The Multifaceted Gene 275 Embedded in the PKS-PTS Gene Cluster Was Involved in the Regulation of Arthrobotrisin Biosynthesis, TCA Cycle, and Septa Formation in Nematode-Trapping Fungus Arthrobotrys oligospora

Jiao Zhou 1, Qun-Fu Wu 1, Shu-Hong Li, Jun-Xian Yan, Li Wu, Qian-Yi Cheng, Zhi-Qiang He, Xu-Tong Yue, Ke-Qin Zhang, Long-Long Zhang and Xue-Mei Niu *

State Key Laboratory for Conservation and Utilization of Bio-Resources & Key Laboratory for Microbial Resources of the Ministry of Education, School of Life Sciences, Yunnan University, Kunming 650091, China * Correspondence: xmniu@ynu.edu.cn; Tel.: +86-871-65032538
† These authors contributed equally to this work.

Abstract: The predominant nematode-trapping fungus Arthrobotrys oligospora harbors a unique polyketide synthase-prenyltransferase (PKS-PTS) gene cluster AOL_s00215g responsible for the biosynthesis of sesquiterpenyl epoxy-cyclohexenoids (SECs) that are involved in the regulation of fungal growth, adhesive trap formation, antibacterial activity, and soil colonization. However, the function of one rare gene (AOL_s00215g275 (275)) embedded in the cluster has remained cryptic. Here, we constructed two mutants with the disruption of 275 and the overexpression of 275, respectively, and compared their fungal growth, morphology, resistance to chemical stress, nematicidal activity, transcriptomic and metabolic profiles, and infrastructures, together with binding affinity analysis. Both mutants displayed distinct differences in their TCA cycles, SEC biosynthesis, and endocytosis, combined with abnormal mitochondria, vacuoles, septa formation, and decreased nematicidal activity. Our results suggest that gene 275 might function as a separator and as an integrated gene with multiple potential functions related to three distinct genes encoding the retinoic acid induced-1, cortactin, and vacuolar iron transporter 1 proteins in this nematode-trapping fungus. Our unexpected findings provide insight into the intriguing organization and functions of a rare non-biosynthetic gene in a biosynthetic gene cluster.

Keywords: Arthrobotrys oligospora; AOL_s00215g275; sesquiterpenyl epoxy-cyclohexenoids (SECs); polyketide synthase-prenyltransferase (PKS-PTS) hybrid pathway; multifunction; retinoic acid induced-1 (Rai1) protein; cortactin and (Vit1) proteins; TCA cycle; mitochondria

1. Introduction

It is not unusual for there to be ‘other’ incongruous genes embedded in a biosynthetic gene cluster that is responsible for the biosynthesis of secondary metabolites. Previous studies have suggested that such ‘other’ genes appear to be unnecessary for the formation of secondary metabolites but display important roles in fungal self-protection mechanisms, including duplication of the metabolite target, detoxification of the metabolites, and efflux of the metabolites [1]. However, limited attention has been paid to the functions of such genes in new biosynthetic gene clusters.

Arthrobotrys oligospora is a typical nematode-trapping fungus that is widespread in diverse terrestrial and aquatic environments and is an ideal natural agent against the detrimental crop parasites known as nematodes [2–4]. Previous studies of A. oligospora have identified a number of unique polyketide–terpenoid hybrid metabolites—including oligosporon, arthrobotrisins, and arthrosporols—involved in the regulation of mycelial
growth and fusion, conidial development, adhesive trap formation, antibacterial and antiviral activities, and soil colonization [5–8]. This class of unique metabolites, which consist of a farnesyl-unit-derived sesquiterpenyl unit linked to a 6-methylsalicylic acid (6-MSA)-derived epoxy polyketide nucleus and belong to the sesquiterpenyl epoxy-cyclohexenoids (SECs), are widely distributed in fungi and include the well-known potent antifungal yanuthones from Aspergillus sp. and Penicillium sp. and the notorious phytotoxic macrophorins from the fungus Macrophoma, which causes fruit rot in apples [9–11]. Among these metabolites, anthrobotrisins A–D, have been regarded as characteristic chemotaxonomic markers for A. oligospora and could be used as chemical indicators for nematode-associated or infected environments [12,13].

A gene cluster $AOL_s00215g$ (215g) containing 10 genes (Figure 1A)—including a 6-MSA polyketide synthase (PKS) gene 283 and a prenyltransferase (PTS) gene 276—is responsible for this class of hybrid metabolites [14–16]. Combination of the construction of the mutants with the deletion of each gene in the 215g gene cluster, along with comparison of the metabolite profiles between the mutants and WT, revealed that genes 273–274 and 276–283 were involved in the biosynthesis of the hybrid metabolites, except for one gene (275) [15]. Moreover, the homologue of gene 275 was only missing in the Yan and Mac gene clusters that are responsible for the biosynthesis of SECs in Aspergillus sp. and Penicillium sp. (Figure 1A). Thus, two questions arise: why the gene 275 is embedded in the 215g cluster of the nematode-trapping fungus, and what kind of role the gene 275 plays in the fungus.

Figure 1. Organization of the genes in the 215g, Yan, and Mac gene clusters for the PKS-PTS hybrid metabolites (A), and phylogenetic analysis of 13 sequences with high coverage and similarity to 275 (B).

In the present study, we found that no conserved domain could be predicted in 275 using routine threshold values for bioinformatics analysis. The protein 275 only shared a limited identity to the retinoic acid induced-1 (Rai1) protein, src substrate cortactin, and vacuolar iron transporter 1 (Vit1) protein (Figure 1B). Next, we constructed two mutants:
one mutant Δ275 with disruption of the gene 275 in A. oligospora, and the other mutant OE-275 with the replacement of a strong promoter PtrpC upstream of the gene 275. Then, we compared the morphology, growth, trap formation, nematicidal activity, resistance to chemical stressors, and transcriptomic and metabolic profiles of the mutants and wild type (WT). The results revealed that the differentially expressed genes (DEGs) were highly enriched in the TCA cycle and diminished in membrane transport and ABC transporters. Interestingly, all of the genes in the 215g cluster responsible for the anthrobotrisin biosynthesis were dramatically upregulated in Δ275 vs. WT. Importantly, the genes involved in period circadian and actin-related proteins were the most downregulated. Metabolic profiles revealed that the TCA cycle and the hybrid metabolites were most dramatically regulated in the mutants. Moreover, Δ275 displayed swollen hyphae, while OE-275 had abnormal septa. A series of detailed analyses revealed that the Δ275 mutants displayed phenotypic and infrastructural traits due to the lack of three putative functions related to Rai1, cortactin, and Vit1. Our results suggest that 275 can perform multiple functions in fungal growth and development via regulating the SEC biosynthesis, TCA cycle, and septa formation.

2. Materials and Methods

2.1. Fungal Strain and Culture Conditions

A. oligospora YMF1.3170 was obtained from the State Key Laboratory for Conservation and Utilization of Bio-Resources and Key Laboratory for Microbial Resources of the Ministry of Education. PDA (potato 200 g/L, glucose 10 g/L, agar 15 g/L), TG (10 g/L tryptone, 10 g/L glucose, 15 g/L agar), TYGA (10 g/L tryptone, 5 g/L yeast extract, 10 g/L glucose, 5 g/L molasses, 15 g/L agar), and YMA (5 g/L yeast extract, 10 g/L malt extract, and 15 g/L agar) media were used for analyzing mycelial growth and related phenotypic traits. For the flask culture, the strain was cultured on PDA medium at 28 °C for 10 days to obtain conidia, and then the conidia were inoculated in 250 mL of liquid YPG medium (5 g of yeast extract, 5 g of peptone, and 20 g of glucose per liter) with a final concentration of 1 × 10⁶ conidia/mL at 28 °C at 180 rpm.

2.2. Sequence and Phylogenetic Analysis of 275 in A. oligospora

Predicted 275 in A. oligospora was annotated by BLAST searches against protein databases and InterProScan searches against protein domain databases. The amino acid sequence of 275 in A. oligospora was downloaded from GenBank, as were the sequences from different fungi. The amino acid sequences from different fungi were analyzed using the DNAman software package, version 5.2.2 (Lynnon Biosoft, Vaudreuil, QC, Canada). A neighbor-joining tree was constructed using the Mega 5.1 software package (PSU, USA). The web-based analysis platform antiSMASH 2.0 (https://fungismash.secondarymetabolites.org/; accessed on 16 April 22) was applied to perform genome mining of the biosynthetic gene clusters. The functional domains were predicted using the default parameters of InterProScan.

2.3. Construction of Mutant Δ275 and OE-275

A modified homologous recombination method was applied to construct the vectors for the mutant Δ275 and OE-275. All primers used in this study are provided in Tables S1 and S2. The DNA fragments (5′ flanks and 3′ flanks) were purified using the RiboEXTRACT Universal DNA Purification Kit (Tsingke, Beijing, China). The purified 5′ flank DNA fragment was inserted into the specific enzyme-digested PUC19-1300-D-HYB vector via the In-Fusion method to produce the PUC19-1300-D-HYB-5′ vector. Then, the 3′ flank DNA fragment was inserted into the specific enzyme-digested PUC19-1300-D-HYB-5′ vector to generate the PUC19-1300-D-HYB-5′–3′ vector. The homologous fragment was amplified and purified as described previously. The gene 275 was inserted into a PUC19 plasmid containing the strong promoter PtrpC and terminator TtrpC from Aspergillus
The plasmid was amplified as described previously. The purified homologous fragments cut from the plasmid were transformed into A. oligospora protoplasts. Transformation colonies were selected after incubation at 28 °C for 2–4 days, and every single colony was transferred to a new plate containing PDA. After incubation at 28 °C for 5 days, genomic DNA of putative transformants was extracted and verified by PCR to check for integration of the target in the genome. The mutant deficient in the target gene was screened out and confirmed by PCR and sequencing analysis.

2.4. Quantitative Real-Time PCR Analysis

The WT strain and OE-275 mutant were cultured in PDA medium at 28 °C for 6 days. The vegetative hyphae were harvested in 9-cm Petri dishes, which were then incubated at 28 °C for 3 and 5 d. After induction, hyphae were collected and frozen immediately in liquid nitrogen. Total RNA was extracted from all samples with the AxyPrep Multisource RNA Miniprep Kit (Axygen, Jiangsu, China). The extracted RNA was then reverse transcribed to cDNA with the FastQuant RT Kit with gDNase (Takara, Kusatsu, Japan). The cDNA was used as the template for a qRT-PCR assay, which was conducted to analyze the transcription levels of candidate genes. The gene-specific qRT-PCR primers were designed with Primer3 software, and the β-tubulin gene (Tub, AOL_s00076g640) was used as an internal standard. The qRT-PCR analysis was performed as previously described [16]. The relative transcription level (RTL) of each gene was calculated as the ratio of the transcription level in the deletion mutant to the transcription level in the WT strain at a given timepoint according to the $2^{-\Delta\Delta CT}$ method [16].

2.5. Ultrahigh-Performance Liquid Chromatography–Mass Spectrometry (UPLC–MS) Analysis

The WT and mutant strains were inoculated in flasks containing 250 mL of potato dextrose broth (PDB) and cultured at 28 °C (180 rpm) for 7 days. The fermentation broth was extracted with an equal volume of ethyl acetate, and the organic layer was evaporated and dissolved in 1 mL of menthol and filtered through a 0.22 μm filter membrane. All samples were then analyzed using a UPLC system coupled with a Q Exactive Focus Orbitrap mass spectrometer (Thermo Fisher, Waltham, MA, USA) and equipped with an Agilent Zorbax ODS 4.6 × 250 mm$^2$ column (Agilent, Santa Clara, CA, USA) using the electrospray ionization (ESI) mode. The metabolites extracted from the WT and mutant strains of A. oligospora were analyzed by UPLC–MS according to the methods described in the literature [15].

2.6. Fungal Colony Growth and Morphology

The WT and mutant strains were cultured on PDA, TYGA, and YMA media at 28 °C, and their growth rates and colony morphology were observed. Hyphal morphology was examined using 5-day-old PDA cultures grown at 28 °C. The fungal strains were stained with calcofluor-white (CFW, Sigma-Aldrich, St. Louis, MO, USA)—a fluorescent dye that naturally binds to cellulose and chitin. The hyphae were collected using a sterilized coverslip. The fungal strains were stained with 20 μg/mL CFW and then observed under an inverted fluorescence microscope (Nikon, Tokyo, Japan).

2.7. Fungal Conidial Production and Germination

The conidial yields of the fungal strains were determined as previously described [15,16]. Three replicates for each strain were initiated by spreading a 100 μL aliquot of a $10^5$ conidia mL$^{-1}$ suspension per CMY plate, followed by 10 days of incubation at 28 °C. Three colony plugs (5 mm in diameter) were taken from each plate culture, and the conidia on each plug were washed off into 1 mL of 0.02% Tween 80 through 10 min of vibration. The conidial concentration in the suspension was determined by microscopic counting in a hemocytometer and converted to the number of conidia per cm$^2$ of plate culture as an estimate of conidial yield. Moreover, conidial formation was observed with
an established side-shot approach. To calculate the conidial germination rates, 50 μL aliquots of 10^5 conidia mL^-1 suspensions of the WT strain and mutants were used to inoculate WA medium. The samples were incubated at 28 °C, and the germinated conidia were counted at 4 h intervals until 24 h post-inoculation. The phenotypic analyses were repeated three times.

2.8. Multi-Stress Assays

Chemical stressors including sorbitol, Congo red, sodium dodecyl sulfate (SDS), and H2O2 were used for the evaluation of fungal stress responses. The colonies of the WT and mutant strains were initiated with 9 cm diameter hyphal disks at 28 °C for 6 days on plates of TG or supplemented with each of the following chemical stressors: NaCl (0.1 mol/L, 0.2 mol/L, 0.3 mol/L) and sorbitol (0.25, 0.5, 0.75 mM) for osmotic stress; Congo red (0.1, 0.2, 0.3 mg/mL) and sodium dodecyl sulfate (SDS) (0.01, 0.02, 0.03%) for cell-wall-disrupting agents; and H2O2 (5, 10, 15 mmol/L) for oxidative stress. The diameter of each colony was measured and then calculated with GraphPad 9.3.0 (San Diego, CA, USA). The experiments were performed with at least three replicates.

2.9. Trap Formation and Nematicidal Activity Assays

WT and mutant strains were cultured on 9 cm agar plates at 28 °C for 3–4 days, and then about 400 nematodes (Caenorhabditis elegans) were introduced to the cultures. After 12 h, traps and captured nematodes per plate were observed under a microscope and counted at specific timepoints.

2.10. Endocytosis Assay

To evaluate endocytosis, hyphae of the WT and Δ275 strains were cultured on PDA plates for 4–5 days. The sterilized coverslips were inserted into the PDA medium at a slant (45° angle), and aerial hyphae were cultured on the coverslip for 5 days. The coverslip with aerial mycelia was incubated with FM4-64 staining solution (SynaptoRed C2, Bio-tium, Fremont, CA, USA; 10 μL of FM4-64 was diluted to a final concentration of 4 μM in 50 mM Tyrode solution) and immediately placed under a fluorescence microscope (Nikon, Tokyo, Japan) for observation [17].

2.11. Transmission Electron Microscopy (TEM)

For TEM analysis, ultrathin sections of hyphal cells were prepared and examined as previously described [16]. The samples of the transmission electron microscope (TEM) were pretreated with 4% paraformaldehyde and then incubated with 2.5% glutaraldehyde (Sigma) in phosphate buffer (pH = 7.4) overnight at 4 °C and post-fixed in 1% OsO4. The samples were dehydrated in a gradient ethanol series, embedded in Spurr resin, and stained with 2% uranyl acetate and Reynold’s lead solution. The samples were examined under an H–7650 transmission electron microscope (Hitachi, Tokyo, Japan).

2.12. Quantitative Analysis of Arthrobotrisins and TCA Metabolites and Amino Acids

Twenty standard amino acids and seven standard compounds of the tricarboxylic acid (TCA) cycle were dissolved in methanol at concentrations of 10 μM. The relative contents of metabolites of TCA and amino acids were calculated by evaluating the peak areas according to standard substances. The metabolites were identified through retention time indices and mass spectra of peaks. Compounds were designated as target metabolites if they were identified with a match of 950 on a scale of 0–1000 with a retention index deviation of 3.0. The semi-quantitative analysis of the main compounds was carried out by internal normalization with the area of each compound.
2.13. Transcriptomic Analysis

The WT and mutant strains were cultured in PDB medium (potato 200 g/L, glucose 10 g/L) at 28 °C for 7 days. Sequencing of mycelial samples was performed by Shanghai Majorbio Bio-pharm Technology Co., Ltd. (Shanghai, China), and the data were analyzed using the Majorbio Cloud Platform (www.majorbio.com (accessed on 10 March 2022)). The transcripts per kilobase million method (TPM) was used to calculate the expression level of each transcript to identify differentially expressed genes (DEGs). Functional enrichment analysis identified the DEGs that were significantly enriched in the GO and KEGG terms ($p \leq 0.05$) through GO and KEGG analysis.

2.14. Protein Modeling and Docking

Protein structural modeling of 275 was performed using AlphaFold v2.0 with Colab-Fold v1.0 (https://deepmind.com/research/open-source; accessed on 24 July 2022) [18,19]. Minimized energy optimization of structures of the precursors, intermediates, and their derivatives in the biosynthetic pathway for anthrobotrinsins in *A. oligospora* was processed using ChemBio3D Draw (version 20.0.0.41, Cambridge Soft, Cambridge, MA, USA). The processing of the 275 protein (hydrogenation, and Gasteiger charge for merging nonpolar hydrogen atoms) was performed using MGLTools (v1.5.6) in AutoDock Vina software (https://autodock.scripps.edu/download-autodock4/; accessed on 24 August 22) [19]. The original pdb file format was converted to the pdbqt file format, which can be recognized by the AutoDock Vina program. Furthermore, the docking active center (including all residues around the original ligand) was set through the grid box function in the software, based on the size of the root-mean-square deviation (RMSD) of the docked ligand molecule and the original ligand molecule with reasonable docking parameter settings. RMSD ≤ 4 was set for the threshold for conformation of the ligand after docking to match the conformation of the original ligand. The best docking mode was selected as the lowest obtained binding energy result.

2.15. Statistical Analysis

All experiments were repeated at least three times. Data are presented as the mean ± standard deviation (SD). GraphPad Prism v9.0.2 software (San Diego, CA, USA) was used for statistical analysis. Comparisons were performed using the two-tailed Student’s *t*-test. Probabilities of $p < 0.05$ were used as the threshold for determining significant differences (* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$).

3. Results and discussion

3.1. Sequence and Phylogenetic Analyses of 275

The gene 275 consists of only one intron and encodes a protein of 275 amino acids with an isoelectric point of 5.3 and a molecular mass of 30.7 kDa. Surprisingly, no conserved domains were identified in 275 using InterProScan and the Conserved Domain Database (CDD). Importantly, only one unknown protein was retrieved from fungi when searching against UniProt (Parameter: E-Threshold < 0.1) with 275. The protein from *Anthrobotrys flagrans* displayed a high similarity to 275 (76% identity). Other retrieved sequences were mainly from mammals, fish, insects, and flies. All of the proteins displayed a low similarity to 275, ranging from 23% to 29%. To illustrate the homology of 13 retrieved sequences with-known protein functions to the 275 protein, a maximum-likelihood (ML) phylogenetic tree under the best-fit amino acid substitution model with 200 bootstrap replications was constructed (Figure 1B). The E value was adjusted to 1 in order to include those protein sequences with higher similarity to 275. The phylogenetic tree revealed that 275 fell into a clade containing retinoic acid induced-1 (Rai1) protein, src substrate cortactin, and vacuolar iron transporter 1 (Vit1) protein.

Analysis of the subcellular localization of the 275 protein with Cell-PLoc indicated that 275 might be located in the nucleus (12), mitochondria (10), cytoplasm (4), and...
cytoskeleton (1). Interestingly, Rai1 was localized to the nucleus, cortactin was localized to the cytoplasm, and Vit1 was localized to the plasma membrane. Moreover, we found that the Rai1 protein covers the first half of 275, while cortactin covers most of the latter half of 275 (Figure 1B), and Vit1 covers almost 80% of the 275 protein. Previous studies suggest that Rai1 is involved in the circadian regulation of gene expression [20,21]. Cortactin acts as an actin-binding protein by binding and activating Arp2/3, playing critical roles in the regulation of the cytoskeleton, cell migration, and endocytosis [22–24]. Vit1 in the plant *Eucalyptus grandis* is a H+ -coupled antiporter for metal ions, which provides the vacuolar H+-ATPase with an important function in metal storage [25,26].

3.2. Function of 275 in Regulating Fungal Growth and Morphology

The mutants Δ275 with the disruption of 275 and OE-275 with the replacement of the strong promoter *PtrpC* were constructed and identified via PCR analyses (Figure 2A,B). qRT-PCR analysis revealed that the transcription level of 275 in OE-275 was 2.68 times and 3.97 times higher than that in the WT at 3 and 5 days, respectively, suggesting that OE-275 displayed a significantly increased transcription level of 275 compared with the wild-type strain (WT) (Figure 2C). Three media—including one nutrient-deficient medium (PDA, which is usually used for trap formation) and two nutrient-rich media (YMA and TYGA)—were evaluated for fungal colony growth and morphology. Compared with the WT, both Δ275 and OE-275 displayed significantly decreased fungal growth on the nutrient-rich media (YMA and TYGA), while no significant difference in fungal growth was observed on the nutrient-deficient medium (PDA) (Figure 2D). Interestingly, at day 6 on all three media, both Δ275 and OE-275 displayed distinct fungal colonies with different morphology from that of WT (Figure 2D). Obviously, the WT displayed much denser and thicker aerial mycelia than both mutants. Moreover, we found that both Δ275 and OE-275 displayed distinct mycelia zones compared to the WT (Figure 2D). These results suggest that the disruption of gene 275 or the enhanced transcription level of 275 could lead to significantly varied fungal colony growth, aerial mycelia, and mycelia zones.
Figure 2. Confirmation of the mutant Δ275 with the disruption of 275 and the mutant OE-275 with the strong promoter replaced, and comparison of the colony growth of these two mutants with the WT on three media: (A,B) PCR analysis confirmed the mutants Δ275 (A) and OE-275 (B). (C) The relative transcriptional level (RTL) of 275 in the mutant OE-275 at 3 and 5 days compared to that in the WT. (D) Comparison of the colony growth of these two mutants with the WT on three media (PDA, TYGA, and YMA).

3.3. Roles of 275 in Conidial Production, Germination, Trap Formation, and Nematicidal Activity

The mutants Δ275 and OE-275 displayed quite opposite conidial production and conidial germination compared with the WT at 28 °C (Figure 3A). The mutant Δ275 showed a 204.4% increase in conidial formation and a 7.6% increased germination rate, while OE-275 displayed significantly decreases in conidial formation and germination rate (by 33.1% and 11.3%, respectively) (Figure 3A,B). Similarly, Δ275 and OE-275 also displayed
quite opposite trap formations compared with the WT. The mutant Δ275 showed slightly decreased trap formation, while OE-275 displayed significantly enhanced trap formation (by 98.4% and 65.0%, respectively), compared with the WT at 24 h and 36 h (Figure 3C,D). The mutant Δ275 exhibited significantly decreased nematocidal activity (by 18.5%) compared with the WT at 24 h (Figure 3D,E). Surprisingly, the mutant OE-275 also exhibited significantly decreased nematocidal activity (by 29.6%) compared with the WT at 24 h (Figure 3E). All of the results suggested that the gene 275 was involved in inhibiting conidial formation, germination, and nematocidal activity but promoting trap production. We found that the decreased trap formation in Δ275 was in stark contrast to the dramatically increased trap formation in the other mutants that were deficient in each biosynthetic gene in the gene cluster 215g [15].

Figure 3. The effects of 275 on conidial production, germination, trap formation, and nematocidal activity: (A–D) Quantitative analysis of conidial production (A), germination rate (B), trap formation (C), and nematocidal activity (D), comparing the mutants Δ275 and OE-275 with the WT. (E) Comparison of trap formation nematocidal activity between the mutants Δ275 and OE-275 and the WT at 24 and 36 h. The arrows indicate the traps.
3.4. The Association of 275 with Multiple Stress Responses

Surprisingly, we found that the mutant Δ275 showed significantly increased resistances to several chemical stressors—including two cell-wall-perturbing agents (SDS and Congo red), two osmotic agents (NaCl and sorbitol), and the oxidant H₂O₂—on the TYGA medium (Figure 4A,B). Δ275 displayed increased colony growth when exposed to Congo red, NaCl, and sorbitol at all of the tested concentrations. Moreover, Δ275 also exhibited distinct aerial mycelia on 15 mM H₂O₂ compared to the WT. Notably, OE-275 also displayed distinct aerial morphology on SDS and enhanced growth on NaCl, sorbitol, and 10–15 mM H₂O₂ (Figure 4A,B). Interestingly, we also found that the inner-ring mycelia of OE-275 became much thinner than those of the WT on Congo red, NaCl, and H₂O₂ (Figure 4A). These results suggest that the disruption of 275 or the enhanced transcription of 275 could exert intensive influences on the fungal resistances and significantly regulate the fungal morphology and responses to all of the tested chemical stressors.

![Figure 4](image_url)

Figure 4. The effects of 275 on the fungal resistances to chemical stressors on TG plates: (A, B) Comparison of the colony growth between the two mutants (Δ275 and OE-275) and the WT under treatment with Congo red, SDS, NaCl, sorbitol, and the oxidant H₂O₂ (*: p < 0.05; **: p < 0.01; ***: p < 0.001).

3.5. Transcriptional Link of 275 to the PKS-PTS Hybrid Pathway, TCA Cycle, Membrane Transport, and Redox Homeostasis

Transcriptomic analysis displayed that there were significant differences between Δ275 and OE-275 and the WT (Figure 5A). There were 227 genes expressed only in the WT, 136 genes only in Δ275, and 114 genes only in OE-275. A total of 9502 genes were co-expressed between the WT, Δ275, and OE-275 (Figure 5B). Compared with the WT, 943 and 902 genes were differentially expressed in Δ275 and OE-275, respectively. Among them, 447 genes were significantly upregulated and 496 genes were significantly downregulated in Δ275 (Figure 5C). In OE-275, 441 genes were significantly upregulated and 461 genes were significantly downregulated (Figure 5D).
Figure 5. Transcriptomic analysis of Δ275, OE-275, and the WT: (A) Plot analysis of Pearson’s correlation (r) among the mutants Δ275 and OE-275 and the WT. (B) Venn analysis for the numbers of significantly differentially regulated genes in Δ275 and OE-275, compared to the WT. (C,D) Volcano
plot analysis of upregulated and downregulated genes in Δ275 and OE-275 vs. the WT, respectively. (E) Comparison of the transcriptional levels of the genes in the PKS-PTS hybrid pathway in Δ275 and OE-275, compared to the WT. (F) Comparison of the transcriptional levels of the key downregulated genes in Δ275 and OE-275 compared to the WT. (G,H) KEGG analysis categorizing all of the significantly upregulated and downregulated genes in OE-275 compared to the WT (relative abundance, log2 scale, p < 0.05). (I,J) KEGG analysis categorizing all of the significantly upregulated and downregulated genes in OE-275 compared to the WT (relative abundance, log2 scale, p < 0.05).

Interestingly, the gene 276, which is immediately adjacent to 275 and encodes a PTS for biosynthesis of the first hybrid precursor, ranked the first among the upregulated genes in Δ275 vs. the WT due to having the lowest Padj value. Moreover, among the top 20 upregulated genes in Δ275 vs. the WT, the gene 276 also ranked the first due to having by far the highest FPKM value (19.6 in the WT vs. 422.7 in Δ275) (Figure 5F, Table S3). Further analysis of the transcriptional levels of all of the genes involved in the PKS-PTS hybrid pathway (273–283) indicated that all of the genes were dramatically upregulated in Δ275. Among them, two genes—279 and 277, which were both involved in the formation of the key epoxy ring in the first PKS-PTS hybrid precursor—displayed similar increases in their transcriptional levels to that of gene 276 (Figure 5E).

Among the top 20 downregulated genes, 79g256, which encodes the period circadian protein, displayed by far the highest FPKM value (1101.44 in the WT vs. 29.41 in Δ275), with a 36.4-fold increase (Figure 5F, Table S4). Moreover, three genes—including 76g531, encoding a putative L-fucose-proton symporter; 4g471, encoding neuromodulin; and 6g248, encoding actin-related protein 2/3 complex subunit 5 (Arp2/3)—were also in the top five FPKM values among these downregulated genes in Δ275 vs. the WT (Figure 5F). Previous studies suggest that neuromodulin is among the most prominent substrates of protein kinase C (PKC) in the brain and that phosphorylation of neuromodulin by PKC functions to allow maximal Ca\(^2+\)/CaM binding and vesicle release in mammals [27,28]. Interestingly, previous studies indicate that Arp2/3 is a protein that can be recruited by cortactin to exist in actin microfilaments, facilitating and stabilizing nucleation sites for actin branching, thereby playing an important role in promoting cell migration and endocytosis [25,26].

KEGG enrichment analysis of all of the upregulated genes in Δ275 vs. the WT indicated that the TCA cycle was the most upregulated in Δ275 (Figure 5G), while arginine biosynthesis, pyruvate metabolism, nitrogen metabolism, and alanine/aspartate/glutamate metabolism ranked the 2nd to the 5th, respectively. KEGG enrichment analysis of all of the downregulated genes in Δ275 vs. the WT exhibited that membrane transport and ABC transporters were the most downregulated in Δ275 (Figure 5H). KEGG analysis of the differentially expressed genes in OE-275 vs. the WT (Tables S5 and S6) indicated that the most upregulated pathways included glycolysis/gluconeogenesis, phenylalanine/tyrosine/tryptophan biosynthesis, pyruvate metabolism, alanine/aspartate/glutamate metabolism, and the TCA cycle (Figure 5I), while the most downregulated pathways were membrane transport, ABC transporters, fatty acid degradation, glycosylphosphatidylinositol (GPI)-anchored proteins, and SNARE interactions in vesicular transport (Figure 5J). A previous study suggested that, in mammalian cells, GPI-anchored proteins are concentrated in lipid rafts that are involved in receptor-mediated signal transduction pathways and membrane trafficking [29].

Further GO analysis of all of the upregulated genes in Δ275 vs. the WT indicated that oxidoreductase, amide transmembrane transporter, and nucleoside monophosphate kinase activities were the most upregulated in Δ275 (Figure 6A). GO analysis of all of the downregulated genes in Δ275 vs. the WT showed that ABC-type transporter activity was the most downregulated in Δ275 (Figure 6B). GO analysis of the differentially expressed genes in OE-275 vs. the WT showed that the amide transmembrane transporter was the most upregulated (Figure 6C) while cysteine-type endopeptidase activity was the most downregulated in OE-275 (Figure 6D). A previous study suggested that cysteine-type endopeptidase activity was mainly involved in antioxidant activity [30]. Vacuolar
processing enzyme (VPE) is a cysteine-type endopeptidase that mediates programmed cell death (PCD) by provoking vacuolar rupture and initiating the proteolytic cascade leading to PCD [30].

![Figure 6](image)

**Figure 6.** GO analysis of the significantly differentially regulated genes in Δ275 and OE-275, compared to the WT: (A,B) GO analysis of the significantly upregulated genes (A) and downregulated genes (B) in Δ275 vs. the WT. (C,D) GO analysis of the significantly upregulated genes (C) and downregulated genes (D) in OE-275 vs. the WT.

3.6. 275 Regulated the Production of Unique PKS-PTS Hybrid Metabolites, TCA Metabolites, and Amino Acids

UPLC-MS analysis of the metabolic profiles of Δ275, OE-275, and the WT was performed. Both Δ275 and OE-275 displayed quite similar metabolic profiles to the WT, except that Δ275 had two more peaks at 13 and 28 min (Figure 7A). Detailed analysis of the hybrid arthrobotrisins showed that Δ275 had 2.1-fold higher arthrosporol levels than the WT, and there was no significant difference in OE-275 (Figure 7B). Furthermore, citrate dramatically increased by 911.4% in Δ275 vs. the WT, while malate decreased by 62.9%. Small significant increases were observed for pyruvate (35.1%) and succinate (7.6%) in Δ275 vs. the WT (Figure 7B). Surprisingly, cis-aconitate, D-isocitrate, and α-ketoglutarate
could be not detected in Δ275. Furthermore, all of the above TCA metabolites—except citrate—could not be detected in OE-275 either (Figure 7C). However, OE-275 displayed quite similar composition of aromatic acids to the WT, except that OE-275 had 438.0% more serine, 67.2% more alanine, and 215.9% more arginine but 70.8% less glutamine and 34.7% less glutamic acid than the WT (Figure 7D). The mutant Δ275 displayed large increases in serine (401.2%), histine (103.5%), alanine (204.6%), and arginine (1127.4%), but significant decreases in threonine (67.8%) and glutamine (56.7%), compared with the WT (Figure 7D).

**Figure 7.** The effects of the gene 275 on the arthrobotrisin production, TCA cycle, and major amino acids: (A) HPLC profiles of the extracts of the liquid cultures of Δ275, OE-275, and the WT. The arrows indicate the extra peaks in Δ275. (B–D) Comparison of the levels of arthrobotrisin (B), major metabolites from the TCA cycle (C), and major amino acids (D) among Δ275, OE-275, and the WT.
3.7. Vital Role of 275 in Endocytosis, Septum Formation, and Mitochondrial Homeostasis

Considering the increased responses of Δ275 to osmotic NaCl and sorbitol, as well as cell-wall-disturbing Congo red and SDS, combined with the most downregulated membrane transport and ABC transporters, endocytosis analysis was performed (Figure 8A). Hypha of both Δ275 and the WT were stained with an endocytic tracer—the lipophilic styryl dye FM4-64, which was gradually internalized and stained endosomes and vacuoles after incubation. Δ275 clearly displayed distinct endocytosis from the WT. The WT exhibited local discontinuities of strong endocytosis, while Δ275 did not, suggesting that the lack of 275 resulted in dramatically decreased endocytosis. Interestingly, Δ275 displayed swollen hyphae and OE-275 had abnormal septa compared with the WT (Figure 8B). In particular, further septum dye assay suggested that septa with larger hollows were observed in the conidia of Δ275 and OE-275 compared to those in the WT, while the conidia of both Δ275 and OE-275 displayed thinner septa than the WT (Figure 8C). Transmission electron microscopy (TEM) analysis displayed that the numbers of mitochondria and vacuoles in Δ275 were dramatically higher than those in the WT (Figure 8D). Moreover, mitochondrial hypertrophy, hyperplasia, and vacuolization were also observed in Δ275 (Figure 8D). In addition, Δ275 displayed amplified rough endoplasmic reticulum, concentric lamellar bodies, and lacked large vacuoles compared to the WT. Importantly, the WT strain consisted of many black tight bodies, whereas Δ275 had few (Figure 8D). In order to evaluate the effects of varied vacuoles on the iron levels in the fungus, the levels of Fe\(^{2+}\), Fe\(^{3+}\), and total iron were also evaluated. However, no significant differences in the levels of Fe\(^{2+}\), Fe\(^{3+}\), and total iron were observed in Δ275 compared to the WT. Interestingly, OE-275 displayed increased levels of Fe\(^{2+}\), Fe\(^{3+}\), and total iron compared to the WT.
Figure 8. The effect of 275 on the fungal endocytosis, septum formation, infrastructure, and iron levels: (A) Comparison of the fungal endocytosis between Δ275 and the WT within 120 s using the lipophilic styryl dye FM-464. (B) Comparison of the fungal mycelial morphology and septum formation between the mutants Δ275 and OE-275 and the WT using CFW staining. (C) Comparison of the conidial septum formation between the mutants Δ275 and OE-275 and the WT. White arrow: septum. (D) TEM analysis of the infrastructures between the mutant Δ275 and the WT. Yellow arrow: vacuole; Blue arrow: mitochondria; red arrow: black particles. (E) Comparison of the Fe$^{2+}$, Fe$^{3+}$, and total iron levels between the mutants Δ275 and OE-275 and the WT.
3.8. Docking of 275 with the PKS-PTS Hybrid Metabolites

To investigate the possible binding ability of the precursors, intermediates, and their derivatives from the PKS-PTS hybrid biosynthetic pathway in *A. oligospora* with the 275 protein, molecular docking analysis was performed. Interestingly, all of the 14 tested compounds displayed binding free energy values ranging from −3.0 to −6.75 kcal/mol (Figure 9). A previous study indicated that binding free energy less than −2 kcal/mol denotes good binding activity between ligands and receptors [31]. It is worth noting that the addition of the farnesyl moiety to toluquinol greatly improved the binding affinity to 275. Moreover, the PKS-PTS hybrid metabolites with the epoxy ring displayed the highest binding affinity (Figures 9 and 10). Molecular visualization showed that the formation of hydrogen bonds between compounds and amino acid residues within the 275 protein can strengthen the interactions (Figure 10).

![Figure 9. The hybrid metabolites derived from the PKS-PTS hybrid biosynthetic pathway with binding free energy values (referring to the binding affinity of SEC metabolites with 275) predicted by molecular docking analysis. All of the metabolites displayed binding free energy values below −3 kcal/mol.](image-url)
4. Discussion

Here, we found that the rare gene 275, embedded in between the two biosynthetic genes 276 and 274 in the gene cluster 215g (Figure 1A), was not involved in the biosynthesis of SEC metabolites in the nematode-trapping fungus. Notably, the gene 276 was responsible for the biosynthesis of the first hybrid precursor farnesyl–toluquinol by transferring a farnesyl unit to toluquinol (Figure 9). The gene 274 is a dehydrogenase that is involved in the oxidation modification of SECs [14]. A previous study suggested that toluquinol could be easily dehydrogenated to a more active toluquinone for the methylation, along with a pair of protons and electrons [14]. Interestingly, we also noted that the PKS-PTS hybrid gene clusters Yan in Aspergillus sp. and Mac in Penicillium sp., without the analogues of gene 275, displayed distinct gene organizations from 215g in A. oligospora. Interestingly, the analogues of gene 276 in the Yan and Mac clusters were just between the PKS gene and the decarboxylase gene that were involved in the biosynthesis of the first and second precursors—6-MSA and m-cresol, respectively—for the SECs. Importantly, all of the oxidoreductase genes including the analogues of gene 274 were arranged on the other side of the PKS gene, suggesting that gene 276 should be apart from 274 or other oxidoreductase genes, and that gene 275 might function to separate gene 276 from gene 274 in the 215g cluster. Solid evidence from the varied resistances to H₂O₂ and GO enrichment analysis indicated that oxidoreductase activity and antioxidant cysteine-type
endopeptidase activity were most regulated in Δ275 and OE-275, respectively. In addition, the gene 276 was the most abundant upregulated gene in Δ275. In fact, all of the genes involved in the PKS-PTS hybrid pathway (273–283) were dramatically upregulated in Δ275. In particular, two oxidation genes—279 and 277, which were both involved in the formation of the key epoxy ring in the first PKS-PTS hybrid precursor—were the most upregulated. The metabolic profiles confirmed the dramatically increased SEC contents. These results suggested that the gene 275 played a key role in regulating the biosynthesis of SEC metabolites.

Bioinformatics analysis indicated that no homologues with high identity to 275 could be found in the known proteins. Detailed phylogenetic trees suggested that 275 shared a very low identity to the Rai1, cortactin, and Vit1 proteins. From the chemical structural point of view, arthrobotrosins shared much similarity to retinol (ligands 1–3 in Figures 9 and 10). Molecular docking analysis suggested strong binding affinity between the epoxy PKS-PTS hybrid metabolites with protein 275, suggesting that 275 might be a Rai1-like protein. A previous study suggested that Rai1 is a transcription factor associated with Smith–Magenis syndrome, where deletion of Rai1 disrupted light-activated daytime melatonin suppression, with consequently high diurnal levels and low nocturnal levels of melatonin, and causing abnormal circadian signaling with genetic disorders [32]. Interestingly, the period circadian protein was by far the most abundant downregulated gene in the fungus with the disruption of 275. A recent study suggested that calcium signaling is pivotal to the circadian clockwork in the suprachiasmatic nucleus—particularly in rhythmic entrainment to environmental light–dark cycles [21]. We also found that the gene encoding neuromodulin—which is reportedly involved in Ca2+/CaM binding—was also among the top five downregulated genes in Δ275 vs. the WT. All of these results suggest that 275 might be a Rai1-like gene.

Interestingly, transcriptional analysis also revealed that the key gene encoding Arp2/3, which was recruited by cortactin to play an important role in the cytoskeleton and endocytosis, was also the most downregulated. Further analysis revealed that the fungal mycelia, the septa of the fungal conidia, and endocytosis became abnormal in both mutants (Δ275 and OE-275), together with distinct aerial morphology on the cell-wall-disturbing agent SDS, consistent with the downregulated gene Arp2/3. All of these results suggest that 275 might also be a cortactin-like gene.

A previous study suggested that Vit1 is a H+-coupled antiporter for metal ions, which provides the vacuolar H+-ATPase with an important function in metal storage [25]. A recent study reported that loss of vacuolar acidification impaired vacuolar cysteine storage, causing oxidative damage to mitochondrial iron–sulfur cluster biosynthesis components and activities, such as aconitase [33]. As a consequence, cells developed an age-related mitochondrial dysfunction characterized by the activation of the iron-deficiency and DNA-damage responses. Notably, a putative L-fucose-proton symporter gene was most downregulated in Δ275 vs. the WT. Moreover, the mutants displayed distinct responses to NaCl and sorbitol, together with fast aging of the aerial mycelia of OE-275 compared to the WT. Importantly, exceptional TCA cycles, mitochondrial dysfunctions, and unusual vacuoles were also observed in Δ275; for example, cis-aconitate, D-isocitrate, and α-ketoglutarate could not be detected in Δ275. These results indicate that 275 might be a Vit1-like gene.

In conclusion, our findings reveal that the gene 275 might function as a separator and as an integrated gene that might perform multiple functions related to three distinct proteins—Rail, cortactin, and Vit1—in the nematode-trapping fungus (Figure 11). Our findings provide insight into the intriguing organization of a rare gene that is likely derived from three genes, which might help to develop effective biological agents for combating nematode pests.
Figure 11. The proposed functions of 275 in the nematode-trapping fungus *A. oligospora*.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/jof8121261/s1. Table S1: The list of the primers for Δ275; Table S2: The list of the primers for OE-275; Table S3: The upregulated genes in Δ275 vs. the WT; Table S4: The downregulated genes in Δ275 vs. the WT; Table S5: The upregulated genes in OE-275 vs. the WT; Table S6: The downregulated genes in OE-275 vs. the WT.


**Funding:** This research was funded by NSFC grants 21977086, 82260691, and 21867018, Program 202201BF070001-012 of “Double tops” from Yunnan Province, and Yunnan University and the Yunnan Innovative Research Team for Discovery and Biosynthesis of Bioactive Natural Products (2018HC012).

**Institutional Review Board Statement:** Not applicable.

**Informed Consent Statement:** Not applicable.

**Acknowledgments:** We would like to thank the Institutional Center for Shared Technologies and Facilities of Kunming Institute of Zoology, Chinese Academy of Sciences, for providing us with transmission electron microscopy. We are grateful to Guo Yingqi for her great help in preparing TEM samples and taking/analyzing TEM images.

**Conflicts of Interest:** The authors declare no conflict of interest.
References

27. Selmon, J.R.; Martzen, M.R. Neurmodulin (GAP-43) can regulate a calmodulin-dependent target in vitro. Biochemistry 1994,