AMPK Is Involved in Regulating the Utilization of Carbon Sources, Conidiation, Pathogenicity, and Stress Response of the Nematode-Trapping Fungus Arthrobotrys oligospora

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ABSTRACT AMP-activated protein kinase (AMPK), a heterotrimeric complex, can sense energy and nutritional status in eukaryotic cells, thereby participating in the regulation of multiple cellular processes. In this study, we characterized the function of the catalytic α-subunit (SNF1) and the two regulatory β- and γ-subunits (GAL83 and SNF4) of AMPK in a representative nematode-trapping fungus, Arthrobotrys oligospora, by gene knockout, phenotypic analysis, and RNA sequencing. The ability of the AMPK complex mutants (including ΔAosnf1, ΔAogal83, and ΔAosnf4) to utilize a nonfermentable carbon source (glycerol) was reduced, and the spore yields and trap formation were remarkably decreased. Moreover, AMPK plays an important role in regulating stress response and nematode predation efficiency. Transcriptomic profiling between the wild-type strain and ΔAosnf1 showed that differentially expressed genes were enriched for peroxisome, endocytosis, fatty acid degradation, and multilipid metabolism (sphingolipid, ether lipid, glycerolipid, and glycerophospholipid). Meanwhile, a reduced lipid droplet accumulation in ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants was observed, and more vacuoles appeared in the mycelia of the ΔAosnf1 mutant. These results highlight the important regulatory role of AMPK in the utilization of carbon sources and lipid metabolism, as well as providing novel insights into the regulatory mechanisms of the mycelia development, conidiation, and trap formation of nematode-trapping (NT) fungi.

IMPORTANCE NT fungi are widely distributed in various ecosystems and are important factors in the control of nematode populations in nature; their trophic mycelia can form unique infectious devices (traps) for capturing nematodes. Arthrobotrys oligospora is a representative NT fungus which can develop complex three-dimensional networks (adhesive networks) for nematode predation. Here, we demonstrated that AMPK plays an important role in the glycerol utilization, conidiation, trap formation, and nematode predation of A. oligospora, which was further confirmed by transcriptomic analysis of the wild-type and mutant strains. In particular, our analysis indicated that AMPK is required for lipid metabolism, which is primarily associated with energy regulation and is essential for trap formation. Therefore, this study extends the functional study of AMPK in NT fungi and helps to elucidate the molecular mechanism of the regulation of trap development, as well as laying the foundation for the development of efficient nematode biocontrol agents.

KEYWORDS Arthrobotrys oligospora, AMP-activated protein kinase (AMPK), utilization of carbon sources, conidiation, trap formation

The SNF1/AMP-activated protein kinase (AMPK) is a phylogenetically conserved kinase which can maintain cellular energy homeostasis in almost all eukaryotic cells (1, 2). The cell activates AMPK to actively phosphorylate downstream targets, and it acts to restore energy balance by initiating ATP synthesis pathways while shutting
down other nonessential biosynthesis processes that consume ATP when the cellular energy is inadequate to meet the requirements of various physiological activities (3, 4). Furthermore, it has been shown that AMPK is structurally similar in most eukaryotes and is made up of three different subunits, including a catalytic α-subunit and two regulatory β- and γ-subunits (1). In mammals, AMPK is activated primarily by phosphorylation at Thr-172, which lies in the α-subunit, where phosphorylation is regulated by AMP or ADP (5, 6). However, it is not directly regulated by AMP in plants and fungi (7).

In *Saccharomyces cerevisiae*, the AMPK complex is composed of α-subunit SNF1; three alternate β-subunits, GAL83, SIP1, and SIP2; and γ-subunit SNF4 (8). Studies have shown that the SNF1 pathway of *S. cerevisiae* is involved in cell cycle regulation, cell proliferation, endocytosis, carbon source utilization, and stress (9). Moreover, AMPK plays a central role in the regulation of virulence, cell growth, and proliferation in several filamentous fungi. In plant-pathogenic fungus *Magnaporthe oryzae*, the AMPK complex consists of a catalytic subunit, SNF1, and two regulatory subunits SIP2 and SNF4, and it is crucial to aerial hyphae development, conidiogenesis, appressorial formation, and morphology. In addition, the defects of ∆Mosn1 were more serious than those of ∆Mosip2 and ∆Mosn4 mutants. Interestingly, these three mutants inoculated on media with nonfermentable carbons as the sole carbon source displayed a significant reduction in hyphal growth (10). Similarly, deletion of the three genes encoding AMPK subunits in *Fusarium graminearum* resulted in defects in virulence, sexual development, and vegetative growth, whereas ∆Fgsnf1 showed considerable vegetative growth defects compared to the ∆Fggal83 and ∆Fgsnf4 mutants (11). In *Alternaria alternata*, deletion of *Aasn1* reduced the number of conidia, mycelial growth rates, pathogenicity, and utilization of carbon sources, while it significantly increased tolerance to cell wall-perturbing agents (12). Taken together, AMPK is closely involved in the regulation of growth and pathogenicity of diverse fungi.

Of particular interest to nematode-trapping (NT) fungi are the special trapping devices (traps) involved in capturing and killing nematodes (13–16). The traps are specifically formed by the vegetative hyphae of NT fungi, divided into adhesive knobs and columns, adhesive networks, and constricting and nonconstricting rings according to their different predatory methods (17–19). *Arthrobotrys oligospora* is one of the best-known species of NT fungi, which can develop adhesive networks for nematode predation, and its whole genome has previously been sequenced (20). Recently, some signaling proteins were identified to be involved in the mycelium development and pathogenicity of *A. oligospora*, including heterotrimeric G proteins (21–23), small GTPases (24, 25), and mitogen-activated protein kinase (MAPK) (26–28). The *A. oligospora* AMPK is composed of a catalytic α-subunit (AoSNF1) and two regulatory β- (AoGAL83) and γ-subunits (AoSNF4) according to the amino acid sequence alignment of orthologous AMPK subunits in *S. cerevisiae*. It has been reported that AMPK participates in the regulation of mycelial growth, sporulation, and pathogenicity in several pathogenic fungi, whereas its function remains unclear in NT fungi. In this study, we characterized the *A. oligospora* AMPK by targeted deletions and multiphenotypic analysis of the three AMPK complex subunits, and we performed transcriptomic analysis between the wild-type (WT) and mutant strains to further probe the regulatory mechanism of AoSNF1 in mycelium development, cellular processes, and trap formation of *A. oligospora*.

**RESULTS**

**Bioinformatics analysis of the AMPK complex in *A. oligospora***. AMPK is a heterotrimer, composed of a catalytic α-subunit and regulatory β- and γ-subunits. In the *A. oligospora* AMPK, the α-subunit AoSNF1 contains more amino acid (aa) residues (822 aa), a greater molecular weight (90.57 kDa), and more functional domains (protein kinase domain, SNF1, ubiquitin-associated domain, AMPK, C-terminal adenylate sensor domain) than the β-subunit AoGAL83 (399 aa; 43.26 kDa; ASC domain and AMPK glycogen-binding domain) and the γ-subunit AoSNF4 (366 aa, 40.93 kDa, CBS domain). Additionally, the conserved domains contained in the orthologs of the AMPK subunits from other fungi are similar to those of *A. oligospora*, whereas the α-subunit of *Aspergillus nidulans* has no C-terminal adenylate sensor domain, and only one CBS domain is detected in the γ-subunit (Fig. 1A). AoSNF1
also contains a Ser/Thr protein kinase active site (Interpro accession no. IPR008271) and a protein kinase ATP-binding site (Interpro accession no. IPR017441), whereas AoGAL83 and AoSNF4 do not. Phylogenetic analysis showed that the three AMPK subunits (α-, β-, and γ-subunits) from various fungi are divided into different branches (Fig. 1B).

Deletion of AMPK impairs the utilization of carbon sources. Three positive transformations for Aosnf1 and two transformations for genes Aogal83 and Aosnf4 were generated, respectively, as described in Materials and Methods, and they were confirmed by PCR and Southern blot analyses (see Fig. S1 in the supplemental material). Because the independent mutants of each gene exhibited similar phenotypic traits, a single mutant for each gene was selected to represent the results of the analysis. Compared with the WT strain, the mycelial growth and morphology of the ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants almost displayed no difference on tryptone-glucose (TG) and potato dextrose agar (PDA) media (Fig. 2A to D), whereas there was a remarkable difference in colony diameter on Czapek-Dox agar (CDA) medium modified with different carbon sources (Fig. 2E). In addition, aerial hyphae were more abundant on PDA and TG medium (Fig. 2A), whereas the hyphae of the WT and mutant strains on CDA medium were very sparse, and almost no aerial mycelia could be observed.

The colony diameters of ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants were smaller on CDA medium without a carbon source than the wild-type (WT) strain, as well as on the CDA medium supplemented with glucose as the only carbon source. However, the colony diameters of the mutant strains were much smaller than those of the WT strain on the CDA medium supplemented with glycerol or sucrose as the carbon source. Specifically, when glycerol (nonfermentative carbon source) was used as the only carbon source, the colony diameters of the ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants were decreased by 36%, 29%, and 30%, respectively (Fig. 2E).

AMPK is required for conidiation. Deletion of Aosnf1, Aogal83, and Aosnf4 resulted in a significant reduction (P < 0.05) in spore production. Compared with the WT strain, the spore yields of ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants were decreased by 81.5%, 74.7%, and 73.9%, respectively. Among them, the ΔAosnf1 mutant produced fewer spores than the WT and other two mutant strains (Fig. 3A and B). However, the spore morphology of all mutants showed no difference from the WT strain (Fig. 3C).

To further probe the function of AMPK in regulating conidiation, the transcript levels of eight sporulation-related genes (veA, flug, abaA, medA, ireA, flbC, vosA, and flbA) (29) in the WT and mutant strains cultured at different growth stages were examined by reverse transcription-PCR (RT-PCR). In the ΔAosnf1 mutant, two genes (flug and abaA) were upregulated and one gene (flbA) downregulated on the third day, one gene (medA) was upregulated and four genes (abaA, flbC, vosA, and flbA) downregulated on the fifth day, and five genes (medA, ireA,
flbC, vosA, and flbA) were downregulated on the seventh day (Fig. 3D). In the ΔAogal83 mutant, three genes (fluG, abaA, and vosA) were upregulated on the third day, and two genes (fluG and vosA) were upregulated and four genes (abaA, lreA, flbC, and flbA) downregulated on the fifth day. On the seventh day, veA, flbC, abaA, and vosA genes were upregulated, and medA, lreA, flbC, and flbA genes were downregulated (Fig. 3E). In the ΔAosnf4 mutant, three genes (medA, vosA, and flbA) were upregulated on the third day, two genes (medA and lreA) were upregulated and three genes (veA, fluG, and abaA) downregulated on the fifth day, and three genes (fluG, abaA, and vosA) were upregulated, and two genes (medA and veA) were downregulated on the seventh day (Fig. 3F).

**AMPK plays a crucial role in trap formation and nematode predation.** We observed significant differences (P < 0.05) in producing traps between the WT and mutant strains after being induced with *Caenorhabditis elegans* (Fig. 4A). At 12 h, the WT and mutant (ΔAosnf1, ΔAogal83, and ΔAosnf4) strains started to produce a few immature traps containing a half or single hyphal loop, and they formed an average of 43, 3, 10, and 3 traps per plate, respectively. All the WT and mutant strains produced a certain number of mature traps at 24 h, but the WT strain produced significantly (P < 0.05) more traps than the three mutant strains, as well as at 36 and 48 h (Fig. 4B). At 12 h, fewer nematodes were captured by the mutants than by the WT strain. At 24 h, 50.0% of the nematodes were captured by the WT strain, whereas 28.1%, 19.9%, and 40.87% of the nematodes were captured by the ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants, respectively. At 36 h, 95.4% of the nematodes were captured by the WT strain, whereas 96.4%, 56.3%, and 61.7% of the nematodes were captured by ΔAosnf1, ΔAogal83, and ΔAosnf4, respectively. However, at 48 h, more than 95% of the nematodes were captured by the WT and mutant strains (Fig. 4C).

According to the size of hydrolytic circles on the milk plate, the ΔAosnf1 mutant strain produced more extracellular hydrolytic enzymes (proteases) than the WT strain,
whereas the hydrolyzed circles of the ΔAogal83 and ΔAosnf4 mutants were almost the same as those of the WT strain (Fig. 4D). Furthermore, we determined the transcription of six genes (76g4, 75g8, 112g42, 78g136, 215g702, and 54g992) encoding serine proteases by RT-PCR. Except for the 76g4 gene, which was upregulated on the third day and downregulated on the seventh day, other genes (75g8, 112g42, 78g136, 215g702, and 54g992) were remarkably upregulated to various degrees in the ΔAosnf1 mutant at different time points. In the ΔAogal83 mutant, four genes (75g8, 112g42, 215g702, and 54g992) and two genes (78g136 and 54g992) were substantially upregulated on the third and fifth days, respectively. On the seventh day, one gene (54g992) was upregulated, and two genes (76g4 and 215g702) were downregulated. Furthermore, in the ΔAosnf4 mutant, four genes (76g4, 75g8, 78g136, and 54g992), four genes (75g8, 112g42, 78g136, and 215g702), and three genes (75g8, 78g136, and 215g702) were remarkably upregulated at the three different time points, whereas one gene (76g4) and two genes (76g4 and 54g992) were downregulated on the fifth and seventh days, respectively (Fig. 4E).

**AMPK regulates stress response.** Compared with the WT strain, the mycelial growth of the ΔAosnf1 mutant was more suppressed when cultured on TG plates supplemented with oxidative agents (H2O2 and menadione), while the ΔAogal83 and ΔAosnf4 mutants were suppressed under 0.07 and 0.09 mM menadione but grew faster under 10 mM H2O2 (Fig. 5A). The relative growth inhibition (RGI) values of the AMPK mutants were significantly increased (∗, P < 0.05) compared to the WT strain in the presence of 0.07 and 0.09 mM menadione, and the RGI value of the ΔAosnf1 mutant was also remarkably increased under 0.05 mM menadione, as well as 5 and 10 mM H2O2. In contrast, the RGI values were decreased in the ΔAogal83 and ΔAosnf4 mutants under 10 mM H2O2 (Fig. 5B to D). Meanwhile, the ΔAosnf1 mutant grew faster than the WT strain under 0.09 mg/mL Congo red and 0.2 M NaCl (Fig. S2A and B). The RGI value of

![FIG 3](image-url) Conidiation and transcript levels of sporulation-related genes of the WT and mutant strains. (A) Sporulation of the WT and mutant strains on PDA medium. Red arrows, conidium. Scale bar, 50 μm. (B) Spore yields. (C) Conidia stained with calcofluor white (CFW) and examined under a confocal laser scanning microscope. Scale bar, 5 μm. (D to F) Relative transcription levels (RTLs) of sporulation-related genes in the WT and mutant strains. ∗, significant difference between the mutant and WT strains (P < 0.05). CK was used as the standard (RTL = 1) for statistical analysis of the RTL of each gene under a given condition.
the ΔAogal83 mutant was considerably decreased under 0.06 mg/mL Congo red and increased under 0.02% SDS, as well as 0.5 and 0.75 M sorbitol (Fig. S2C and D), whereas that of ΔAosnf4 was remarkably decreased under 0.06 and 0.09 mg/mL Congo red and increased under 0.3 M NaCl and 0.25 M sorbitol (Fig. S2E and F).

We further analyzed the expression of five genes (glr, gld, per, thi, prx, and cat) associated with oxidative stress response in the WT and mutants (Fig. 5E). In the ΔAosnf1 mutant, with the exception of the glr and cat genes, which were upregulated on the third and fifth days, respectively, the expression levels of the other genes were remarkably downregulated at different time points. In the ΔAogal83 mutant, all five genes were downregulated on the third and seventh days, one gene was downregulated and two genes (thi and cat) upregulated on the fifth day. In the ΔAosnf4 mutant, two genes (glr and per) were downregulated and two genes (thi and cat) upregulated on the third and seventh days, respectively. Additionally, one gene (cat) was upregulated, and gene per was downregulated on the fifth day.

Transcriptomic insight into the regulatory role of AoSNF1. The WT and ΔAosnf1 mutant strains were induced with nematodes for 0 and 12 h, and then the mycelial samples were collected for RNA sequencing. Quality control data showed that the error rate of transcriptome data was less than 0.1%, while the Phred-like quality scores of the samples at the Q20 and Q30 levels were both higher than 90%, and the GC content was 48.23 to 48.82% (Table S1). The quality assessment of the transcriptome showed that the saturation quality of the sequencing was high, the sequencing depth was sufficient to cover most expressed genes, and the sequences obtained by sequencing were evenly distributed on the genes. In the principal-component analysis (PCA), the proportions of PC1 and PC2 that could distinguish samples were 73.81% and 8.31%, respectively. Moreover, the four samples were distributed in different regions.
indicating high similarity and good reproducibility of the three replicates (Fig. S3). To verify the transcriptome data, the transcription levels of 11 selected genes related to peroxisome, glycerolipid metabolism, and endocytosis were examined by RT-PCR. The results showed that expression levels of all six genes related to peroxisome were up-regulated, whereas genes related to glycerolipid metabolism and endocytosis (except 210g5) were downregulated at 0 and 12 h. These results were highly consistent with transcriptome analysis (Fig. S4).

There were 3,312 differentially expressed genes (DEGs) in the ΔAosnf1 mutant compared with the WT strain at 0 h, of which 662 were specifically expressed. At 12 h, 3,476 DEGs were found, of which 826 were specifically expressed (Fig. 6A). Among the total DEGs, 1,452 genes were upregulated, and 1,860 genes were downregulated at 0 h. At 12 h, 1,430 genes were upregulated, and 2,046 were downregulated (Fig. 6B). In gene ontology (GO) enrichment analysis, upregulated genes were enriched for 65 GO terms, and downregulated genes were enriched for 76 GO terms at 0 h, while upregulated genes were enriched for 93 GO terms, and downregulated genes were enriched for 89 GO terms at 12 h (Fig. S5A). The top 15 GO terms enriched by upregulated and downregulated DEGs at different time points almost all belonged to biological processes. The upregulated GO terms included coenzyme biosynthetic process, pantothenate biosynthetic process, and pantothenate metabolic process at 0 h and peroxisome, oxidoreductase activity, and oxidation-reduction process at 12 h (Fig. S5B and C). The downregulated GO terms included response to oxidative stress, nitrate assimilation, nitrate metabolic process, response to nutrient, and

**FIG 5** Stress response between the WT and mutant strains. (A) Growth of the WT and AMPK mutant strains on media supplemented with oxidants. (B to D) Relative growth inhibition (RGI) of the WT and mutant strains (ΔAosnf1, ΔAoga83, and ΔAosnf4) cultured on TG plates supplemented with H2O2 (5 and 10 mM) and menadione (0.05, 0.07, and 0.09 mM). (E) RLTs of the genes related to oxidative stress response in the WT and mutants cultured at different time points. The red line indicates the standard (RTL = 1) for statistical analysis of the RTL of each gene under a given condition. *, significant difference between the mutant and WT strains (Tukey’s HSD, P < 0.05).
regulation of generation of precursor metabolites and energy at 0 h and nitrate assimila-
tion, nitrate metabolic process, response to nutrient, NADPH activity, and oxidoreductase
activity at 12 h (Fig. S5D and E).

In the Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis, there
were 15 and 16 enriched KEGG pathways at 0 and 12 h, respectively (Fig. 6C). Speci-
fically, the KEGG pathways enriched for upregulated genes were the same at different time points,
including peroxisome, sulfur relay system, pantothenate and CoA biosynthesis, sphingolipid
metabolism, fatty acid degradation, and linoleic acid metabolism (Fig. 6D and Fig. S6). In
addition, KEGG pathways enriched for downregulated genes were very similar at 0 and 12
h, including glycerophospholipid metabolism, galactose metabolism, glycerolipid metabo-
lism, ether lipid metabolism, nitrogen metabolism, alpha-linolenic acid metabolism, linoleic
acid metabolism, and glycosphingolipid biosynthesis (globo and isoglobo series), but not
glyoxylate and dicarboxylate metabolism at 0 h and not endocytosis and amino sugar and
nucleotide sugar metabolism at 12 h (Fig. 6D and Fig. S6). On the basis of the KEGG analysis,
we performed cluster analysis of peroxisome-related genes that were enriched (Fig. 6E). The
results showed that the expression levels of genes related to peroxisome in the \( \Delta Aosnf1 \)
mutant were higher than those of the WT strain, and the gene upregulation at 0 h was
more remarkable than that at 12 h. Of these, three genes (4g547, 215g40, and 215g205)
were involved in peroxisome biosynthesis, and four genes (215g56, 215g212, 210g122, and
215g200) were involved in fatty acid oxidation.

**AMPK plays a role in lipid metabolism and cellular components.** By staining the
lipid droplets (LDs) in mycelia, the difference in LD volume between the WT and mu-
tant strains was observed under a confocal laser scanning microscope. Compared with
the WT strain, the volume of LDs was decreased in the \( \Delta Aosnf1 \), \( \Delta Aogal83 \), and
\( \Delta Aosnf4 \) mutants (Fig. 7A). In addition, the transmission electron microscopy (TEM)
results showed that dividing mitochondria and endocytic vesicles could be observed
in the WT strain. In contrast, more vacuoles and lipid droplets (LDs) were observed in
the \( \Delta Aosnf1 \) mutant (Fig. 7B). Meanwhile, the expression levels of genes involved in
lipid metabolism and endocytosis were visualized using heatmaps (Fig. 7C and D).
Compared to the WT strain, the expression levels of the gene encoding fatty acid oxygenase PpoC (AOL_s00215g31) and four genes (AOL_s00215g212, AOL_s00215g56, AOL_s00103g190, and AOL_s00215g200) involved in fatty acid \( \beta \)-oxidation were upregulated in the \( \Delta \)Aosnf1 mutant, whereas 193g190, 210g156, 193g104, 210g100, and 210g115 were downregulated (Fig. 7C). The expression levels of genes (AOL_s00210g80, AOL_s00210g4, and AOL_s00210g5) involved in endocytosis were downregulated in the \( \Delta \)Aosnf1 mutant, whereas they were upregulated in the WT strain (Fig. 7D).

**DISCUSSION**

AMPK, also known as sucrose nonfermenting (SNF1) protein kinase, is a key cellular energy sensor (30). Previous studies showed that it is involved in the regulation of carbon source utilization, sporulation, pathogenicity, and stress response in some filamentous fungi and yeasts (9, 10, 12). In this study, we found that AMPK plays a conserved role in carbon source utilization, sporulation, and stress response, along with a crucial role in trap formation and lipid metabolism in NT fungus *A. oligospora*.

Compared with the WT strain, the mycelial growth of AMPK complex mutant strains (\( \Delta \)Aosnf1, \( \Delta \)Aogal83, and \( \Delta \)Aosnf4) cultured on PDA and TG was not affected, but it was significantly affected on CDA medium with different carbon sources, especially when a nonfermentable carbon source (glycerol) was used as the only carbon source. In addition, the aerial hyphae of all strains on CDA medium were very sparse. In *S. cerevisiae*, SNF1 was involved in meiosis and invasive growth (8). Deletion of the *snf1* and *snf4* genes resulted in the mutants being unable to grow on fermentable carbon source sucrose and nonfermentable carbon source glycerol or ethanol (30–32). Similarly, in *M. oryzae*, the mycelia of \( \Delta \)Mosnf1, \( \Delta \)Mosip2, and \( \Delta \)Mosnf4 mutants grew normally, whereas growth was inhibited when inoculated on the medium with nonfermentable carbon as the sole carbon source (10). In *A. alternata*, the hyphal growth of the \( \Delta \)Aasnf1 mutant was defective, and the colony diameter and mycelia radial growth were decreased on minimal medium supplemented with...
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glucose or glycerol (12). These findings showed that AMPK plays a varied role in the mycelial growth of *A. oligospora* and other fungi, with a key role in the utilization of carbon sources, especially in the use of nonfermentable carbon sources.

Deletion of *Aosnf1*, *Aagal83*, and *Aosnf4* resulted in a remarkable reduction in conidia yield compared with the WT strain, which was consistent with previous reports in filamentous fungi, including *Gibberella zeae* (33), *M. oryzae* (10), *Beauveria bassiana* (34, 35), *Pestalotiopsis microspora* (36), and *A. alternata* (12). Furthermore, the spore morphology of *A. oligospora*, *B. bassiana*, and *A. alternata* exhibited no change, whereas the spores of *G. zeae* became shorter, and spores of *M. oryzae* were abnormal. In addition, compared with the WT strain, the transcription levels of several sporulation-related genes were varied in the AMPK mutants; for example, the transcription of *abaA* was downregulated in three mutants during the middle stage of conidiation (fifth day), whereas the *medA* was upregulated in mutants (Δ*Aosnf1* and Δ*Aosnf4*). In particular, *abaA*, along with *brlA* and *wetA*, forms a central regulatory pathway governing sporulation in filamentous fungi and regulates the transcription of genes with important developmental functions (36). In *A. nidulans*, deletion of *abaA* and *medA* resulted in abnormal conidial morphology (37–39). In *F. graminearum*, *abaA* deletion mutants completely abolished sporulation (40). These results suggest that AMPK plays a conserved role in the spore production of diverse filamentous fungi.

Previous studies have shown that AMPK protein kinase plays an important role in the pathogenicity of different fungi. In *M. oryzae*, the disruption of *Mosnf1*, *Mosp2*, and *Mosnf4* genes reduced the virulence of fungi, especially Δ*Mosnf1*, which almost completely lost virulence (10). In *Metarhizium acridum*, the pathogenicity of Δ*Mosnf1* was decreased, which may be due to the reduction in appressorium formation and the decline in growth rate in insect hemolymph (41). In *Penicillium digitatum*, deletion of the *Pdsnfl* gene reduced the virulence of the mutant strain and the expression of many genes related to cell wall-degrading enzymes (42). The formation of traps is crucial for NT fungi to capture and kill nematodes (21). In this study, traps produced by the AMPK mutants were remarkably reduced, and more extracellular enzymes were detected in the Δ*Aosnf1* mutant than the WT strain. In addition, the nematode predatory efficiency was decreased in the Δ*Aagal83* and Δ*Aosnf4* mutants at 24 h and 36 h, whereas the nematocidal activity of the Δ*Aosnf1* mutant was decreased at 24 h, exhibiting no difference from the WT strain at 36 h. Previous studies have proven that extracellular serine protease is involved in the nematode predation of NT fungi by degrading the proteinaceous components of the nematode cuticle (43, 44). The increased proteolytic activity of Δ*Aosnf1* mutant may have resulted in the increase in predatory efficiency at 36 h. This was further confirmed by the variation of the transcription level of the genes encoding serine proteases in mutants compared with the WT strain. Thus, AMPK is involved in pathogenicity by regulating the development of infectious structures in *A. oligospora* and other pathogenic fungi, and *Aosnf1* may also participate in the regulation of production of serine proteases.

In yeast, oxidative stress resulted in the phosphorylation of Thr-210 and an increase in the catalytic activity of SNF1, and upstream kinases were required to phosphorylate Thr-210 to resist hypertonic stress (1 M NaCl) (45). Similarly in *Pestalotiopsis microspora*, the vegetative growth of the Δ*snf1* mutant was markedly inhibited under 1 M NaCl, 1 M sorbitol, 2 mM H$_2$O$_2$, and 0.02% Congo red compared to the controls (36). In *A. alternata*, the Δ*Aasnf1* mutant had reduced resistance to cell wall inhibitors SDS and Congo red, whereas there was no obvious difference in the growth rate between the WT and mutant strains supplemented with sorbitol, KCl, NaCl, and H$_2$O$_2$ (12). In this study, the Δ*Aosnf1* mutant strain had reduced resistance to oxidative stress (H$_2$O$_2$ and menadione) compared to the WT strain, and the Δ*Aagal83* and Δ*Aosnf4* mutants also had increased sensitivity to menadione. Accordingly, the downregulated transcription levels of most genes related to oxidative stress response were associated with their increased sensitivity to oxidative stress. In addition, the AMPK mutants had increased resistance to cell wall inhibitors (0.06 or 0.09 mg/mL Congo red), whereas the Δ*Aagal83* and Δ*Aosnf4* mutants showed increased
sensitivity to osmotic stress (NaCl and sorbitol). These results indicate that AMPK is involved in the stress response, but its role may vary in different fungi.

Transcriptome analysis of the ΔAosnf1 mutant strain showed that upregulated DEGs were remarkably enriched into the KEGG pathways of peroxisome and fatty acid degradation during vegetative growth (0 h) and trap formation (12 h). Fatty acid degradation in most organisms occurs primarily via the β-oxidation cycle; β-oxidation occurs in both mitochondria and peroxisomes in mammals, whereas only the peroxisomes harbor the β-oxidation cycle in plants and most fungi (46). In fungi, the peroxisome plays an important regulatory role in the metabolism of nonfermentable compounds such as glycerol and fatty acids (47). In this study, the transcription of four enzymes involved in fatty-acid β-oxidation, namely, carnitine O-acetylttransferase (AOL_s00215g212), long-chain acyl-CoA synthetase (AOL_s00215g200), enoyl-CoA hydratase (AOL_s00215g56), and 3-ketoacyl-CoA ketothiolase (AOL_s00210g122), was upregulated. Meanwhile, fatty acid oxygenase PpoC (AOL_s00215g31) was also upregulated in the ΔAosnf1 mutant. It has been reported that PpoC is involved in the growth and development of fungi. In Aspergillus fumigatus, the ΔpooC mutant produced fewer conidia, which were oval-shaped and larger, germinated faster in shaking cultures and slower in stationary cultures, and showed increased survival at both room temperature and 37°C, than the WT strain. In particular, the deletion of ppoC increased phagocytosis and killing by alveolar macrophages (48). Similarly, in A. nidulans, deletion of ppoC reduced conidia, increased ascospore production, and delayed conidiophore formation. PpoC was also shown to be involved in the production of lipo- genic signal molecule psI8 oxylipins (49). Recently, two peroxisome biogenesis genes, pex1 and pex6, were characterized in A. oligospora; deletion of gene pex1 or pex6 resulted in a failure to produce traps and conidia, as well as a reduction in nematode predation (50). In M. oryzae, deletion of the snf1 gene resulted in the absence of peroxisome structure, and mutant strains could not survive on medium with nonfermentable carbon sources (10). In fact, in the present study, the AMPK mutants showed a reduction in glycerol utilization capacity, LD accumulation, and conidial yield.

In addition, fatty acids can serve as the sole sources of carbon and energy; acetyl-CoA must be converted to C4 compounds via the glyoxylate bypass, comprising the enzymes isocitrate lyase and malate synthase, allowing gluconeogenesis (51). In this study, glyoxylate and dicarboxylate metabolism was downregulated in the ΔAosnf1 mutant at 0 h. Previous studies have proven that deletion of Aoal8 (encoding malate synthase) resulted in a reduction in conidiation and trap formation, as well as failure to utilize fatty acids and sodium acetate for growth; furthermore, its conidia were unable to germinate on medium supplemented with sodium oleate (52). In Candida albicans, deletion of Caicl1 (encoding isocitrate lyase) resulted in a failure to grow on the medium supplemented with citrate or glycerol as the sole carbon source, and the idc1 gene was required for the full virulence of this fungal pathogen (53). Furthermore, the glyoxylate cycle is essential for the pathogenicity of bacterial pathogens, such as Mycobacterium tuberculosis, Rhodococcus equi, and Pseudomonas aeruginosa (53). Here, deletion of Aosnf1, Aogal83, and Aosnf4 genes resulted in reduced glycerol utilization capacity and trap formation, as well as decreased pathogenicity at 24 h, consistent with the downregulated nitrogen metabolism, regulation of energy, and reduced glyoxylate and dicarboxylate metabolism. In addition, the downregulated terms of response to nutrient and oxidative stress were likely consistent with the mycelial growth and stress response of the mutants.

SNF1 plays a major role in lipid biosynthesis in S. cerevisiae, with deletion of snf1 increasing lipid production (54, 55). Similarly, in Yarrowia lipolytica, the Δsnf1 mutant accumulated more fatty acids than the WT strain (55). In Mortierella alpina, knockout of Masip2 contributed to fatty acid synthesis, whereas Masip2 overexpression increased lipid production, and knockout of Masip2 increased fatty acid unsaturation (56, 57). In our study, the LD volume of the ΔAosnf1, ΔAogal83, and ΔAosnf4 mutants was decreased compared with the WT strain, consistent with the expression of genes involved in lipid metabolism. In addition, more vacuoles and LDs were observed in the ΔAosnf1 mutant than the WT strain by transmission electron microscopy (TEM). Moreover, endocytic vesicles were observed in the WT strain but not in ΔAosnf1; consequently, the associated downregulated endocytosis.
pathway was enriched in ΔAosnf1. Taken together, our findings suggest that AMPK may play different roles in the lipid metabolism of different fungi, and it may affect the biosynthesis of cellular components.

To sum up, we constructed the ΔAosnf1, ΔAogal83, and ΔAosnf4 mutant strains for the first time, and our results showed that three subunits of AMPK play a similar role in phenotypic traits of A. oligospora. A combination of phenotypic and transcriptome analyses between the WT and mutant strains suggested that AMPK is involved in the peroxisome, lipid metabolism, glyoxylate and dicarboxylate metabolism, regulation of energy, nitrogen metabolism, response to nutrient, and response to oxidative stress, ultimately regulating the carbon source utilization, sporulation, trap formation, pathogenicity, and stress response in A. oligospora (Fig. 8). It is worth noting that the trap formation process requires energy (20), whereas AMPK can sense cellular energy and maintain cellular energy homeostasis (2). Deletion of Aosnf1, Aogal83, and Aosnf4 genes resulted in a significant reduction in the traps of A. oligospora. Therefore, further exploring the mechanisms underlying how AMPK influences trap formation and development through energy regulation will provide new insights into the lifestyle switching of NT fungi.

MATERIALS AND METHODS

Strains and culture conditions. The WT strain A. oligospora (ATCC 24927) and the mutant strains ΔAosnf1, ΔAogal83, and ΔAosnf4 were cultivated on PDA at 28°C. S. cerevisiae (FY834) was cultured in yeast extract-peptone-dextrose (YPD) medium and inoculated on synthetic complete medium lacking uracil (SC-Ura) for screening the recombinational clones (58). The Escherichia coli strain DH5α (TaKaRa, Shiga, Japan) was maintained on Luria-Bertani (LB) medium for storing plasmids pCSN44 and PRS426. PDA, TG, water agar (WA), and cornmeal-molasses-yeast (CMY) medium were used for various phenotypic analyses, and Czapek-Dox agar medium was used to compare the utilization of carbon sources between the WT and mutant strains (12). Caenorhabditis elegans (strain N2) was incubated at 26°C on oatmeal medium for trap induction of A. oligospora.

Sequence and phylogenetic analyses of AMPK complex in A. oligospora. The sequences of SNF1 (AOL_s00173g192), GAL83 (AOL_s00215g614), and SNF4 (AOL_s00004g493) were retrieved from A. oligospora using the homologous sequences of S. cerevisiae as a query. The NCBI database was used to search the homologous sequences of the A. oligospora AMPK subunits in different fungi. A neighbor-joining tree was constructed using MEGA 5.0 software (59). The InterProScan database was used to predict and analyze the conserved domains and functional sites (60).

Deletion of Aosnf1, Aogal83, and Aosnf4 genes. The Aosnf1, Aogal83, and Aosnf4 genes were deleted by homologous recombination as described previously (61). The 5′ and 3′ flanking sequences of the target genes and the hygromycin resistance gene cassette (hph) were amplified from A. oligospora and pCSN44 using primers (see Table S2 in the supplemental material), respectively. These three DNA fragments and linearized pRS426 vector were cotransformed into S. cerevisiae FY834 by use of electroporation. Then, the complete fragments for targeted gene disruption were individually transformed into A. oligospora protoplasts (62). For selection of positive transformants, PDA supplemented with 0.6 M...
sucrose (PDAS) plates supplemented with 200 mg \( \cdot \text{mL}^{-1} \) hygromycin were used, and PCR and Southern blot analyses were used to further substantiate them (Fig. S1).

**Comparison of mycelial growth, utilization of carbon sources, and conidiation.** The WT and mutant (\( \Delta \text{Aosnf1}, \Delta \text{Aogal81}, \) and \( \Delta \text{Aosnf4} \)) strains were inoculated on PDA and TG plates at 28°C for 6 days, respectively, and we recorded colony growth and measured colony diameter. By use of Czapek-Dox agar medium as the base medium, various carbon sources (glycerol, sucrose, and glucose) were added or not to test the carbon source utilization capacity of the WT and mutant strains, and the colony diameter was measured after 6 days of incubation (12). After 14 days of incubation on CMY, 20 mL of sterile double-distilled water (ddH2O) was added to elute and collect the spore suspension, followed by counting under a microscope (63). The spores were stained with 20 \( \mu \)g/mL calcofluor white (CFW; Sigma-Aldrich), and the morphology of the conidia was visualized under a light microscope (64).

**Trap development and pathogenicity assays.** The above-described spore suspension was spread on the WA plates (2 \( \times \) 10^4 conidia/plate) and cultured at 28°C for 3 to 4 days. *Caenorhabditis elegans* (cultured on oatmeal medium) was washed with sterile ddH2O, and then about 400 nematodes were added to the WA plates (64). At 12, 24, 36, and 48 h, the number of traps and captured nematodes were counted, and the percentages of captured nematodes of the WT and mutant strains were calculated.

**Extracellular protease activity analysis.** Skim milk was added to the PDA medium, and then the fungal strains were inoculated for 7 days at 180 rpm at 28°C. The fermentation broth was collected, and the extracellular enzyme activity was determined on the skim-milk plates as previously described (22).

**Stress response.** Using TG solid medium, the response of fungal strains to environmental stress was detected. The WT and mutant strains were inoculated in TG medium containing osmotic stressors (sorbitol and NaCl), cell wall-perturbing agents (Congo red and SDS), and oxidative stressors (H2O2 and menadione) at 28°C for 6 days. Different concentrations of chemical stressors were added as described in previous reports (26).

**Transcriptome and lipid staining analysis.** The WT and \( \Delta \text{Aosnf1} \) mutant strains were incubated onto PDA plates with cellophane on the surface at 28°C. After 3 days of incubation, equal amounts of nematodes were added for induction for 0 and 12 h, and mycelia were collected. Three independent biological replicates were used for each sample, and hyphal samples (12 samples) were sent to the Shanghai Meiji Biological Company (Shanghai, China) for RNA sequencing and analysis (65). GO and KEGG analyses were performed using the free online Majorbio Cloud Platform (http://www.majorbio.com).

Lipid droplets were stained with 10 mg/mL BODIPY (Sigma-Aldrich, St. Louis, MO, USA) for 30 min and observed under a confocal laser scanning microscope.

**RT-PCR analysis.** The WT and mutants were inoculated on CMY or PDA plates at 28°C, and mycelia were collected at different time points. The total RNA of mycelial samples was extracted and reverse transcribed to cDNA as previously described (64). Primer pairs for the target genes related to conidiation, oxidative stress response, serine proteases, peroxisome, glycerolipid metabolism, and endocytosis were designed (Table S3); the transcriptional levels of genes were detected by the LightCycler 480 SYBR green I master mix standard. The relative transcription level of each gene was calculated using the threshold cycle (2^-DDCt) method (66).

**Statistical analysis.** All experiments were biologically replicated three times, and the data were expressed as mean values \( \pm \) standard deviations (SD). Statistical analysis was performed using Prism 8 (GraphPad Software, San Diego, CA, USD), and Tukey's honestly significant difference (HSD) test was used to analyze the significance of the differences among treatments.

**Data availability.** All data generated or analyzed during this study are included in the published paper and the associated supplementary files. The raw sequence was deposited to the GEO under accession number GSE205188.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, PDF file, 2.7 MB.

**ACKNOWLEDGMENTS**

We are grateful to the Microbial Library of the Germplasm Bank of Wild Species from Southwest China for preserving and providing experimental strains and Guo Yingqi (Kunming Institute of Zoology, Chinese Academy of Sciences) for her help in taking and analyzing TEM images.

Funding for this study was provided by the National Natural Science Foundation of China (no. 31960556) and the Applied Basic Research Foundation of Yunnan Province (no. 202001BB050004).

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