Siderophore Biosynthesis but Not Reductive Iron Assimilation Is Essential for the Dimorphic Fungus Nomuraea rileyi Conidiation, Dimorphism Transition, Resistance to Oxidative Stress, Pigmented Microsclerotium Formation, and Virulence

Yan Li, Zhongkang Wang, Xuee Liu, Zhangyong Song, Ren Li, Changwen Shao and Youping Yin*

Chongqing Engineering Research Center for Fungal Insecticides, School of Life Science, Chongqing University, Chongqing, China

Iron is an indispensable factor for the dimorphic insect pathogenic Nomuraea rileyi to form persistent microsclerotia which can replace conidia or blastospores for commercial mass production. There are two high affinity iron acquisition pathways in N. rileyi, siderophore-assisted iron mobilization and reductive iron assimilation systems. Transcription of the two iron uptake pathways related genes is induced under iron-limiting conditions. Stage-specific iron uptake-related genes expression during microsclerotia development shows siderophore-mediated iron acquisition genes are rigorously upregulated specifically during the formation and mature period while reductive iron assimilation related genes just display a higher expression at the late maturation period. Abrogation of reductive iron assimilation, by the deletion of the high affinity iron permease (NrFtrA), has no visible effect on microsclerotia biogenesis in N. rileyi. In sharp contrast, N. rileyi L-ornithine-N5-monooxygenase (NrSidA), required for synthesis of all siderophores, is absolutely necessary for the development of pigmented microsclerotia. In agreement with the lower intracellular iron contents of microsclerotia in ΔNrSidA strains, not only the pigments, but both the number and the biomass are also noticeably reduced. Certain concentration of ROS is required for promoting microsclerotia biogenesis. Combined with expression pattern analysis of related genes and quantitative of intracellular iron or extracellular siderophore in WT and mutants, these data demonstrate the lack of adequate intracellular iron caused by the loss of the siderophore results in the deficiency of ROS detoxication. Furthermore, ΔNrSidA strains show significantly increased sensitivity to hydrogen peroxide. Besides, NrSidA, but not NrFtrA, play a crucial role in vegetative growth under iron-limiting conditions, conidiation, and dimorphic switching. Remarkably, the slower growth of the ΔNrSidA strains in vivo due to a reduced capacity for iron acquisition.
INTRODUCTION

To control the damage of pests to agriculture, widespread and continuous use of chemical insecticides causes environmental problems and leads to the development of insect resistance. Microbial insecticides such as entomopathogenic fungi can provide an alternative, more environmentally friendly option. Unlike insect-pathogenic viruses, bacteria, and protozoa, which must be ingested to infect a host, most entomopathogenic fungi infect by penetrating the host cuticle (Srisukchayakul et al., 2010; Thakre et al., 2011). However, lack has a potential for development into mycoinsecticide in the rhizosphere (Zhong et al., 2014; Song et al., 2015). MS granules have been tested for the control of pests such as the sugar beet root maggots, *Tetanops myopaeformis*, Asian Longhorned Beetles (Coleoptera: Cerambycidae), and black-legged ticks, *Ixodes scapularis* Say (Jaronski and Jackson, 2008; Jackson and Jaronski, 2009; Behle et al., 2013; Goble et al., 2015). Although the ability to produce dry granules instead of conidia or blastospores represents an innovation for commercial interests, the mechanisms underlying the key developmental events of entomopathogenic fungi remain unresolved.

**MS development is controlled by a network of signaling pathways, which must occur in concert with other complex signaling events (Song et al., 2013).** To study the mechanisms of MS development, comparative transcriptome analysis of MS development in *N. rileyi* had been performed, showing that oxidative stress occurs during the physiological process (Song et al., 2013). Multiple genes of *N. rileyi* have been identified as involved in MS development (Song et al., 2013; Jiang et al., 2014; Liu et al., 2014; Zhou et al., 2015). **RacA** and **Cdc42**, small GTPases, are required for MS formation through the control of ROS generation (Jiang et al., 2014). Both NADH: flavin oxidoreductase/NADH oxidase gene (Nox) and alternative oxidase gene were proved to play a role in MS differentiation by regulating intracellular **H2O2** concentration (Liu et al., 2014; Zhou et al., 2015). Silencing mutants of two transmembrane proteins, Sho1p and Shn1p, resulted in severely reduced MS yields and conidiation production (Song et al., 2015). In addition, iron was found to be an indispensable for the MS formation. Furthermore, the yield of the MS increased with the rising iron concentrations of the fermentation broth within a certain range (Song et al., 2014). On the basis of previous results, further studies should be focused on the roles of the iron and related genes play in the MS formation.

As an essential nutrient, iron is required for various metabolic processes, including electron transport and redox reactions in nearly all organisms (Haas et al., 2008). Fungi have evolved various strategies, often used in parallel, to acquire iron. More detailed information on the iron uptake mechanisms is available in the model fungus, *Aspergillus fumigatus*. *A. fumigatus* possesses two high affinity iron uptake mechanisms: reductive iron assimilation (RIA) and siderophore-mediated iron uptake (Schrettl et al., 2004). **RIA** involves reduction of ferric to ferrous iron and subsequent uptake of ferrous iron by the *P*trA/FsEC complex, an activity that is blockable with the...
ferrous iron-specific chelator bathophenanthroline disulphonate (BPS) (Haas, 2003; Schrettl et al., 2007). Siderophores are low molecular mass, ferric iron-specific chelators, which are excreted during iron starvation (Haas, 2003). The majority of fungal produce hydroxamate-class siderophores (Haas, 2014) but are able to take up other types of siderophores as well, such as Saccharomyces cerevisiae (Phippott et al., 2002). A. fumigatus produces four hydroxamate siderophores all based on L-ornithine-N\textsuperscript{3}-monooxygenase (SidA) (Moore, 2013), which catalyzes both intra- and extracellular siderophore biosynthesis at initial biosynthetic step (Schrettl et al., 2007). And then the formation of the hydroxamate group is accomplished by N\textsuperscript{5}-acylation of N\textsuperscript{5}-hydroxornithine with the N\textsuperscript{3}-transacylases (SidF) (Schrettl et al., 2007; Haas et al., 2008). SidF homologs can be found among hydroxamate-producing fungi and numerous bacterial species, for example, the Escherichia coli homolog IucB for synthesis of siderophore aerobactin (de Lorenzo et al., 1986; Schrettl et al., 2007; Haas et al., 2008) and the potential siderophore-biosynthetic transacylase Fer5 from Ustilago maydis (Eichhorn et al., 2006). After the addition of an acyl moiety to N\textsuperscript{5}-hydroxornithine, the hydroxamates are covalently linked via ester or peptide bonds to form the final siderophore accomplished by nonribosomal peptide synthetases (NRPSs) (Haas et al., 2008). Different NRPSs appear to be responsible for producing different siderophores. In A. fumigatus, SidC or SidD takes part in the biosynthesis of intracellular siderophores FC or extracellular siderophores TAFc, respectively (Schrettl et al., 2007). Siderophore-iron chelates are usually taken up through transporters of the UMF/SIT subfamily of the major facilitator superfamily (Haas et al., 2008). In S. cerevisiae, the transporter Sit1p/Arn3p recognizes the bacterial hydroxamate ferrioxamine B, coprogen, and a variety of ferrichromes. S. pombe Str2 or Str1 has been suggested to transport ferrichrome and ferrioxamine B or ferrichrome, respectively (Ardon et al., 2001; Pelletier et al., 2002).

The function of siderophore-mediated iron acquisition and RIA in entomopathogenic fungi is not well understood. In N. rileyi, iron stimulates the formation of MS at some range of concentrations and transcriptome analysis revealed that most of the genes involved in iron uptake were found to be upregulated during MS development (Song et al., 2013, 2014). Hence, a deeper understanding of the role of iron and its uptake related genes play in the corresponding physiological process will facilitate to guide the liquid fermentation of MS in N. rileyi. In present study, we mainly analyzed the roles of two essentially important genes (NrSidA and NrFtrA) played in MS formation, spore production, dimorphism transition and the virulence against Spodoptera litura and the influence on the transcriptional regulation of several genes implicated in MS formation.

**MATERIALS AND METHODS**

**Strains, Media, and Culture Conditions**

N. rileyi CQNnr01 was isolated from cadavers of Spodoptera litura infected naturally and stored at the Engineering Research Center for Fungal Insecticides, Chongqing University, China. The fungal strain and engineered strains generated in this study were grown on Sabouraud maltose agar with 1% yeast extract (SMAY) under continuous illumination at 25°C for 14 days to produce conidia. The conidia suspensions were prepared in sterile 0.5% TWEEN 80 (Sigma-Aldrich, USA). The spores of N. rileyi was incubated in liquid-amended medium (AM) (containing 40 g/L of glucose, 2.5 g/L of peptone, 5 g/L of yeast extract, 4.0 g/L of KH\textsubscript{2}PO\textsubscript{4}, 0.8 g/L of CaCl\textsubscript{2}·2H\textsubscript{2}O, 0.6 g/L of MgSO\textsubscript{4}·7H\textsubscript{2}O, 0.1 g/L of FeSO\textsubscript{4}·7H\textsubscript{2}O, 37 mg/L of CoCl\textsubscript{2}·6H\textsubscript{2}O, 16 mg/L of MnSO\textsubscript{4}·H\textsubscript{2}O, and 14 mg/L of ZnSO\textsubscript{4}·7H\textsubscript{2}O) at 28°C under shaking at 250 rpm for MS production. Phenotypic characterizations were carried out using strains grown on Aspergillus minimal medium (AMM) amended with 4% (wt/vol) maltose as carbon source, 20 mM glutamine as nitrogen source, and 10 μM FeSO\textsubscript{4}·7H\textsubscript{2}O or FeCl\textsubscript{3}·7H\textsubscript{2}O when required. For iron-depleted conditions, iron was omitted.

**Gene Cloning and Bioinformatic Analyses**

To study the potential functions of NrSidA and NrFtrA on the MS development, the full cDNA sequences were acquired using fusion primer and nested integrated PCR (FPNI-PCR) and RACE based on the ESTs from the transcriptome (Liu and Chen, 2007; Wang et al., 2011; Song et al., 2013). Both deduced amino acid sequences were used for Blastp searches against GenBank. The homologs of the two genes from other fungal species were collected from the NCBI database for multiple sequence alignment analysis separately with ClustalX (Larkin et al., 2007) and a neighbor-joining tree was generated with 1000 bootstrap replicates using the program MEGA 7.0 (Tamura et al., 2013).

**Generation of NrSidA and NrFtrA mutants**

For inactivation of NrSidA and NrFtrA through homologous recombination, respective flanking regions were obtained from genomic DNA by FPNI-PCR (Wang et al., 2011) with the primers listed in Table S1 and inserted in to pMD19-T vector for sequencing.

To generate the NrSidA and NrFtrA disruption vectors, upstream and downstream flanking sequences were obtained by PCR with primers listed in Table S1. The resulting upstream flanking sequence of the NrSidA or NrFtrA was digested with EcoRI/XhoI or BamHI/XbaI and with XbaI/HindIII or PstI for the downstream flanking sequence and then inserted into pPZP-Hph vector harboring a hph cassette to yield pPZP-Hph-NrSidA or pPZP-Hph-NrFtrA, respectively. The resulting clones were introduced into Agrobacterium strain AGL1 for fungal transformation. Agrobacterium tumefaciens mediated transformation (ATMT) of N. rileyi CQNnr01 with pPZP-Hph-NrSidA or pPZP-Hph-NrFtrA was performed as described elsewhere (Shao et al., 2015).

Transformants resistant to the hygromycin were selected. Initial screening of transformants for the disruption of the NrSidA or NrFtrA gene was carried out by PCR with the primers listed in Table S1. Gene deletion was further assessed in single conidial progenies by Southern analysis according to the instruction of the DIG High Prime DNA Labeling and Detection Starter Kit I (Roche, Mannheim, Germany). All primers used in construction and verification of engineered strains are listed in Table S1.
Quantification of Intracellular Iron in MS and LIP Determination in Yeast-Spores

Intracellular iron concentration in MS was measured using the BPS-based colorimetric assay (Epsztejn et al., 1997; Tamarit et al., 2006) with modifications. Briefly, 3.5-day-old MS was harvested by filtration, washed three times with ddH$_2$O, ground with liquid nitrogen, weighed after vacuum freeze drying and resuspended in 1000 µL of 3% nitric acid. Suspensions were boiled for 8 h and centrifuged at 12,000 g for 15 min to discard cell debris. A total of 100 µL of supernatant was diluted 10-folds (except the MS cultivated in AFM-Me medium) and then 400 µL of supernatant diluent was mixed with 160 µL of sodium ascorbate, 320 µL of BPS, and 126 µL of ammonium acetate, and reactions were incubated at room temperature for 5 min. OD$_{535}$ of the BPS-Fe complex was measured with the DU®640 Spectrophotometer (Beckman Coulter, USA). To eliminate the nonspecific absorbance, OD$_{680}$ was subtracted from OD$_{535}$. Iron was quantified taking into account a calibration curve using seven solutions with known FeSO$_4$ concentrations (0.02–0.16 mM; y = 0.127x-0.003; R$^2$ = 0.996), and the results were expressed as µmol g$^{-1}$ dry weight. Three replicates were performed, and each replicate was measured in triplicate.

The labile iron pool (LIP) in yeast-spores is measured by using calcein, a fluorescent probe that binds to both Fe$^{2+}$ and Fe$^{3+}$ (Epsztejn et al., 1997). The binding of Fe quenches the fluorescence of calcein. After cellular uptake of calcein-AM, intracellular esterases convert the molecule into a non-permeable acid-form of calcein (Cabisco et al., 2002). N. rileyi strains yeast-spores were obtained after growing conidiophores on the SMAY for 4 days. Yeast-spores were washed twice with phosphate-buffered saline (PBS), incubated for 3 h at 37°C in 5 µM Calcein-AM (a final concentration) in PBS, and then washed twice for 5 min with PBS to remove unincorporated dye. Fluorescence was observed (excitation and emission at 485 nm and 530 nm, respectively) under a Nikon fluorescence microscope (Nikon, Japan).

Chrome Azurol S (CAS) Liquid Assay and Quantification of Siderophores

Extracellular siderophores were determined using the chrome azurol S liquid assay as described (Schwn and Neilands, 1987; Payne, 1994; Louden et al., 2011). One hundred microliter of the CAS solution (10x) was mixed with 900 µl the culture filtrates from the 3.5-day-old MS growing in AM. Siderophore production was further measured by a Csaky test (Csaky, 1948). 0.5 mL of culture filtrates were mixed with 2.5 mL of iron (III) perchlorate (5 mM) dissolved in 0.1 M perchloric acid and measured spectrophotometrically for absorbance at 480 nm.

Purification and Measurement of MS Pigments

The MS melanin/pigment was isolated from 3.5-day-old cultures with 2%NaOH. The collected MS was ground with liquid nitrogen, suspended in NaOH solution (1 mL) and boiled at 100°C for 2 h. The solution was acidified to pH 2.0 with 5 M HCl and centrifuged at 6000 g for 15 min. The resulting melanin/pigment was dissolved in 2% NaOH, and the solution was measured spectrophotometrically for absorbance at 459 nm (Babitskaya et al., 2000).

Phenotypic Analyses

The sensitivity to NaCl, sorbitol, sodium dodecyl sulfate (SDS), calcofluor white (CFW), congo red (CR) and various high concentration metal ions including CaCl$_2$, MnSO$_4$, ZnSO$_4$, and CuSO$_4$ was tested in AMM plus 1.8% agarose as the solidifying agent. Conidial suspensions (5 × 10$^5$ conidia) were pipetted onto plates and incubated at 25°C and inspected regularly for 14 days for colony morphology and diameter. Heat tolerance was determined according to Liu et al. (2010).

The sensitivity of conidia to killing by hydrogen peroxide was assayed under different consideration iron. As mentioned above, 5 × 10$^5$ conidia of WT and mutants were point inoculated and radial growth was measured after 14 days at 25°C on AMM medium without iron (−Fe) or with 10 µM FeSO$_4$ (Fe) or 1.5 mM FeSO$_4$ (hFe), respectively, and with the addition of 0.25 mM bathophenanthroline disulfonate (BPS) or/and 2 mM H$_2$O$_2$. The AMM medium without H$_2$O$_2$ was as a control.

The sensitivity of conidia to hydrogen peroxide with different concentrations was assayed as described by Schrätti et al. (2007). Briefly, conidial suspensions approximating 10$^9$ conidia ml$^{-1}$ were incubated for 30 min at room temperature with varying concentrations of hydrogen peroxide. To determine the number of surviving conidia, the spore suspensions were diluted 50-fold and plated on SMAY. Following incubation for 6 days at 25°C, colonies were counted and normalized to that without hydrogen peroxide treatment. The sensitivity of hyphae to hydrogen peroxide was estimated by using a modification of the protocol of Kawasaki et al. (1997). Approximately 100 conidia were plated on SMAY medium and grown at 25°C for about 8 day for the countable colonies formation and at this time each colony grew out the fungal mycelia. Subsequently, the plates were overlaid with 8 ml of the same medium as top-agar but containing the indicated concentration of hydrogen peroxide. After further incubation for 5 days at 25°C, colonies able to resume growth were counted as survivors and normalized to the number before hydrogen peroxide treatment.

Sporulation rates were tested using 5 × 10$^5$ conidia on AMM amended with 0, 10 µM and 1.5 mM FeCl$_3$ or FeSO$_4$, respectively. The plates were incubated at 25°C for 14 days for sporulation and the numbers were counted. Besides, the sporulation rates were also recorded at intervals of 3–11 days using an equal number conidia on SMAY.

Germination tests were conducted using 1 × 10$^5$ conidia on SMAY or AMM amended with (250 µM) BPS, 0 or 30 µM FeSO$_4$. Cultures were incubated at 25°C and the conidial germination rates at various time-points were determined by counting 150 conidia. The experiment was performed three times; each involved three replicates.

N. rileyi displays a dimorphic switching, using the yeast cell form to avoid phagocytosis and the cytotoxic environment of the phagolysosomal system (Wanchoo et al., 2009; Boyce and Andrianopoulos, 2015). In vitro, N. rileyi can grow in yeast-cell form at the early stage (3–6 days) on SMAY and then...
gradually transform into filamentous form. Switching rates from yeast-cell to hyphal were studied using $1 \times 10^5$ yeast-cells on SMAY and regularly observed for colony morphology. In addition, approximately 80 simple yeast-cells were plated on SMAY medium, grown at 25°C and the switching rates at various time-points were recorded. The median transition rates time (TT$_{50}$) of the mutants were compared with WT.

**Insect Virulence Assays**

Virulence of the WT, $\Delta NrSidA$ and $\Delta NrFtrA$ was assayed against newly emerged second-instar *Spodoptera litura* larvae. Conidia from SMAY plates were applied topically on the abdominal dorsum in a mineral oil suspension containing $1 \times 10^8$ conidia ml$^{-1}$. Each treatment had three replicates with 15 insects each, controls were treated with mineral oil only, and the experiments were repeated three times. For injection assays, each insect was injected from the second proleg with $10^7$ conidia ml$^{-1}$. The number of dead insects was recorded daily for 10 days. The median lethal time (LT$_{50}$) was calculated using a probit analysis with the SPSS program.

Hyphal bodies represent a yeast-like fungal cell type produced in the insect hemolymph after hyphal penetration through the integument to evade the insect immune system (Wanchoo et al., 2009). For collection of hemolymph, infected and control larvae were anesthetized on ice and the rear leg was cut off with a scissor at 72, 96, and 132 h after injection with $10^7$ conidia ml$^{-1}$. The exuded hemolymph from the wound was collected. Fungal hyphal bodies present in the insect hemolymph were observed via bright-field microscopy (Olympus, Japan), and the number of hyphal bodies was determined by direct counting using a hemocytometer. Three replicates were performed for each treatment, and each replicate contained three randomly picked larvae.

**Gene Expression by Quantitative Reverse Transcription Polymerase Chain Reaction (qRT-PCR)**

In order to induce iron starvation, conidia from the WT were grown in liquid AMM lacking iron ($-\text{Fe}$) for 2 days, and then transferred to fresh AMM containing 0, 0.03 or 1.5 mM FeSO$_4$, or 0.25 mM BPS for an additional day. Subsequently, the mycelia were harvested for RNA extraction. For time-specific expression patterns during MS development, $10^9$ conidia of the WT, $\Delta NrSidA$ and $\Delta NrFtrA$ were inoculated in AM and the cultures were collected by filtration and washed three times with sterile distilled water at 36, 48, 60, 72, 84, 98, 120, and 144 h for RNA extraction. To study the effects of the different consideration iron on growth and formation of MS, $10^9$ conidia of the WT, $\Delta NrSidA$ and $\Delta NrFtrA$ were inoculated in AM for 3.5 days under different iron concentrations and collected for RNA extraction. To explore the impacts of the hydrogen peroxide on the pathways of the iron uptake in MS, 3.5-day-old MS was treated with varying concentrations of hydrogen peroxide for 30 min and then collected for transcriptional analysis.

Total RNA from *N. rileyi* was isolated using Trizol reagent (TAKARA, Japan) and then treated with DNase I (Thermo Fisher Scientific, USA). cDNA of each sample was generated using RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher Scientific, USA). Amplification mixtures (20 µl) for qRT-PCR contained 2 µl template cDNA, 10 µl 2 × IQ SYBR Green Supermix (Bio-Rad, USA) and 1 µl of each primer (PF/PB, 10 µmol l$^{-1}$). The reaction profile was performed as an initial 95°C for 3 min, followed by 42 cycles of 95°C for 10 s, appropriate annealing temperature for 20 s and 72°C for 20 s. PCR reaction was performed using the iCycler system (CFX, Bio-Rad) and primers listed in Supporting Information Table S1. qRT-PCR efficiency was determined by 10-fold gradient dilutions of cDNA of each target sequence for standard curve production. Under the optimal annealing temperature, the calculated efficiency of all primers was 95–105%. Melt curve analyses were carried out to confirm the absence of the nonspecific products. Relative expression of each gene was normalized against the expression of a $\beta$-tubulin gene (TUB) and a transcription elongation factor (TEF). The resulting data were analyzed with the CFX qPCR software. All PCR amplifications were conducted in triplicate, and trials were repeated three times.

**Data Analysis**

Unless otherwise indicated, all data were analyzed by one-way ANOVA, followed by Duncan’s Multiple Range test using the SPSS 17 program (SPSS 17.0, SPSS Inc., USA). Uppercase or lowercase letters indicated the statistically significant level at $P < 0.01$ or $P < 0.05$, respectively. Means indicated by the same letter are not significantly different from one another.

**RESULTS**

**Molecular Characterization of *NrSidA* and *NrFtrA***

Two fragments that showed strong sequence similarity to fungal genes encoding L-ornithine-N$^3$-monooxygenase (SidA) or a high-affinity iron permease (FtrA) separately were identified from the expression profiles due to the up-regulated expression in the transcriptome of MS. Both entire cDNA sequences were amplified by FPNI-PCR and RACE and designated as *NrSidA* (GenBank accession number: KX181541) or *NrFtrA* (accession number: KX181542), respectively. The putative coding sequences of *NrSidA* and *NrFtrA* were 1863 and 1080 bp, encoding 620 and 359 amino acid residues with a calculated molecular weight of 66.97 and 38.84 kDa and an isoelectric point of 5.85 and 8.89, respectively (http://expasy.org/tools/protparam.html). The comparison of cDNA and genomic sequences revealed the presence of two introns in both genes. Both *NrSidA* and *NrFtrA* expressions are under the SreA-mediated regulation (Oberegger et al., 2001) as the promoter regions contained several HGATAR motifs (data not shown), which represent putative binding sites for GATA-factors such as SreA.

The amino acid sequence of *N. rileyi* NrsidA showed very high identity to sidA from *M. anisopliae* (81%) and *B. bassiana* ARSEF 2860 (65%) and also displayed 50 and 48% identity to SidA from *A. fumigatus* and *A. nidulans*, respectively (Figure S1A).
The *N. rileyi* NrSidA sequence contained the three signature sequences typical of amino acid hydroxylase enzymes. The first of these is the flavin adenine dinucleotide binding (FAD) motif GXGXGG (*N. rileyi* CQNr01 159CVGFGP 164) and the last glycine in this motif of NrSidA was exchanged for proline, which is a typical feature of siderophore biosynthetic enzymes (Stehr et al., 1998). Another typical feature for siderophore biosynthetic genes was a putative nicotinamide adenine dinucleotide phosphate (NADP)-binding motif GXGXX(G/A) (365GAGQSAA 371). Besides, the conserved motif D(X)5(L/F)ATGY(X)4(H/P) (512DLV1ATGYQRN AH 525) was proposed to be involved in substrate binding (Eisenindle et al., 2003). Phylogenetic analysis indicated that NrSidA protein had the closest relationship with that from the entomopathogenic fungus *Metarhizium* spp. (Figure S1D).

NrFtrA showed typical features of a high affinity iron permease protein. NrFtrA protein, the same as the *S. cerevisiae* iron permease Ftr1, contained seven transmembrane domains (Figure S1C; Kwok et al., 2006) and two motifs of REXXE, Askwith et al., 1994; Stearman et al., 1996; Kwok et al., 2006). The glutamic acid residue in REXXE motif, which was interact directly with iron, was conserved in the selected FTR protein sequences from other organisms (Figure S1B). With respect to amino acid similarity, NrFtrA displayed 59% and 48% identity to FtrA from *A. fumigatus* and Ftr1 from *R. oryzae*, respectively. Phylogenetic analysis indicated that NrFtrA protein was closely related to that from the entomopathogenic fungus *Metarhizium* spp., while having rather distant phylogenetic relationships to *R. oryzae* (Figure S1E).

The transcriptional response of two high affinity iron uptake pathways related genes of *N. rileyi* to iron availability was investigated by qRT-PCR in the AMM liquid medium. Under iron-limiting conditions (—Fe or BPS), *N. rileyi* wild-type strain CQNr01 induced the expression of genes involved in the biosynthesis of siderophores (*NrSidA*, *NrSidC*, *NrSidF* and *NrSidD*), uptake of siderophores (*NrSit1p* and *NrStr3*; Heymann et al., 2002; Pelletier et al., 2002) and RIA (*NrFtrA* and *NrFet3*), a multicopper oxidase; Ashkith et al., 1994; Stearman et al., 1996; Kwok et al., 2006; Figure S1F). High iron concentrations (30 µM or 1.5 mM FeSO4) efficiently down-regulated *NrSidD* and RIA genes transcripts at concentrations but led to increased transcript levels of the *NrSidA* and *NrSidC* genes required for synthesis of intracellular storage siderophores and control of intracellular iron homeostasis. As expected, under iron depletion conditions, transcripts of the GATA transcription factor NrSre, a putative repressor of genes involved in iron uptake (Chao et al., 2008; Haas, 2012), were present at low concentration while bZIP-type regulator NrHapX, required for adaption to iron starvation (Schrett et al., 2010; Gisaller et al., 2014), showed high transcript levels (Figure 1). Increasing concentrations of FeSO4 to 30 µM or 1.5 mM increased transcript abundance of *NrSre* and decreased transcript concentration of *NrHapX*. *NrSre* and *NrHapX* had a strong response to iron in exactly the opposite way: iron abundance induced *NrSre* expression in wild-type *N. rileyi*, whereas iron-poor conditions induced *NrHapX* expression when grown in the AMM liquid medium (Figure 1).

**Identification of Deletion Mutants**

Through homologous recombination and *Agrobacterium tumefaciens* mediated transformation, *NrSidA* and *NrFtrA* were individually disrupted by replacement with the hygromycin b phosphotransferase (hph) cassette (Figures S2A,B). Several mutant candidates for each of the targeted genes were identified by PCR screening. After single spore isolation, single integration of the replacement constructs were verified by PCR and Southern blot analysis (Figures S2C,D). Finally, two disruption mutants were obtained for *NrsidA* and three for *NrFtrA*. Both Δ*NrSidA* strains displayed the same phenotype and all three Δ*NrFtrA* strains showed the same features.

**NrSidA Is Required for the Biogenesis of Pigment MS by Regulating the Siderophore Production and Iron Uptake**

To examine whether both *NrSidA* and *NrFtrA* participate in growth and MS production, conidia from the WT and mutants were inoculated in AM for MS biogenesis. After 3.5 days of induced culturing, the WT and Δ*NrFtrA* strains generated MS normally and accumulated melanin in MS, whereas the Δ*NrSidA* strains developed few MS and accumulated less melanin (Figure 2A). To ascertain whether a defect of the iron uptake caused by *NrSidA* deletion led to deficiencies in MS, conidia of the WT and mutants were inoculated in AM at different iron supplies for 3.5-days induction culture as before. Under iron deletion condition (AM-Fe), both WT and Δ*NrFtrA* mutants formed few MS and accumulated less melanin. By contrast, the Δ*NrSidA* mutants were not only pigment deficient, accumulating less melanin than WT and Δ*NrFtrA* mutants (Figures 2A,B,E) but also failed to form compact aggregations of hyphae and accumulate significant biomass (Figure 2B). With a increase in the iron concentration of AM, although the hyphae of MS in the Δ*NrSidA* strains was compact aggregations and the biomass were comparable to that of WT, the number of MS was still distinctly less than that in WT and Δ*NrFtrA* (Figures 2C,D). Furthermore, the pigments in the WT and Δ*NrFtrA* mutants could fully recover when supplemented with 100 mg/L FeSO4, and increased as the concentrations of iron rose. However, increasing the concentrations of FeSO4 to 400 or 600, even to 800 mg/L in AM just restored a portion of pigments in Δ*NrSidA* (Figures 2A,B,E), suggesting that *NrSidA* might play a vital role in the MS development especially in pigment biosynthesis.

To understand how the *NrSidA* and the *NrFtrA* genes affect the MS formation, extracellular siderophore and intracellular iron contents were measured. Under iron depletion condition, a large amount of extracellular siderophore was detected in both the Δ*NrFtrA* and WT, and then a decline appeared followed by the increase of the concentrations of iron in AM. In sharp contrast, no siderophore production was evident in the Δ*NrSidA* strains regardless of whether the iron was sufficient or deficient (Figures 3A,B). Remarkably, a much larger amount was detected in the Δ*NrFtrA* strains under iron deficient condition compared with WT (Figure 3B), indicating that Δ*NrFtrA* strains had to excreted more siderophores to remedy the missing of RIA. Meanwhile, the intracellular iron contents in different stages of
Li et al. Siderophore Is Essential for Nomuraea

FIGURE 1 | Relative transcript abundance of iron uptake-related genes. Relative transcript abundances of iron uptake-related genes at different iron supplies were measured by qRT-PCR. TEF and TUB transcripts were used as reference. Uppercase or lowercase letters indicated the statistically significant level at $P < 0.01$ or $P < 0.05$, respectively; the same letter were not significantly different from one another. Error bars are standard deviations of three trials.

the MS development and in formation period (3.5 days) with different iron concentrations were measured to evaluate the loss of siderophore on the ability of WT and mutants to acquisition and store of iron. As the MS of $N.rileyi$ was developing, the intracellular iron contents in the $\Delta NrFtrA$, as well as in the WT strains, gradually rose whereas the increase in the $\Delta NrSidA$ strains was slower (Figure 3C). Likewise, the intracellular iron contents in the $\Delta NrFtrA$ and WT strains rose dramatically as the iron concentrations increased whereas $\Delta NrSidA$ strains showed a slower increase (Figure 3D). A analogous result was found in the labile iron pool determination by the fluorescent iron dye calcein that much less free iron was tested in the yeast-spores of the $\Delta NrSidA$ rather than in the $\Delta NrFtrA$ strains (Figure S3).

Loss of $NrSidA$, but Not $NrFtrA$, Leads to Defects in Mycelial Growth and Conidiation, Delaying Conidial Germination and the Dimorphic Transition

When grown on AMM amended with 0, 10 µM FeCl$_3$ or FeSO$_4$, the $\Delta NrSidA$ mutants were defective in sporulation while the $\Delta NrFtrA$ mutants behaved like WT (Figure 4A). The inability to sporulate only partially restored by the addition of 1.5 mM FeSO$_4$, but not FeCl$_3$. But the amount was still 100 folds less than that of WT and $\Delta NrFtrA$ mutants. Meanwhile, the dimorphic transition of these strains on SMAY medium was observed. The $\Delta NrFtrA$ mutants were comparable to that of WT: yeast-cell began to switch to mycelial and sporulated on the 5th day (Figures 4B,C). In comparison, the $\Delta NrSidA$ mutants showed a much slower switching: the conversion of yeast-cell to hyphae was present on the 9th day and mycelial phase started to sporulate on the 11th day, resulting in prolonging the yeast-to-mold conversion (Figures 4B,C). To eliminate the effects of the initial inoculum size on the morphologic switch, single yeast cell of the WT and mutants was grown on the SMAY medium. The similar results were showed in Figures 4D,E that the differences between WT (TT$_{50} = 7.2 \pm 0.1$ days) and $\Delta NrSidA$ mutants (TT$_{50} = 9.1 \pm 0.4$days) were significant with no obvious differences between WT and $\Delta NrFtrA$ mutants ($P < 0.001$), which proved once more that a delay in the dimorphic transition was indeed caused by $NrSidA$ deletion.

The effects of $NrSidA$ and $NrFtrA$ genes disruption on conidial germination were investigated on SMAY or AMM medium. $\Delta NrSidA$ exhibited a significantly decreased germination either on SMAY or AMM compared with that of WT (Figure 5). When grown on SMAY medium, the germination efficiency was distinctly delayed while the rate of $\Delta NrFtrA$ was the same as that in WT (Figure 5A). After 24 h of incubation, about 20% of $\Delta NrSidA$ conidia had germinated compared with 64 or 68% of WT or $\Delta NrFtrA$ conidia, respectively. Similarly,
after 56 h on AMM-Fe media, the germination rate was about 18% for ΔNsidA conidia but 65 or 45% for WT or ΔNftrA conidia, respectively (Figure 5C); when supplemented with 30 μM FeSO₄, the germination rates at 56 h were increased to 29% for ΔNsidA conidia while 74% or 60% for WT or ΔNftrA conidia, respectively (Figure 5D). Nevertheless, the ΔNsidA conidia under iron depletion in the presence of BPS nearly failed to germinate and after 56 h, the germination rates only reached 4% compared to 55 or 43% for WT or ΔNftrA, respectively (Figure 5B). These observations suggested that siderophore-mediated iron storage or utilization of intracellular iron is required for efficient growth initiation under iron limitation.

**ΔNsidA Mutants, but Not ΔNftrA Are Hypersensitive to Iron Starvation, Oxidative Stress and Other Abiotic Stresses**

The resistance of WT and mutants to iron starvation was studied under iron-sufficient and deficient conditions. In all assays performed as before, the growth rates of both genes mutants were comparable to that of WT except that the growth
rate in ΔNrSidA was reduced significantly under iron-depleted conditions (AMM+BPS or AMM+Fe+BPS) (Figure 6A).

The effects of the other high concentration metal ions on the growth of the WT and mutants were also investigated. The susceptibility of high concentration CaSO₄, MnSO₄ and CuSO₄ ions to both genes mutants as well as to WT except zinc (Figure S4B). When treated with 10 mM ZnSO₄, ΔNrSidA was significantly more sensitive to zinc ion than ΔNrFtrA. As the concentration of ZnSO₄ increased 2.5-fold or 5-fold, these mutants were as sensitive as WT (Figure S4B), suggesting that zinc appeared to affect iron uptake by interfering with RIA pathway.

In addition, the resistance of NaCl, sorbitol and heat to these mutants was as well as to WT when grown on normal medium and non-biotic stress but the ΔNrSidA mutants were slightly sensitive to the cell wall disturbing agent (SDS, CFW and CR) (Figure S4).

Inappropriate iron storage can catalyze formation of reactive oxygen species (ROS) and detoxification of hydrogen (peroxide) depends on iron because catalases and peroxidases require heme as cofactor (Schrettl et al., 2007). Therefore, the impact of siderophores on the resistance to oxidative stress was studied. As showed in (Figures 6A,B) deficiency of siderophores (ΔNrSidA) caused hypersensitivity to H₂O₂ during iron-depleted growth. With the increase of extracellular iron availability, the defect was cured slightly, indicating the major role of siderophore is efficient iron utilization rather than iron detoxification (Figures 6A,B).

To further verify the speculation as mentioned above, the effects of hydrogen peroxide on conidial and hyphal survival were tested with these mutants in more details (Figures 6C,D). Whether conidia or hyphae of N. rileyi strains killed by hydrogen peroxide, ΔNrSidA mutants were significantly more sensitive than WT, implying that siderophore played an important role in the oxidative-stress resistance of N. rileyi conidia and hyphae.

**NrSidA Is Required for Full Virulence Against S. litura Host**

To examine the effects of NrSidA and NrFtrA deletion on the pathogenic ability of N. rileyi, insect bioassays by both topical infection and injection of fungal spores into S. litura larvae were performed. The values of median lethal time (LT₅₀) were estimated and compared among the WT, ΔNrSidA and ΔNrFtrA mutants. After topical infection, a significant difference was found between the WT (LT₅₀ = 5.97 ± 0.29 days) and ΔNrSidA (LT₅₀ = 11.52 ± 0.85 days) (P < 0.001) but not
Li et al. Siderophore Is Essential for Nomuraea

**FIGURE 4** | Quantitative analysis and comparison of conidia and the dimorphic transition of *N. rileyi* WT and mutants. (A) Quantitative analysis of the conidia produced by *N. rileyi* WT and mutants on AMM medium at different iron supplies. (B) Conidial production on SMAY medium every 2 days. (C) The dimorphic transition of the *N. rileyi* WT and mutants grown on the SMAY. Equal yeast-cells of the WT and mutants were pipetted onto plates and recorded every 2 days. Bar = 5 mm. (D) Quantitative analysis of the dimorphic transition (from yeast-cell to hypha) rate with approximately 80 single yeast-cells plated on the SMAY medium. The growth morphology were observed every half day or a day. (E) The median transition rates time (TT$_{50}$) of *N. rileyi* WT and mutants were compared. TT$_{50}$ was calculated using a probit analysis with the SPSS program.

between the WT and ΔNrFtrA (LT$_{50}$ = 6.55 ± 0.24 days) (Figures 7A, C). Similar results were observed in injection assays that the differences between the WT (LT$_{50}$ = 4.89 ± 0.22 days) and ΔNrSidA (LT$_{50}$ = 9.32 ± 0.52 days) were significant with no obvious differences between the WT and ΔNrFtrA (P < 0.001) (Figures 7B, D), demonstrating that it is NrSidA, but not NrFtrA that is absolutely required for *N. rileyi* virulence.

*S. litura* hemolymph was investigated microscopically and hyphal body production was also quantified over time after inoculation with the mutants and WT conidia. Both ΔNrFtrA and WT conidia escaped from insect immune and started yeast-type budding to multiple hyphal-bodies 72 h following the injection of fungal spores. However, the ΔNrSidA spores were continuously attacked and encapsulated by insect haemocytes and only a few yeast-like cells were observed (Figure 8). After 96 h, ΔNrSidA evaded insect immune defense and started to multiply in the insect haemocoel (Figure 8A). After 132 h, ΔNrSidA hyphal-bodies in insects increased rapidly but the average number remained 20-fold less than that of WT and ΔNrFtrA mutants (Figure 8B). In both bioassays, almost all the *S. litura* larvae died of the infection of the WT and ΔNrFtrA strains were before pupation. By contrast, most death by the ΔNrSidA strains were in the stage of the pupae (Figure 7E). Furthermore, insect cadavers were fully covered by the WT and ΔNrFtrA fungal spores or mycelia post-insect death, but these were not observed in ΔNrSidA-treated insects. Thus, abrogation of *N. rileyi* siderophore biosynthesis by NrSidA deletion prevents initiation of *S. litura* infection, which cannot be supported by RIA alone. Consistently, inactivation of RIA by deletion of NrFtrA is inconsequential for virulence.

**Loss of Both NrSidA and NrFtrA Alters Iron-Related Genes Expression, but Only the NrSidA Deletion Changes the ROS Detoxification at Transcription Level during MS Development**

To better understand the contribution of two high affinity iron uptake pathways required by MS production, we investigated stage-specific expression pattern analysis of related genes during the MS development in WT. The expression trends of the siderophore biosynthesis genes (NrSidA, NrSidC, NrSidF, and
Li et al.  
Siderophore Is Essential for Nomuraea

**FIGURE 5** | Germination rates of *N. rileyi* WT and mutants grown on SMAY medium (A), or AMM-Fe medium (C), AMM-Fe medium supplied with BPS (B) or with 30 µM FeSO₄ (D).

*NrSidD* were consistent with that of siderophore transporter genes (*NrSit1p* and *NrStr3*) during the MS formation. During coalescing hyphae time (36 and 48 h), the expression abundances were low, started to increase at MS initiation stage (60 h), peaked at the MS formation stage (72 and 84 h) or MS maturation stage (96 h) and then declined at the late stage (120 and 144 h; **Figure 9A**). Interestingly, the *NrSidC* expression was highest at MS maturation stage, which was in agreement with the finding that the greatest intracellular iron content in WT was present at 96 h. The analogous expression profiles were found in the RIA genes (*NrFtrA* and *NrFet3*), but the difference was that the higher expression levels occurred at late maturation and at MS maturation stage, implying that RIA mainly took part in iron supplement at the maturation and late maturation. To dive deep into the iron homeostasis by the gene regulation during MS formation, the expression patterns of two iron regulatory genes (*NrHapX* and *NrSre*) were analyzed in the WT and mutants. The transcript abundance of bZIP-type regulator *NrHapX* was enhanced as the MS matured (**Figure 9B**). It was also worth mentioning that a more significant increase in the *ΔNrSidA* mutants rather than in WT or *ΔNrFtrA* mutants because of a defect of iron uptake caused by the siderophore missing under iron depletion (**Figure 3C**). As expected, the transcript concentrations of the GATA transcription factor *NrSre* also increased with the augmentation of intracellular iron contents in WT and mutants. In spite of this, a dramatcal decreased expression occurred at maturation and late maturation stage of MS in *ΔNrSidA* rather than in *ΔNrFtrA* compared with the WT due to the consumption of iron in AM. As oxidative stress occurred during MS development (Song et al., 2013), the transcript abundances of reactive oxygen species (ROS) detoxification genes (*NrSOD* and *NrCata*) were analyzed in the WT and mutants. The forming of MS increased the transcripts of the *NrCata* and *NrSOD* among the WT and mutants, validating that the production of ROS is along with the formation of MS (**Figure 9B**). Notably, a dramatically higher transcript level of *NrSOD* appeared in *ΔNrSidA* mutants at both initiation and formation stage of MS compared to the WT and *ΔNrFtrA* mutants, suggesting that a decrease of the intracellular iron by *NrSidA* deletion resulted in a rise of ROS.

The transcriptional levels of the iron uptake related genes and two iron regulatory genes were further investigated under different iron concentrations for MS formation in the WT and mutants. Expression of the siderophore biosynthesis and RIA genes in WT and mutants was strongly repressed under iron supplemented conditions except the intracellular siderophore biosynthesis *NrSidC* which showed a high expression under the high iron concentrations (**Figure 10A**). Under iron starvation condition, the loss of *NrSidA* led to an obviously increased expression of the RIA related genes in *ΔNrSidA* mutants whereas the *NrFtrA* deletion resulted in evidently higher transcription levels of the siderophore biosynthesis genes in *ΔNrFtrA* mutants.
Li et al. Siderophore Is Essential for Nomuraea

FIGURE 6 | Impact of siderophores on resistance to iron limitation and oxidative stress. (A) $5 \times 10^5$ conidia of N. rileyi WT and mutants were point inoculated on minimal medium lacking iron ($-\text{Fe}$), containing 30 $\mu$M FeSO$_4$ (Fe), or 1.5 mM FeSO$_4$ (hFe), or 0.25 mM bathophenanthroline disulfonate (BPS), or 2 mM H$_2$O$_2$, respectively. (B) Colony diameter of the WT and mutants were measured after 14 days at 25°C. The data represent the means ± standard deviations of results from three independent experiments. (C, D) Analysis of hydrogen peroxide sensitivity of conidia (C) and hyphae (D) was determined as described in Materials and Methods. The conidia used were harvested from SMAY medium. Samples were prepared in triplicate.

(FIGURE 10A). As expected, transcripts of the regulator $NrHapX$ were present at higher concentration in $\Delta NrSidA$ mutants under iron deletion compared to WT and $\Delta NrFtrA$ mutants. The repressor $NrSre$ was typically negatively regulated by iron. To explore the role of the iron played in the ROS detoxification, the expression levels of the $NrSOD$, $NrCata$, glutathione synthase ($NrGS$) and glutathione S transferase ($NrGST$) genes were also further detected. Under iron-limiting conditions, a significantly higher transcript concentration was detected in $\Delta NrSidA$ mutants rather than in $\Delta NrFtrA$ mutants compared to WT. Increasing concentrations of iron to 400, 600, or 800 mg/L rapidly decreased the transcript concentrations of the $NrSOD$, $NrCata$, $NrGS$, and $NrGST$ (FIGURE 10B) in WT and mutants, suggesting that high levels of intracellular iron might promote the detoxification of ROS during MS formation.

To test the hypothesis that iron acquisition deficiency by lack of siderophores led to the observed oxidative stress sensitivity and the higher expression of the ROS detoxification-related genes in $\Delta NrSidA$ mutants, the expression profiles of the iron uptake related genes were further analyzed when treatment with different concentrations of hydrogen peroxide. The expression of selected genes did not associate with the change of hydrogen peroxide (data not show) with the exceptions of the siderophore transporter genes ($Nrsit1p$ and $NrStr3$). They were present at higher concentration only in the WT and $\Delta NrFtrA$ mutants as the levels of H$_2$O$_2$ increased (FIGURE S5).

**DISCUSSION**

**Siderophore-Mediated Iron Uptake, but Not RIA, Is Indispensable for Pigmented MS Development in N. rileyi**

The necessity of the iron involving in MS biogenesis and the up-regulated expression of some iron acquisition related genes in comparative transcriptome raised the questions of what are the main acquisition strategies to achieve the requirement of iron during MS biogenesis and what roles the iron plays. Firstly, we examined the contribution of two high affinity iron uptake pathways played in this physiological process at transcription level. Stage-specific expression pattern analysis showed that the expression of siderophore biosynthesis related genes, as well as the siderophore transporter genes were strongly upregulated during MS development of entomogenous fungi N. rileyi while the higher transcript abundances of RIA related genes were present at the maturation and late maturation stage (FIGURES 3, 9A). These data along with quantitative analysis of intracellular iron in WT indicated that both iron uptake pathways take part in the MS production but the siderophore-mediated iron
acquisition pathway meets the most requirement of iron during the formation and mature period of MS while the RIA just partly compensates the iron at the late maturation.

To understand how two high affinity iron uptake pathways affect the MS formation, NrSidA, which catalyzes the first committed step of hydroxamate-type siderophore biosynthesis, and the high affinity iron permease NrFtrA, the key permease gene for the reductive iron assimilation, were deleted. The loss of the NrSidA led to the defects in pigmented MS development and the similar phenotype also appeared in WT and ΔNrFtrA mutants under iron deletion condition (Figure 2A). Combined with the absence of NrSidA and changes in intracellular iron and extracellular siderophore of MS demonstrated the lack of the total siderophore caused by NrSidA missing impaired cellular iron uptake (Figure 3) and the reduction of the intracellular iron contents in the ΔNrSidA mutants contributed to the defect in MS production. These results were further validated with the differences in the expression levels of iron regulatory genes among the WT and both genes mutants. The increased expression levels of NrHapX during the MS differentiation and the decreased transcript concentrations of NrSre at the maturation and late maturation of MS in the ΔNrSidA mutants is a consequence of the lack of adequate intracellular iron (Figure 9B). The increase in the iron concentrations of AM medium fully restored altered biomass; however, it was not enough to overcome reduced pigment (Figures 2D,E). The pigment in the ΔNrSidA mutants could be basically recovered by adding of the high concentrations of ferrichrome into AM (results not shown). Likewise, whether in the maize (Zea mays) pathogen Cochliobolus heterostrophus or the citrus fungal pathogen Alternaria alternata, the loss of the extracellular siderophore by a nonribosomal peptide synthetase (NPS6) deletion is defective in pigmentation (Oide et al., 2006; Liu et al., 2012). Loss of SIT1, a siderophore transporter gene, in Cryptococcus neoformans exhibits altered melanin deposition and laccase activity (Tangen et al., 2007). It remains uncertain the role of the siderophore plays in the biosynthesis of secondary metabolites. And a reduction in melanin biosynthesis by ΔNrSidA mutants could be at least a result of a reduction in iron availability for some melanin biosynthesis enzymes require metal repletion for full activity (Tangen et al., 2007; Choi et al., 2012).

The role of the iron in MS development is suggested by the studies that Fe^{2+} promotes MS biogenesis by catalyzing the formation of hydroxyl radicals that induce sclerotal biogenesis (Georgiou et al., 2006; Papapostolou and Georgiou, 2010; Song et al., 2013, 2014). Nevertheless, detoxification of hydrogen peroxide depends on iron because catalases and peroxidases require heme as cofactor (Haas et al., 2008). The decreased transcript abundances of the NrCata, NrSOD, NrGS, and NrGST induced by increased iron concentration in AM medium at the 3.5-days old MS indicated that the major role of iron is efficient ROS detoxification rather than the formation of ROS. Furthermore, the higher transcript concentrations of these genes in the ΔNrSidA mutants suggested that the lack of
Remarkably, Chen et al. also demonstrated in the citrus fungal pathogen *Alternaria alternata* that siderophore-mediated iron acquisition has profound effects on ROS detoxification by regulating the NADPH oxidase, the redox activating yes-associated protein 1 regulator, and the high-osmolarity glycerol 1 mitogen-activated protein kinase (Chen et al., 2014). Here, our findings show that the major role of siderophore is collaborated with iron for the ROS detoxification during the MS biogenesis.

The expression patterns of MS after the treatment of hydrogen peroxide is further evidence that the increase of the ROS requires more iron for detoxification by the upregulated expression of the siderophore transporter genes (Figure 5S). However, a different expression pattern is present in *A. nidulans* (Eisendle et al., 2006) and there is no obvious effects on the expression of siderophore transporter mirB after treatment with hydrogen peroxide. The difference may result from that the MS biogenesis is induced by oxidative stress (Georgiou et al., 2006; Song et al., 2013; Liu et al., 2014) and the requirements of iron for ROS detoxification during MS biogenesis is more than the normal for growth (Song et al., 2013).

**Siderophore-Mediated Iron Uptake Is Required for Sporulation, Germination and Dimorphic Switching**

In *N. rileyi*, like in the vast majority of fungus analyzed so far, high-affinity iron uptake systems are indispensable not only for hyphal growth under iron-limited conditions, but also for sporulation (Eichhorn et al., 2006; Oide et al., 2006; Schrettl et al., 2007; Figure 4). Under iron-sufficient conditions, vegetative growth rates of these mutant strains did not significantly differ from that of WT but under iron-limiting conditions in the presence of BPS, the $\Delta NrSidA$ mutants grew significantly slowly (Figure 6A). In contrast, $\Delta sidA$ of *A. fumigatus* failed to grow under iron deletion conditions with BPS (Schrettl et al., 2007). With respect to conidiation of *N. rileyi*, the $\Delta NrSidA$ mutants had a marked reduction either on AMM or SMAY medium and was only rescued by addition of a high concentration of iron (1.5 mM FeSO$_4$) (Figure 4A). Similarly, both $\Delta sid1$ mutants of *Colletotrichum graminicola* and $\Delta SIDA$ mutants of *A. fumigatus* are unable to produce conidia under iron starvation and high iron or corresponding siderophores fully restored conidiation (Schrettl et al., 2004; Wallner et al., 2009; Albarouki et al., 2014). Besides, *SidA* of *A. nidulans* was found to be significantly upregulated during asexual sporulation (Garzia et al., 2013). Thus, growth and sporulation of *N. rileyi* depends on iron steady supplies and siderophore is particularly crucial for iron trafficking. Noteworthily, sporulation in the $\Delta NrSidA$ could be stimulated only with 1.5 mM FeSO$_4$, but not FeCl$_3$, indicating the preference on Fe$^{2+}$ in a reductive iron assimilatory system of *N. rileyi*. However, under iron-sufficient or deficient conditions, the $\Delta NrFtrA$ mutants behaved like WT, suggesting

![Image](image_url)

**FIGURE 8** | (A) Fungal development in insect haemocoels. *S. litura* larvae were injected with the spores of the WT and mutants and bled for microscopic examination of fungal development at 72, 96, and 132 h. (B) Concentration of hyphal bodies in insect haemocoels. Fungal cells were counted using a haemocytometer. HC, haemocyte; HB, hyphal body. Bar = 20 μm. Error bars are standard deviations of three trials. Means indicated by the same uppercase letter are not significantly different from one another, $P < 0.01$.

adequate intracellular iron by the *NrSidA* missing leads to a defects in ROS effective detoxification. The hypothesis is also supported by the hypersensitivity of the conidia and the hypha in the $\Delta NrSidA$ mutants to oxidative stress (Figures 6C,D). Some similar findings are also observed in *A. fumigatus*, *Cochliobolus heterostrophus* and *Magnaporthe grisea* that *NrSidA* homologs missing makes fungi be hypersensitive to oxidative stress (Schrettl et al., 2007; Hof et al., 2009; Condon et al., 2014). So iron is crucially involved in the destruction of ROS. As iron can potentiate oxidative stress, siderophore might play an instrumental role to avoid iron toxicity, which is proven by that the sharp increase in the transcripts of *NrSidA*, an intracellular siderophore synthetase, were followed by an augment of intracellular iron contents at MS formation stage (Figures 3C, 9A). Iron metabolism is interconnected with the regulatory production of ROS. In *A. fumigatus*, Brandon et al. predicted blocking siderophore-mediated iron uptake reduces resistance to oxidative stress by constructing a Boolean network model (Brandon et al., 2015). Remarkably, Chen et al. also demonstrated in the citrus fungal pathogen *Alternaria alternata* that siderophore-mediated iron acquisition has profound effects on ROS detoxification by regulating the NADPH oxidase, the redox activating yes-associated protein 1 regulator, and the high-osmolarity glycerol 1 mitogen-activated protein kinase (Chen et al., 2014). Here, our findings show that the major role of siderophore is collaborated with iron for the ROS detoxification during the MS biogenesis.
that the loss of reductive iron uptake system was compensated fully by the siderophore-mediated iron acquisition.

In addition, the NrSidA deletion in the dimorphic fungus *N. rileyi* prolonged the conversion from yeast-cell to hypha on SMAY medium ([Figures 4C,D](#)). The conversion from yeast-cell to hypha contributes to environmental survival and transmission to new hosts (Gauthier, 2015). Several evidences have explained the role of iron acquisition in the yeast-to-hyphal transition. Cell-type-specific transcriptional profiles analysis reveals *sidF* (*NrSidA* orthologue) and *sidD* (*NrSidD* orthologue) are highly expressed in *Penicillium marneffei* yeast cells (Pasricha et al., 2013). Besides, the loss of a vacuolar ATPase, *VMA1*, involved in iron homeostasis in *H. capsulatum*, impairs the mycelial transition at ambient temperature (Hilty et al., 2008). Collectively, iron limitation by the *NrSidA* deletion should have a drastic effect on the morphologic switch.

Conidial siderophore storage was an important germination factor, as germination of conidia fails or is greatly delayed unless a suitable siderophore is supplied due to the loss of cellular siderophores in *A. nidulans*, *A. fumigatus* and *N. crassa* (Eisendle et al., 2006; Schrettl et al., 2007). Hence, a delay of the germination under iron-sufficient or deficient conditions ([Figures 5C,D](#)) and a failure under depletion condition in the presence of BPS in Δ*NrSidA* mutants ([Figure 5B](#)) results from the missing of siderophore-mediated iron storage or utilization of intracellular iron (Figure S3B).

**Loss of the NrSidA, but Not NrftrA Is Required for Full Virulence of the *N. rileyi* Against Spodoptera litura**

The research on the roles of siderophores in fungal pathogenicity to host was widespread studied in pathogenic fungi of plant
and human with the differences among different pathosystems, while the studies about entomogenous fungal pathogenicity to insect are scarce. In this study, we found that the siderophore biosynthesis, but not reductive iron assimilation is essential for virulence in the \textit{N. rileyi} against \textit{S. litura}. In both topical infection and injection assay, the \(\Delta\text{NrSidA}\) mutants showed less virulence in \textit{S. litura} than WT while the \(\Delta\text{NrFtrA}\) mutant was as virulent as the wild-type strain (Figure 7). Further investigation revealed that both germination and proliferation of the \(\Delta\text{NrSidA}\) mutants conidia in insect haemocoel slowed dramatically (Figure 8). Disruption of siderophore production resulted in iron-dependent growth during the infection and the growth defect in \textit{S. litura} ultimately led to the loss of virulence. By contrast, in \textit{A. fumigatus}, no germination of \(\Delta\text{sidA}\) conidia was observed in vivo, leading to the total loss of the virulence caused by the \textit{SidA} missing (Schrettl et al., 2004).

In other fungal pathogens, one mode of high-affinity iron acquisition is also favored over the other. In \textit{A. fumigatus},

FIGURE 10 | Relative transcript abundances of iron uptake-related genes (A) and ROS detoxification related genes (B) in 3.5-day-old MS of the WT and mutants at different iron supplies. TEF and TUB transcripts were used as reference. Uppercase or lowercase letters indicated the statistically significant level at \(P < 0.01\) or \(P < 0.05\), respectively and the same letter were not significantly different from one another. Error bars are standard deviations of three trials.
siderophore biosynthesis is essential for virulence, but RIA is neither necessary nor sufficient for normal growth and survival in the host (Schrettl et al., 2007). Similar results are reported in the phytopathogens Cochliobolus miyabeanus, Alternaria alternata, Magnaporthe grisea, Fusarium graminearum, Colletotrichum graminicola, and human pathogen Histoplasma capsulatum that extracellular siderophore production is required for full virulence (Greenshields et al., 2007; Hwang et al., 2008; Hof et al., 2009; Chen et al., 2013; Albarouki et al., 2014). However, the loss of the SIDI gene in U. maydis does not affect virulence in maize, instead, RIA is required for virulence (Mei et al., 1993). Analogous observations have also been found in Candida albicans and Cryptococcus neoformans, where high-affinity iron permease genes are essential for systemic infection (Ramanan and Wang, 2000; Haas et al., 2008). Our findings indicate that N. rileyi prefers high-affinity siderophore iron acquisition over RIA.

In summary, our studies revealed siderophore-mediated iron uptake is more important than reductive iron uptake. Even though both are induced upon iron deprivation, NrSidA, but not NrFirA, is required for pigmented-MS morphogenesis, oxidative-stress resistance, sporulation, phase transition, germination and full virulence of N. rileyi against S. litura. As a cofactor, iron is necessary for ROS detoxification instead of catalyzing the formation of ROS during the MS formation; siderophore is very important to assist iron mobilization and meanwhile avoid iron toxicity. In view of these findings, a certain amount of iron is indeed an essential part for the MS induction. Thus, a complete understanding of iron uptake strategies during the MS formation in N. rileyi at the molecular level provides a valuable insight into the molecular mechanism of MS differentiation and serves as a guidance in industrialized mass production of N. rileyi.

AUTHOR CONTRIBUTIONS

YL conducted the main experiments and wrote the original manuscript. XL provided a lot of help throughout the experiment. RL provided great help to screen the stable transformants. CS provided assistance with fungal transformation. ZS, ZW, and YY provided technical oversight and critical manuscript review and editing. All authors discussed the results and implications and commented on the manuscript at all stages.

ACKNOWLEDGMENTS

This research was financially supported by the National Natural Science Foundation of China (Grant No.31570073) and the National Transformation Fund for Agricultural Science and Technology Achievements (Project No. CSTC2014JCSF-NYCGZHBB0011).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.00931

REFERENCES

Albarouki, E., Schafferer, L., Ye, F. H., Wirén, N., Haas, H., and Deising, H. B. (2014). Biotrophy-specific downregulation of siderophore biosynthesis in Colletotrichum graminicola is required for modulation of immune responses of maize. Mol. Microbiol. 92, 338–355. doi: 10.1111/mmi.12561

Ardon, O., Bussey, H., Philpott, C., Ward, D. M., Davis-Kaplan, S., and Veronneau, S., et al. (2001). Identification of a Candida albicans ferrichrome transporter and its characterization by expression in Saccharomyces cerevisiae. J. Biol. Chem. 276, 43049–43055. doi:10.1074/jbc.M108701200

Askwith, C., Eide, D., Van Ho, A., Bernard, P. S., Li, L., Davis-Kaplan, S., et al. (1994). The Fet3 gene of Saccharomyces cerevisiae encodes a multicopper oxidase required for ferrous iron uptake. Cell 76, 403–410. doi:10.1016/0092-8674(94)90346-8

Babitskaya, V. G., Scherba, V. V., Filimonova, T. V., and Grigorchuk, E. A. (2000). Melanin pigments from the fungi Paeilomyces varioti and Aspergillus carbonarius. Appl. Biochem. Microbiol. 36, 128–133. doi: 10.1007/BF02759067

Behle, R. W., Jackson, M. A., and Weiler, L. B. (2013). Efficacy of a granular formulation containing Metarhizium brunneum F52 (Hypocreales: Clavicipitaceae) microsclerotia against nymphs of Ixodes scapularis (Acari: Ixodidae). J. Econ. Entomol. 57, 56–63. doi:10.1603/EC12226

Boyce, K. J., and Andrianopoulos, A. (2015). Fungal dimorphism: the switch from hyphae to yeast is a specialized morphogenetic adaptation allowing colonization of a host. FEMS Microbiol. Rev. 39, 797–811. doi:10.1093/femsre/fuv035

Brandon, M., Howard, B., Lawrence, C., and Laubenbacher, R. (2015). Iron acquisition and oxidative stress response in Aspergillus fumigatus. BMC Syst. Biol. 9:19. doi:10.1186/s12918-015-0163-1

Cabisco, E., Belli, G., Tamarit, J., Echave, P., Herrero, E., and Ros, J. (2002). Mitochondrial Hsp60 to oxidative stress, and the labile iron pool are closely connected in Saccharomyces cerevisiae. J. Biol. Chem. 277, 44531–44538. doi:10.1074/jbc.M206525200

Chao, L. Y., Marletta, M. A., and Rine, J. (2008). Srel, an iron-modulated GATA DNA-binding protein of iron-uptake genes in the fungal pathogen Histoplasma capsulatum. Biochemistry 47, 7274–7283. doi:10.1021/bi800066x

Chen, L. H., Lin, C. H., and Chung, K. R. (2013). A nonribosomal peptide synthetase mediates siderophore production and virulence in the citrus fungal pathogen Alternaria alternata. Mol. Plant Pathol. 14, 497–505. doi:10.1111/mpp.12021

Chen, L. H., Yang, S. L., and Chung, K. R. (2014). Resistance to oxidative stress via regulating siderophore-mediated iron acquisition by the citrus fungal pathogen Alternaria alternata. Microbiology 160, 970–979. doi:10.1099/micro.0.076182-0

Choi, J. N., Kim, J., Kim, J. Y., Jung, W. H., and Lee, C. H. (2012). Influence of iron regulation on the metabolome of Cryptococcus neoformans. PLoS ONE 7:e4165. doi:10.1371/journal.pone.0041654

Condon, B. J., Oide, S., Gibson, D. M., Krasnoff, S. B., and Turgeon, B. G. (2014). Reductive iron assimilation and intracellular siderophores assist extracellular siderophore-driven iron homeostasis and virulence. Mol. Plant Microbe Interact. 27, 793–808. doi:10.1094/MPMI-11-13-0328-R

Gajty, T. Z. (1948). An estimation of bound hydroxylamine in biological materials. Acta Chem. Scand. 2, 450–454. doi:10.3891/acta.chem.scand.02-0450

de Lorenzo, V., Binderfeir, A., Paw, B. H., and Neilands, J. B. (1986). Aerobactin biosynthesis and transport genes of plasmid CoV-K30 in Escherichia coli K-12. J. Bacteriol. 165, 570–578.

Eichhorn, H., Lessing, F., Winterberg, B., Schirawski, J., Kämper, J., Müller, P., et al. (2006). A ferrooxidation/ permeation iron uptake system is required for virulence in Ustilago maydis. Plant Cell. 18, 3332–3345. doi:10.1105/tpc.106.043588

Eisendle, M., Oberegger, H., Zadra, I., and Haas, H. (2003). The siderophore system is essential for viability of Aspergillus nidulans: functional analysis of two genes encoding l-ornithine N 5-monooxygenase (sidA) and a nonribosomal

AUTHOR CONTRIBUTIONS

YL conducted the main experiments and wrote the original manuscript. XL provided a lot of help throughout the experiment. RL provided great help to screen the stable transformants. CS provided assistance with fungal transformation. ZS, ZW, and YY provided technical oversight and critical manuscript review and editing. All authors discussed the results and implications and commented on the manuscript at all stages.

ACKNOWLEDGMENTS

This research was financially supported by the National Natural Science Foundation of China (Grant No.31570073) and the National Transformation Fund for Agricultural Science and Technology Achievements (Project No. CSTC2014JCSF-NYCGZHBB0011).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.00931
peptide synthetase (sidC). Mol. Microbiol. 49, 359–375. doi: 10.1046/j.1365-2958.2003.03586.x

Eisenfeld, M., Schrettl, M., Kragl, C., Müller, D., Illmer, P., and Haas, H. (2006). The intracellular siderophore ferricinocin is involved in iron storage, oxidative-stress resistance, germination, and sexual development in Aspergillus nidulans. Eur. J. Biochem. 263, 1218–1226. doi: 10.1111/j.1432-1033.2005.04576.x

Epstein, S., Kakhlon, O., Glickstein, H., Breuer, W., and Capanich, I. (1997). Fluorescence analysis of the labile iron pool of mammalian cells. Anal. Biochem. 248, 31–40. doi: 10.1006/abio.1997.2126

Garzia, A., Etxebeste, O., Rodríguez-Romero, J., Fischer, R., Espeso, E. A., and Ugaleda, U. (2013). Transcriptional changes in the transition from vegetative cells to asexual development in the model fungus Aspergillus nidulans. Eukaryot Cell 12, 311–321. doi: 10.1128/EC.00274-12

Gauthier, G. M. (2015). Dimorphism in fungal pathogens of Mammals, plants, and Insects. PLoS Pathog. 11:e1004608. doi: 10.1371/journal.ppat.1004608

Giorgiou, C. D., Patsoyukis, N., Papapostolou, I., and Zervoudakis, G. (2006). Siderolysis in filamentous fungi is induced by oxidative stress. Integr. Comp. Biol. 46, 691–712. doi: 10.1093/iceb/icc034

Goble, T. A., Hajek, A. E., Jackson, M. A., and Gardesca, S. (2015). Microsclerotia of Metarhizium brunneum F52 Applied in Hydromulch for control of Asian Longhorned Beetles (Cerambyx cerambycidae). J. Econ. Entomol. 2, 433–443. doi: 10.1093/jee/tov013

Greenshields, D. L., Liu, G., Feng, J., Selvaraj, G., and Wei, Y. (2007). The siderophore biosynthetic gene SID1, but not the ferroxidase gene FET3, is required for full Fusarium graminearum virulence. Mol. Plant Pathol. 8, 411–421. doi: 10.1111/j.1364-3703.2007.00401.x

Gaßler, F., Hortschansky, P., Beattie, S. R., Klammer, V., Tuppatsch, K., Lechner, B. E., et al. (2014). The Janus transcription factor HapX controls fungal adaptation to both iron starvation and iron excess. EMBO J. 33, 2261–2276. doi: 10.15252/embj.201489468

Haas, H. (2003). Molecular genetics of fungal siderophore biosynthesis and uptake: The role of siderophores in iron uptake and storage. Appl. Microbiol. Biotechnol. 62, 316–330. doi: 10.1007/s00253-003-1335-2

Haas, H. (2012). Iron–A key nexus in the virulence of Aspergillus fumigatus. Annu. Rev. Phytopathol. 50, 113–139. doi: 10.1146/annurev.phyto.45.062806.094338

Heinkekamp, T., Thywissen, A., Macheleidt, J., Keller, S., Valiante, V., and Brakhage, A. A. (2013). Aspergillus fumigatus melanos: interference with the host endocytosis pathway and impact on virulence. Front. Microbiol. 3:440. doi: 10.3389/fmicb.2012.00440

Heymann, P., Gerads, M., Schaller, M., Dromer, F., Winklmann, G., and Ernst, J. F. (2002). The siderophore iron transporter of Candia albicans (Sit1p/Arn1p) mediates uptake of ferrichrome-type siderophores and is required for epithelial invasion. Infect. Immun. 70, 5246–5252. doi: 10.1128/IAI.70.12.5246-5252.2002

Hilty, J., S mutants, A. G., and Newman, S. L. (2008). The Histoplasma capsulatum vacular ATPase is required for iron homeostasis, intracellular replication in macrophages, and virulence in a murine model of histoplasmosis. Mol. Microbiol. 70, 127–139. doi: 10.1111/j.1365-2958.2008.06395.x

Hof, C., Eisel, K., Antelo, L., Foster, A. J., and Anke, H. (2009). Siderophore synthesis in Magnaporthe grisea is essential for vegetative growth, conidiation and resistance to oxidative stress. Fungal Genet. Biol. 46, 321–332. doi: 10.1016/j.fgb.2008.12.004

Hwang, L. H., Mayfield, J. A., Rine, J., and Sil, A. (2008). Histoplasma Requires SID1, a member of an iron-regulated siderophore gene cluster, for host colonization. PLoS Pathog. 4:e1000044. doi: 10.1371/journal.ppat.1000044

Jackson, M. A., Dunlap, C. A., and Jaronski, S. T. (2010). Ecological considerations in producing and formulating fungal entomopathogens for use in insect biocontrol. BioControl 55, 129–145. doi: 10.1007/s10526-009-9240-y

Jackson, M. A., and Jaronski, S. T. (2009). Production of microsclerotia of the fungal entomopathogen Metarhizium anisopliae and their potential for use as a biocontrol agent for soil-inhabiting insects. Mycol. Res. 113, 842–850. doi: 10.1016/j.mycres.2009.03.004
Schrettl, M., Bignell, E., Kragl, C., Joechl, C., Rogers, T., Arst, H. N. Jr., et al. (2004). Siderophore biosynthesis but not reductive iron assimilation is essential for *Aspergillus fumigatus* virulence. *J. Exp. Med.* 201, 213–219. doi: 10.1084/jem.20041422

Schrettl, M., Bignell, E., Kragl, C., Sabiba, Y., Loss, O., Eissenle, M., et al. (2007). Distinct roles for intra- and extracellular siderophores during *Aspergillus fumigatus* infection. *PLoS Pathog.* 3:e128. doi: 10.1371/journal.ppat.0030128

Schwyn, B., and Neilands, J. B. (1987). Universal chemical assay for the detection and determination of siderophores. *Anal. Biochem.* 160, 47–56. doi: 10.1016/0003-2697(87)90612-9

Shanthakumar, S. P., Murali, P. D., Malarvannan, S., Prabavathy, V. R., and Srisukchayakul, P., Wiwat, C., and Pantuwatana, S. (2005). Studies on the physiology of two *Nomuraea rileyi* strains against tobacco caterpillar, *Spodoptera litura* Fabricius (*Noctuidae: Lepidoptera*) and its safety to *Trichogramma sp.* *J. Biopesticides* 3, 132–137.

Shao, C. W., Yin, Y. P., Qi, Z. R., Li, R., Song, Z. Y., Li, Y., et al. (2015). *Agrobacterium tumefaciens* mediated transformation of the entomopathogenic fungus *Nomuraea rileyi*. *Fungal Genet. Biol.* 83, 9–25. doi: 10.1016/j.fgb.2015.08.002

Tamarit, J., Irazusta, V., Moreno-Cermeño, A., and Ros, J. (2006). Colorimetric assay for the quantitation of iron in yeast. *Anal. Biochem.* 351, 49–151. doi: 10.1016/j.ab.2005.12.001

Tamura, K., Stecher, G., Peterson, D., Filipski, A., and Kumar, S. (2013). MEGA6: molecular evolutionary genetics analysis version 6.0. *Mol. Biol. Evol.* 30, 2725–2729. doi: 10.1093/molbev/mst197

Song, Z. Y., Yin, Y. P., Jiang, S. S., Liu, J. J., and Wang, Z. K. (2014). Optimization of culture medium for microsclerotia production by *Nomuraea rileyi*. *F. Microbiol. Egypt.* 160, 47–56. doi: 10.1016/j.ab.2005.12.001

Song, Z. Y., Shen, L., Yin, Y. P., Tan, W. Y., Shao, C. W., Xu, J. M., et al. (2015). Role of two *Nomuraea rileyi* transmembrane sensors Sho1p and Sin1p in adaptation to stress due to changing culture conditions during microsclerotia development. *World J. Microbiol. Biotechnol.* 31, 773–485. doi: 10.1007/s11274-015-1801-x

Song, Z. Y., Yin, Y. P., Jiang, S. S., Liu, J. J., Chen, H., and Wang, Z. K. (2013). Comparative transcriptome analysis of microsclerotia development in *Nomuraea rileyi*. *BMC Genomics*. 14:411. doi: 10.1186/1471-2164-14-411

Stehr, M., Diekmann, H., Smau, L., Seth, O., Ghisla, S., Singh, M., et al. (1998). A hydrophobic sequence motif common to N-hydroxylating enzymes. *Trends Biochem. Sci.* 235, 6–57.

Tang, K. L., Jung, W. H., Sham, A. P., Lian, T., and Kronstad, J. W. (2007). The iron-and CAMP-regulated gene *SIT1* influences ferrioxamine B utilization, melanisation and cell wall structure in *Cryptococcus neoformans*. *Microbiology* 153, 29–41. doi: 10.1099/mic.0.009927-0

Thakre, M., Thakur, M., Malik, N., and Ganger, S. (2011). Mass scale cultivation of entomopathogenic fungus *Nomuraea rileyi* using agricultural products and agro wastes. *J. Bioprocess. Sci.* 4, 176–179.

Wallner, A., Blatter, M., Schrettl, M., Sarg, B., Lindner, H., and Haas, H. (2009). Ferricrocin, a siderophore involved in intra- and extracellular iron distribution in *Aspergillus fumigatus*. *Appl. Environ. Microbiol.* 75, 4194–4196. doi: 10.1128/AEM.00479-09

Wanchoo, A., Lewis, M. W., and Keyhani, N. O. (2009). Lectin mapping reveals stage-specific display of surface carbohydrates in *in vitro* and haemolymph-derived cells of the entomopathogenic fungus *Beauveria bassiana*. *Microbiology* 155, 3121–3133. doi: 10.1099/mic.0.029157-0

Wang, H., Lei, Z., Reitz, S., Li, Y., and Xu, X. (2013). Production of microsclerotia of the fungal entomopathogen *Lecanicillium lecanii* (Hyphocreales: Cordycipitaceae) as a biological control agent against soil-dwelling stages of *Frankliniella occidentalis* (Thysanoptera: Thripidae). *Biocontrol. Sci. Tech.* 23, 234–238. doi: 10.1080/09583157.2012.745482

Wang, Z., Ye, S. F., Li, J. J., Zheng, B., Bao, M. Z., and Ning, G. G. (2011). Fusion primer and nested integrated PCR (FPNI-PCR): a new high-efficiency strategy for rapid chromosome walking or flanking sequence cloning. *PMC Biotechnol.* 2011, 11:109. doi: 10.1186/1476-7565-11-109

Wang, Z. Y., Huang, S., Song, Z. Y., and Wang, Z. K. (2012). Microsclerotia Artificial Inductions of *Nomuraea rileyi* CQNtr01. *Sci. Agric. Sin.* 45, 4801–4807. doi: 10.3864/j.issn.0578-1752.2012.23.006

Zhong, J., Frases, S., Wang, H., Casadevall, A., and Stark, R. E. (2008). Following fungal melanin biosynthesis with solid-state NMR: biopolymer molecular structures and possible connections to cell-wall polysaccharides. *Biochemistry* 47, 4701–4710. doi: 10.1021/bi702093r

Zhao, G. L., Song, Z. Y., Yin, Y. P., Jiang, W., and Wang, Z. K. (2015). Involvement of alternative oxidase in the regulation of hypha growth and microsclerotia formation in *Nomuraea rileyi* CQNtr01. *World J. Microbiol. Biotechnol.* 31, 134–1352. doi: 10.1007/s11274-015-1877-3

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

Copyright © 2016 Li, Wang, Liu, Song, Li, Shao and Yin. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.