGCGCTGGCGCTTAC, aa: SAGAY); 3) the fragment of 1×sGFP-Tef-1-HYGB was cloned from the above primary plasmid; and 4) these four fragments were simultaneously inserted into the pUC19 plasmid to generate the final HITI plasmid containing the triple sGFP tag, which was named as pUC19-HITI-FoCHS5-3×sGFP. The detailed N-terminal FoSSO2 gene tagging vector for HITI was constructed as follows: 1) a ~800 bp sequence in front of the FoSSO2 gene start codon was
cloned with a sgRNA site; 2) 1 × mCherry was cloned from plasmid ss3591; 3) the FoSSO2 coding sequence including introns and a ~450 bp 3-UTR region were amplified from genomic DNA; 4) the hygromycin cassette was obtained from plasmid pCWHyg1; 5) the above four fragments were simultaneously inserted into the pUC19 plasmid, which was later labeled as pUC19-HITI-mCherry-FoSSO2.

The FoSSO1 C-terminal tagging plasmids contained a mCherry tag fused with a Tef-1 terminator and hygromycin resistance cassette. The detailed HDRI vector construction was carried out as follows: 1) a ~400 bp DNA template fragment prior to the gene stop codon was cloned from the genomic DNA (upstream sequence); 2) the other DNA fragment after the sgRNA cleavage site, ~400bp in length, was obtained using PCR (downstream sequence); 3) the mCherry was cloned from the plasmid ss3591 and the Tef-1 terminator fused with hygromycin cassette was cloned from pUC-HITI-FoCHS5-3×sGFP as constructed above; 4) these four fragments were assembled in the following order, upstream sequence, mCherry, Tef-1 fused with the hygromycin cassette, and downstream sequence, into pUC19 (pUC19-HDRI-FoSSO1-mCherry). All primers used in this study are listed in Table 3.3.

3.2.4 sgRNA selection and in vitro Cas9 nuclease assay.

The sgRNA sites were located after the stop codon between 0 and 200 bp for the HDRI strategy (FoSSO1-mCherry), while those for C-terminal tagging of the FoCHS5 gene and N-terminal tagging of the FoSSO2 gene were located prior to the gene stop or start codon, respectively. A ~2kb fragment was cloned for use as a template for the in vitro Cas9 nuclease assay to confirm the Cas9 endonuclease cleavage activity. The in vitro Cas9 nuclease assay was performed as previously described (Wang et al., 2018). In brief, a mixture of 200 ng Cas9, ~40
ng sgRNA, 1×Cas9 Nuclease Reaction Buffer and ~75 ng DNA template in a 20 μL total volume was incubated at 37°C for 1 h. The cleaved products were evaluated on a 1% agarose gel. All sgRNA were able to effectively cleave PCR fragments with the Cas9 nuclease (Figs. 3.5C and 3.6B).

3.2.5 Fungal transformation and transformant identification.

Transformation of *F. oxysporum* was carried out as previously described (Wang et al., 2018). For the HDRI strategy, donor PCR fragments were amplified from the donor plasmids (pUC19-HDRI-FoSSO1-mCherry and pUC19-HDRI-FoSSO2-mCherry) with the primers, NA_FoSSO1upF/NA_FoSSO1downR and NA_FoSSO2upF/NA_FoSSO2downR, and used to conduct the transformations. The desired transformants will contain the gene of interest fused with the different marker tags under the native promoter, and were screened by three PCR reactions to determine the location of the integrated fragment in the fungal genome. Detailed illustrations of correct transformants are shown in Fig. 3.5A and 3.6A. The first pair of primers, Tf and Tr, were used to assess the full length near the sgRNA cleavage site after the fragments were integrated. Another two pairs of primers were used to determine the upstream and downstream sequences of the inserted fragments. All primers used are listed in Table 3.3.

3.2.6 Confocal microscopy observation.

Transformant hyphae were picked and inoculated into 50 mL YG media and cultured for 4 days at room temperature on a rotary shaker at 150 rpm. The 500 μL hyphae-conidia mixture of the *F. oxysporum* transformants were inoculated into 50 mL fresh YG media and incubated on a shaker at 18°C at 220 rpm for 16 h. One mL of the fungal culture was centrifuged at 13,000 rpm for 1 min and the supernatant removed. The resulting pellet was suspended in 500 μL of
4% paraformaldehyde and the mixture was incubated at room temperature at least 15 min, followed by at least two washes of PBS, pH 7.4. Transformants were suspended in a final volume of 300 μL of PBS, pH 7.4 and imaged by confocal microscopy using an inverted agar method under 60 × oil magnification (Nikon A1 Confocal Microscopy) (Hickey and Read, 2009). For live cell imaging, the fungus was cultured in a four-chambered coverglass (Thermo Scientific™ Nunc™ Lab-Tek™ Chambered Coverglass) or the live fungal cells were dispensed on the surface of a 0.5 cm × 0.5 cm piece of agar on a slide and covered with the coverglass. The sGFP fluorescent marker was excited at 488 nm while the mCherry fluorescent marker was excited at 561 nm.

3.3 Results and Discussion

3.3.1 Identification of FoChs5, FoSso1, and FoSso2

Orthologs to several well-characterized genes in other filamentous fungi were selected in *F. oxysporum* to develop an EGT system to study subcellular localization. The well characterized localization of the transmembrane chitin synthases (CHSs) and their importance in production of the fungal cell wall component chitin, make this group of enzymes ideal to use in the development of an EGT system. Nine CHS genes reside in the *F. oxysporum* genome and the presence of multiple copies of CHSs suggest functional complexity exists in regards to chitin biosynthesis in these fungi (Kong et al., 2012). Among these CHSs, the class V CHS (CHS5) usually contain several protein domains including a Myosin head, Cyt-b5, Chitin_synth_2, and DEK C domains (Fig. 3.1A); importantly, this class of CHS enzymes have been shown to be involved in pathogenesis and vesicle transportation (Madrid et al., 2003;
Among these domains, the myosin domain is involved in traversing on actin filaments and microtubules while the chitin synthase domain participates in chitin synthesis (Schuster et al., 2016; Treitschke et al., 2010). To characterize the subcellular localization of FoChs5 in different fungal structures and stages of development, the orthologous gene (FOTG_09933) in the *F. oxysporum* genome was identified based on the *CHS5* genes from other fungi. FoChs5 shares 83.3% and 52.8% amino acid similarity to Chs5 in *N. crassa* and Mcs1 in *U. maydis*, respectively; supporting Chs5 performs similar functions among these fungi.

In addition to *FoCHS5*, we chose the duplicated genes *SSO1* and *SSO2*, that encode homologues of syntaxin 1, as representative genes to develop this EGT system based on their known localization to membranes (Kienle et al., 2009). Syntaxins are Qa SNARE transmembrane proteins that play essential roles in membrane fusion events (Salaun et al., 2004). In the yeast *S. cerevisiae*, the *SSO1* and *SSO2* genes are involved in the fusion between secretory vesicles and the plasma membrane. Deletion of *SSO1* halted vesicle fusion and decreased sporulation efficiency, while mutation of *SSO2* resulted in increased sensitivity to chemicals and temperature (Jantti et al., 2002; Nakanishi et al., 2006). Syntaxin 1 and its homologues contain, in their cytosolic region, an N-terminal syntaxin domain followed by a TMD-proximal SNARE domain (Fig. 3.1A), and the syntaxin Sso1 has been used for characterization of the plasma membrane in some fungi (Kilaru et al., 2017; Schuster et al., 2016; Taheri-Talesh et al., 2008; Valkonen et al., 2007). To identify the homologous genes of syntaxin 1, several previously described syntaxin 1 genes from *S. cerevisiae, U. maydis*, and *Z. tritici* were used as a query for the *FoSSO1* and *FoSSO2* gene sequences (Kilaru et al., 2017).
FoSso1 and FoSso2 contained the two typical domains (syntaxin and SNARE) and shared ~44% amino acid sequence similarity between the two proteins. Although some strains of *Aspergillus* species only contain a single copy of the SSO genes (Kienle et al., 2009), phylogenetic analysis revealed that a majority of the ascomycete fungi encode two copies of the SSO genes, and can be resolved into two distinct phylogenetic clades suggesting that these two genes diverged some time ago and have since evolved to function differently (Fig. 3.1B). Both copies of the SSO genes were highly conserved among *Fusarium* spp. and were clustered in the phylogenetic tree supporting that the presence of two SSO genes is ancestral in this genus.

3.3.2 HITI-mediated endogenous gene tagging at the C-terminus of *FoCHS5*

A homology-independent targeted integration (HITI) strategy was developed utilizing CRISPR/Cas9 technology (Wang et al., 2018), which enables long DNA fragments to be inserted into the genome using the NHEJ repair mechanism. The main advantage of HITI is that it can occur in non-dividing cells, and has been validated in many cell types including human cell lines and zebrafish (Auer et al., 2014; Suzuki et al., 2016). The HITI strategy was used to insert a 3×sGFP (a variant of eGFP) tag before the *FoCHS5* stop codon (C-terminal tagging), generating a fusion protein consisting of FoChs5 and a strong green fluorescent signal for protein subcellular localization. The donor plasmid DNA was composed of the following: a ~700 bp homologous fragment containing the sgRNA cleavage site before the *FoCHS5* stop codon was fused with a 3×sGFP sequence, a *Tef-1* terminator sequence, and the hygromycin resistance cassette sequence (Fig. 3.2A). During the transformation, both the genome and donor plasmid sequences can be simultaneously cleaved by the Cas9 RNPs. The activated DNA repair
mechanism allows “cross-ligation” between the chromosome and the donor plasmid at some frequency (C1--P2 and P1--C2, Fig. 3.2A). The desired transformants will contain the complete \textit{FoCHS5} endogenous gene sequence fused in frame with the 3\times s\textit{GFP} tag followed by the remaining portion of the donor plasmid containing the hygromycin resistance cassette for selection of the desired transformants (Fig. 3.2A).

Previous evidence showed the HITI strategy allowed perfect repair of DNA at a higher frequency, and that error-prone DNA repair occurred infrequently (Auer et al., 2014; Suzuki et al., 2016). We hypothesized if the Cas9 RNP transformation of \textit{F. oxysporum} protoplasts only allowed a miniscule amount of the Cas9 RNPs into the cell and the Cas9 RNPs carried out the cleavage a short time before their degradation, some proportion of the transformants would have perfect DNA repair occurring between the cleavage sites of the chromosome and donor plasmid. After transformation, two of the four transformants were used for PCR analysis to confirm that the whole plasmid (~ 8 kb) was inserted into the genome at the sgRNA cleavage site (Fig. 3.5D). In theory, the HITI-mediated gene editing may generate short losses or additions of nucleotides at the repair sites (Auer et al., 2014; Suzuki et al., 2016). However, sequencing of the sgRNA target sequence site of the two transformants showed identical nucleotide sequence with the wide-type strain (Fig. 3.5B), demonstrating that the NHEJ DNA repair mechanism for some \textit{F. oxysporum} protoplast cells would be error free, which was consistent with previous publications (Auer et al., 2014; Suzuki et al., 2016).

The subcellular localization of FoChs5 revealed that the protein had variable distribution and accumulated in some areas (Fig. 3.2B). In conidia, FoChs5-3\times s\textit{GFP} accumulated at both ends of the spore; however in \textit{F. oxysporum} germlings FoChs5-3\times s\textit{GFP} was distributed in the
cell membrane primarily at the newly formed hyphal tips emerging from the conidium showing that FoChs5, as a class V myosin chitin synthase, was strongly polarized. In addition, confocal microscopy showed that FoChs5 localized to the hyphal tips and septa (Mov. 1, Fig 3.7). As FoChs5 is polarized and contains a myosin motor domain, it may have a similar function with *U. maydis* Mcs1 and participate in the transportation of vesicles (Treitschke et al., 2010).

3.3.3 HDRI-mediated endogenous gene C-terminal tagging of *FoSSO1*

Our previous study showed Cas9 RNPs significantly improved the efficiency of homologous recombination (Wang et al., 2018). *FoSSO1* was tagged with a *mCherry* peptide at the 3′-terminus of the genes by a homology-dependent recombination integration (HDRI) strategy. The sgRNA site was located after the gene stop codons (Fig 3.3A). The donor DNA was constructed by the following: an upstream homologous fragment, ~400 bp in length, prior to the gene stop codon was cloned and fused with the mCherry-*Tef-1-HYGB* and a ~400 bp downstream homologous fragment after the sgRNA cleavage site. During the transformation with the donor DNA fragments and Cas9 RNPs, two homologous recombination events were expected to occur at the sgRNA cleavage site (Fig. 3.2B). The desired transformants will express the Sso-mCherry fusion protein and can be selected for by hygromycin resistance.

Since the transformation efficiency was significantly increased with the CRISPR/Cas9 RNPs (Wang et al., 2018), three potential *FoSSO1-mCherry* transformants were randomly selected for further analysis. All of the transformants possessed the desired fusion (Fig. 3.6C) and fluorescence signal when excited at a wavelength of 561 nm under confocal microscopy. The previous study showed when the donor plasmid was directly used for the protoplast
transformation, the maximum correct integrated frequency was ~50% (Wang et al., 2018). However, all our selected HDRI transformants contained the correctly integrated fragments at the expected sites, suggesting that PCR fragment transformation might have a higher integration efficiency in F. oxysporum than transformation with plasmid donors. In conidia, FoSso1-mCherry primarily accumulated at the septum and the cell plasma membrane (Fig 3.3B, a and b). FoSso1-mCherry had no specific accumulation except for the septum in the germinating conidia and showed a weak signal in the newly-formed hyphal tip (Fig 3.3B, c and d). Conversely, FoSso1-mCherry accumulated in well-formed hyphae distant from the fungal tips, instead of newly formed hyphae and fungal tips (Fig 3.3B e and f). To further confirm the FoSso1-mCherry subcellular localization results, the Z-series scanning function on the confocal microscopy was used to observe the fungus at a fluorescent range of 3.2 μm with a step of 0.4 μm (Mov. 2, Fig 3.8). The Z-series stack indicated that all three fungal tips showed weak fluorescent signals while the septa and hyphae distal from the fungal tips had strong fluorescence accumulation.

3.3.4 HITI-mediated endogenous gene tagging at the N-terminus of FoSSO2

Previous studies have tagged SSO2 orthologs at the N-terminus to avoid the fluorescent protein residing in the lumen of exocytic vesicles, which can result in improper folding of the protein tag (Taheri-Talesh et al., 2008; Valkonen et al., 2007). To further confirm the subcellular localizations of FoSso2 proteins, we conducted with N-terminal tagging to FoSSO2 gene using the HITI strategy. The donor DNA was composed of a ~800 bp fragment prior to the FoSSO2 start codon containing a sgRNA cleavage site which was fused with the mCherry coding
sequence fused in frame with the FoSSO2 coding sequence, including the introns and the 3′-UTR sequence, followed by the hygromycin resistance cassette (Fig 3.4A). During the transformation the entire plasmid is intended to be integrated at the sgRNA cleavage site, and the desired transformants will contain an intact endogenous FoSSO2 gene fused with a mCherry marker gene at the N-terminus (Fig 3.4A). The number of transformant colonies generated using the Cas9 RNPs was much greater than those without the Cas9 RNPs (Fig 3.4B, Table 3.2), indicating that the addition of the Cas9 RNPs significantly enhanced transformation efficiency. A total of 49 independent colonies were selected and screened for detection of mCherry fluorescence, and only five transformants failed to produce any detectable fluorescent signal under confocal microscopy (positive fluorescent efficiency was ~90% (44/49)). Nine randomly selected mCherry-FoSSO2 transformants were further assessed for precise integration at the expected site in the genome by PCR results (data not shown) and sequencing the locus. Sequencing revealed eight of the transformants were correct, having the desired mCherry-FoSSO2 gene fusion, while one transformant contained a 3-bp nucleotide deletion at the sgRNA cleavage site, although the resulting transformant was still fluorescent.

The mCherry fluorescent marker for FoSso2 indicated that the Qa SNARE protein was localized to the fungal plasma membrane (Fig 3.4C). To further investigate the mCherry-FoSso2 plasma membrane localization, confocal microscopy Z-series stacks of live cells were used to further observe key structures involved in F. oxysporum conidiation (Fig 3.4C d, Fig 3.9). mCherry-FoSso2 was localized to an extensive array of fungal structures including conidia, phialides, septa, germlings, hyphae, and fungal tips (Fig 3.4C). The localization of FoSso2 to the plasma membrane of a broad range of fungal structures indicates it may serve as
a better marker for the characterization of the plasma membrane than FoSso1.

Clearly the differences in subcellular localization between FoSso1 and FoSso2 indicate divergence during their evolution. While FoSso1-mCherry showed a weak fluorescent signal in germlings and at subapical regions of the hyphae, mCherry-FoSso2 showed strong plasma membrane localization in conidia, fungal germlings, and in hyphae at the apical fungal tips. Qa SNARE proteins encode a single type II transmembrane domain which allows it to be localized to the plasma membrane (Sharpe et al., 2010). Previously, the single Qa SNARE protein SSOA from Aspergillus nidulans was shown to localize to the plasma membrane of the hyphal tip with some accumulation at the apical plasma membrane (Taheri-Talesh et al., 2008). Alternatively, in the fungus Trichoderma reesei the site of SNARE complex formation between SNCI and SSOI or SSOII was spatially segregated (Valkonen et al., 2007). Both Qa SNARE proteins from T. reesei was localized to the plasma membrane; however the SNCI-SSOII complexes were mainly localized to the growing hyphal apical regions while SSOI and SNCI co-localized to subapical regions.

This EGT system is a rapid and efficient method for fungal protein subcellular localization studies. There are several advantages of this system over traditional methods including: 1) conventional methods for protein subcellular localization in fungi required cloning and fusing the promoter region with the gene coding region and fluorescent marker sequences (Kilaru et al., 2015). However, in this CRISPR/Cas9-based EGT system, one or two short homologous fragments around the sgRNA sites were cloned. This avoids the usually time-consuming process to clone large gene coding regions such as the FoCHS5 gene from cDNA; 2) this method does not require the promoter region of a gene to be predicted to generate a construct,
which avoids inaccurate promoter prediction or incorrect expression; 3) the HITI strategy allows a large DNA fragment to be integrated into the fungal genome. This is a potential advantage to simultaneously expressing multiple genes from plasmids in fungi, to be tested in the future; 4) the EGT system could accurately insert DNA fragments at the desired site(s) in the genome, and has a clear advantage over traditional transformation methods since it limits the frequency of random insertion; 5) the desired transformant(s) can be obtained from a small quantity of transformants due to the high efficiency of the system. Despite these advantages, using this EGT system might be difficult for genes with low expression under the native promoter due to low fluorescent signal detection during microscopy.

In this study, we used two different strategies for tagging endogenous genes in situ with fluorescent markers in the plant pathogenic filamentous fungus *F. oxysporum*. Both methods allowed the gene of interest to be expressed with a fluorescent signal under the control of the native promoter. The HITI strategy allowed a long DNA fragment to be inserted into the fungal chromosome and was used for the C-terminal tagging of *FoCHS5* and the N-terminal tagging of *FoSSO2*. In addition, the other Qa SNARE syntaxin 1 homologous gene, *FoSSO1*, was tagged by an HDRI strategy. All these tagging transformants carried the desired fluorescent signals. Overall, this CRISPR/Cas9 RNP-mediated transformation method for EGT may facilitate the study of genetic engineering and the elucidation of gene function(s) in *F. oxysporum* and other fungi.


Lev, S., et al., 2012. The Crz1/Spl transcription factor of Cryptococcus neoformans is activated by calcineurin and regulates cell wall integrity. PLOS ONE. 7, e51403.


Figure 3.1 Identification of FoChs5, FoSso1 and FoSso2 proteins in *F. oxysporum*.

(A) The protein domain structures of the chitin synthase FoChs5, and the two syntaxin proteins FoSso1 and FoSso2. (B) The maximum-likelihood phylogenetic tree of Sso1 and Sso2 amino acid sequences from selected filamentous ascomycete fungi. The tree was generated with a bootstrap value of 500.
Figure 3.2 *FoCHS5* gene C-terminal tagging with a $3\times sGFP$ fluorescent marker based on the HITI strategy.

(A) The homology-independent targeted integration (HITI) strategy scheme for the C-terminal tagging of *FoCHS5* gene. A bait sequence, ~700 bp in length containing the sgRNA cleavage site, was cloned prior to gene stop codon and fused with a $3\times sGFP$ marker and the hygromycin resistance cassette sequences. During transformation, two DNA cross–ligation events will occur between the donor plasmid and the fungal chromosome (C1-P2 and P1-C2). The whole plasmid will be inserted into the sgRNA cleavage site on the fungal chromosome in the desired transformants. The C1-C2 fragment is homologous to the P1-P2 fragments. B) *FoChs5*-3×sGFP subcellular localization in different fungal structures. a and b: conidia, c and d: conidia with the emerging germination tubes, e: fungal tips, f: hyphae. Scale bars, 10 μ
Figure 3.3  FoSSO1 gene C-terminal tagging with a mCherry fluorescent marker based on the HDRI strategy.

(A) An illustration of the homology-dependent recombination integration (HDRI) strategy for the C-terminal tagging of FoSSO1 and FoSSO2 genes. A homologous upstream fragment was amplified prior to the gene stop codon and fused with mCherry, the hygromycin resistance cassette, and a homologous downstream fragment that is immediately after the sgRNA cleavage site. During the transformation, two homologous recombination events will be expected to occur and mCherry-Tef-1-HYGB fragment will be integrated at the 3′-terminus of the endogenous gene. (B) the FoSso1-mCherry subcellular localization in different fungal structures. a and b: conidia, c and d: germlings, e and f: fungal tips and hyphae. Gradient red arrows weaken from strong to weak, suggesting different distribution of the FoSso1 protein. Scale bars, 10 μm.
Figure 3.4  *FoSSO2* gene tagging with a N-terminal *mCherry* fluorescent marker based on the HITI strategy.

(A) A ~800 bp homologous sequence containing the sgRNA cleavage site in front of the *FoSSO2* gene start codon was cloned and fused with a 1×*mCherry* marker, *FoSSO2* endogenous gene, a 3′-UTR region of the *FoSSO2* gene and the hygromycin resistance cassette sequences. During the transformation, the whole plasmid can be inserted into the sgRNA site and a N-terminal *mCherry* fusion gene for the *FoSSO2* can be generated. (B) Comparisons of the transformation efficiency with or without Cas9 RNPs for the *FoSSO2* N-terminal tagging. (C) *mCherry*-FoSso2 subcellular localization in different fungal structures. a: conidia, b: fungal tips, c: a germling, d: fungal conidiation, *: phyllides.
Figure 3.5  Screen of the correct HITI transformants for generation of FoCHS5-3×sGFP.  
(A) the illustration of the chromosomal region of a correct HITI transformant after the integration of the entire plasmid. Tf and Tr: a pair of primers located outside of the bait sequence and can be used to clone a ~2.1 kb fragment, from the wild-type strain DNA, which was used for the in vitro Cas9 nuclease assay. Three PCR products, Up, Down and Full, were used to confirm the location of the integrated plasmid at the correct target site. (B) The sequencing results of DNA fragments containing the sgRNA sites for the wild-type strain and two transformants. (C) In vitro Cas9 RNPs cleavage assay to assess sgRNA target sequence. (D) agarose gel electrophoresis for the three DNA fragments based on the three pairs of primers from (A).
Figure 3.6  Screen of the correct HDRI transformants for generation of *FoSSO1-mCherry* (A) the illustration of the chromosomal region for correct HDRI transformants after homologous recombination. Tf and Tr: a pair of primers located outside of the homologous arms and can be used to amplify a fragment from the wild-type strain DNA for the *in vitro* Cas9 nuclease assay. Three fragments, Up, Down and Full, were used to determine the location of the integrated fragment at the desired positions. The two orange arrows represent different primers located in the region of the *mCherry-Tef-1-HYGB*. B) *In vitro* Cas9 RNPs cleavage assay for the *FoSSO1* gene. C) The results of agarose gel electrophoresis for three DNA fragments based on three pairs of primers from A).
**Figure 3.7**  Each frame of the FoChs5-3×sGFP Z-series stack
Figure 3.8  Each frame of the FoSso1-mCherry Z-series stack
Figure 3.9  Each frame of the mCherry-FoSso2 Z-series stack
Table 3.1 The sgRNAs and associated transformants of *F. oxysporum* in Chapter 3

<table>
<thead>
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<th>sgRNA</th>
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<th>Targeted gene</th>
<th>Gene No.</th>
<th>Genotype of transformants</th>
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<td>AGG</td>
<td><em>FoCHS5</em></td>
<td>FOTG_09933</td>
<td><em>FoCHS5</em>-3×3GFP&lt;sup&gt;EGT&lt;/sup&gt;, hyg&lt;sup&gt;R&lt;/sup&gt;</td>
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<td>ATCTCAGCG</td>
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<td>GAGCATCAAACA</td>
<td>CGG</td>
<td><em>FoSSO1</em></td>
<td>FOTG_02772</td>
<td><em>FoSSO1</em>-mCherry&lt;sup&gt;EGT&lt;/sup&gt;, hyg&lt;sup&gt;R&lt;/sup&gt;</td>
</tr>
<tr>
<td>TCATAACAG</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GATCAAAGAGA</td>
<td>GGG&lt;sup&gt;RCS&lt;/sup&gt;</td>
<td><em>FoSSO2</em></td>
<td>FOTG_04307</td>
<td>mCherry-<em>FoSSO2</em>&lt;sup&gt;EGT&lt;/sup&gt;, hyg&lt;sup&gt;R&lt;/sup&gt;</td>
</tr>
<tr>
<td>AGAATGATC&lt;sup&gt;RCS&lt;/sup&gt;</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

*hyg*<sup>R</sup>, hygromycin B resistance; *EGT*, endogenous gene tagging; *RCS*: reverse complement sequence.
Table 3.2 The transformant amounts of the N-terminal tagging of *FoSSO2* gene

<table>
<thead>
<tr>
<th>Transformation plate</th>
<th>Cas9 RNP</th>
<th>Transformation amounts</th>
</tr>
</thead>
<tbody>
<tr>
<td>FoSSO2_C1</td>
<td>-</td>
<td>7</td>
</tr>
<tr>
<td>FoSSO2_C2</td>
<td>-</td>
<td>4</td>
</tr>
<tr>
<td>FoSSO2_T1</td>
<td>+</td>
<td>59</td>
</tr>
<tr>
<td>FoSSO2_T2</td>
<td>+</td>
<td>46</td>
</tr>
<tr>
<td>FoSSO2_T3</td>
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<td>61</td>
</tr>
<tr>
<td>FoSSO2_T4</td>
<td>+</td>
<td>37</td>
</tr>
<tr>
<td>FoSSO2_T5</td>
<td>+</td>
<td>54</td>
</tr>
<tr>
<td>FoSSO2_T6</td>
<td>+</td>
<td>43</td>
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</table>
### Table 3.3 The sequences of oligonucleotides used in Chapter 3

<table>
<thead>
<tr>
<th>Primer names</th>
<th>Oligomer sequences</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>NA_PV_sGFPf</td>
<td>AGTGAATTCGAGCTCGGTACCCGGGA TGGTAGCAAGGGCGAG</td>
<td>The construction of the plasmid primary vector (sGFP - Tef-1 - HYGB)</td>
</tr>
<tr>
<td>NA_PV_sGFPr</td>
<td>ATCGAATGTCGCTTACCTTTGTACACG TCGTCCATG</td>
<td></td>
</tr>
<tr>
<td>NA_PV_Tef1f</td>
<td>GCTGTACAAGTAAGCGGACATTCGAT TTATGC</td>
<td></td>
</tr>
<tr>
<td>NA_PV_Tef1r</td>
<td>TCCATTAATATATTTGGTGCGATGTG</td>
<td></td>
</tr>
<tr>
<td>NA_PV_HYGBf</td>
<td>ATCATGCCCACCAATATATTAATGGGAAG GGTATATACCACGC</td>
<td></td>
</tr>
<tr>
<td>NA_PV_HYGBr</td>
<td>AACAGCTATGACCATGATTACGCCAA GGGTTGCGAGGTCCAATG</td>
<td></td>
</tr>
<tr>
<td>CHS5TF1</td>
<td>CAACGCAAAGGGCTCTTTCA</td>
<td>1. <em>FoCHS5</em> <em>in vitro</em> cleavage template; 2. transformant PCR detection (Full)</td>
</tr>
<tr>
<td>CHS5TR1</td>
<td>CGCACTCGTTTGCTACGTCT</td>
<td></td>
</tr>
<tr>
<td>NA_CHS5hitiF</td>
<td>AGTGAATTCGAGCTCGGTACCCGGGA TGGTAGCAAGGGCGAG</td>
<td>The construction of the plasmid pUC19-HITI- <em>FoCHS5</em>-3×sGFP. (HITI bait -3×sGFP - Tef-1 - HYGB)</td>
</tr>
<tr>
<td>NA_CHS5hitiR</td>
<td>ATCGAATGTCGCTTACCTTTGTACACG TCGTCCATG</td>
<td></td>
</tr>
<tr>
<td>NA_CHS5hiti-3×egfpF1</td>
<td>GAGGAG</td>
<td></td>
</tr>
<tr>
<td>NA_CHS5hiti-3×egfpR1</td>
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</tr>
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<tr>
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</tr>
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<tr>
<td>NA_CHS5hiti-3×egfpF2</td>
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<td>NA_CHS5hiti-3×egfpF3</td>
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<tr>
<td>HyB_R3</td>
<td>TCCTCGGCCCCAAGCATCAG</td>
<td>Detection of Up fragment with TF primer (CHS5TF1 and FoSSO1TF1)</td>
</tr>
<tr>
<td>H_outF</td>
<td>GTCGATGCGACGCAATCGT</td>
<td>Detection of Down fragment with TR primer (CHS5TR1 and FoSSO1TR1)</td>
</tr>
<tr>
<td>FoSSO1TF1</td>
<td>CCCGCCATTGCTTTCTCTGTTT</td>
<td>1. <em>FoSSO1</em> <em>in vitro</em> cleavage template; 2. PCR detection (Full fragment) of the transformants;</td>
</tr>
<tr>
<td>FoSSO1TR1</td>
<td>TTCGTCAAGCCCTGTCCGAT</td>
<td></td>
</tr>
<tr>
<td>NA_FoSSO1upF</td>
<td>AGTGAATTCGAGCTCGGTACCCCGGGC</td>
<td>The construction of the plasmid pUC19-HDRI-FoSSO1-mCherry (up arm - mCherry - Tef-I – HYGB – down arm)</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>NA_FoSSO1upR</td>
<td>GCTATCGCGATCACATTG</td>
<td></td>
</tr>
<tr>
<td>NA_FoSSO1mChH YBF</td>
<td>CACCATACATAAAGGTTGCGAGGT</td>
<td></td>
</tr>
<tr>
<td>NA_FoSSO1mChH YBR</td>
<td>GGCAGG</td>
<td></td>
</tr>
<tr>
<td>NA_FoSSO1downF</td>
<td>ATCCCTGCAACCTTTATGTTTGTG</td>
<td></td>
</tr>
<tr>
<td>NA_FoSSO1downR</td>
<td>ATACCATGGAAATCTCTACTC</td>
<td></td>
</tr>
<tr>
<td>FoSSO2TF2</td>
<td>GGATAGTGCCGGTGATTGCC</td>
<td>1. FoSSO2 N-terminal tagging template for in vitro cleavage; 2. PCR detection (Full fragment) of the transformants;</td>
</tr>
<tr>
<td>FoSSO2TR2</td>
<td>TCCATAACCCTGAGCGG</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2hiti1F</td>
<td>AGTGAATTCGAGCTCGGTACCCCGGGC</td>
<td>The construction of the plasmid pUC19-HITI-mCherry-FoSSO2 (HITI bait -1×mCherry – SSO2 ORF(3-UTR)- HYGB)</td>
</tr>
<tr>
<td>NA_mCS2hiti1R</td>
<td>CATGACGATGCGAACCTTG</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2mcherryF</td>
<td>ATCCCTGCAACCTTTATGTTTGTG</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2mcherryR</td>
<td>ATCCCTGCAACCTTTATGTTTGTG</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2SSO2F</td>
<td>GGATAGTGCCGGTGATTGCC</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2SSO2R</td>
<td>TCCATAACCCTGAGCGG</td>
<td></td>
</tr>
<tr>
<td>NA_mCS2HYGbf</td>
<td>ATCCCTGCAACCTTTATGTTTGTG</td>
<td></td>
</tr>
<tr>
<td>FoCHS5CTsgRNA</td>
<td>AAGCTAATACGACTCTACTATAGGATG</td>
<td>sgRNA DNA template synthesis</td>
</tr>
<tr>
<td>FoSSO1CTsgRNA</td>
<td>TCACGAATCTCAGCGGTTTTTAGAGCT</td>
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</tr>
<tr>
<td>FoSSO2NTsgRNA</td>
<td>AGAAATAGCAAG</td>
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</tr>
</tbody>
</table>


Chapter 4.

The extracellular superoxide dismutase Sod5 from *Fusarium oxysporum* is localized in response to external stimuli and contributes to fungal pathogenicity

4.1 Introduction

The ascomycete fungus, *Fusarium oxysporum* (Fo), is an important pathogenic fungus which can infect and cause disease on a wide range of hosts including plants, animals, and humans (Gordon, 2017). Over 100 formae speciales have been described based on the ability of these fungi to cause disease on different host plants. For example, *F. oxysporum* f. sp. *lycopersici* is regarded as a tomato pathogen causing tomato wilt disease while *F. oxysporum* f.sp. *cubense* is the causal agent of banana wilt disease (Lagopodi et al., 2002; Ploetz, 2006). *F. oxysporum* f. sp. *vasinfectum* (*Fov*), is responsible for Fusarium wilt of cotton, a significant disease that reduces cotton yields. This soil-borne fungus invades vascular tissue via host roots and rapidly spreads to the aboveground portion of the host. Typical field symptoms of Fusarium wilt include yellowing, chlorotic leaves, dark-brown/necrotic xylem (discoloration), and wilting eventually leading to plant death (Davis et al., 2006).

Reactive oxygen species (ROS) produced by host cells play an important role in defense
against various pathogens. These small ROS molecules are highly toxic to the infective agents and are able to directly kill them. At the interaction between the host and pathogen, host cells rapidly synthesize and accumulate large quantities of ROS, during a process called the oxidative burst (Heller and Tudzynski, 2011). In plants, the oxidative burst may result in a hypersensitive response (HR) phenomenon which can inhibit the spread of the pathogen to surrounding tissue (Zurbriggen et al., 2010). Additionally, ROS serve as an important signal which can trigger a series of other plant defense responses or the production of plant hormones (Sauer et al., 2001). These processes effectively stimulate plant hosts to re-program the expression of genes involved in defense, leading to the production of antimicrobial compounds such as phytoalexins, callose deposition, and systemic acquired resistance (SAR); collectively impeding pathogens from spreading and causing disease (Forman et al., 2010; Sauer et al., 2001).

Despite the extensive production of ROS in plant defense, successful pathogens are usually able to survive in a high ROS environment and have evolved strategies to overcome these detrimental effects (Aguirre et al., 2006; Fones and Preston, 2012) as it is essential for pathogens to tolerate ROS during the infection process. The group of enzymes referred to as superoxide dismutases (SOD) are well-known to participate in catalyzing the partitioning of superoxide radicals. SOD enzymes are divided into several families according to structure of the enzymes and the binding of a metal cofactor, and include the copper and zinc SODs (Cu/Zn SOD), the iron or manganese SODs (Fe/Mn SOD), and the nickel SODs (Miller, 2012). The cytosolic Cu/Zn SODs are the most common SOD enzymes in eukaryotic cells where there is at least one Cu/Zn SOD encoding gene in a genome. In *Saccharomyces cerevisiae*, loss of
SOD1 can lead to slow growth and increased sensitivity to ROS when exposed in ROS-generating compounds (Moradas-Ferreira and Costa, 2000). Additionally, SOD1 mutants showed attenuated virulence in some fungal pathogens. For example, the SOD1 mutant strain of Cryptococcus neoformans showed less virulence than the wide-type strain and showed significantly decreased growth in macrophages (Cox et al., 2003). Additionally, loss of function of SOD1 gene can alter normal fungal physiology. For instance, SOD1 mutants of the ericoid mycorrhizal fungus Oidiodendron maius had reduced production of conidia and the capacity for mycorrhization (Abba et al., 2009). Fe/Mn SODs mainly exist in prokaryotes, protists, or mitochondria/chloroplasts, suggesting that Fe/Mn SODs are an ancient family of SODs (Miller, 2012). Only a few prokaryotic microorganisms use nickel SODs for reducing its substrates (Youn et al., 1996).

A class composed of extracellular SODs has recently been described in various pathogens (Robinett et al., 2018). In bacteria, highly virulent E. coli serotype O157:H7 obtained additional secreted SodC through horizontal prophage gene transfer and is highly resistant to degradation by host proteases (D'Orazio et al., 2008). In fungi, extracellular SODs are a typical glycosylphosphatidylinositol (GPI) protein where they usually contain an N-terminal secreted signal peptide and a C-terminal GPI anchor attachment site. The mature GPI anchor enables proteins to be located to the cell membrane or cell wall by covalent attachment (Mayor and Riezman, 2004; Robinett et al., 2018). Current experimental evidence has indicated that some fungal extracellular SODs can be secreted out of the cell and participate in catalyzing the partitioning of superoxide radicals produced by host cells allowing fungi to survive in a high ROS environment (Youseff et al., 2012). The acquirement of extracellular SODs during the
evolution of fungal pathogens largely improves their pathogenicity (D'Orazio et al., 2008; Gleason et al., 2014; Youseff et al., 2012). *Candida albicans* can secret a SOD (Sod5), a Cu-only co-factor type, and this gene is required for pathogen defense (Gleason et al., 2014). In addition, the secreted Sod5 rapidly obtains the copper co-factor from the host to compromise the copper toxicity to *C. albicans* (Li et al., 2015). Similarly, an extracellular superoxide dismutase in the dimorphic fungus *Histoplasma capsulatum*, contributes to resistance to host-derived oxidative stress in yeast cells (Youseff et al., 2012). In *Puccinia striiformis*, the causative agent of stripe rust on wheat, a secreted extracellular Zn-only SOD contributes to enhanced resistance to oxidative stress during the interaction between the wheat host and the fungus (Liu et al., 2016). All these results showed extracellular SODs played an important role in pathogenicity to hosts and reduction of oxidative stress.

In *S. cerevisiae*, Sod1 has been shown to be localized to the nucleus when yeast cells were exposed to elevated H$_2$O$_2$ (Tsang et al., 2014). Once shuttled into the nucleus, Sod1 acts as a transcription factor binding DNA promoter regions and regulating gene expression (Tsang et al., 2014). Extracellular SODs are phylogenetically conserved throughout fungi and are predicted to have evolved from the canonical Sod1 (Robinett et al., 2018), and are therefore likely to share a similar biological function. However, little research has focused on the functional role of the homologous gene in other filamentous fungi. In this study, the extracellular SOD (FoSod5) from *Fusarium oxysporum* f. sp. *vasinfectum* was characterized and shown its subcellular localization is dependent on environmental conditions. During normal growth, FoSod5 was mainly localized to the fungal phialides. When fungi were exposed to a high ROS environment, FoSod5 was rapidly re-localized to the septum and cell wall. In a
lacZ reporter assay, the results indicate FoSod5 was a GPI protein that was rapidly induced in alkaline conditions and the GPI site was required for correct protein subcellular localization. Loss of the FoSOD5 gene led to a reduction in fungal virulence on cotton.

4.2 Materials and Methods

*F. oxysporum* strain, *E.coli* strain, plant materials, and growth conditions

*Fusarium oxysporum* f. sp. *vasinfectum* (*Fov*), a strain highly virulent on cotton, was obtained from the Fungal Genetics Stock Center (FGSC #10442) and used in this study. The wild-type *Fov* isolate and all mutant and complemented strains were cultured at room temperature (25 °C) on Potato Dextrose agar (PDA) medium or minimum nutrient medium (M-100). All *F. oxysporum* strains used in this study are listed in the Table 4.1. The mutant and complemented strains were screened on M-100 agar medium containing hygromycin B or G418 at a final concentration of 150 μg/mL or 60 μg/mL, respectively. For evaluation of fungal sensitivity to various chemical stressors, a 5 mm diameter mycelial plug was taken from the edge of a four-day-old culture grown on M-100 agar medium and put onto the new agar medium containing the chemical. The diameter of the colony was evaluated after seven days. Each treatment was repeated three times.

*E.coli* DH5α was used for construction and propagation of the transformation vectors. *Agrobacterium tumefaciens* strain AGL-1 was used for fungal transformation (Mullins et al., 2001). Cotton cultivar FM1944 was chosen for pathogenicity assays, and was cultured in a growth chamber with a period of 16 h light 30 °C and 8 h darkness 25 °C.
Bioinformatics analysis

The SOD protein family in *F. oxysporum* was identified using BLASTP software in the Ensembl Fungi database (https://fungi.ensembl.org/index.html) with several SOD query sequences (*S. cerevisiae* Sod1: YJR104C and Sod2: YHR008C; *C. albicans* Sod5: XP_719507). Protein domains were predicted using PFAM (El-Gebali et al., 2019) and BLASTP software. The subcellular localization peptide sequences were identified with SeqNLS and WoLF PSORT (Horton et al., 2007; Lin and Hu, 2013). Secreted signal peptides were predicted with SignalP 4.1 (Petersen et al., 2011). Multiple sequence alignments were conducted with the online software TCOFFEE (Notredame et al., 2000), and the phylogenetic tree was generated with PhyML 3.1 (Guindon et al., 2010). The 3D protein structure of FoSod5 was predicted by the SWISS-MODEL server (Gleason et al., 2014; Waterhouse et al., 2018).

Plasmid constructions

The *FoSOD5* gene complementary plasmid, pCom-G418<sup>R</sup>-*FoSOD5*, was constructed based on the background plasmid of pCAMBIA1302 as the following: 1) the hygromycin cassette under the 35S promoter between the XhoI and KpnI restriction sites was replaced with a neomycin phosphotransferase II cassette (trpC promoter) amplified from the pII99 plasmid; 2) the *FoSOD5* gene locus, ~4 kb in length, was cloned from genomic DNA using the primers FoSOD5cassF and FoSOD5cassR, and ligated into the above constructed plasmid between the LB- and RB- T-DNA sequences with the NEBuilder<sup>®</sup> HiFi DNA Assembly Cloning Kit (New England Biolabs).

The HITI Cas9 RNP transformation plasmid, pUC19-HITI-*FoSOD5C2*, was constructed
using the NEBuilder® HiFi DNA Assembly Cloning Kit and four PCR fragments were simultaneously assembled into the pUC19 plasmid between the BamHI and the HindIII restriction sites in the following order, the first fragment contained a ~1kb upstream sequence and partial amino acid coding region (AA: 1-53) (primers: NA_FoSod5C1hitiF and NA_FoSod5C2uphitiR), the second fragment was the sGFP coding region (primers: NA_FoSod5C2midsGFPF and NA_FoSod5C2midsGFPF), the third fragment contained the partial amino acid coding region (AA: 186-end containing the GPI site) and the predicted native terminator of the FoSOD5 gene (~800 bp) (primers: NA_FoSod5C2downF and NA_FoSod5C2downR), and the forth fragment was the hygromycin cassette amplified from plasmid pUC19-HDRI-FoSSO1-mCherry (primers: NA_FoSod5C2HYGBF and Universal_NA_forHITITR) (Wang and Coleman, 2019). The plasmid pUC19-HITI-FoSOD5C2 contained the FoSOD5 gene with the catalytic SOD domain replaced by sGFP. The plasmids pUC19-HITI-FoSOD5C1 and pUC19-HITI-FoSOD5C3 were generated in a similar way. Plasmid pUC19-HITI-FoSOD5C1 did not contain the amino acid sequences after the SOD domain, while in pUC19-HITI-FoSOD5C3, the ORF of the FoSOD5 gene was completely replaced by the β-galactosidase (lacZ) coding region cloned from the plasmid, pCYC-lacZ. The three different plasmids were used to generate variants of FoSod5 enabling the study of the function of the FoSod5 protein, including protein secretion, protein subcellular localization, and protein expression. All PCR primers used are listed in Table 4.2

**Target gene replacement and gene complementation**

The target gene, FoSOD5, was replaced with the hygromycin B resistance cassette using
a previously described split-marker approach with a few modifications (Goswami, 2012). Briefly, \( \sim 10^7 \) conidia were inoculated into 100 mL of fresh PDB liquid medium and grown on a rotary shaker at 30 °C 180 rpm for 12-14 h. The fungal germlings were collected and treated by a mixture of protoplasting enzymes, 10 mg/mL Driselase™ from *Basidiomycetes sp.* (Sigma), 15 mg/mL β-Glucanase from *Trichoderma longibrachiatum* (Sigma), and 5 mg/mL Lysing enzymes from *T. harzianum* (Sigma) for 2-3 h. The resulting protoplasts were filtered through Nitex Nylon mesh, centrifuged, collected, and washed with SuTC buffer. The protoplasts were suspended in SuTC buffer to a concentration of \( 2 \times 10^7 \) protoplasts/mL. The DNA fragments were generated by overlapping fusion PCR (Goswami, 2012). The protoplast suspension (200 μL) was mixed with a total of 10 μg of the DNA fragments (5 μg of the upstream fragment and 5 μg of the downstream fragment) and were incubated at room temperature for 20 min. One mL of PSTC was slowly added to the above protoplast mixture, mixed, and left at room temperature for 20 min. After that time, 3 mL of TB3 liquid medium with 25 μg/mL Amp antibiotics were added. The transformed protoplasts were cultured on a rotary shaker (160 rpm) for 20 h. Once the cell wall was regenerated, \( \sim 15 \) mL of melted bottom agar (\( \sim 60 ^\circ \text{C} \)) with 100 μg/mL hygromycin B were mixed with the transformed protoplasts and poured onto plates. After 10 h, a top layer of agar medium with 150 μg/mL hygromycin B was poured over the bottom medium. Resistant colonies emerged within 3 ~ 5 days, and the resistant colonies are transferred to M-100 medium with hygromycin B for further evaluation.

For the FoSOD5 gene complementation, a *Agrobacterium*-mediated transformation method was used according to a previous publication with a few modifications (Mullins et al., 2001). The ΔFoSOD5 strain was inoculated into fresh PDB medium at 160 rpm, room
temperature (25 °C) for three to five days. A. tumefaciens strain AGL-1 containing the plasmid pCom-G418\textsuperscript{R}-FoSOD5 was grown at 28 °C, 180 rpm for two days in LB liquid medium with appropriate antibiotics. The AGL-1 was diluted to an OD\textsubscript{600} value of 0.2 in induction medium containing 100 μM acetosyringone (Sigma) and associated antibiotics. The induction time length was about 6 h, after which a 100 μL conidial suspension (1 × 10\textsuperscript{7} / mL) was mixed with an equal volume of the above AGL-1 induction medium. The total mixture was plated on the surface of a 0.45 μm nitrocellulose filter on co-cultivation medium. The plates were set in the dark at 28 °C for two days. One mL of sterile water was used to wash the conidia from the surface of the nitrocellulose filter, and each aliquot of 200 μL of the conidia-containing water mixture was plated on M100 selective medium containing 50 μg/mL of G418 and 300 μg/mL of cefotaxime (to counter select the AGL-1 cells). All selective plates were incubated at room temperature for 5-7 days. After the G418 resistant colonies appeared on the surface of selective medium, they were transferred to new selective medium for several rounds of screening. The mutants and transformants in this experiment were further confirmed by PCR and Southern blots. All PCR primers are listed in the Table 4.2.

**In vitro Cas9 nuclease assay and Cas9 RNP transformation**

Before Cas9 RNP transformation, an in vitro Cas9 nuclease assay was conducted to confirm the Cas9 RNP cleavage activity. This assay was performed as described previously (Wang et al., 2018). Briefly, a total 20 μL reaction system composed of 200 ng of the Cas9 protein, 40 ng of sgRNA, 1×Cas9 Nuclease Reaction Buffer, and 50 ~ 100 ng of the DNA template, was mixed and incubated at 37°C for 1 hr. The cleaved DNA products were resolved
through a 1% agarose gel. The Cas9 RNP transformation was conducted as previously described (Wang et al., 2018), and the amount of donor plasmid did not exceed 6 µg per transformation plate. After transformation, the positive colonies were isolated on M100 selective medium containing 150 µg/mL of hygromycin. Three pairs of PCR primers were used to confirm the location of the integrated plasmids. Since three different FoSod5 variants were generated using this HITI strategy, the cleavage site at the 5′-terminus (SS, Fig. 4.3A) was sequenced.

**β-Galactosidase (lacZ) activity assay**

*FoSOD5* expression and regulation was investigated under various conditions. For different pH values, M100 agar medium was adjusted to pH values ranging from 4 to 8 with a final X-gal concentration of 200 ng/mL. Regulation of *FoSOD5* under the influence of different carbon sources was investigated using M100 medium with the glucose replaced by 1% (w/v) of various carbon sources (glycerol, carboxymethylcellulose, starch, sucrose, sorbitol, and mannitol). Conidia (~ 1 × 10⁵) were pipetted onto the center of agar plates with or without X-gal. For quantification of β-Galactosidase enzymatic activity, an O-Nitrophenyl-B-D-Galactopyranoside (ONPG) assay was conducted. FoSod5C3 was grown to the exponential stage in YG liquid medium and different chemicals (0.05% H₂O₂, 0.5 mM diamide, 0.2 mM menadione, 20 µg/mL xanthine oxidase/0.1mM hypoxanthine, and 150 µM CuSO₄) were added to the YG medium with the fungi and cultured for an extra 3 hrs. Total protein was extracted under liquid nitrogen, and 5 µg of the total protein (diluted to 75 µL) from each treatment and 40 µL of 4 mg/mL ONPG was added to 700 µL of the reaction buffer (8.5 mg/mL
Na₂HPO₄, 5.5 mg/mL Na₂H₂PO₄·H₂O, 750 ng/mL KCl and 246 ng/mL MgSO₄·7H₂O, pH 7.0) and incubated at 30 °C for 25 min. After which, 100 μL of 1M Na₂CO₃ was used to stop the above reaction. The β-Galactosidase enzymatic activity was determined by spectroscopy at an absorbance of 420 nm (Cytation 3, BioTek).

RNA extraction of Fov-infected cotton roots and qRT-PCR

Cotton seedlings that were ~10-days-old were uprooted from soil, washed with sterile water, placed in a water-culturing box containing ~ 500 mL of nutrient buffer (1/10 Murashige & Skoog basal medium with vitamins (MS) with 0.1% sucrose, PhytoTechnology Laboratories), and allowed to acclimate for 3 days in a growth chamber. Conidia (~ 2×10⁷) were inoculated into the nutrient buffer in a water-culturing box and mixed well by gently shaking. The Fov-infected cotton roots were collected at different time-points (0h, 4h, 12h, 24h, 36h, 2d, 3d, 5d and 7d post inoculation) for total RNA extraction. Total RNA of the cotton roots was extracted using the E.Z.N.A.® Total RNA Kit I (OMEGA, Bio-tech), according to the manufacturer’s instructions. Reverse transcription was immediately conducted to generate first-strand cDNA using the RETRO Script kit (Ambion) with the random oligos. The transcript expression of FoSOD5 was determined by qRT-PCR on a Bio-Rad CFX96 instrument for the different time points of fungal infection using the primers qFoSOD5F and qFoSOD5R. The F. oxysporum elongation factor 1 alpha gene (EF1α) was used as the internal reference gene using the primers qEF1αF and qEF1αR. The relative expression levels were calculated by the ΔΔCt method (Livak and Schmittgen, 2001). Three cotton roots were combined for extracting total RNA and three technical replicates were performed in this experiment. The genes and primers sequences...
are listed in Table 4.2.

6 × His-FoSod5 protein purification and SOD activity assay

The 6×His-FoSod5 protein purification was conducted using a previously published method (Gleason et al., 2014). Briefly, the signal peptide (N-terminus) and GPI-anchor sites (C-terminus) of the FoSOD5 gene sequence were removed and the middle amino acid sequence was inserted into the pHis-Parallel1 vector (Sheffield et al., 1999). The recombinant plasmid was transformed into E. coli Rosetta™(DE3) competent cells. Protein production was induced by the addition of 0.5 mM IPTG to a culture at an OD₆₀₀ value between 0.5 and 0.6 and allowed to grow for 4 h at 37 °C. Protein purification was performed under denaturing conditions with the prepared denaturing buffer (50 mM Tris-HCl (pH 8.0), 8 M urea, and 1.5 mM reduced glutathione) using a Ni-NTA Purification System (Thermo Fisher). A series of dialysis buffers with different concentrations of urea (50 mM Tris-HCl (pH 8.0), 6/4/2/0 M urea and 0.25 mM oxidized glutathione) were used for FoSod5 protein refolding with a 10 kDa dialysis tube. SOD proteins usually require different metals to fold into the correct structure for function. Different metal ions (ZnCl₂, FeCl₃, CuSO₄, and MnSO₄) were provided to the 6×His-FoSod5 at the final dialysis overnight. The final 6×His-FoSod5 protein was concentrated to 2 mg/mL with 3K protein concentrator tubes (Millipore Sigma).

Two methods, the nitro-blue tetrazolium (NBT) method and the SOD-WST assay, were used for assessing the enzymatic activity of the purified 6×His-FoSod5. The NBT method reflects SOD enzyme efficiency through inhibition of the reduction of NBT when superoxide is present. When NBT is reduced by superoxide, formazan is produced causing a proportional
color change to dark blue that can be monitored. Three mL of the freshly prepared reaction buffer (10 μM riboflavin, 45 μM EDTA, 350 μM NBT, 60 μM methionine, and potassium phosphate with a pH value of 7.8) was mixed to obtain different concentrations (0, 4, 10, 20 and 40 μg) of the purified 6×His-FoSod5 protein and immediately incubated under the light (~4000 Lx) for 20 min at 28°C. The color change of the reaction solution is an indication of formazan production, and is proportional to the amount of SOD enzyme activity. The SOD-WST assay was conducted using the EnzyChrom™ Superoxide Dismutase Assay Kit (BioAssay Systems) and conducted according to the manufacture instructions. One μg of the FoSod5 protein that was reconstituted in the presence of different metal cations was added to the WST-1 reaction buffer and incubated at room temperature for 1 h. The results were measured by spectroscopy at the the absorbance A_{440 nm}.

Secreted protein extraction and western blotting

A trichloroacetic acid (TCA) precipitation method was used to extract fungal secreted proteins. The fungal isolates were cultured in YG liquid medium for three days at room temperature. The resulting fungal culture was centrifuged at 13000 g for 10 min and filtered through a 0.22 μM filter to remove hyphae and conidia. TCA was added to a final concentration of 10%, and the mixture chilled at 4 °C overnight. The solution was centrifuged at 13000 g for 10 min, and the resulting pellets were washed with ice-cold acetone at least twice. The final protein pellets were dissolved in 50 mM Tris-HCl buffer (pH=7.5), and the protein concentration was determined using a Qubit 3.0 Fluorometer with the Qubit™ Protein Assay Kit (Thermo Fisher Scientific).
The western blotting was conducted with a total of 5 μg of the secreted proteins which were separated by 10% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and stained with Coomassie blue staining solution (0.1% Coomassie Brilliant Blue R-250, 50% methanol and 10% glacial acetic acid). The proteins in the SDS-PAGE gel were transferred to a nitrocellulose filter membrane (GE Healthcare) using the Mini Trans-Blot® Electrophoretic Transfer Cell (BioRad). Anti-GFP (Rockland Immunochemicals) and the ECL chemiluminescent detection kit (GE Healthcare) were used to detect the resulting proteins.

**Confocal microscopy observation**

Confocal microscopy was conducted with a Nikon A1 Confocal Microscope with an excitation wavelength of 488 nm for detecting the sGFP fluorescence. The observation was performed according to a previously described protocol (Wang and Coleman, 2019). For different chemical treatments, a mixture of hyphae and conidia was inoculated into YG medium and cultured for three days. One mL of the hyphal-conidial mixture was transferred into fresh YG medium and cultured on a rotary shaker at 18 °C at 220 rpm for 16 h. Different chemicals (0.03% H₂O₂ and 20 μg/mL xanthine oxidase/0.1mM hypoxanthomine) were added to the YG medium and cultured for an additional 2 h. At this time, samples were aliquoted, fixed with 4% paraformaldehyde, and observed by confocal microscopy using an inverted agar method (Hickey and Read, 2009).

**Pathogenicity assay**

Two different infection assays were used to assess fungal virulence. In the cotton root
lesion assay, cotton seeds (cultivar FM1944) were washed and surface sterilized with 20% bleach followed by 75% ethanol, washed in sterile water, and germinated on wet filter paper for 3~5 days. Cotton seedlings with cotton roots 4~6 cm in length were chosen for the infection assay and at least ten cotton roots were used for evaluating the virulence of a single fungal strain. Conidia were collected and dispersed into sterile water at a concentration of $1 \times 10^6$ per mL. Ten $\mu$L were dropped onto the root surface and incubated in a dark, wet box for 48 h. Virulence was quantified by the length of the lesions caused by *F. oxysporum*.

For the cotton root infection assay, cotton seeds (cultivar FM1944) were planted in sterile soil and placed in a growth chamber with a 16 h 28 °C light / 8 h 24 °C dark cycle. After 2 weeks, cotton plants at a two-to-three true leaf stage were used for the infection assay, where the cotton seedlings were carefully uprooted and immersed in sterile water to remove adhering soil from the root. A set of eight seedlings were transplanted to a water-culturing box which contained nutrient solution (1/10 MS with 0.1% sucrose) to sustain the growth of the cotton plant and allowed to acclimate for four days in the growth chamber. During this time, the wild-type and mutant strains of *Fov* were inoculated to PDB and allowed to grow at room temperature for 5 days on a rotary shaker. The conidia were collected from the cultures, washed in sterile water, and suspended at a concentration of $1 \times 10^7$ conidia/mL in sterile water. Two mL of the conidia were pipetted into the water-culturing box containing the cotton plants and mixed well with gentle shaking. Each week the remaining nutrient solution in the plant culturing box was replaced with fresh nutrient solution. After 5-8 weeks, disease symptoms were recorded by photo and the cotton vascular tissue was evaluated. Based on *F. oxysporum* disease symptoms on cotton, four disease index categories were developed: 0 – no symptoms,
1 – light yellowing of the leaves and black root tips, 2 – extensive yellowing of the leaves, wilting evident, and brownish discolored vascular tissue, 3 – severely wilted plant with dark discolored vascular tissue or whole plant death. Pathogenicity assays for each strain were repeated at least twice.

4.3 Results

4.3.1 Identification of an extracellular GPI SOD protein (FoSod5) from *F. oxysporum*

Sod proteins from different fungal species were used as queries to identify five Sod proteins encoded in the *F. oxysporum* genome, which included two Fe/Mn SOD genes (FoSOD3: FOTG_10379; FoSOD4: FOTG_02058) and three Cu/Zn SOD genes (FoSOD1: FOTG_01421; FoSOD2: FOTG_16882; FoSOD5: FOTG_08628) (Fig 4.1A). Subcellular localization prediction analysis indicated these five Sod proteins had varied in subcellular localization where FoSod1 (a Cu/Zn SOD) was localized to the fungal cytosol while FoSod3 (a Fe/Mn SOD) was localized to the mitochondrion, in agreement with the subcellular localization of homologous proteins from other fungi (Luk et al., 2003; Yao et al., 2016). However, FoSod2 (a Cu/Zn SOD) was predicted to be localized to the peroxisome or the nucleus while FoSod4 (a Fe/Mn SOD) contains a predicted secretion signal peptide and could represent a secreted SOD enzyme. FoSod5, the final SOD in *F. oxysporum*, contains a N-terminal secreted signal (AA: 1-21) and a C-terminal glycosphatidyl inositol (GPI) attachment site (AA: 240), which suggests that this protein is secreted and anchored to the fungal cell wall or membrane (Fig. 4.1A). Sequence alignment of select Cu/Zn SODs revealed that like CaSod5, FoSod5 lacked two histidine sites involving in zinc binding, suggesting
FoSod5 was a Cu-only SOD enzyme (Fig. 4.8) (Gleason et al., 2014). Although the amino acid sequence similarity between CaSod5 and FoSod5 was only 33%, the predicted structure of FoSod5 was highly similar to the Cu-only CaSod5, suggesting functional similarities between the two proteins (Fig. 4.1B). Phylogenetic analysis showed that most fungi in the Ascomycota phylum contained a single ortholog of the SOD5 gene (Fig. 4.1C). Interestingly, all the isolates of Aspergillus lost the GPI anchor site in the Sod5 protein during their evolution. Isolates within Fusarium carried a single phylogenetically conserved copy of the SOD5 gene, which likely had diverged from the well-characterized extracellular SOD from C. albicans, CaSOD5 (Fig. 4.1C).

4.3.2 FoSod5 was a functional SOD protein.

To assess the SOD activity for FoSod5, the N-terminus secretion signal peptide and the C-terminus GPI site were removed, and the middle amino acid sequence harboring the SOD domain was fused with a N-terminus 6 × His tag for protein purification using E. coli Rosetta™(DE3) (Fig. 4.2B). Two SOD activity detection methods, the nitro-blue tetrazolium (NBT) method and the SOD-WST1 assay, were used for determining the enzymatic activity of the purified 6 × His-FoSod5 and the specificity for different metal ions to serve as a cofactor for FoSod5 activity.

In the SOD-WST1 assay, of all the heavy metals tested, only copper could efficiently confer SOD activity, demonstrating that FoSod5 was a copper-only SOD (Fig. 4.2D), consistent with a previous publication (Gleason et al., 2014). The traditional NBT-riboflavin method was also used to confirm SOD activity of the 6 × His-FoSod5. Under illumination, the riboflavin-methionine mixture was able to produce superoxide, and NBT was reduced to blue formazan.
which can be inhibited by SOD activity. A series of the 6 × His-FoSod5 protein at an increasing quantity (0 to 40 μg) were added to the reaction buffer, and as the concentration of the 6 × His-FoSod5 protein increased, a decrease in the production of formazan was evident (Fig. 4.2C) indicating that FoSod5 was a SOD enzyme able to inhibit NBT reduction. In addition, a radial growth ROS stress assay showed the ΔFoSOD5 mutant had increased sensitivity to hydrogen peroxide and diamide, a ROS-generating compound (Fig. 4.1D). Collectively, these results indicate that FoSod5 is a functional copper specific SOD enzyme that may have an important role in overcoming ROS damage.

4.3.3 FoSOD5 is up-regulated during infection of cotton.

Many fungal virulence factors have increased expression during the infection of host plants. For example, two LysM effectors from Zymoseptoria tritici are expressed during infection of wheat where they protect fungal hyphae against plant-derived hydrolytic enzymes (Marshall et al., 2011). In addition, the effector SIX6, a virulence factor from Fusarium oxysporum f. sp. lycopersici, is expressed during the early and late stages of infection of tomato and its expression is triggered by living host cells (Gawehns et al., 2014). As FoSod5 is a SOD and many of these enzymes are known to be involved in virulence in other fungal pathogens, the expression profile of FoSOD5 during infection of cotton was investigated by qRT-PCR. The expression of FoSOD5 was low over the initial 12 hours of infection, but this gene was drastically induced and expressed from 24 h to 7 d after infection (Fig. 4.2A). At all-time points 36 hr after infection, there was at least a >50 fold increase in FoSOD5 transcripts, peaking at 3 d dpi (Fig. 4.2A).
4.3.4 *FoSOD5* was required for full virulence of *Fov*

The WT, *FoSOD5* mutant (*ΔFoSOD5*), and complemented (*ΔFoSOD5/FoSOD5*) isolates were used to investigate the role of FoSod5 in pathogenicity on cotton. In a cotton root lesion assay, *Fov* conidia inoculated on the surface of the cotton seedlings are able to infect the root tissue and cause a lesion within 48 h. While the *FoSOD5* mutant was able to cause a lesion, it was significantly smaller in size than the lesions formed by the wild-type isolate (Fig. 4.7A and 4.7B), supporting the involvement of *FoSOD5* in pathogenicity. As *Fov* is responsible for wilting symptoms, a second virulence assay was conducted on whole plants to confirm the role of FoSod5 in infection. While the wild-type isolate was able to colonize the xylem tissue in the roots and cause some necrosis, the *ΔFoSOD5* mutant had reduced necrosis (Fig. 4.7C). Overall, the cotton plants inoculated with the *FoSOD5* mutant displayed less wilting symptoms, had less yellowing of the leaves, and there was less necrosis of the root xylem when compared to cotton plants inoculated with the WT and the *ΔFoSOD5/FoSOD5* complement isolates (Fig 4.7C and 4.7D).

4.3.5 Expression of *FoSOD5* is regulated by various carbon sources and repressed in a nutrient rich-medium.

The function of cell wall proteins (CWPs) can vary and sometimes be species-specific, as they contribute to the diverse properties of the cell wall and enable fungi to adapt to harmful environments (Gow et al., 2017). The expression of these CWPs is highly regulated; for example, *C. albicans* significantly alters the composition of CWPs in response to the availability of a carbon source, iron limitation, or hypoxia (Chaffin, 2008; Ene et al., 2012;
Sorgo et al., 2013). A lacZ reporter construct was generated under the control of the native FoSOD5 promoter to further investigate the expression of FoSOD5 in various environments. The lacZ gene, encoding the beta-galactosidase protein converting X-gal to the blue product which was easily seen by naked eyes, is commonly used as a reporter gene for the in vivo analysis of gene regulation in various organisms. After transformation, eight independent colonies were selected for analysis, and seven of these transformants produced the blue pigment when grown on M100 medium containing X-gal, indicating the lacZ gene can be a suitable reporter for the in vivo analysis of gene regulation in F. oxysporum. Sequencing results of six of the seven transformants revealed that error free DNA repair occurred at the sgRNA cleavage region without any nucleotide additions or deletions (Fig. 4.3A, SS sites), whereas the remaining transformant had a 77 bp nucleotide deletion (Fig. 4.10).

The FoSOD5-lacZ reporter was used to investigate the expression pattern of FoSOD5 when grown on various carbon sources. LacZ activity was evident using this strain when the fungus was grown on a minimal nutrient medium (M-100) for four days. When a agar plug (5 mm in diameter) was removed from the lacZ inducing M-100 plate and placed on a rich nutrient medium (TB3), the resulting hyphae that grew on the TB3 medium failed to have significant LacZ activity after 4 days, and the hyphae were primarily white in color (Fig. 4.4C). Even after one week, there was no significant color change of the mycelia on TB3 media. When an agar plug of the FoSOD5-lacZ reporter mycelia that was grown on TB3 was placed on the minimal M-100 medium, the mycelia had LacZ activity. This regulation of LacZ activity by the FoSOD5 promoter indicates that the superoxide dismutase is dependent on the available nutrient(s).

Since the available carbon sources between the M-100 and TB3 media were different, it
was hypothesized that the carbon source in the medium may regulate *FoSOD5* gene expression. The 1% (w/v) glucose content in the M-100 agar medium was replaced with 1% (w/v) of various other carbon sources (sucrose, mannitol, glycerol, CMC, starch, and sorbitol). Four days after inoculation the mycelia of the *FoSOD5-lacZ* reporter indicated LacZ activity was evident for all carbon sources investigated, but there was a clear difference between nutrients. Interestingly, the LacZ activity of the mycelia in the presence of CMC and starch were higher than those for the other carbon sources (Fig. 4.4B), suggesting that plant-derived carbon sources may facilitate *FoSOD5* gene expression. In addition, *FoSOD5* gene regulation was investigated when in the presence of different chemical stimuli. After the addition of different chemicals including hydrogen peroxide and the ROS generating compounds diamide and menadione, the LacZ activity of the mycelia was increased to various levels, suggesting *FoSOD5* expression is up-regulated under different stimuli (Fig. 4.4D).

4.3.6 *FoSOD5* is rapidly induced under an alkaline environment.

Many virulence factors of phytopathogenic fungi, including those in *F. oxysporum*, are dependent on the pH of the surrounding environment, and therefore, we hypothesized that the pH of the medium may play a role in the regulation of *FoSOD5*. Conidia of the *FoSOD5-lacZ* reporter strain were placed on M-100 plates that have been adjusted to pH’s ranging from 4 to 8 and monitored over time. In a neutral or alkaline environment (pH= 7 or 8), *FoSOD5* was induced within the first 24 hours; however, mycelia at a pH 6 did not display LacZ activity until 48 hours, and at a pH of 4 or 5 it required 3 days for LacZ activity to be evident (Fig. 4.4A).
4.3.7 FoSod5 is only associated with fungal cells

Typical glycoprophosphatidyl inositol (GPI) proteins contain both the N-terminus secreted signal peptide and the C-terminus GPI site (Mayor and Riezman, 2004). These GPI proteins are attached to either the fungal cell wall or the plasma membrane. Previously Sod3, a homologous GPI protein from *H. capsulatum*, could be associated with yeast cells and secreted into the culture medium (Youseff et al., 2012). To assess the N-terminal secretion signal peptide of FoSod5, two FoSod5 variants under the control of the native promoter were generated (Fig. 4.3B). The homologous-independent targeted integration (HITI) strategy was used to insert an entire plasmid at the *FoSOD5* endogenous gene locus (Fig. 4.3A). The FoSod5C1 variant was constructed by replacing the nucleotide sequences coding for the SOD domain and GPI site (AA: 53 - 263) with sGFP; while the FoSod5C2 variant replaced the internal catalytic SOD domain (AA: 53 - 186) with sGFP (SP+sGFP+GPI) (Fig. 4.3B). After transformation, the loci of interest of three FoSod5C1 and two FoSod5C2 transformants were sequenced, confirming that all five transformants do not contain indels or other undesired alterations (Fig. 4.10). These FoSod5 variants and the wild type isolate were grown in liquid culture and after several days the supernatant analyzed for the presence of sGFP by western blot. Using anti-GFP antibody only the sGFP of FoSod5C1 was able to be detected in the liquid culture medium (Fig. 4.3C), confirming FoSod5 contained a N-terminal functional secretion peptide. When the GPI anchor is included (FoSod5C2) the protein is unable to be detected in liquid culture medium, even when up to 25 μg of the extracted protein was used for the western blot and increasing the exposure time for detection. Therefore, unlike Sod3 from *H. capsulatum*, FoSod5 was only associated with the fungal cells and was not secreted into the liquid medium.
4.3.8 Subcellular localization of FoSod5

Bioinformatic analysis of FoSOD5 indicates a GPI anchor resides after the SOD catalytic domain towards the C-terminus, and therefore this protein could be attached to the cell wall although little is known about the subcellular localization of this protein in detail. Two different modifications, a sugar modification or a lipid modification, can occur at GPI sites, and the mature protein will be sorted with a C-terminal attachment to the vesicle membrane (Mayor and Riezman, 2004). GPI proteins can be secreted and attached to the fungal cell wall under the direction of a N-terminal cleavable secreted signal peptide. Since mature proteins pass through the secretory pathway, the SOD domain was replaced with a sGFP coding sequence (FoSod5C2) which was used for subcellular localization studies (Fig. 4.3B). Surprisingly, most of FoSod5 protein were specifically localized to the fungal phialides in YG medium, a medium that is less rich in nutrients than TB3 (Fig. 4.5A). However, a weak fluorescent signal was present in conidia, fungal tips, septa, and hyphae indicating that FoSod5 also accumulated in these structures (Fig. 4.5A and 4.5B). When this isolate was grown in minimal nutrient M-100 medium, FoSod5 was localized to the conidia, hyphae, and septa, indicating a significant alteration of the subcellular localization (Fig. 4.5C and 4.5D). Interestingly, large amounts of FoSod5 was accumulated in the hyphal tips (Fig. 4.5E). The in vivo FoSod5 protein regulation appears to be more complex in YG medium when compared to growth in M-100 medium. A Z-series stack using confocal microscopy indicated that strong fluorescence was present in the cell wall of conidia, hyphae, and phialides (Fig. 4.11).
4.3.9 The GPI anchor is required for proper FoSod5 subcellular localization

The GPI anchor is known to play an important role in subcellular localization for many proteins. Comparison of the two sGFP constructs, FoSod5C1 and FoSod5C2 which only differed by the presence of the GPI site in FoSod5C2, demonstrated that the FoSod5C1 transformants without the GPI site had no fluorescent signal at the phialides under confocal microscopy (Fig. 4.6A and 4.6C). This finding, in conjunction with the previous result that sGFP was detected by western blot of the supernatant of the FoSod5C1 transformant, confirms that the GPI site of FoSod5 is required for proper physiological localization (Fig. 4.6A and 4.6C).

4.3.10 FoSod5 could be localized to the cell wall and septum in the presence of ROS

Increased expression of FoSOD5 was observed when the fungus was treated with various chemicals, including H$_2$O$_2$ and ROS generating compounds. Since FoSod5 preferentially accumulates to the phialides in YG medium but accumulates at the fungal cell walls/septa in M-100 media, we hypothesized that some FoSod5 proteins would be localized to the cell wall after the fungal cell recognizes the appropriate stimuli. Two different ROS-generating chemicals, H$_2$O$_2$ and xanthine oxidase/ hypoxanthine, which generates superoxide, were used to treat the FoSod5C2 transformants in YG medium. While the sGFP localization was previously found primarily at the phialides (Fig. 4.5A), in the presence of H$_2$O$_2$ and xanthine oxidase/ hypoxanthine treatments the sGFP fluorescent signal accumulated at the cell walls and septa of the FoSod5C2 transformants (Fig. 4.6A and 4.6B), suggesting that FoSod5 was localized to the fungal cell wall and septa when challenged with ROS. Statistical analysis
indicated that the frequency of FoSod5 localization to the septa was significantly different between the two treatments ($P = 0.002$ and 0.021; Fig. 4.6D).

### 4.3 Discussion

A vast amount of evidence supports the importance of ROS in fungal-host interactions. Successful fungal pathogens must overcome the cellular damage from ROS during infection (Heller and Tudzynski, 2011). In the present study, the GPI-protein Sod5 from *F. oxysporum* was shown to be a virulence factor. *FoSOD5* was gradually up-regulated during infection, suggesting that the FoSod5 protein may participate in ROS scavenging during the infection process. Mutation of *FoSOD5* led to increased sensitivity to ROS and attenuated virulence on cotton.

In fungi, the number of *SOD* encoding genes in a genome varies greatly. For example, *Saccharomyces cerevisiae* only encodes two *SOD* genes (Sod1 and Sod2) while some filamentous fungi, including *F. oxysporum*, encode five or more *SOD* encoding genes, suggesting that the expansion of the *SOD* gene family was a complex, selective process. Usually protein family expansion leads to divergence of protein function. Bioinformatic analysis indicated that the five SOD enzymes of *F. oxysporum* have a different subcellular localization. FoSod2 and FoSod5 are predicted to have evolved from FoSod1 (a Cu/Zn SOD), but they are hypothesized to be localized to the peroxisome/nucleus and cell wall, whereas FoSod1 is in the cytosol. FoSod4, which is most closely related to the mitochondrial localized SOD FoSod3, has a secretion signal peptide at its N-terminus. These results indicate that Sod proteins are translocated into different organelles/localizations which may have an influence
on the enzymatic function. Increasing evidence has shown that Sod proteins have various subcellular localizations and are re-localized under stressful conditions. In *S. cerevisiae*, the Cu/Zn SOD enzyme (Sod1) is mainly distributed in the cytosol and the Mn SOD enzyme is localized in the mitochondrion. Interestingly, Sod1 could rapidly relocate into the nucleus in response to high endogenous and/or exogenous ROS, aiding to maintain genomic stability (Tsang et al., 2014). In addition, Sod1 as a nuclear transcription factor is able to bind DNA promoter regions of oxidative resistance and repair genes and regulate the gene expressions. These results clearly indicate that the function of Sod1 is far more complex than only a ROS scavenging enzyme. In addition, some Sod1 proteins are able to be localized into the mitochondrial intermembrane space and the peroxisome (Fischer et al., 2011).

The FoSod5 protein accumulates to high levels at the fungal phialides in YG medium, suggesting that a large amount of ROS might be produced at the phialides and FoSod5 might act as a ROS scavenging enzyme during *F. oxysporum* conidiation. In addition, regulation of FoSod5 appears to be more complex in M100 medium as it was found in many locations. Although the FoSod5 protein was localized to the cell walls of conidia and hyphae in M100 medium, a strong accumulation of FoSod5 was present at the fungal tip. As rapidly growing regions, fungal tips have been shown to produce large amounts of ROS in *M. oryzae* (Egan et al., 2007). Fungal tip-localized FoSod5 might cooperate with other cytosol Sod proteins to scavenge ROS. Interestingly, when *F. oxysporum* was treated with superoxide and hydrogen peroxide in YG medium, FoSod5 was translocated to the hyphal cell wall and septa, indicating FoSod5 was not restricted to a single location in the fungal cell for function, but instead it could be translocated into other cellular regions in response to environmental stimuli.
Usually multiple virulence factors from a pathogen might cooperate to facilitate the infection process of a host plant. Other isolates of *F. oxysporum* secrete peptides which induce alkalization in plants and enhance fungal virulence during infection (Fernandes et al., 2017; Masachis et al., 2016). MAPK-mediated fungal growth on cellophane is more invasive at pH 7 than that at pH 5 (Masachis et al., 2016). In addition, alkaline treatment can increase root damage and inhibit seedling growth inhibition with ROS accumulation (Masachis et al., 2016). As the infection process is dynamic, fungi must evolve relevant strategies to overcome these pressures. *FoSOD5* was rapidly induced under an alkaline environment, suggesting that an alkaline condition may be a key induction factor during early fungal infections. The alkalization-inducing expression profiles might help fungi to overcome ROS damage at the alkaline environments.

The *SOD5* gene regulation in *F. oxysporum* was different than the regulation of homologous genes of human pathogenic fungi including *C. albicans*. For instance, the *SOD5* gene in *C. albicans* was shown to be a hyphae-induced factor and was gradually up-regulated during the hyphal induction (Heilmann et al., 2011; Martchenko et al., 2004). However, the *SOD5* gene from *F. oxysporum* was found to be regulated by external environments, since the *lacZ* assay showed there was nearly no expressions of *FoSOD5* under the rich-nutrient medium TB3. When cultured in minimum nutrient medium, the fungus showed high expression of *FoSOD5*. These results indicated that *SOD5* gene regulations largely diverged during fungal evolution. In addition, *FoSOD5* was involved in fungal pathogenicity during the infection. However, the previous evidence showed that *FgSOD5* from the closely-related fungus, *F. graminearum*, did not contribute to fungal infection (Rittenour and Harris, 2013). qRT-PCR results showed that
FgSOD5 was expressed at a low level and could not be highly induced during the infection of wheat (Yao et al., 2016). This divergence may suggest the various Sods have a different importance in virulence and could be dependent on fungus and/or host plants. Above all, FoSOD5 expression is highly regulated by environmental factors rather than fungal development, and could be highly up-regulated during infection and contribute to virulence on cotton.

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Figure 4.1  Bioinformatic analysis of the FoSod5 protein and stress assay
A) Identification of the Sod protein family in *F. oxysporum* and subcellular localization predictions of FoSod proteins. NLS: nucleus localization sequence; SP: secreted peptide
B) The predictive structure of FoSod5 and comparison with that of CaSod5.
C) The maximum-likelihood phylogenetic tree of Sod5 amino acid sequences from the ascomycete fungi. The tree was generated with a bootstrap value of 500.
D) The evaluation of fungal sensitivity to different chemical stresses. The indicated strains were inoculated on the minimal medium (M-100).
**Figure 4.2**  *FoSOD5* gene was up-regulated during infection and FoSod5 was a functional SOD protein.

A) The expression of *FoSOD5* during infection was measured by quantitative real-time RT-PCR (qRT-PCR) using cDNA from infected plant tissue. B) SDS-PAGE gel depicting protein expression and purification conditions. Lane 1 represents total protein before inducing; Lane 2 total protein after addition of 0.5 mM IPTG for 4 h; Lane 3 represents purified 6xHis-FoSod5 protein. C) the nitro-blue tetrazolium (NBT) method was used for testing purified 6xHis-FoSod5 enzyme activity. The production of formazan decreases as the 6xHis-FoSod5 protein increases. D) Statistics analysis of 6xHis-FoSod5 enzyme activity with different mental ions using SOD-WST assay.
Figure 4.3 CRISPR/Cas9 RNPs-based transformant generation
A) The homologous-independent targeted integration (HITI) strategy was illustrated and the desired transformant contained the donor plasmid integrated into endogenous FoSOD5 locus with the hygromycin cassette as the selective marker. B) Schematic diagram of three different FoSod5 constructs. C1: using sGFP to replace the SOD domain and GPI site (SP+sGFP); C2: using sGFP to replace the SOD domain (SP+sGFP+GPI); C3: using lacZ to replace the whole ORF of FoSOD5 gene. C) Western blotting analysis showed that FoSod5 could be only associated with fungal cells. Coomassie protein staining indicated the same volume of loading samples.
lacZ as a suitable reporter gene was used for *in vivo* analysis of FoSOD5 expression under different environments in *F. oxysporum*.

A) The pH value influences FoSOD5 gene expression. The minimum medium (M-100) with X-gal and a series of different pH values were used and FoSod5C3 transformant conidia were dropped onto the M-100 medium. The agar color changes were monitored. Under the alkaline environment (pH = 7 or 8), was FoSOD5 gene could be rapidly induced. B) Expression of FoSOD5 was regulated by various carbon sources C) the nutrient medium TB3 could repress FoSOD5 gene expression. D) the lacZ gene showed variously up-regulated expression under different chemical stimuli.
**Figure 4.5** FoSod5C2 subcellular localization under confocal microscopy

A-B) FoSod5C2 subcellular localization in YG medium
C-E) FoSod5C2 subcellular localization in M-100 medium.
**Figure 4.6**  GPI site was required for FoSod5 subcellular localization and ROS treatments allowed FoSod5 to localize to cell walls and septa in YG medium.

A) phialide localization of FoSod5C1 and FoSod5C2; B) septum localization of FoSod5C2 and FoSod5C2 with treatments of H$_2$O$_2$ and superoxide (xanthine oxidase / hypoxanthine); C) Statistical analysis of phialide localization among FoSod5C1 and FoSod5C2; D) Statistical analysis of septum localization among FoSod5C2 and FoSod5C2 with treatments of H$_2$O$_2$ and superoxide (xanthine oxidase / hypoxanthine)
Figure 4.7  Assessment of the role of the FoSOD5 gene in virulence on the natural host.
A) Pathogenicity assay using cotton seedlings B) Statistical analysis of the size of lesions caused by
WT, ΔFoSOD5, and ΔFoSOD5/FoSOD5 strains. C) The cotton symptoms caused by infections of WT,
ΔFoSOD5, and ΔFoSOD5/FoSOD5 strains and control group (Mock) D) Statistical analysis of the
disease index for WT, ΔFoSOD5, and ΔFoSOD5/FoSOD5 strains.
Figure 4.8 Amino acid multiple sequence alignment of the SOD domain from FoSod5, CaSod5, CaSod1 (C. albicans), FoSod1, and ScSod1 (Saccharomyces cerevisiae). Two ∆H sites represented the missing zinc binding histidine residues and the Δel represents the missing Sod1 electrostatic loop, suggesting FoSod5 is a Cu-only binding enzyme. R181 is a conserved site required for the activity of SODs.
Figure 4.9  The FoSOD5 gene replacement and complementation.
A) The ~1 kb upstream and ~1 kb downstream flanking sequences of FoSOD5 were amplified. These two fragments were then ligated adjacent to the hygromycin phosphotransferase (HPH) by overlapping PCR. B) Four pairs of primers were used to assess the correct inserted position of the HPH cassette. C) Southern blotting was used to confirm a single integration event occurred of the HPH cassette in the F. oxysporum genome. D) Southern blotting was used to confirm the single FoSOD5 gene cassette integrated into a ΔFoSOD5 strain.
**Figure 4.10** The DNA sequences of the target region of FoSod5C1, FoSod5C2, and FoSod5C3 transformants.

The red arrow represents the cleavage site of the Cas9 RNP.
Figure 4.11  The Z-stacks analysis of FoSod5C2 on M-100 medium under confocal microscopy.

c: conidia; p: phialide; h: hyphae; s: septum
### Table 4.1 Strains of *F. oxysporum* used in Chapter 4

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<tr>
<th>Strain</th>
<th>Genotype description</th>
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<td>Wild type (WT)</td>
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<td>ΔFoSOD5</td>
<td><em>FoSOD5</em> gene deletion from WT</td>
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<td>ΔFoSOD5/FoSOD5</td>
<td>The complementation strain for ΔFoSOD5</td>
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<td>EGT&lt;sup&gt;CRISPR&lt;/sup&gt;-based transformation of WT expressing SP+sGFP+GPI</td>
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<td>FoSod5C3</td>
<td>EGT&lt;sup&gt;CRISPR&lt;/sup&gt;-based transformation of WT expressing <em>lacZ</em> under the native <em>FoSOD5</em> promoter</td>
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SP: secreted peptide; GPI: glycosylphosphatidylinositol sites; EGT<sup>CRISPR</sup>: CRISPR/Cas9-mediated endogenous gene tagging system.
<table>
<thead>
<tr>
<th>Primer names</th>
<th>Oligomers sequences</th>
<th>Usage</th>
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