

SCIENTIFIC REPORTS



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Nox Complex signal and MAPK cascade pathway are cross-linked and essential for pathogenicity and conidiation of mycoparasite *Coniothyrium minitans*

Received: 30 October 2015

Accepted: 24 March 2016

Published: 12 April 2016

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The NADPH oxidase complex of a sclerotial mycoparasite *Coniothyrium minitans*, an important biocontrol agent against crop diseases caused by *Sclerotinia sclerotiorum*, was identified and its functions involved in conidiation and mycoparasitism were studied. Gene knock-out and complementary experiments indicated that *CmNox1*, but not *CmNox2*, is necessary for conidiation and parasitism, and its expression could be significantly induced by its host fungus. *CmNox1* is regulated by *CmRac1-CmNoxR* and interacts with *CmSlt2*, a homolog of *Saccharomyces cerevisiae* *Slt2* encoding cell wall integrity-related MAP kinase. In Δ *CmNox1*, *CmSlt2*-GFP fusion protein lost the ability to localize to the cell nucleus accurately. The defect of conidiation in Δ *CmRac1* could be partially restored by over-expressing *CmSlt2*, indicating that *CmSlt2* was a downstream regulatory factor of *CmNox1* and was involved in conidiation and parasitism. The expressions of mycoparasitism-related genes *CmPks1*, *Cmg1* and *CH1* were suppressed in the knock-out mutants of the genes in *CmNox1-CmSlt2* signal pathway when cultivated either on PDA. Therefore, our study infers that *CmRac1-CmNoxR* regulates *CmNox1-CmSlt2* pathway in regulating conidiation and pathogenicity of *C. minitans*.

Sclerotinia sclerotiorum (Lib.) de Bary is a significant necrotrophic pathogen that could infect more than 400 plant species worldwide and cause huge economic losses every year^{1,2}. *Coniothyrium minitans* is an important mycoparasite of *Sclerotinia* spp.; it could parasitize and destroy both sclerotium and hypha of its hosts. It is an effective agent for controlling crop diseases caused by *S. sclerotiorum*³⁻⁵. Like its hosts, *C. minitans* also occurs widely⁶. It shares the same growing season as its hosts, and its growth and proliferation are tightly related to the hosts. Without host fungi, *C. minitans* stays dormant for seasons in soil⁷. *C. minitans* is one of the commercialized fungal agents for biological control of crop diseases⁸.

Understanding the parasitism and conidiation of *C. minitans* at molecular level could help us to utilize this biological control agent more efficiently. Studies indicated that the primary cell wall degrading enzymes secreted by *C. minitans* are β -1,3 glucanase (encoded by *Cmg1*) and chitinase (encoded by *CH1*). The expression level of *Cmg1* was enhanced when *C. minitans* parasitized *S. sclerotiorum*, indicating that β -1,3 glucanase played important roles in the parasitism process⁹. *C. minitans* also could secrete antifungal substances and degrade oxalic acid, an essential pathogenicity factor of *S. sclerotiorum*^{10,11}. The parasitism of *C. minitans* is likely very complicated¹², and shares some signal transduction pathways with conidiation. Fungal cell wall integrity-associated MAP kinase cascade, fatty acid beta-oxidation, reactive oxygen and nitrogen species, and possibly other unknown pathways in peroxisomes are required for both conidiation and mycoparasitism of *C. minitans*^{13,14}. Recent research revealed that

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C. minitans can regulate ambient pH by degrading oxalic acid to facilitate mycoparasitism of *S. sclerotiorum*^{15,16}. *C. minitans* requires a large amount of L-arginine during conidiation and L-arginine-derived nitric oxide was likely to be involved in conidiation with cyclic GMP functions as a second messenger^{17,18}. Qin *et al.* found that phosphoribosylamidotransferase is essential for conidiation of *C. minitans* via adenosine related molecules, and *C. minitans* is able to obtain adenosine or related components from its host during parasitization¹⁹.

The NADPH oxidase complexes are conserved and play important roles in the life cycle of filamentous fungi²⁰. In rice blast pathogen *Magnaporthe oryzae*, NADPH oxidase-derived reactive oxygen species (ROS) is essential for pathogenicity²¹, and further study found that NADPH oxidases are necessary for septin-mediated reorientation of the F-actin cytoskeleton to facilitate cuticle rupture and plant cell invasion²². NADPH oxidases are involved in sclerotial formation and pathogenicity of necrotrophic fungal pathogen *Botrytis cinerea* and *S. sclerotiorum*^{22–25}, and fusion of conidial anastomosis tubes of *B. cinerea*²⁶. ROS is critical in maintaining a mutualistic interaction between *Epichloë festucae* and perennial ryegrass^{27–31}. In saprophytic fungi, such as *Aspergillus* spp., *Neurospora crassa* and *Podospora anserina*, NADPH oxidases are required for growth, cell differentiation, conidiation, and sexual reproduction^{32–36}. *Trichoderma harzianum* NADPH oxidases are involved in the antagonism against *Pythium ultimum*³⁷.

Besides NADPH oxidases mitogen-activated protein kinases (MAPKs) also play critical roles in pathogenicity and in fungal development^{38–47}. In endophytic fungi, a stress-activated mitogen-activated protein kinase (sakA) of *E. festucae* is essential to maintain mutualistic symbiosis with perennial ryegrass. Deletion of sakA converted this endophytic fungus to a pathogen of its host⁴⁸. MAP kinase cascade also is involved in the mycoparasitism and development of hyperparasitic fungi, such as *C. minitans*, *Stachybotrys elegans* and *T. artoviride*^{13,49,50}. The phenotype of Δ sakA mutants are very similar to that of mutants whose genes in the Nox complex were disrupted, suggesting that there is a possible link between ROS signal and MAPK signal pathway on the maintenance of mutualistic symbiosis²⁹. Similar phenomena were observed in tangerine pathogen *Alternaria alternata*⁵¹; Medina-Castellanos *et al.* found that extracellular ATP could promote the Nox1-derived ROS and activate a MAPK pathway in *T. artoviride*⁵². Recently, Jaimes-Arroyo *et al.* found that SrkA kinase could regulate stress responses and development in *A. nidulans*, and H₂O₂ could induce mitochondrial fragmentation and relocalize SrkA at the presence of SakA⁵³.

Previously, we investigated a fungal cell wall integrity-associated MAP kinase cascade in *C. minitans* and found that this cascade was required for conidiation and mycoparasitism¹³. In this study, we analyzed the function of NADPH oxidases (Nox1/Nox2) complex of *C. minitans*, and found that *CmNox1* played critical roles in conidiation and mycoparasitism, but not *CmNox2*. In *C. minitans*, *CmNoxR* interacts with *CmRac1* to activate *CmNox1*, *CmNox1* could also interact with *CmSlt2* and thus adjust its location to the cell nucleus, and deliver the signal of conidiation and mycoparasitism.

Results

NADPH oxidases (Nox1/Nox2) in *C. minitans*. Two NADPH oxidase genes, *CmNox1* and *CmNox2*, were isolated from *C. minitans*. The deduced amino acid sequence of *CmNox1* (GenBank Accession No: KJ596434) shows high similarity to Nox1 homologs from other filamentous fungi, including *Alternaria alternata* (AaNox1, BAK52527.1, 89% identity), *Curvularia lunata* (ClNOX1, AHC53982.1, 89% identity), and *Pyrenophora tritici-repentis* (PrNOX1, XP_001935118.1, 88% identity). *CmNox2* (GenBank Accession No: KJ596435) is highly similar to Nox2 homologs from other filamentous fungi. The multiple alignment analysis showed that both *CmNox1* and *CmNox2* contained NOX family signature regions. Phylogenetic analysis of NADPH oxidases in several fungi placed *CmNox1* and *CmNox2* homologs into two different clades based on the amino acid sequences (Supplementary Figure S1).

***CmNox1* is essential for ROS production and conidiation.** To study the function of *CmNox1* in *C. minitans*, a replacement vector p3300neo*CmNox1* (see Supplementary Figure S2a) was constructed and transformed into strain ZS-1 to disrupt *CmNox1*. Twenty transformants were obtained, and three of which, Δ *CmNox1-1*, Δ *CmNox1-6*, and Δ *CmNox1-107* were selected randomly as candidates for further analyses. Furthermore, a complement vector pNox1 was transformed into mutant Δ *CmNox1-6*. The deletion and complementary events were confirmed by RT-PCR (see Supplementary Figure S2b) and Southern blot analysis (see Supplementary Figure S2c). Using the same methods, a replacement vector p3300neo*CmNox2* was constructed (see Supplementary Figure S2a) and transformed into strain ZS-1 to disrupt *CmNox2*. Three deletion mutants Δ *CmNox2-20*, Δ *CmNox2-323* and Δ *CmNox2-347* were also confirmed by Southern blot analysis.

Colony staining with NBT solution showed that superoxide production was decreased significantly in Δ *CmNox1-6*, compared to Δ *CmNox2-323* and the wild-type strain ZS-1 (Fig. 1a). The conidiation of the wild-type strain ZS-1, *CmNox1* deletion mutants, *CmNox1* complemented mutants, and *CmNox2* deletion mutants were determined after incubating for 15 days on PDA (Fig. 1b). *CmNox1* deletion mutants completely lost the ability to produce conidia (Table 1). In contrast, the wild-type strain ZS-1, *CmNox1* complemented mutants and *CmNox2* deletion mutants were normal in conidiation under the same condition (Table 1). Unlike strain ZS-1 and Δ *CmNox2-323*, which could form matured pycnidia and conidia, Δ *CmNox1-6* could only form a few pycnidial primordia that could not further develop to mature pycnidia, and no conidium was produced (Fig. 1b). These data indicated that *CmNox1*, but not *CmNox2*, played significant roles in conidiation and production of superoxide. The experiments also suggested *CmNox1* and *CmNox2* are not essential for hyphal growth of *C. minitans* on PDA.

***CmNox1* is essential to parasitize sclerotia of *S. sclerotiorum*.** To determine whether *CmNox1* is related to sclerotial parasitizing, the parasitic ability of *CmNox1* mutants and the other strains to sclerotia of *S. sclerotiorum* were examined. The hyphae of Δ *CmNox1-6* was inoculated on to sclerotia and incubated for 30

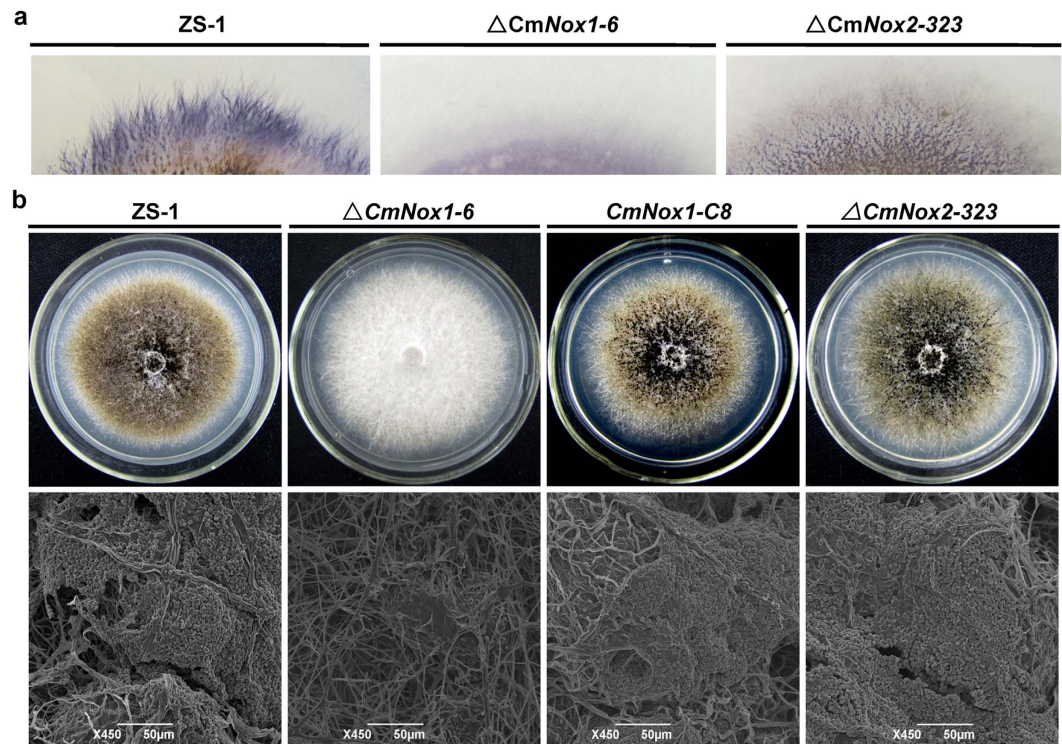


Figure 1. Production of superoxide and conidia of *Coniothyrium minitans* *CmNox1* and *CmNox2* mutants. (a) Detection of superoxide using colony staining with NBT solution (0.05% w/v, 30 min) of cultures of *C. minitans* wild-type strain ZS-1, *CmNox1* deletion mutant $\Delta CmNox1-6$, *CmNox2* deletion mutant $\Delta CmNox2-323$ on PDA. (b) Pycnidial and conidial production of *C. minitans* wild-type strain ZS-1, *CmNox1* deletion mutant $\Delta CmNox1-6$, complementary transformant *CmNox1-C8*, and *CmNox2* deletion mutant $\Delta CmNox2-323$ 12 days after incubation on PDA. Compared to ZS-1, *CmNox1-C8* and $\Delta CmNox2-323$, $\Delta CmNox1-6$ lost the ability to produce pycnidia.

days at 20 °C, and no pycnidia and conidia were observed on either the surface or interior of the sclerotia (Fig. 2a, Table 1). The inoculated sclerotia were surface sterilized and then incubated on PDA containing 50 μ g/ml hygromycin, and no *C. minitans* colony emerged from the sclerotia, suggesting that $\Delta CmNox1-6$ did not invade the inner of sclerotia (Fig. 2b, Table 1). Meanwhile, *CmNox1-C8*, $\Delta CmNox2-323$ and the wild-type strain ZS-1 could degrade sclerotia and produce mature pycnidia there. The results demonstrated *CmNox1* played a significant role in sclerotial mycoparasitism.

Expression of *CmNox1* is highly induced by *S. sclerotiorum*. When dual cultured with *S. sclerotiorum*, the expression of *CmNox1* was highly induced. Compared to growing on PDA, the expression of *CmNox1* peaked at 12 hpi, and the high expression was maintained till 24 hpi, and then sharply declined to undetectable level at 36 hpi (Fig. 3a). This phenomenon suggested that *CmNox1* played an important role in parasitism. Expression of *CmNox2* also was up-regulated slightly by *S. sclerotiorum* at 12 hpi and 24 hpi, but compared to culture on PDA, the induction was much lower (Fig. 3b). The results showed that *S. sclerotiorum* could induce the expression of *CmNox1* and *CmNox2*, with stronger induction to *CmNox1*.

***CmRac1* interacts with *CmNoxR* together to regulate *CmNox1* and to control conidiation in *C. minitans*.** It is reported that Bem1 and Cdc24 are components of the NADPH oxidase complex in filamentous fungi, and NoxR interacting with RacA *in vitro* together regulates ROS production and control hyphal branching and patterning in *E. festucae*^{27,29}. Since *CmNox1* is involved in the production ROS in *C. minitans*, we identified *C. minitans* homologs of NoxR, GTPase Rac1 and Bem1, named *CmNoxR* (GenBank Accession No: KJ596436), *CmRac1* (GenBank Accession No: KJ596436) and *CmBem1*, respectively. The interaction between *CmRac1* and *CmNoxR*, *CmBem1* and *CmNoxR* were confirmed by Yeast two-hybrid (Fig. 3c,d).

To investigate the role of *CmNoxR*, *CmBem1* and *CmRac1* in *C. minitans*, replacement vector for *CmNoxR* (p3300neo*CmNoxR*), *CmRac1* (p3300neo*CmRac1*), and *CmBem1* (p3300neo*CmBem1*) were constructed and transformed into ZS-1. The *CmNoxR* deletion mutants ($\Delta CmNoxR-5$ and $\Delta CmNoxR-107$), *CmBem1* deletion mutant ($\Delta CmBem1-344$) and *CmRac1* deletion mutant ($\Delta CmRac1-79$) were obtained and confirmed by RT-PCR. In contrast to the wild-type strain ZS-1, the *CmNoxR* deletion mutants and the *CmRac1* deletion mutant lost the ability to produce conidia and to parasitize *S. sclerotiorum*. In addition to the conidiation and parasitizing phenotype, growth rate also was reduced significantly in the *CmRac1* deletion mutant. The growth rate of $\Delta CmRac1-79$ was less than 0.6 mm/day on average on PDA, while it was about 3mm/day for strain ZS-1 under the same

Strains	Growth rate ^a (mm/d)	Conidiation ^b (x10 ⁷ conidia/plate)	Rot index ^c (%)
ZS-1	2.9 ± 0.1 ^A	128 ± 7.2 ^A	82 ± 3 ^A
Δ <i>CmNox1-1</i>	2.8 ± 0.1 ^A	0 ^B	0 ^B
Δ <i>CmNox1-6</i>	2.8 ± 0.1 ^A	0 ^B	0 ^B
Δ <i>CmNox1-107</i>	2.9 ± 0.1 ^A	0 ^B	0 ^B
<i>CmNox1-C1</i>	2.9 ± 0.1 ^A	115 ± 4.8 ^A	80 ± 2 ^A
<i>CmNox1-C7</i>	2.8 ± 0.1 ^A	119 ± 4.8 ^A	81 ± 2 ^A
<i>CmNox1-C8</i>	2.8 ± 0.1 ^A	121 ± 4.8 ^A	80 ± 2 ^A
Δ <i>CmNox2-20</i>	2.9 ± 0.1 ^A	120 ± 5.4 ^A	81 ± 3 ^A
Δ <i>CmNox2-323</i>	2.8 ± 0.1 ^A	122 ± 5.4 ^A	82 ± 3 ^A
Δ <i>CmNox2-347</i>	2.9 ± 0.1 ^A	125 ± 5.4 ^A	82 ± 3 ^A
Δ <i>CmRac1-79</i>	0.5 ± 0.1 ^B	0 ^B	0 ^B
Δ <i>CmRac1-95</i>	0.5 ± 0.1 ^B	0 ^B	0 ^B
Δ <i>CmRac1-102</i>	0.4 ± 0.1 ^B	0 ^B	0 ^B
<i>CmRac1-C1</i>	2.9 ± 0.1 ^A	121 ± 9.8 ^A	81 ± 4 ^A
<i>CmRac1-C4</i>	2.9 ± 0.1 ^A	119 ± 9.8 ^A	80 ± 4 ^A
<i>CmRac1-C7</i>	2.8 ± 0.1 ^A	120 ± 9.8 ^A	81 ± 4 ^A
<i>OVS-Rac1-1</i>	0.6 ± 0.1 ^B	0.6 ± 5 ^A	40 ± 3 ^A
<i>OVS-Rac1-3</i>	0.5 ± 0.1 ^B	0.6 ± 5 ^A	45 ± 3 ^A
<i>OVS-Rac1-4</i>	0.6 ± 0.1 ^B	0.5 ± 5 ^A	39 ± 3 ^A
Δ <i>CmNoxR-5</i>	2.9 ± 0.1 ^A	0 ^B	0 ^B
Δ <i>CmNoxR-107</i>	2.8 ± 0.1 ^A	0 ^B	0 ^B
Δ <i>CmNoxR-160</i>	2.8 ± 0.1 ^A	0 ^B	0 ^B
Δ <i>CmBem1-344</i>	2.9 ± 0.1 ^A	124 ± 14.2 ^A	84 ± 2 ^A
Δ <i>CmBem1-359</i>	2.9 ± 0.1 ^A	125 ± 14.2 ^A	85 ± 2 ^A
Δ <i>CmBem1-370</i>	2.9 ± 0.1 ^A	120 ± 14.2 ^A	80 ± 2 ^A

Table 1. Comparison of hyphal growth rate, conidial production, and parasitic ability among mutants and the wild-type strain ZS-1 of *C. minitans*. ^aGrowth rate was detected by measuring the colony diameter of cultures incubated at 20 °C for 7 days. ^bConidia produced by 14-day-old cultures and counted with a haematocytometer. ^cRot index of sclerotia was calculated after infected by *C. minitans* for 30 days¹⁴. ^dDifferent letters in the same column indicated statistically significant differences (P = 0.05). Means and standard errors were calculated from three replicates.

conditions. However, there was no significant difference in the growth rate between *CmNoxR* deletion mutants and ZS-1, and the deletion mutant displayed a phenotype similar to *CmNox1* deletion mutant (Fig. 3e, Table 1). Δ*CmBem1-344* shared the same phenotype as strain ZS-1. It suggests that *CmRac1* and *CmNoxR* are required to activate *CmNox1* to adjust conidiation and parasitism, while *CmBem1* is not involved.

CmSlt2 is *CmNox1* effector required for conidiation and parasitism. To understand the mechanism of *CmNox1*-mediated conidiation and parasitism in *C. minitans*, we further investigated the functional relationship between *CmNox1* and *CmSlt2*. We reported previously that *CmSlt2* (cell wall integrity-related MAP kinase) is essential for conidiation of *C. minitans*. Δ*CmSlt2* lost the ability to produce pycnidia and conidia, and had dramatically reduced sclerotial mycoparasitism¹³. The phenotype of Δ*CmSlt2* was similar to that of Δ*CmNox1*, thus *CmSlt2* and *CmNox1* might have somewhat interaction directly or indirectly. To identify whether *CmSlt2* interact with *CmNox1*, full length gene of *CmSlt2* and *CmNox1* were cloned into yeast two-hybrid vector pGADT7 and pGBDT7 to test the possible interaction, and the results showed that *CmSlt2* could interact with *CmNox1* (Fig. 4a). Furthermore, we generated fusions of *CmSlt2* and *CmNox1* with N- and C-terminal domains of YFP and assessed their interaction in tobacco (*Nicotiana benthamiana*) using bimolecular fluorescence complementation (BiFC) (Fig. 4c). To confirm this interaction in *C. minitans*, a transformant with TrpC-*CmNox1*-Flag and TrpC-*CmSlt2*-GFP and a transformant with TrpC-*CmSlt2*-GFP and Vector-Flag in ZS-1 were constructed. In Western blot analysis with total protein, the anti-Flag and anti-GFP antibodies detected a 65-kDa and an 80-kDa band, respectively. In proteins eluted from anti-Flag agarose, the 80-kDa *CmSlt2*-GFP band was detected with an anti-GFP antibody in transformant with *CmSlt2* and *CmNox1*, but not in transformant with *CmSlt2* only (Fig. 4b). The results showed that *CmSlt2* and *CmNox1* interacted directly in *C. minitans*.

We expressed the *CmSlt2*-GFP fusion in the wild-type strain ZS-1 and Δ*CmNox1*, and found that over-expressing *CmSlt2*-GFP could not rescue the deficiency of conidiation and parasitism in Δ*CmNox1* (Fig. 5a,b). In the wild-type strain ZS-1, *CmSlt2* was located in the cytoplasm during the hyphal growth period (3-day-old hyphae), and was located in the nucleus in 40% of the hyphae at the later stage of conidiation (5-day-old hyphae) observed under a fluorescence microscope; While in Δ*CmNox1*, no fluorescence signal of *CmSlt2*-GFP was observed in the nucleus at any stages (Fig. 5c). The results suggested that *CmSlt2* was a downstream regulatory factor of *CmNox1* and was involved in conidiation and parasitism.

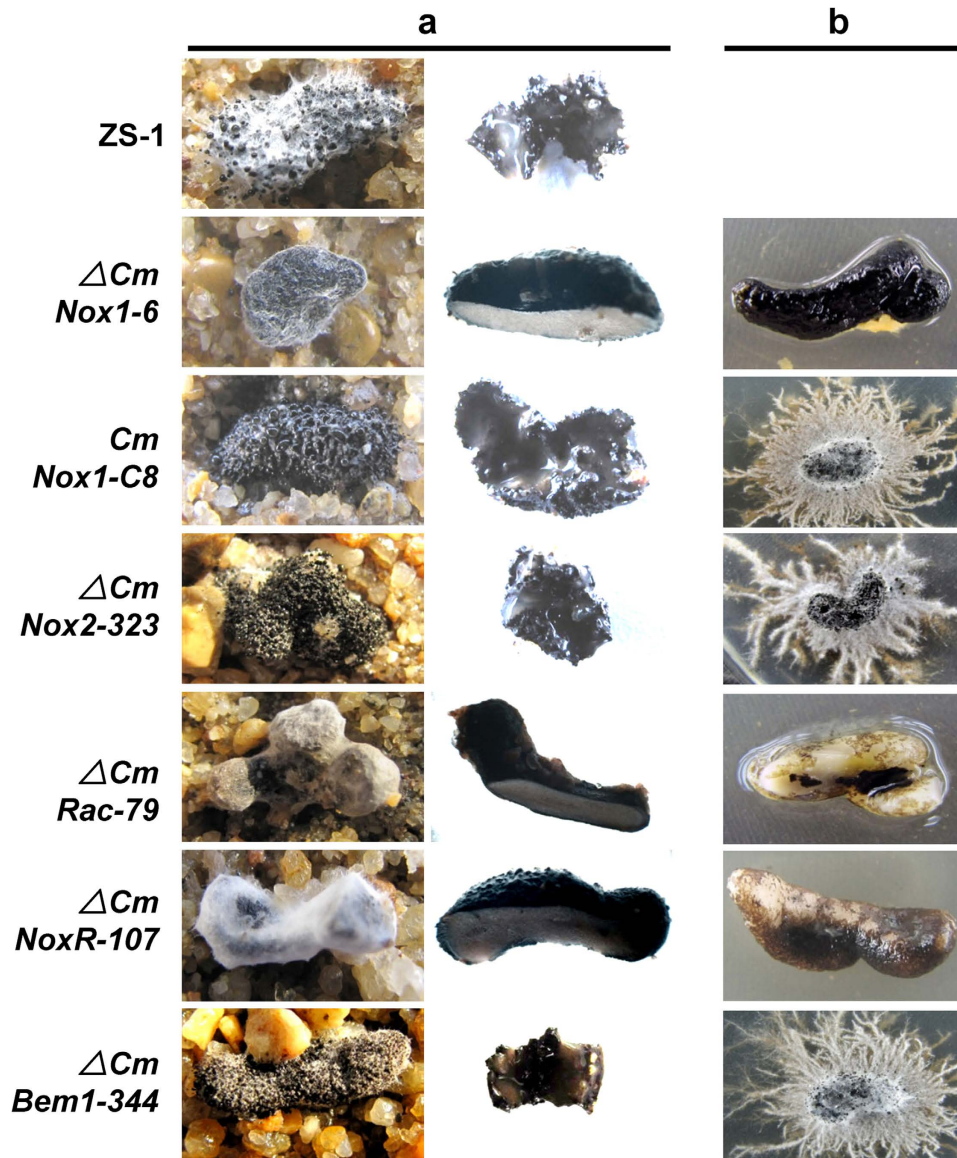


Figure 2. Parasitism of *C. minitans* mutants to sclerotia of *S. sclerotiorum*. (a) Microscopic observation of the surface and interior of sclerotia infected by *C. minitans*. (b) Colony formed by all inoculated sclerotia. All inoculated sclerotia were surface sterilized and then placed on hygromycin (50 µg/ml) amended PDA for 7 days.

Over expressing *CmSlt2* in $\Delta CmRac1$ could partially restore conidiation and parasitism. Since *CmRac1* interacts with *CmNoxR* together in regulating *CmNox1*, and *CmNox1* mediates the localization of *CmSlt2* in nucleus, we suspected that *CmRac1* is likely to regulate the expression of *CmSlt2*. qRT-PCR analysis showed that the expression of *CmSlt2* was obviously decreased in $\Delta CmRac1$ by 0.5 fold, compared to the wild-type strain (Fig. 6a). We fused *CmSlt2* with a *trpC* promoter from *Aspergillus* and transformed into $\Delta CmRac1$, and found that over-expressing of *CmSlt2* could partially restore conidiation and parasitic ability of $\Delta CmRac1$ (Fig. 6b, and Table 1), in spite of no significant improvement of the hyphal growth.

The expression of *CmPks1* was suppressed in *CmNox1* signal pathway mutants. *C. minitans* produces dark pigment and black pycnidia during the late stage of growth. However, all $\Delta CmRac1$, $\Delta CmNoxR$, $\Delta CmNox1$, and $\Delta CmSlt2$ produce little dark pigment, and colonies were whitish. It is likely that the expression of *CmPks1*, a melanin biosynthesis associated polyketide synthase-encoding gene, was suppressed in those mutants. The transcript profile of *CmPks1* was monitored by qRT-PCR, and the result showed that the expression of *CmPks1* peaked at 96 hpi in the wild type ZS-1 and was significantly suppressed in $\Delta CmNoxR$, $\Delta CmRac1$, $\Delta CmNox1$ and $\Delta CmSlt2$, whereas there was no obvious difference in the transcript level between $\Delta CmNox2$ and ZS-1 (Fig. 7a). Further experiments proved that the expression of *CmPks1* in ZS-1 was enhanced by 300–400 fold when dual cultured with *S. sclerotiorum* than cultured on PDA alone at 24 hpi and 36 hpi (Fig. 7b), while it could not be induced in $\Delta CmRac1$ and $\Delta CmNox1$ (Fig. 7c). These results indicated that the expression of *CmPks1* could

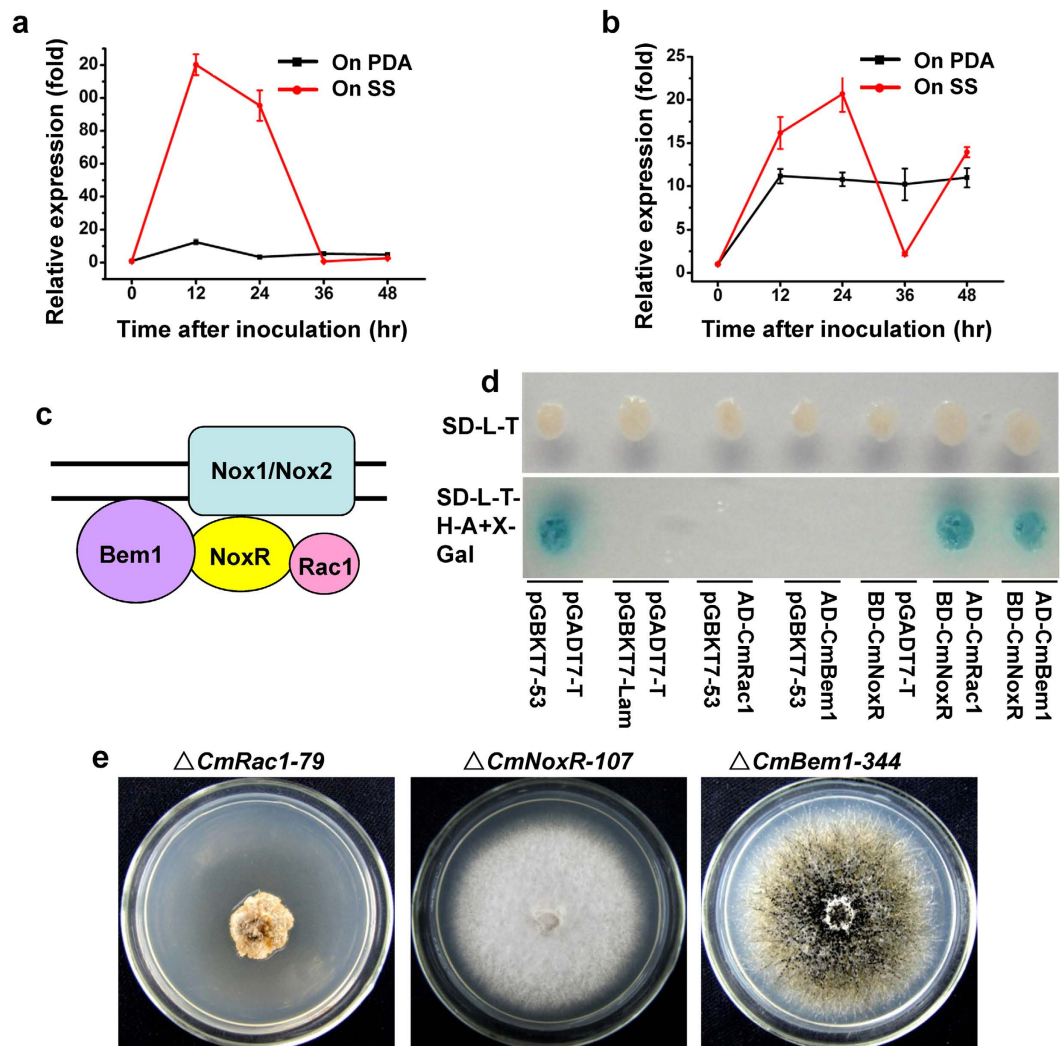


Figure 3. NADPH oxidase complexes in *C. minitans*. Expression analysis of *CmNox1* (a) and *CmNox2* (b). Total RNA was isolated from mycelia of ZS-1 cultured alone or dual-incubated with *S. sclerotiorum* on PDA, and used for cDNA synthesis. The transcript level of *CmActin* was used to normalize different samples. Bars represent means and standard deviations (three replications). (c) Schematic drawing showing the NADPH oxidase complexes in *C. minitans*. (d) Yeast two-hybrid assay of interaction between *CmNoxR*, *CmBem1*, and *CmRac1*. (e) Colony morphology of *CmRac1* deletion mutant $\Delta CmRac1-79$, *CmNoxR* deletion mutant $\Delta CmNoxR-107$ and *CmBem1* deletion mutant $\Delta CmBem1-344$ grown on PDA at 20 °C for 12 days.

be induced at the early stages of interaction with *S. sclerotiorum*, while could not in all of the *CmNox1* signal pathway mutants either cultured on PDA or interacted with *S. sclerotiorum*. It suggested that the *CmNox1* signal pathway is involved in regulating the expression of *CmPks1*.

The *CmNox1* signal pathway is involved in regulating cell wall degrading enzyme genes. The parasitic ability of all mutants in *CmNox1* signal pathways decreased obviously and it is also reported that the cell wall degrading enzyme *Cmg1* (β -1,3 glucanase-encoding gene) and *CH1* (chitinase gene) play important roles in the process of parasitizing sclerotia of *S. sclerotiorum*⁹. Our results of qRT-PCR demonstrated that the transcript levels of both *Cmg1* and *CH1* in ZS-1 were enhanced significantly at 24 hpi and 36 hpi when dual cultured with *S. sclerotiorum* (Fig. 7e,h), while were suppressed in $\Delta CmNoxR$, $\Delta CmRac1$, $\Delta CmNox1$ or $\Delta CmSlt2$ when incubated on PDA compared to that in ZS-1 (Fig. 7d,f,g,i). The results indicated that *Cmg1* and *CH1* were involved in parasitism and could be regulated by the *CmNox1* signal pathway.

Discussion

The mycoparasitism system of *C. minitans*/*S. sclerotiorum* is unique and important to probe fungi and fungi interaction. In this study, the role of NADPH oxidases complex of *C. minitans* on growth, conidiation and mycoparasitism were studied. We found that *CmNox1*, but not *CmNox2*, played essential roles on conidiation and mycoparasitism. Furthermore, we found that *CmRac1* could interact with *CmNoxR*, and *CmNox1* could interact with *CmSlt2*. Expression of *CmSlt2* in $\Delta CmRac1$ could partially restore the conidiation and parasitism. These

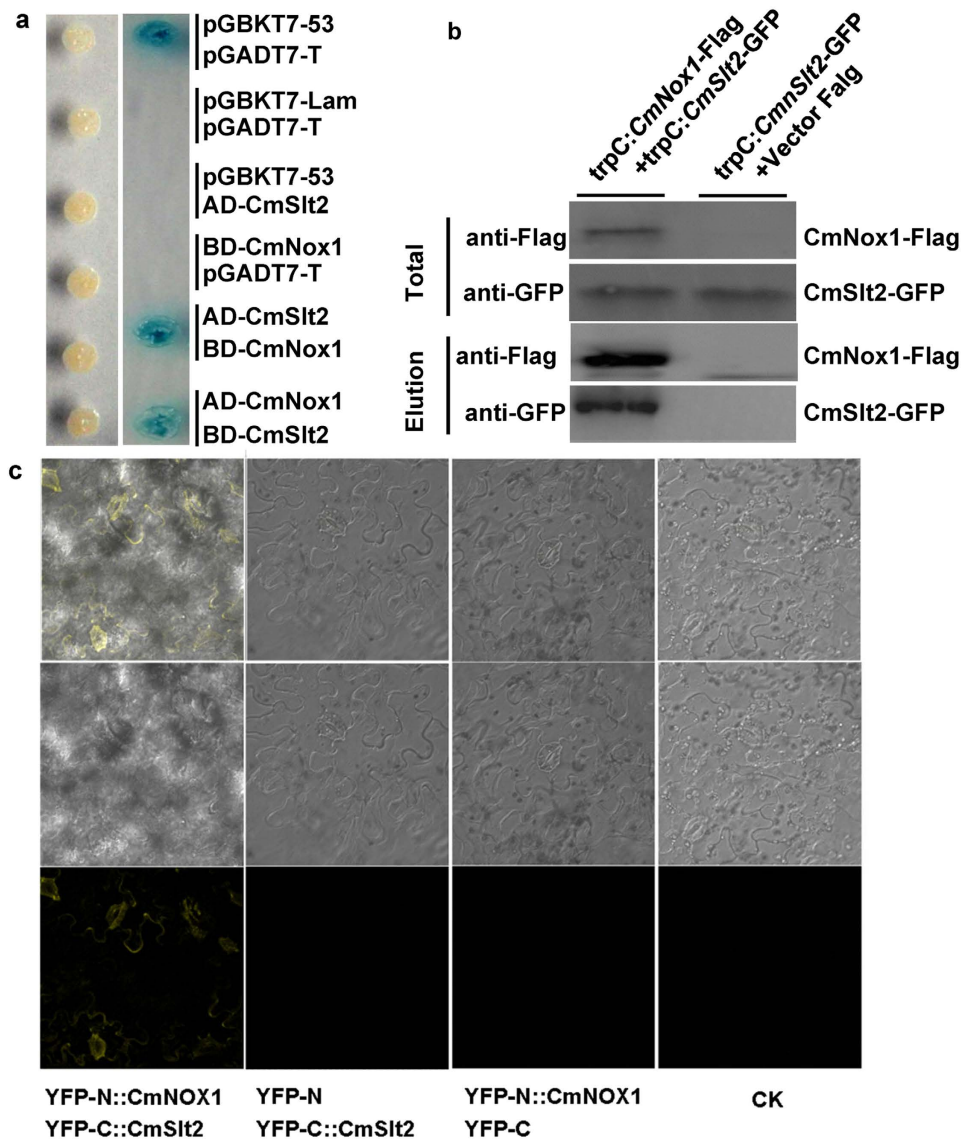


Figure 4. Interaction of CmNox1 with CmSl2. (a) Yeast two-hybrid assay of interaction between CmNox1 and CmSl2. (b) Co-IP assays. Western blots of total proteins and proteins eluted from anti-Flag agarose from transformant (*CmSl2*-GFP and vector Flag) and transformant (*CmSl2*-GFP and *CmNox1*-Flag) were detected with anti-Flag or anti-GFP antibodies. (c) Bimolecular fluorescence complementation (BiFC) analysis of the interaction between CmNox1 and CmSl2 tagged with YFP in *Nicotiana benthamiana*.

findings broadened and strengthened our knowledge of mycoparasitism, and supplied a possible link between Nox complex signal and MAPK cascade signal in fungi.

Fungal Nox isoforms have different roles in fungi lifecycle. In *M. oryzae*, both Nox1 and Nox2 are essential for pathogenicity²¹. Nox1 is required for penetration, hyphal elongation, and Nox2 is required for assembly of a toroidal F-actin network during the penetration peg formation²²; In *S. sclerotiorum*, Nox1 is required for virulence, oxalic acid production and sclerotial development, while, Nox2 might be involved in sclerotial development²⁴. In *P. anserine*, *PaNox1* mutants are impaired in the differentiation of fruiting bodies from their progenitor cells, and deletion of the *PaNox2* specifically blocked ascospore germination^{33,54}. New Nox proteins were also identified from *P. anserine*, a Nox isoform Nox3 was found to play a minor role³⁶. In *N. crassa*, Nox1 is required for female sterility, and involved in asexual development and hyphal growth, while Nox2 might be involved in ascospore germination³⁴. In *E. festucae*, both Δ *NoxA* and Δ *NoxB* mutants could produce conidia, while NoxA, but not NoxB, is essential for hyphal polarized growth and hyphal fusion, NoxB did not affect conidiation²⁹. In *Sordaria macrospora*, Nox1 is required for fruiting body formation, normal hyphal growth, and hyphal fusion, while Nox2 is involved in strict melanin-dependent ascospore germination⁵⁵. In *C. purpurea*, CpNox1 is essential for infection, but CpNox2 is not essential for infection. Interestingly, Δ *cpnox2* mutants converted endophytic lifestyle to pathogenic lifestyle, and CpNox2 functioned in the infection process and moderates damage to the host⁵⁶. In this study, the Nox1 of

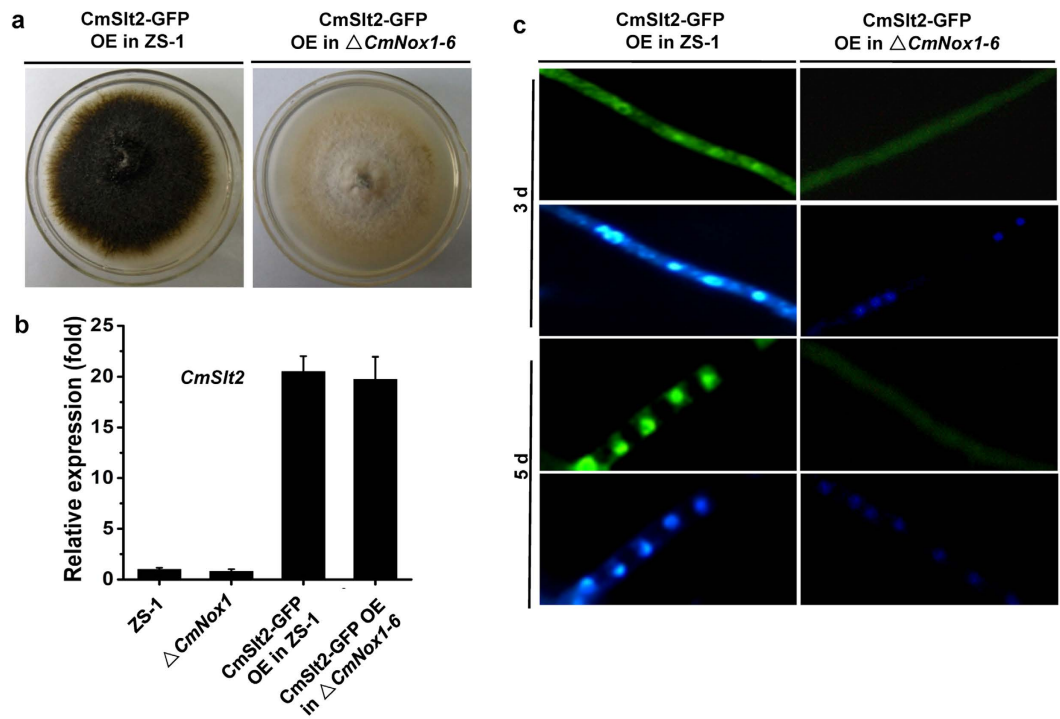


Figure 5. CmNox1 controls the nucleus location of CmSlt2. (a) Colony morphology of CmSlt2-GFP-overexpressed ZS-1 and Δ CmNox1-6. (b) Relative expression of CmSlt2 in CmSlt2-GFP-overexpressed ZS-1 and Δ CmNox1-6. The expression of ZS-1 was set as level one. (c) CmSlt2-GFP nuclear localization in CmSlt2-GFP-overexpressed ZS-1 and Δ CmNox1-6.

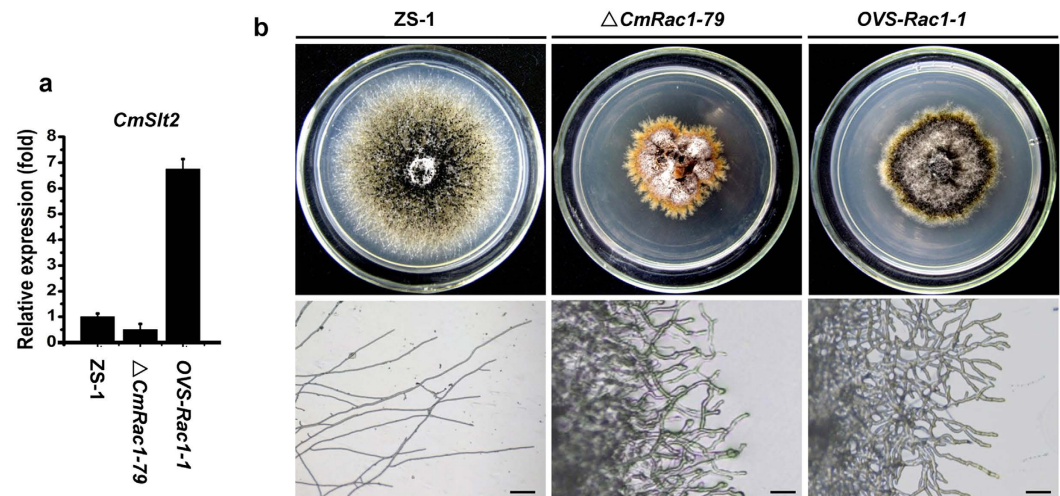


Figure 6. Over-expression of the MAP kinase gene CmSlt2 partially restored the conidiation of Δ CmRac1-79. (a) The relative expression of CmSlt2 in the wild type strain ZS-1, Δ CmRac1-79 and OVS-Rac1-1 (CmSlt2 over-expressing in Δ CmRac1-79). (b) Comparison of colony morphology and tip hyphae branching pattern among the wild-type strain ZS-1, Δ CmRac1-79, and OVS-Rac1-1 grown on PDA medium at 20 °C for 20 days.

C. mitans is essential for mycoparasitism, conidiation and pigmentation, while deletion of Nox2 did not affect these bioprocesses. However, how Nox1 regulates these bioprocesses is necessary to be unraveled.

Rac1, a member of the Rho-family GTPases, widely exists in eukaryotes and in fungi as well. Sequence alignment of RAC1 homologs from fungi revealed significant conservation in amino acid composition and RAC1 from *M. grisea* could rescue the conidiation, parasitization and growth in Δ Rac1 mutant of *C. mitans* (data not shown). Fungal Rac1 is crucial for the growth, virulence and development in many fungi, such as *M. grisea*, *U. maydis*, *A. fumigatus*, *Claviceps purpurea* and *F. graminearum*⁵⁷⁻⁶¹. In *M. grisea*, MgRac1 is essential

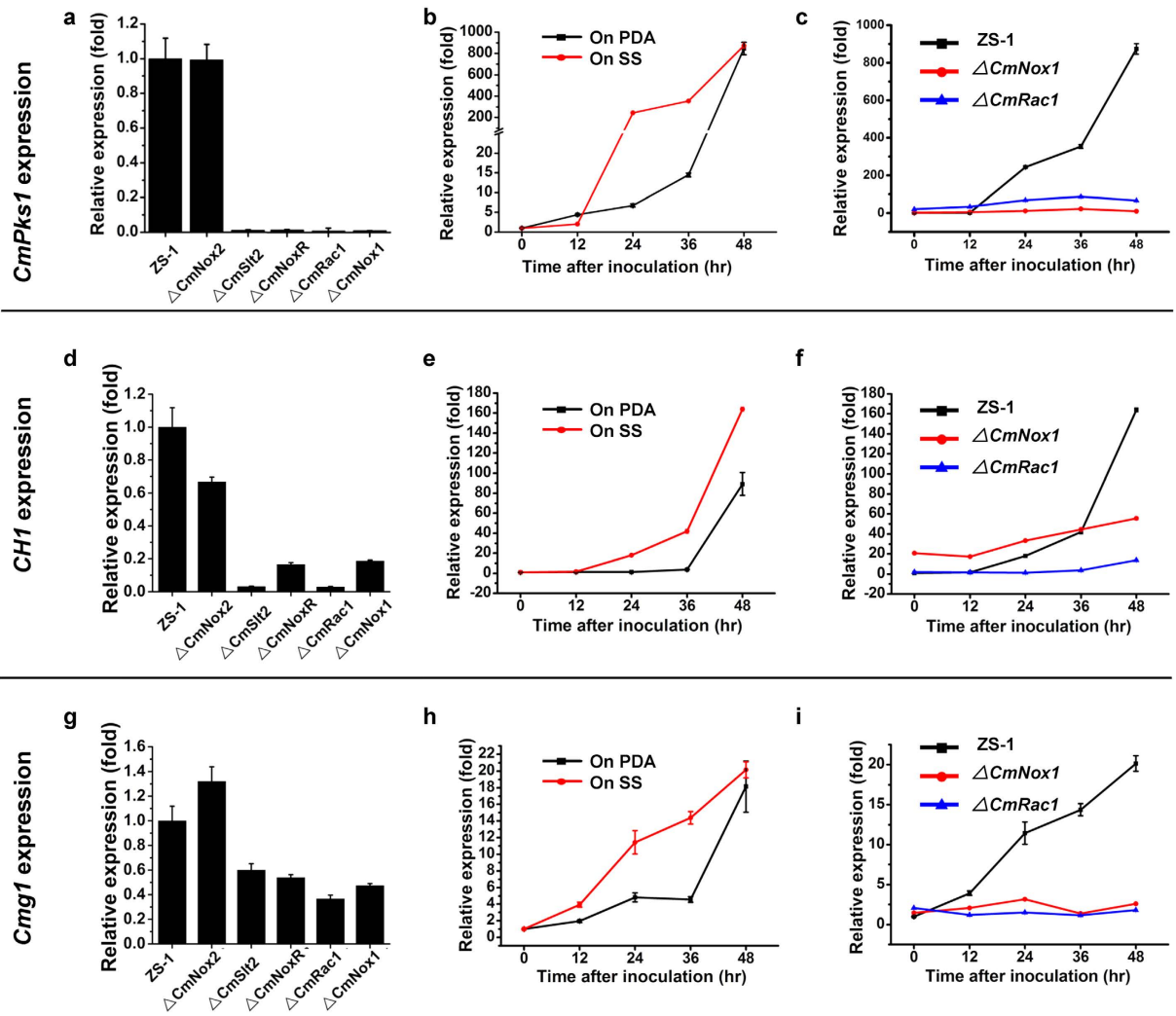


Figure 7. Gene expression analysis in the wild-type strain ZS-1 and deletion mutants of *C. minutans*.

Relative transcript accumulations of *CmPks1* (a), *CH1* (d) and *Cmg1* (g) detected with qRT-PCR amplification in ZS-1, Δ *CmNox2*, Δ *CmSlt2*, Δ *CmNoxR*, Δ *CmRac1* and Δ *CmNox1* after growing on PDA for 96 hr. The expression of ZS-1 was set as level one. Relative transcript patterns of *CmPks1* (b), *CH1* (e) and *Cmg1* (h) in ZS-1 after contacting with *S. sclerotiorum* (red) or growing on PDA (black) for 0–48 hr. The gene expression of ZS-1 inoculated in plate at 0 hr was set as level one. Relative transcript patterns of *CmPks1* (c), *CH1* (f) and *Cmg1* (i) genes in ZS-1 (black), Δ *CmNox1* (red) and Δ *CmRac1* (blue) after growing on PDA for 0–48 hr. The gene expression of ZS-1 at 0 hr was set as level one. The relative level of transcript was calculated by the comparative Ct method. The level of *CmActin* transcript was used to normalize different samples. Bars represent means and standard deviations (three replications).

for conidiogenesis, and contributes to the formation of appressorium and pathogenicity through activating its downstream factors: the PAK kinase Chm1 and NADPH oxidases⁵⁸. In *Candida albicans*, Rac1 is an upstream regulatory factor of the MAP kinase Cek1 and Mkk1, but the control mechanism is still unclear⁶². In this study, the qRT-PCR assay demonstrated that the expression level of *CmSlt2* was obviously decreased in Δ *CmRac1*. When *CmSlt2* was over expressed in Δ *CmRac1*, the conidiation could be partially restored, further confirming that Rac1 was an upstream regulatory factor of MAPK cascade signal pathway.

The fact that the Nox complex mutants and the MAPK cascade pathway mutants of fungi sharing highly similar phenotypes suggests that these two signal pathways are cross-linked. This study provides further evidence supporting this cross-link since *CmNox1* could interact with *CmSlt2*, and *CmRac1* could interact with *CmNoxR*. Lalucque *et al.* reported that PaNox1 acted on the PaASK1/PaMkk1/PaMpk1 MAPK module by promoting nuclear translocation of the PaMpk1 MAP kinase in *P. anserina*⁶³. More recently, H₂O₂ was found to induce the relocalization of a putative MAPK-activated protein kinase SrkA to nuclei and mitochondria under the presence of Saka in *A. nidulans*⁵³. We further found that *CmNox1* could regulate the nucleus location of a cell wall integrity-associated MAP kinase (*CmSlt2*). In Δ *CmNox1*, *CmSlt2* could not move into fungal nuclei, suggesting that Nox complex-derived ROS may function on the localization of *CmSlt2*.

The Nox complex signal and MAPK cascade signal affect fungal gene expression globally to regulate complicated processes in physiology, pathogenicity and development. In *S. macrospora*, Nox1 affects the expression of genes involved in cytoskeleton remodeling, hyphal fusion, metabolism, and mitochondrial respiration⁵⁵. In *P. anserine*, the expressions of 15% genes were modified in $\Delta PaMpk1$, $\Delta PaMpk2$ and $\Delta PaNox1$, and about 1000 genes were regulated similarly in these three mutants⁶⁴. We attempted to explain how the Nox complex signal and MAPK cascade signal regulate mycoparasitism by monitoring the expression levels of *CmPks1*, *Cmg1* and *CHI* in *C. minitans*. We found that expression of these genes was obviously enhanced when interacted with its host fungus *S. sclerotiorum*, while could not be induced in $\Delta CmNox1$, $\Delta CmRac1$ and $\Delta CmSlt2$, indicating that these three genes were regulated by the Nox complex signal and MAPK cascade signal pathways, thus to affect parasitism. Mycoparasitism of *C. minitans* is a very complicated process. There are many other genes that are also involved in the mycoparasitism and remain to be investigated.

Briefly, we analyzed the function of Nox complex on mycoparasitism and conidiation of *C. minitans*, and we found that it is *CmNox1*, but not *CmNox2*, that is required for mycoparasitism and conidiation. We further found that *CmNox1* interacts with MAP kinase *CmSlt2* and effects the location of *CmSlt2*. Our finding suggests that Nox complex signal pathway and MAPK cascade signal pathway are cross-linked via *CmNox1*.

Methods

Strains and cultural conditions. The *C. minitans* wild-types train ZS-1 (CCAM 041057) was isolated from garden soil at Zhushan County, Hubei Province, P R China⁶⁵. Strain Ep-1PNA367 was a virulent and virus-free strain of *S. sclerotiorum*, derived from a single ascospore of the hypovirulent strain Ep-1PN⁶⁶. The strains used in this research were maintained and cultured on PDA at 20–22 °C and stored in PDA slants at 4 °C^{17,66}. Cultures for genomic DNA and RNA isolation were conducted on PDA at 20–22 °C for 4 d. Conidia were prepared from 15-day-old cultures grown on PDA. The selective PDA was supplemented with 50 µg/ml of hygromycin B (Sigma) or 80 µg/ml of G418 (Sigma), depending on the selection marker in the plasmid vector.

Analysis of colony morphology, growth rate, conidiation and parasitic ability. Colony morphology was observed on PDA after incubating at 20 °C for 12 days. Observation of the pycnidial formation was performed by following the method described¹⁴. To characterize the biological properties of the mutants, growth rate, conidial production and parasitic ability were examined as described^{13,14}. For all the transformants obtained, three individuals were examined (Table1).

ROS detection assay. Production of superoxide was evaluated with NBT using the method modified from Chen *et al.*⁵⁸. The wild-type ZS-1, mutants $\Delta CmNox1$ and $\Delta CmNox2$ were grown on PDA for 7 days. Mycelia were incubated in 0.05 M sodium phosphate buffer, pH 7.5, containing 0.05% (w/v) NBT (Sigma-Aldrich). After 1 h of incubation, the culture was fixed in ethanol to stop the reaction. The stained sample was examined with a compound microscope at ×400 magnification.

Isolation of *CmNox1/CmNox2*. A PCR primer pair 1F1/1R1 (Table S1) was designed based on the *C. minitans* genome database. *CmNox1* was amplified from the ZS-1 genomic DNA by a 32-cycle PCR reaction (94 °C, 1 min; 58 °C, 1 min; 72 °C, 2 min), followed by a 10 min extension at 72 °C. The PCR product was cloned into the pMD18-T vector (TaKaRa) and confirmed by DNA sequencing. The cDNA of *CmNox1* was isolated by RT-PCR from total RNA of *C. minitans* with the primer pair 1F1 and 1R1, followed by cloning into the pMD18-T vector and direct DNA sequencing. The same method was used to clone *CmNox2* in *C. minitans* with the primer pair 2F1/2R1 (Table S1).

Vector construction and Agrobacterium-mediated transformation. The *CmNoxR*, *CmRac1*, *CmNox1*, *CmNox2* and *CmBem1* replacement constructs were generated according to Qin *et al.*¹⁹. The *CmNox1* (primer pairs 9F1/9R1 and 10F1/10R1), *CmNox2* (primer pairs 11F1/11R1 and 12F1/12R1), *CmNoxR* (primer pairs 13F1/13R1 and 14F1/14R1), *CmRac1* (primer pairs 15F1/15R1 and 16F1/16R1), and *CmBem1* (primer pairs 17F1/17R1 and 18F1/18R1) deletion constructs were made by PCR amplification of the 5' - and 3' - flanks of the respective ORFs (0.8–1.2 kb) with genomic DNA of the wild-type strain ZS-1 as template. The fragments were cloned upstream and downstream of *hph* cassette in pMD18 respectively, and then the structures were cloned into the corresponding sites of vector pneo-P3300III, which carries a neomycin resistance gene cassette (neo) as the second selection marker.

In order to construct the complementary vector of *CmNox1*, a 2.7 kb fragment containing the native promoter and ORF of *CmNox1* was amplified by PCR with primer pair 19F1/19R1 (Table S1) and cloned into vector pneoP3300. The complementary mutant *CmNox1-C* was generated by introducing the vector into $\Delta CmNox1-6$, followed by screening for neomycin resistance and RT-PCR confirmation.

cDNA of *CmSlt2* was amplified by RT-PCR with primer pair 20F/20R (Table S1) and cloned into the *HindIII/SmaI* sites of pCIT vector, which contains the *trpC* promoter and terminator, and then the gene cassette was cloned into vector pneoP3300 to get the *CmSlt2* over-expression vector pOVSlt2. The over-expression mutant *OVS-Rac1-1* was generated by introducing pOVSlt2 into $\Delta CmRac1-79$, followed by screening for neomycin resistance and qRT-PCR confirmation.

To tag *CmSlt2* with the GFP, cDNA of *CmSlt2* without stop codon was amplified by RT-PCR with primer pair 20F/20R (Table S1) and cloned into the *HindIII/SmaI* sites of pCIT vector. The GFP coding sequence was excised from plasmid pEGFP-1 with *SmaI/BamHI*, and the corresponding fragment was purified. Both the *CmSlt2* and GFP coding sequences were cloned into vector pneoP3300 digested with *XhoI* to yield plasmid pSGFP1. Mutants were generated by introducing of pSGFP1 into $\Delta CmNox1-6$ and the wide-type strain ZS-1, followed by screening for neomycin resistance. GFP signal was observed under a Nikon Eclipse 80i fluorescent microscope (Nikon, Japan).

DNA extraction and Southern blot analysis. The genomic DNA of the wild-type strain ZS-1 and generated mutants in this study was extracted according to the standard protocols⁶⁷. Southern blot analysis was performed according to Gong *et al.*¹⁷. Genomic DNA (ZS-1 and CmNox1 mutants) aliquots of 15 µg were digested with *Hind*III, separated by electrophoresis on 0.8% agarose gel and transferred onto a Hybond N+ membrane (Amersham Pharmacia Biotech). Interior probe was amplified with the primer pair 30F and 30R (Fig. 2c, Table S1), while the exterior probe was amplified with the primer pair 31F and 31R (Fig. 2c, Table S1). *CmNox2* mutants were also confirmed by Southern blot, with the probe amplified using primer pair 32F and 32R.

RNA manipulation and qRT-PCR analysis. The total RNA sample of fungal strain was isolated with TriZOL reagent (Invitrogen, USA) according to the manufacturer's protocols and potential DNA contamination was removed by DNase I treatment (RNase Free) (TaKaRa, Dalian, China). The first-strand cDNA was synthesized with RevertAid™ First Strand cDNA Synthesis Kit (MBI Fermentas, Lithuania) by following the manufacturer's instructions. In quantitative Real-time PCR, *CmPks1* (primer pair 21F/21R), *CmRac1* (primer pair 22F/22R), *CmNox1* (primer pair 23F/23R), *CmNox2* (primer pair 24F/24R), *CmNoxR* (primer pair 25F/25R), *Cmg1* (primer pair 26F/26R), *CH1* (primer pair 27F/27R) and *CmSlt2* (primer pair 34F/34R) were amplified by using the respective primer pairs (Table S1). As an endogenous control, a 154-bp amplicon of *actin* was amplified with primer pair 28F and 28R (Table S1).

Yeast two-hybrid system. Yeast two-hybrid analysis was carried out using a GAL4-based yeast two-hybrid system-Matchmaker™ Gold Systems (Clontech, Palo Alto, CA). cDNA of *CmNox1* (1F1/1R1), *CmSlt2* (2F2/2R2), *CmRac1* (3F/3R) and *CmNoxR* (4F/4R), were amplified with the respective primer pairs (Table S1) and inserted into the yeast vector pGADT7. To test the specificity of the interaction, the bait plasmid and the prey plasmids were co-transformed into yeast strain Y2HGOLD. The transformants were assayed on SD (synthetic dropout)/-Trp-Leu-His-Ade plates and SD/-Trp-Leu-His-Ade plates with X-α-gal for β-galactosidase test.

Bimolecular fluorescence complementation experiments. For bimolecular fluorescence complementation experiments (BiFC), *CmNox1* and *CmSlt2* were tagged with separated halves of YFP, based on Walter *et al.*⁶⁸, with the following modifications. To fuse YFP-N to *CmSlt2*, *CmSlt2* cDNA without stop codon was amplified with primer pair 7F and 7R and cloned into the *Spe*I/*Sma*I sites of pUC-SPYNE1. To fuse YFP-C to *CmNox1*, *CmNox1* cDNA without stop codon was amplified with primer pair 8F and 8R and cloned into the *Spe*I/*Sma*I sites of pUC-SPYCE1. The two vectors were co-transformed into tobacco (*Nicotiana benthamiana*). Leaf samples were collected 2 days after agro-infiltration. YFP Fluorescence signal was observed under a Nikon Eclipse 80i fluorescent microscope (Nikon, Japan).

In vivo Co-IP. Agrobacterium strain EHA105 carrying the TrpC-*CmSlt2*-GFP and TrpC-*CmNox1*-Flag expression vectors was coinfiltrated into *C. minitans*. The over-expression mutant was generated by screening for neomycin/hygromycin resistance. Mutant transformed with TrpC-*CmSlt2*-GFP and empty Vector-Flag was generated and used as a control. For *in vivo* Co-IP, 8-day-old mutant total protein was extracted in IP buffer (50 mM Tris pH 7.4, 150 mM NaCl, 1% Triton, 1 mM EDTA, 1 mM MgCl₂, and protease inhibitor cocktail). A 15 µl aliquot of anti-Flag agarose (Beyotime) was added to the samples, and the mixtures were incubated for 4 h at 4 °C with shaking. The immunoprecipitated proteins were washed three times with IP buffer and eluted with SDS sample buffer. Eluted sample were loaded on the protein gels for immunoblot analysis using anti-GFP antibody or anti-Flag antibody (Sigma).

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Acknowledgements

The research was financially supported by the National '973' Basic Research Program of China (Grant No. 2012cb114000), the National Natural Science Foundation of China (grant 31371895), the Programme of Introducing Talents of Discipline to Universities in China (the 111 Project no. B14032), and the Program for Changjiang Scholars and Innovative Research Team in University of China (IRT1247).

Author Contributions

W.W., D.J., W.C. and Y.F. designed the research and wrote the paper; W.W., W.Z. and J.C. executed the experiments. J.C., J.X., G.L. and Y.F. performed the data and bioinformatics analyses. All authors read and approved the final manuscript.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Wei, W. *et al.* Nox Complex signal and MAPK cascade pathway are cross-linked and essential for pathogenicity and conidiation of mycoparasite *Coniothyrium minitans*. *Sci. Rep.* **6**, 24325; doi: 10.1038/srep24325 (2016).



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