Genome sequence and transcriptome profiles of pathogenic fungus *Paecilomyces penicillatus* reveal its interactions with edible fungus *Morchella importuna*

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**Abstract**

*Paecilomyces penicillatus* is one of the pathogens of morels, which greatly affects the yield and quality of *Morchella* spp. In the present study, we *de novo* assembled the genome sequence of the fungus *P. penicillatus* SAAS_ppe1. We analyzed the transcriptional profile of *P. penicillatus* SAAS_ppe1 infection of *Morchella importuna* at different stages (3 days and 6 days after infection) and the response of *M. importuna* using the transcriptome. The assembled genome sequence of *P. penicillatus* SAAS_ppe1 was 39.78 Mb in length (11 scaffolds; scaffold N50, 6.50 Mb), in which 99.7% of the expected genes were detected. A total of 7.48% and 19.83% clean transcriptional reads from the infected sites were mapped to the *P. penicillatus* genome at the early and late stages of infection, respectively. There were 3,943 genes differently expressed in *P. penicillatus* at different stages of infection, of which 24 genes had increased expression with the infection and infection stage, including diphthamide biosynthesis, aldehyde reductase, and NAD (P)H-hydrate epimerase (*P* < 0.05). Several genes had variable expression trends at different stages of infection, indicating *P. penicillatus* had diverse regulation patterns to infect *M. importuna*. GO function, involving cellular components, and KEGG pathways, involving glycerolipid metabolism, and plant-pathogen interaction were significantly enriched during infection by *P. penicillatus*. The expression of ten genes in *M. importuna* increased during the infection and infection stage, and these may regulate the response of *M. importuna* to *P. penicillatus* infection. This is the first comprehensive study on *P. penicillatus* infection mechanism and *M. importuna* response mechanism, which will lay a foundation for understanding the fungus-fungus interactions, gene functions, and variety breeding of pathogenic and edible fungi.

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**1. Introduction**

The genus *Paecilomyces* was established by Bainier (1907) and was initially considered to be similar to *Penicillium* in morphology and physiology. Luangsa-Ard et al. [1] demonstrated that *Paecilomyces* was polyphyletic across two subclasses (Sordariomycetidae and Eurotiomycetidae) based on phylogenetic analysis of the 18S rRNA sequence. Inglis and Tignano [2] supported the polyphyly of the genus through phylogenetic analysis of the 5.8S rRNA and internal transcribed spacer (ITS1 and ITS2) sequences from selected *Paecilomyces* species. The diverse species of *Paecilomyces* inhabit a wide range of environments. Some *Paecilomyces* species are human pathogens that can cause onychomycosis [3], pneumonia [4], and rhinosinusitis [5]. Other *Paecilomyces* species cause diseases of apple [6], pistachio [7], and other trees [8]. *P. hepiali* is an endoparasitic fungus of *Cordyceps sinensis* and has various pharmacological activities [9,10].

*P. penicillatus* (white mold) is a pathogen of morels (*Pezizomycetes, Ascomycota*) [11], which are valuable edible fungi [12–14]. Morel cultivation is important in China [12,15,16] and is a major income source for many growers. However, white mold is a serious disease affecting the cultivation of *Morchella* spp. [11]. White mold reduces production and quality of morels during cultivation, storage, and transportation [11]. White mold is highly adaptable, and has the ability of rapid colonization and transmission [11,17], which made it difficult to control without affecting...
mellor production. The pathogenic mechanism of \textit{P. penicillatus} and the related resistance mechanism of morells have not been fully understood [17].

The availability of high-throughput DNA sequencing platforms, including Roche 454, SOLiD, Illumina, Ion Torrent, and PacBio, can aid the study of genomic features [18], evolution, environmental adaptation, pathogenicity [19], and secondary metabolite synthesis of pathogenic fungi. The genomes of four 	extit{Paecilomyces} species (\textit{P. penicillatus} CCMJ2836 [17], \textit{P. variotii} CBS144490 HYG1 [20], \textit{P. variotii} CBS 101075 [20], and \textit{P. niveus} 07 [21]) have been published. However, different fungal strains varied greatly in genome size, gene number and gene sequence. In order to more accurately analyze the pathogenic mechanism of \textit{P. penicillatus} and the resistance mechanism of \textit{M. importuna} spp., we further sequenced and assembled the \textit{P. penicillatus} genome based on two sequencing platforms.

We sequenced the genome of \textit{P. penicillatus} using Illumina and PacBio technologies. PacBio sequencing produces long read lengths (>10 Kb) and greatly improves the continuity of genome assembly. Transcription profiles of \textit{P. penicillatus} infecting \textit{Morchella importuna} (an important cultivated morrel species in China) at different stages were detected by transcriptome sequencing to reveal the pathogenic mechanisms of \textit{Paecilomyces} and the resistance mechanisms of \textit{M. importuna}. This study is the first report on pathogenic mechanism of \textit{Paecilomyces} and resistance mechanism of \textit{M. importuna} from the transcriptional level, which will promote our understanding of the genomic feature, fungi-fungi interaction, pathogenicity, and genome evolution of \textit{P. penicillatus} and \textit{M. importuna}.

2. Materials and methods

2.1. Fungal strains and experimental treatment

\textit{P. penicillatus} mycelia were obtained from the Institute of Plant Protection, Sichuan Academy of Agricultural Sciences (No. SAAS_ppe1) [22]. They were isolated from the fruiting bodies of \textit{M. importuna} with white mold disease and verified by Koch’s law. The \textit{P. penicillatus} mycelia were cultured on potato dextrose agar medium (PDA, potato, 200 g/L; glucose, 20 g/L; agar, 15 g/L) at 28°C for 7 d. Morel cultivation strains were purchased from a local company in Chengdu. \textit{M. importuna} was cultivated in a field in Pujiang, Sichuan province. When the fruiting bodies of \textit{M. importuna} reached 7 cm in height, we inoculated them with cultured \textit{P. penicillatus} mycelia. Four groups of samples were collected for transcripome analysis, and each group contained three replicates (Fig. 1). The groups were pure cultured \textit{P. penicillatus} mycelia in PDA (Ppe); healthy fruiting bodies of \textit{M. importuna} (Mim); \textit{M. importuna} pathogenic sites (1 cm²) infected by \textit{P. penicillatus} for 3 days (early stage of infection) (MP1); \textit{M. importuna} pathogenic sites (1 cm²) infected by \textit{P. penicillatus} for 6 days (late stage of infection) (MP2).

2.2. DNA and RNA extraction

Pure cultured \textit{P. penicillatus} mycelia were harvested for DNA extraction. Genome DNA of \textit{P. penicillatus} was extracted using a fungal DNA kit (Omega Bio-Tek, Norcross, GA, USA) according to manufacturer’s instructions. The extracted DNA was detected using 1% agarose gel electrophoresis. DNA quality was evaluated using a Nanodrop spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA), with the optical density ratio of 260/280 nm and 260/230 nm greater than 1.8 and 2.0, respectively. Samples from the four groups (Ppe, Mim, MP1, and MP2) were collected and frozen in liquid nitrogen for RNA extraction. Total RNA was extracted using TransZol reagent (TransGen Biotech, Beijing, China) according to manufacturer’s instructions. We used the Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA) to evaluate RNA quality. Truseq Stranded Total RNA Sample Preparation Kit (Illumina) was used to prepare paired-end libraries for each RNA-seq sample. The obtained libraries were subjected to quality control steps and then submitted to the Illumina HiSeq 2500 platform for sequencing (Personal Biotechnology, Shanghai).

2.3. Genome sequencing and assembly

We constructed two different gDNA libraries for the Illumina HiSeq system and PacBio system. Illumina sequencing libraries were constructed using a library construction kit (Illumina, San Diego, CA, USA) with 400 bp inserts. The PacBio libraries were constructed according to the manufacturers’ instructions of the PacBio system with a 20-kb-insert size. Generated gDNA libraries were subjected to Illumina HiSeq and PacBio RS II platforms for genomic sequencing by Personal Biotechnology (Shanghai, China). Long reads from the PacBio platform were used to de novo assemble the genome. Short reads from the Illumina platform were used to conduct a genome survey and base level correction after the assembly [23]. We used the Illumina sequencing data to estimate genome size with the K-mer-based method. The frequency of each K-mer from the short sequencing data was calculated using jellyfish (v2.1.3) [24]; FALCON package [25] was used to assemble the long sequencing reads generated from the PacBio RS II platform into contigs. Using the PacBio long reads, we polished the assembled genome sequences through two rounds of polishing with Quiver. Pilon [26] was used to perform genome-wide base-level correction with the Illumina short sequencing reads. Then, we used the GapCloser program [27] to fill gaps and obtained a final draft genome.

2.4. Repetitive element and non-coding RNA annotation

Both homologous comparison and ab initio prediction were applied to annotate repeat elements in the \textit{P. penicillatus} genome [28]. For homologous repeat annotation, RepeatMasker v4.0.5 [29] and RepeatProteinMask [29] were used for known repeat element types by searching against the Repbase database [30]. For ab initio repeat annotation,LTR_FINDER [31], RepeatScout [32], and RepeatModeler v1.0.4 (http://repeatmasker.org/RepeatModeler/) were used to construct a de novo repetitive element database, and the RepeatMasker [29] (http://repeatmasker.org/RMDown load.html) was used to annotate repeat elements with the database. The tRNA and rRNA genes in the \textit{P. penicillatus} genome were predicted using tRNAscan-SE v1.3.1 [33] and Rfam v1.2 [34], respectively. Other non-coding RNAs in the \textit{P. penicillatus} genome were annotated by the internal tool using Rfam database [35].

2.5. Protein-coding gene prediction and functional annotation

Exonerate software [36] was used to predict protein-coding genes in the \textit{P. penicillatus} genome using protein sequences from closely related species. Augustus [37], GlimmerHMM [38], and SNAP [39] were used for ab initio prediction with the optimized parameters that were trained using high-quality proteins derived from the PASA gene models [40]. We combined gene models predicted by the two methods using EvidenceModeler [40]. Carbohydrate-active enzymes in the \textit{P. penicillatus} genome were annotated using hmmScan v3.1b2 [41] against the Carbohydrate-Active enzymes Database [42] (http://www.cazy.org) with an E-value threshold of 1E-5. Public biological function databases of SwissProt [43], evolutionary genealogy of genes: Non-supervised Orthologous Groups (eggNOG) [44], NR from NCBI, Gene Ontology (GO) [45], and Kyoto Encyclopedia of Genes and Genomes (KEGG)
were used for functional annotation of the predicted genes. Secondary metabolite encoding gene clusters were predicted using AntiSmash [47].

2.6. Analyses of RNA-Seq data

Data qualities were checked by FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/). We used the FASTX-Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/) program to filter out rRNA reads, short-fragment reads, sequencing adapters, and other low-quality reads from raw reads of the sequences. We mapped the remaining clean reads onto the \textit{P. penicillatus} and \textit{M. importuna} genomes using Bowtie2 software [48] based on a local alignment algorithm. We judge whether the RNA reads belongs to \textit{M. importuna} or \textit{P. penicillatus} according to the similarities between the reads and corresponding genome sequences. The expression levels were normalized by calculating the fragments per kilobase million reads (FPKM) value. DESeq R package (version 1.22.0) was used to calculate differentially expressed genes (DEGs). The resulting P values were adjusted using Benjamini and Hochberg's approach for controlling the false discovery rate. Genes with an adjusted P-value < 0.05 as detected by DESeq were designated as differentially expressed [49]. GO enrichment analysis of DEGs was implemented by the GOseq R packages that were based on a Wallenius non-central hyper-geometric distribution [50]. KOBAS [51] was used to test the statistical significance of enrichment of DEGs in KEGG pathways [52].

2.7. Gene expression validation by RT-qPCR

To validate the gene expression calculated by transcriptome, we conducted RT-qPCR analysis of 19 genes identified by RNA-Seq. About 2 \(\mu\)g of total RNA from each sample was reverse-transcribed by reverse transcriptase M–MLV (TaKaRa). We designed gene-specific primers based on the gene sequences and used the 5.8S rRNA genes as endogenous loading controls for testing the validity of the template preparation [53]. At least three rounds of independent RT-PCR reactions were conducted to confirm the expression of each gene.

2.8. Data availability

The Illumina short-read sequencing data have been deposited to NCBI Sequence Read Archive (SRA) (Data Citation 1: NCBI Sequence Read Archive SRR9051197). The PacBio long-read sequencing data have been deposited to NCBI Sequence Read Archive (SRA) (Data Citation 2: NCBI Sequence Read Archive SRR9051168). The transcriptome data are available through the NCBI SRA (Data Citation 3: NCBI Sequence Read Archive SRR9682527-SRR9682529). The assembled genome version is available at GenBank (Data Citation 4: GenBank VCCP00000000.1). The annotation gff3 file of the assembled genome is available at Figshare (Data Citation 5: Figshare https://doi.org/10.6084/m9.figshare.8146202.v3). The reference genome of \textit{M. importuna} can be found in JGI database (https://mycocosm.jgi.doe.gov/Morimp1/Morimp1.home.html) [16].

3. Results

3.1. Genome assembly of \textit{P. penicillatus}

After removing adaptors in the raw reads, we obtained about 7.70 Gb (coverage of 187.65\(\times\)) long sequencing subreads from the PacBio platform. The subreads N50 and the longest subreads were 10.64 kb and 63.81 kb, respectively. To improve the accuracy of reads for the \textit{P. penicillatus} genome, approximately 4.05 Gb short sequencing reads were generated from the Illumina platform using paired-end sequencing libraries with insert lengths of 400 bp. Reads with residual adapter sequences and low-quality values (<20) were filtered out. Approximately 3.75 Gb (coverage of 91.25\(\times\)) clean reads were obtained for the k-mer analysis and
base correction of the genome (Fig. S1). We used the GapCloser program [27] to fill gaps and obtained a final 39.78 Mb draft genome. The *P. penicillatus* draft genome contained 11 scaffolds with a scaffold N50 length of 6.50 Mb (Table 1). All 11 scaffolds were over 10 kb, with the longest scaffold being 7.29 Mb. The GC content of the *P. penicillatus* genome was 44.19%, which was the lowest among the five *Paecilomyces* genomes. The single copy orthologs (BUSCO, version 3.0) [54] were used to evaluate the completeness of the assembled genome, and 99.7% of the 290 fungal BUSCO genes were identified in the final assembly. This indicated that the assembly of the *P. penicillatus* genome was of a high quality.

### 3.2. Annotation of the *P. penicillatus* genome

A total of 8,619 protein-coding genes (PCGs) with a mean of 2.6 exons per gene were annotated in the *P. penicillatus* genome based on de novo prediction and a homology-based search (Table 1). The average length of PCGs identified in the *P. penicillatus* genome was 1,593 bp. There were 8,056 (96.48%) PCGs similar to the sequences in the NCBI nr database, 7,844 (93.94%) homologous to sequences in eggNOG, 5,759 (68.97%) mapped to GO, 5,089 (60.95%) classified in Swiss-Prot, and 3,276 (39.23%) assigned to KEGG (Fig. S2).

We identified 50 secondary metabolite encoding gene clusters in the *P. penicillatus* genome using AntiSmash. A total of 6.07 Mb repetitive elements were identified in the *P. penicillatus* genome, accounting for 15.26% of the whole genome. A total of 374 tRNA and 143 rRNA genes were annotated in the *P. penicillatus* genome. We also predicted 32 other ncRNAs in the *P. penicillatus* genome. Comparative genomic analysis showed that the assembly of the *P. penicillatus* SAAS_ppe1 genome had a longer scaffold N50 value than the other four *Paecilomyces* genomes. All scaffolds were longer than 10 kb, indicating that the assembly of *P. penicillatus* SAAS_ppe1 was more complete than that of the other three *Paecilomyces* genomes (Table 1). *P. penicillatus* SAAS_ppe1 had a larger genome size than the other two *Paecilomyces* species (one species has two strains) but smaller than *P. penicillatus* CCMJ2836. The genome of *P. penicillatus* SAAS_ppe1 had the least number of gene models (Table 1). Carbohydrate-active genes in the *P. penicillatus* SAAS_ppe1 genome were more than that in *P. penicillatus* CCMJ2836, *P. variotii* CBS144490 HYG1, and *P. variotii* CBS 101,075 genomes, but less than *P. penicillatus* CCMJ2836. The genome of *P. penicillatus* SAAS_ppe1 had a larger genome size than the other two *Paecilomyces* species (one species has two strains) but smaller than *P. penicillatus* CCMJ2836. The genome of *P. penicillatus* SAAS_ppe1 had the least number of gene models (Table 1).

### 3.3. Overview of the RNA-Seq data

To study the transcriptional regulation mechanism of *P. penicillatus* during infection of *M. importuna*, as well as the transcriptional response of *M. importuna* to infection by *P. penicillatus*, we detected the expression profiles of pure cultured *P. penicillatus* mycelia, healthy *M. importuna* fruiting bodies, and infected samples at different stages using an RNA-Seq technique. An average of 9.08 Gb of raw data (with an average of 60,135,493 raw reads) was generated for each sample, with a Q20 and Q30 of 97.43% and 93.50%, respectively (Table S1). After quality control, we generated an average of 9.01 Gb clean data (with an average of 59,820,429 clean reads) from the raw data; about 0.53% of the raw reads were discarded. All the clean reads were mapped to a corresponding reference genome. The average mapped rates were 78.39% and 95.49% of the samples Mim and Ppe to corresponding genomes, respectively. A total of 86.34% and 74.49% clean reads from the infected sites were mapped to the genome of *M. importuna* at early and late stages of infection, respectively. However, 7.48% and 19.83% clean reads from the infected sites were mapped to the *P. penicillatus* genome at early and late stages of infection, respectively. An average of 5.93% clean reads from the infected sites with mixed *M. importuna* fruiting bodies and *P. penicillatus* mycelia were not mapped to the two reference genomes.

### 3.4. Genes up-regulated or down-regulated with infection stage in *P. penicillatus*

Of the 8,466 genes expressed in pure culture and infected mycelia of *P. penicillatus*, 3,943 genes were differently expressed between the *P. penicillatus* samples at different stages of infection (Fig. 2). Of the 3,943 differently expressed genes, the expression of 24 genes increased with the infection and infection stage (Fig. 3a), and they included diphthamide biosynthesis, aldehyde reductase, NAD(P)H-lydrate epimerase, and GPI anchored protein (*P < 0.05*). A total of 25 genes were down-regulated with the infection and infection stage (*P < 0.05*). The down-regulated genes were related to NAD-dependent epimerase/dehydratase, glyoxalase-like domain-containing protein, putative histidine--RNA ligase, and acetyltransferase (Fig. 3b).

### 3.5. Genes differentially expressed with infection stage in *P. penicillatus*

A total of 47 genes were first up-regulated and then down-regulated with infection in the infection stage in *P. penicillatus* (*P < 0.05*). These were related to fumitremorgin C mono-oxygenase-like protein, glycosyltransferase family 31 protein, ferric-chelate reductase, concanava A-like lectin/glucanase, and cupin 2 domain-containing protein (Fig. 4a). A total of 125 genes, including mannitol-1-phosphate 5-dehydrogenase, phosphatidylinositol 4,5-bisphosphate-binding protein SLM1, glucose repressible protein Grg1, reductase, peptidase activity, and peptidase inhibitor 178 (Fig. 4b and 4c), were first down-regulated and then up-regulated with the infection and infection stage in *P. penicillatus* (*P < 0.05*).
3.6. Functional enrichment analysis in P. penicillatus

GO functional enrichment analysis showed that when the fruiting body of M. importuna was infected by P. penicillatus at the early stage, four GO terms, related to membrane, membrane part, intrinsic component of membrane, and integral component of membrane, were significantly enriched (P < 0.05). The four GO terms could be assigned to the cellular component category (Table S2). Only one GO term was significantly enriched in P. penicillatus samples at the late infection stage compared with samples at the early infection stage (P < 0.05). This was related to carbohydrate metabolic process in the category of biological processes.

Fig. 2. Volcano plot of differentially expressed genes (DEGs) between samples (M. importuna). The X-axis represents the fold change of DEGs in different experimental groups; -log10 (p-value) in the Y-axis represents DEGs that were more significant as the value increased. The dots represent the genes; the grey dots indicate non-differentially expressed genes; the red dots represent up-regulated differentially expressed genes; the blue dots represent down-regulated differentially expressed genes. Ppe, pure cultured P. penicillatus mycelia in PDA medium; MP1, M. importuna fruiting bodies infected by P. penicillatus for 3 days; MP2, M. importuna fruiting bodies infected by P. penicillatus for 6 days. Each treatment had three biological replicates. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 3. Genes up- (a) or down- (b) regulated with infection and infection stages in P. penicillatus. The black number in the color block indicates the fragments per kilobase million reads (FPKM) value of different genes. The gene expression level increased with color from yellow to blue. Ppe, pure cultured P. penicillatus mycelia in PDA medium; MP1, M. importuna fruiting bodies infected by P. penicillatus for 3 days; MP2, M. importuna fruiting bodies infected by P. penicillatus for 6 days. Each treatment had three biological duplicates. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
KEGG pathway analysis showed that 35 pathways were significantly enriched between the pure culture and the early infection stage of *P. penicillatus* mycelia (*P* < 0.05). These were involved in “N-Glycan biosynthesis,” “protein processing in endoplasmic reticulum,” “cardiac muscle contraction,” and “glycerolipid metabolism” (Fig. 5). Metabolism pathways relating to “glyoxylate and dicarboxylate metabolism” and “carbon fixation in photosynthetic organisms,” and organismal systems relating to “plant-pathogen interaction,” and “adrenergic signaling in cardiomyocytes” were also over-expressed at the early infection stage of *P. penicillatus* (*P* < 0.05). Ten KEGG pathways were significantly enriched in *P. penicillatus* mycelia at the late infection stage compared with those at the early stage (*P* < 0.05). These were related to starch and sucrose metabolism, amino sugar and nucleotide sugar metabolism, and galactose metabolism in the metabolism category, prolactin signaling pathway, and vitamin digestion and absorption in the category of organismal systems.

### 3.7 Genes up- or down-regulated with infection stage in *M. Importuna*

A total of 10,955 genes were expressed in healthy and diseased *M. importuna* fruiting bodies, of which 1,576 genes were significantly differently expressed between healthy and diseased *M. importuna* fruiting bodies at different stages (*P* < 0.05) (Fig. 6). There were 142 genes differently expressed between healthy *M. importuna* fruiting bodies and diseased *M. importuna* fruiting bodies at the early stage, and 1,349 genes differently expressed between diseased *M. importuna* fruiting bodies at the early and the late stages (*P* < 0.05). Of these differently expressed genes, the expression of 10 genes increased with the infection and infection stage (*P* < 0.05). These
included cyclin-dependent protein kinase inhibitor and nine genes with unknown functions (Fig. 7a). No gene was always down-regulated with the infection and infection stage.

### 3.8. Genes differentially expressed with infection stage in *M. importuna*

A total of 15 genes were first up-regulated and then down-regulated with the infection and infection stage in *M. importuna* (*P* < 0.05), and they were related to zinc finger MYND domain-containing protein, aquaporin-like protein, and 13 genes with unknown functions (Fig. 7b). Twenty-eight genes, including YTH-domain-containing protein, retrotransposable element Tf2, cdc42 effector protein, mitochondrial 54S ribosomal protein YmL16, and 24 hypothetical proteins were first down-regulated and then up-regulated with the infection and infection stage in *M. importuna* (*P* < 0.05).

### 3.9. Functional enrichment analysis in *M. importuna*

GO functional enrichment analysis showed that no GO term was significantly enriched between healthy and diseased *M.
importuna fruiting bodies at different stages. KEGG pathway analysis showed that nine pathways were significantly enriched between the healthy and diseased *M. importuna* fruiting bodies at the early stage (*P* < 0.05) (Fig. 8). Organismal systems pathways relating to “longevity regulating pathway,” “PPAR signaling pathway,” and “GABAergic synapse,” metabolism pathway relating to “fatty acid metabolism,” “fatty acid biosynthesis,” and “biosynthesis of unsaturated fatty acids,” genetic information processing pathway relating to “fanconi anemia,” and environmental information processing pathway relating to “FoxO signaling” were over-expressed at the early infection stage of *M. importuna* (*P* < 0.05).

Four KEGG pathways were significantly enriched between diseased *M. importuna* fruiting bodies at the early and late stages (*P* < 0.05). These were related to glycan degradation, carbon metabolism, and pentose phosphate pathway in the category of metabolism and mineral absorption in the organismal systems category.

3.10. Validation of the DEG results by qPCR analysis

We validated gene expression profiles obtained by transcriptome analysis using qPCR analysis. This confirmed the expression levels of 19 selected genes, with 5.8S rRNA serving as the reference gene (Table S3). All the 19 identified genes were successfully amplified and resulted in a single band. Three genes were up-regulated in *M. importuna* at the early infection stage compared with healthy *M. importuna* fruiting bodies at the early and late stages (*P* < 0.05). These results were related to glycan degradation, carbon metabolism, and pentose phosphate pathway in the category of metabolism and mineral absorption in the organismal systems category.

### 4. Discussion

*P. penicillatus* is a pathogenic fungus that greatly reduces the yield and quality of morels [11]. However, the pathogenicity mechanism of *P. penicillatus* has not been fully understood. Fungal genomes and mitogenomes are variable in size, structure, intron and gene number [55–57,78]. In order to more accurately analyze the pathogenic mechanism of *P. penicillatus* and the resistance mechanism of *Morchella* spp., we further sequenced and assembled the *P. penicillatus* SAAS_ppe1 genome to improve the integrity of genome assembly. We obtained the genome of *P. penicillatus* SAAS_ppe1 based on PacBio and Illumina sequencings. The genome of *P. penicillatus* had a longer scaffold (N50) than the other four *Paecilomyces* species, and all its scaffolds were greater than 10 kb long. BUSCO assessment results showed that 99.7% of the 290 fungal BUSCO genes exist in the assembled genome of *P. penicillatus* SAAS_ppe1, which was higher than the other four *Paecilomyces* genomes. The combined application of PacBio and Illumina sequencing methods produced a *P. penicillatus* genome that was of higher quality than other species in the genus [58,59]. The other four *Paecilomyces* genomes were assembled based on only one sequencing method. A high-quality *P. penicillatus* genome enabled...
analysis of its pathogenicity mechanism. We were able to successfully distinguish the transcripts of *P. penicillatus* and *M. importuna* in the pathogenic sites, and analyze pathogenic mechanisms and transcriptional response mechanisms. In addition, compared with the other *Paecilomyces* species, *P. penicillatus* has fewer gene models and more carbohydrate-active genes belonging to auxiliary activities (AAs), carbohydrate esterases (CEs), and polysaccharide lyases (PLs). The increased number of carbohydrate-active genes may contribute to the strong adaptation and infection abilities of *P. penicillatus* [60,61].

We obtained the transcriptional profiles of a pure culture of *P. penicillatus* mycelia, healthy fruiting bodies of *M. importuna*, and mixed transcripts of *P. penicillatus* and *M. importuna* at early and late infection stages, from pathogenic sites. During the early stage of infection, 86.34% and 7.48% of the mixed transcripts were mapped to the genome of *M. importuna* and *P. penicillatus*, respectively. While in the late stage of infection, 74.49% and 19.83% of the mixed transcripts were mapped to the genome of *M. importuna* and *P. penicillatus*, respectively. These results showed that *M. importuna* was dominant in the pathogenic site, but its biomass and transcripts decreased over time, indicating that healthy *M. importuna* tissues were gradually infected by *P. penicillatus*. The biomass of *P. penicillatus* gradually increased with the infection and infection stage. Differentially expressed gene analysis showed that 24 and 25 genes were first up-regulated and then down-regulated, respectively, with the infection and infection stage, indicating that these genes are involved in the process of *P. penicillatus* infection. The up-regulated genes included diphthamide biosynthesis, aldehyde reductase, NAD (P) h-hydroxy epimerase, and GPI anchored protein. The biological function of diphthamide is not well understood. Previous studies showed that diphthamide modification plays an important role in regulating the sensitivity of cells to bacterial toxins and maintaining translation fidelity [62,63]. Our results suggest that diphthamide plays a role in *P. penicillatus* infection. Aldehyde reductase is a member of the aldo–keto reductase superfamily, which catalyzes the NADPH-dependent reduction of a variety of aldehydes to their corresponding alcohols [64,65]. NAD (P) h-hydroxy epimerase can help repair NAD(P)H hydrates [66]. GPI anchored proteins act as membrane anchors of many eukaryotic cell surface proteins [67]. These genes are important in the physiological regulation of eukaryotes. Our study provides the first information on the regulatory roles of these genes in the process of *P. penicillatus* infection.

We found 47 genes that were first up-regulated and then down-regulated with the infection and infection stage in *P. penicillatus*. These genes were related to glycosyltransferase family 31 proteins and ferric-chelate reductase. Glycosyltransferases are important proteins for the synthesis of complex carbohydrates [68]. Ferrichelate reductase participates in the synthesis of sesquiterpenoids with important biological activities in fungi [69]. A total of 125 genes were first down-regulated and then up-regulated with the infection and infection stage in *P. penicillatus*, including mannitol-1-phosphate 5-dehydrogenase and glucose repressible protein Grgl1. Mannitol-1-phosphate 5-dehydrogenase participates in the mannitol cycle for NADPH regeneration [70]. Glucose repressible protein Grgl1 helps fungi utilize a wide range of carbon sources except for glucose [71]. GO functional enrichment showed that, in the early infection stage, cellular component function was significantly enriched in *P. penicillatus*, showing its important role in early infection of *P. penicillatus*. In the late infection stage, carbohydrate metabolic process was over expressed in *P. penicillatus*. These results show that *P. penicillatus* has diverse regulation patterns that it uses to infect *M. importuna* in different infection stages. KEGG pathways, including “glycerolipid metabolism” and “plant-pathogen interaction” were over-expressed at the early infection stage of *P. penicillatus*. Glycerolipid metabolism helps regulate lipid synthesis and metabolism and influences stress responses [72]. Plant-pathogen interactions can play an important role in the process of pathogens infecting plants [73]. Our study revealed that metabolic pathways have an important regulatory role in the early stage of fungus-fungus interactions. Ten KEGG pathways were significantly enriched in *P. penicillatus* mycelia at the late infection stage compared to the early infection stage. These pathways were related to amino sugar and nucleotide sugar metabolism and galactose metabolism, which are involved in the regulation of cell morphology and plant-fungus interactions [74,75].

*M. importuna* is cultivated in China and other countries [76,77]. However, the yield of *M. importuna* is reduced by pathogens. *P.
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**Author Contributions**

Conceived and designed experiments: C.C., R.F., J.W., and Q.L.. Analyzed the data: C.C., X.L., and X.C.. Wrote and review the paper: C.C., and Q.L.. Funding acquisition and project administration: D.L.

**Declarations of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Appendix A. Supplementary data**

Supplementary data to this article can be found online at [https://doi.org/10.1016/j.csbj.2021.04.065](https://doi.org/10.1016/j.csbj.2021.04.065).

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[1] Luangsa-ard JJ, Hywel-Jones NL, Samson RA. The polyphyletic nature of M. importuna that can cause stalk and cap rot [11]. However, the transcriptional response mechanism of M. importuna to pathogens was previously unknown. We found that the expression levels of 10 genes increased with the infection and infection stage in M. importuna and included cyclin-dependent protein kinase inhibitor and nine genes with unknown functions. Fifteen genes were first up-regulated and then down-regulated with the infection and infection stage in M. importuna. The results indicated that these genes had roles in regulating the response of M. importuna to P. penicillus infection. Fatty acid biosynthesis and metabolism pathways were significantly enriched at the early infection stage of M. importuna. Fatty acid biosynthesis and metabolism plays an important role in the plant-fungus interaction [73]. We found that this pathway participated in the fungus-fungus interaction. The functions of many proteins from M. importuna are not understood. This study shows the transcriptional response mechanism of M. importuna and the data will also be useful in the functional analysis of M. importuna proteins.

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