Genome Analysis of a Zygomycete Fungus *Choanephora cucurbitarum* Elucidates Necrotrophic Features Including Bacterial Genes Related to Plant Colonization

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A zygomycete fungus, *Choanephora cucurbitarum* is a plant pathogen that causes blossom rot in cucurbits and other plants. Here we report the genome sequence of *Choanephora cucurbitarum* KUS-F28377 isolated from squash. The assembled genome has a size of 29.1 Mbp and 11,977 protein-coding genes. The genome analysis indicated that *C. cucurbitarum* may employ a plant pathogenic mechanism similar to that of bacterial plant pathogens. The genome contained 11 genes with a Streptomyces subtilisin inhibitor-like domain, which plays an important role in the defense against plant immunity. This domain has been found only in bacterial genomes. Carbohydrate active enzyme analysis detected 312 CAZymes in this genome where carbohydrate esterase family 6, rarely found in dikaryotic fungal genomes, was comparatively enriched. The comparative genome analysis showed that the genes related to sexual communication such as the biosynthesis of β-carotene and trisporic acid were conserved and diverged during the evolution of zygomycete genomes. Overall, these findings will help us to understand how zygomycetes are associated with plants.

The kingdom Fungi has a critical role in the carbon cycle¹ as well as various interactions with other living organisms. Zygomycetes diverged in the early stage of fungal evolution²,³ and have various ecological types, which encompass plant and animal pathogens⁴–⁹, saprobes¹⁰, and parasites of other fungal species¹¹. Mucorales is the largest and best-studied clade in zygomycetes.

The Mucorales fungus *Choanephora cucurbitarum* is a necrotrophic plant pathogen, which causes fruit and blossom rot in cucurbits and other plants, including eggplant, squash, ice plant, okra, pumpkin, and petunia, with varying degrees of severity¹²–¹⁷. The infection caused by this fungus is frequently found in warm and humid climates such as tropical and subtropical regions¹⁸. Several dozen Mucorales genomes have been sequenced, including the opportunistic human pathogenic *Rhizopus delemar*¹⁹, oleaginous *Mucor circinelloides*²⁰, human pathogenic *Lichtheimia corymbifera*²¹, and thermophilic *Rhizomucor miehei*²². There are some Mucorales fungi reported as plant pathogen including *Mucor circinelloides* (wound pathogen of noni fruit)¹⁷, *Rhizopus microsporus* (pathogen of maize, sunflower, and rice)⁴, and *Rhizopus stolonifer* (pathogen of sweet potato)²³, but none of their genomic features have been analyzed and reported in previous studies.

Fungal plant pathogens are classified by three different lifestyles: necrotrophs, biotrophs, or hemibiotrophs²⁴. Fungal pathogens have developed different plant colonization strategies depending on their ecological niches and physiological characteristics. Thus, it is important to understand various forms of plant pathogenicity in a genomic scale, and it can be done by comparing the functional repertoires that shape fungal pathogens’ lifestyles. General approaches for understanding plant pathogenicity include identifying functional domains related to host infection, such as adhesion, detoxification, secondary metabolism, and signal conduction²⁵, which can be inferred using Pfam²⁶ or Gene Ontology (GO)²⁷. In addition, the carbohydrate-active enzyme (CAZyme)²⁸ profile can

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be used to characterize the lifestyles of fungi. Global CAZyme investigations of a kingdom of fungi showed that necrotrophic pathogens have more CAZymes than other ecotypes such as biotrophs and saprobes. Interestingly, biotrophic Ustilago maydis has an extremely decreased number of total CAZymes, but instead the gene clusters of secreted virulence factors were found in its genome. This suggests that CAZymes are not the only factors that determine plant pathogenicity in fungi. Secreted effectors have also been investigated because plant pathogenic fungi interact with host cell death machinery via these effectors.

In pathogen–host interaction, genome evolution via sexual communication is important to host adaptation. β-carotene derivatives, particularly trisporoids, have been identified to be responsible for partner recognition and early sexual differentiation in zygomycetes. Three genes, tsp1, tsp2, and tsp3, are associated with trisporic acid biosynthesis that encode 4-dihydromethyltrisporate dehydrogenase, 4-dihydrotrisporin dehydrogenase, and carotene oxygenase, respectively. AcaA encodes an additional carotene cleavage oxygenase acting on the cleavage product of β-carotene made by tsp3-encoded oxygenase in Phycomyces. The biosynthesis of β-carotene is well-conserved in Mucorales and is mediated by two genes, carRA and carB.

Here, we report the genome sequence of the plant pathogenic Mucorales fungus Choanephora cucurbitarum.

Results and Discussion

Genome assembly and gene prediction. We sequenced and assembled the genome of Choanephora cucurbitarum KUS-F28377 isolated from green squash in Korea. On the basis of high-quality reads (18.8 million reads with 4.3 billion bases), we assembled a genome of 29.1 Mbp with 2,814 scaffolds. The estimated genome size based on the k-mer frequency was 29.2 Mbp, thereby indicating that 99.8% of the entire genome was covered by the assembly. The N50 values for the contigs and scaffolds were 24.2 kbp and 27.9 kbp, respectively, and the sequence coverage was 81.3-fold. The read-depth coverage and GC-content profile for each scaffold showed no indication for the sequences of contaminants, symbionts, or parasites in the final assembly.

| Assembly statistics          |          |
|------------------------------|----------|
| Total contig length (Mbp)    | 28.89    |
| Total scaffold length (Mbp)  | 29.10    |
| Average base coverage (fold) | 86.2     |
| Number of scaffolds          | 2,814    |
| N50 contig length (kbp)      | 24.2     |
| N50 scaffold length (kbp)    | 27.9     |
| G+C content (overall) (%)    | 37.41    |
| G+C content (coding region) (%) | 41.65 |
| G+C content (non-coding region) (%) | 34.38 |

| Predicted protein-coding genes       |          |
|--------------------------------------|----------|
| Predicted genes                      | 11,977   |
| Percent coding (%)                   | 49.17    |
| Average coding sequence size (nt)    | 1,194    |
| Gene density (genes/Mb)              | 411.62   |
| Total exons                          | 47,510   |
| Total introns                        | 35,533   |
| Number of introns per gene (median)  | 3        |
| Number of exons per gene (median)    | 3        |
| Average exon length (nt)             | 301.16   |
| Average intron length (nt)           | 71.39    |

Table 1. Genome features of C. cucurbitarum.
Orthologous and orphan genes. Orthologous gene families were inferred on the basis of the proteomes of C. cucurbitarum, ten Mucorales, and two Glomerales genomes. Among the 19,598 inferred gene families, 2,904 (14.82%) families were shared by all 13 genomes (single-member gene families were not counted). The C. cucurbitarum genome shared the most gene families with the M. circinelloides genome (6,383 families, Supplementary Figure S4). The degree of gene sharing was consistent with the relationships in the phylogenetic tree shown in Fig. 1 (Supplementary Table S2 shows the counts for the shared gene families in all 13 genomes). The number of members per gene family provided little evidence of the whole-genome duplication events, as found in Rhizopus genomes19, which was consistent with the smaller genome size of the C. cucurbitarum genome (Supplementary Figure S4 and Supplementary Table S3).

The C. cucurbitarum genome contained 1,326 unique genes (11.11% of overall genes) where their orthologs were absent in any of the comparative genomes used in this study, and they were designated as “orphan genes”. Among 1,326 orphan genes, 1,113 genes belonged to single-member gene families and 213 genes belonged to multi-membered gene families. The second largest gene family for the orphan genes comprised 31 members containing the F-box domain (PF12937), which is related to the ubiquitin proteolysis mechanism36. Five orphan genes were annotated with CAZymes: CE4 (acetyl xylan esterase, EC 3.1.1.72), CBM14 (chitin-binding module), GH31 (α-glucosidase, EC 3.2.1.20), GH37 (α,α-trehalase, EC 3.2.1.28), and AA6 (1,4-benzoquinone reductase, EC. 1.6.5.6) modules. These unique CAZymes may determine the host specificity of this organism. Interestingly, the orphan proteins had shorter lengths than those in the overall proteome, where the median lengths of the orphan proteins and all proteins were 107 aa and 318 aa, respectively (Table 2 and Supplementary Figure S5). This bias still needs to be explained. GO annotations showed that the functions of “protein binding,” “oxidation–reduction process,” “ATP binding,” “integral component of membrane,” “serine-type endopeptidase inhibitor activity,” and “metal ion binding” were frequent in these orphan genes.

To identify effector candidates, we predicted 657 genes containing a signal peptide for transport outside of cells. In particular, 97 secreted protein-encoding genes were orphan genes that could potentially function as host-specific effectors, where 84 of these lacked any known functional domain, which is consistent with the current rule for defining effector genes as secreted proteins without known functional domains30. The other 13 genes included protease inhibitor domain- and catalase domain-containing genes involved in defense against plant immunity.

Comparative genome analysis showed a various number of orphan genes, ranging from 188 to 8,396, which represents the diverse lifestyles of Mucorales fungi adapting to ecological niches. Orphan genes were often highly duplicated. For instance, M. ambiguus genome had an orphan gene family with 69 members. Majority of duplicated orphan genes were not annotated with Pfam or CAZymes. The ortholog groups and the functional
names corresponding to the abbreviations are listed in the methods section. This observation was also identified in the comparative genomes except in R. irregularis DAOM 181602 and U. isabellina NBRC 7884. The orphan genes and statistical tests for secreted orphan proteins are summarized in Table 2 and Supplementary Table S6.

### Pfam domain annotation and bacterial Streptomyces subtilisin inhibitor (SSI)-like domain.

We used Pfam annotations to identify unique or enriched functional domains in the *C. cucurbitarum* genome. We identified 12,767 Pfam domains within 7,936 genes, which was lower than the numbers in the comparative genomes (11,810–28,947 domains, Supplementary Table S7). However, the numbers of unrepeated Pfam entries were similar (3,504 for *C. cucurbitarum* and 3,249–3,676 for the others, Supplementary Figure S6). Indeed, around 65% of the Pfam entries (2,284 entries) were shared by all 13 genomes. In addition, the most frequent Pfam domains were common in all 13 genomes: “protein kinase domain,” “WD domain,” “RNA recognition motif,” “major facilitator superfamily,” and “helicase conserved C-terminal domain” (Supplementary Figure S7). These observations suggest that they share most functions and that small variations determine their lifestyles. The *C. cucurbitarum* genome contained 14 unique, 11 enriched, and 18 depleted Pfam entries (P < 0.01, Fisher’s exact test) compared with the other 12 genomes (Fig. 2).

The most notable enriched domain was the SSI-like domain (PF00720), where 11 copies were present (Fig. 2). It has been reported that only Actinobacteria harbor this domain, which is found mostly in *Streptomyces* spp.37. We found that only one other Mucorales fungus, *Absidia glauca*, contains three SSI-like proteins among all the fungal genomes deposited in NCBI. The lengths of the *C. cucurbitarum* SSI-like proteins were relatively short, ranging from 109 aa to 138 aa, and they were encoded by a single exon, except for one containing two exons. Only 6/11 SSI-like genes were supported by mRNA-seq in the hyphal state, which indicates that they might be differently regulated. Eight of the SSI-like proteins had extracellular localizations in the same manner as bacterial homologs (6/8 bacterial SSI-like proteins investigated). The three non-extracellular SSI-like proteins were destined for the endoplasmic reticulum, mitochondria, and cytosol, respectively, thereby suggesting that they may have various roles in cells.

There were clade-specific conserved positions as well as universally conserved positions in SSI-like protein sequences. We aligned a total of 19 SSI-like protein sequences (11 from *C. cucurbitarum* and eight from bacteria) and identified 35 conserved positions in all 19 sequences. The 16 and 12 positions were conserved only in *C. cucurbitarum* and bacteria, respectively (Fig. 3a). Four cysteine positions that form two disulfide bonds were well conserved in all sequences (positions 107, 122, 144, and 175 in Fig. 3a), which indicates that these disulfide bonds are important for correct structure formation in both groups.

We examined the phylogenetic relatedness between *C. cucurbitarum* and bacterial SSI-like domains by building phylogenetic gene trees. The neighbor-joining tree showed the nestedness among the SSI-like domains of *C. cucurbitarum* and three bacterial species: *Thermobifida*, *Actinosynnema*, and *Nocardia* (Fig. 3b and Supplementary Figure S8). The maximum-likelihood tree built with the same dataset also supported similar grouping pattern (Supplementary Figure S9). Despite that the bootstrap values of branching nodes were weak in the trees to pinpoint the exact phylogenetic relationship, the tree analysis implied that SSI-like domains of *C. cucurbitarum* have been horizontally transferred. These SSI-like paralogs probably help to neutralize the host’s immunity because subtilisin proteases have functions in pathogen recognition in plants.

### Distributions of CAZymes and the CE6 family.

The *C. cucurbitarum* genome contained fewer CAZymes in total than dikaryotic necrotruchs. This genome contained 312 CAZymes whereas other dikaryotic necrotruchs...
contained 419–865 CAZymes with an average of 605 as previously reported. This number is lower than the average number of CAZymes in hemibiotrophs (542 CAZymes) and is similar to that found in symbiotic and biotrophic plant pathogens (328 and 313 CAZymes, respectively). Plant pathogens have been known to have more CAZymes than saprobes and biotrophs because they need to degrade plant biomass to facilitate their invasion. Thus, this suggests that not all plant pathogens have expanded CAZyme modules and different plant pathogenicity factors, such as secreted effector proteins, would lead to dissimilar CAZyme profiles. Interestingly, a decreased number of total CAZymes was also observed in other Mucorales with an average of 342 CAZymes, except for *R. microsporus* with 585 CAZymes, which might be attributable to whole-genome duplication.

The carbohydrate esterase family 6 (CE6, acetyl xylan esterase) was enriched in the *C. cucurbitarum* genome with seven copies, whereas the other Mucorales had one or two copies of each. This family is only conserved in zygomycetes among all fungal clades, but it is also found in bacterial and plant genomes. The CE6 family is annotated with “acetyl xylan esterase” and it improves the saccharification of stem lignocellulose by deacetylating xylan. The gene tree of CE6 family indicated that the fungal CE6 genes were more diverged from the common ancestor of plants and the bacterial CE6 family.

Pathogen–host interactions and bacterial virulence genes. The genes involved in pathogen–host interactions (PHIs) were inferred by identifying their homologs in the PHI-base database. Among 11,977 predicted genes, 2,277 (19.01%) genes shared similarity with the PHI genes. In particular, 1,504 genes were related to plant hosts (monocots or eudicots) and there were fewer of these genes than those in the comparative genomes (1,362–2,967 genes, Table 3 and Supplementary Figure S11) and two *Colletotrichum* genomes (2,395 and 2,972 PHI genes in *C. graminicola* and *C. higginsianum*, respectively), which may be attributable to its small genome size. In total, 226 PHI entries were shared by all 16 genomes (*C. cucurbitarum*, ten Mucorales, two Glomerales, two *Colletotrichum*, and *Ustilago*). These common virulence genes determine the core plant-infecting factors, and they are not specific to hosts. Four known effectors were found in the *C. cucurbitarum* genome: two copies of PHI:2703 (*Xanthomonas axonopodis*-derived citrus canker causing effector), one copy of PHI:2347...
Figure 3. Sequence alignment and gene tree of SSI-like proteins from *C. cucurbitarum* and bacterial genomes. (a) Multiple sequence alignment of SSI-like domain-containing protein sequences from *C. cucurbitarum* and bacterial genomes. Conserved positions in all proteins are highlighted in light green, and positions conserved only in *C. cucurbitarum* and bacteria are highlighted in orange and light purple, respectively. GenBank accession numbers are labeled for each sequence: WP_020666218 from *Amycolatopsis* nigrescens, WP_055506216 from *Nonomuraea* sp. NBRC 110462, WP_042258841 from *Nocardia brasiliensis*, WP_043813828 from *Allokutzneria albata*, WP_01836711 from *Streptomyces sp.* NRRL B-24720, and WP_051772499 from *Saccharothrix* sp. NRRL B-16314. (b) Gene tree of SSI-like domains built by neighbor-joining method. Clustered nodes were collapsed for clarity. Each node is labeled with UniProt ID with corresponding organism name in brackets. The tree was built using FastTree with the default option.
(grey mould-causing effector), and one copy of PHI:4506 (Epichloe festucae-derived effector). Genes involved in host-interactions in *C. cucurbitarum* still need to be fully elucidated, so we expect that many genes were missed in this investigation.

**Sexual communication genes in the Mucorales and Glomerales genomes.** The analysis of trisporoids and β-carotene biosynthesis-related genes revealed the conservation and divergence of these genes over Mucorales and Glomerales genomes. As shown in Fig. 5, we investigated four genes related to trisporoids biosynthesis (*tsp1*, *tsp2*, *tsp3*, and *acaA*) and two genes related to β-carotene biosynthesis (*carB* and *carRA*). *Tsp3*, which encodes carotene oxygenase catalyzing β-carotene cleavage, was well-conserved in the Mucorales genomes as a single copy with the exception of two copies in the *R. microsporus* genome. The *acaA* gene had high sequence similarity with *tsp3* (33% identity and 50% similarity in *Phycomyces* proteins), therefore the ortholog-finding method was applied to differentiate the two genes. Interestingly, *tsp1* (4-dihydromethyltrisporate dehydrogenase) and *tsp2* (4-dihydrotrisporin dehydrogenase) had various degrees of duplications over the genomes ranging from one to eight copies. The phylogenetic tree showed that *tsp1* genes were divided into three groups (Supplementary Figure S12), suggesting that *tsp1* genes were duplicated for functional divergence in contrast to *tsp2* genes (Supplementary Figure S13). Two *R. irregularis* genomes lacked both *tsp3* and *acaA* genes, but contained *tsp1* and *tsp2* genes. This observation suggests the synthesis of signal compounds by non-β-carotene pathway, which is also supported by an absence of *carB* and *carRA* orthologs in these genomes.

**Conclusion**

Here we report the genome sequence of *C. cucurbitarum* KUS-F28377, a pathogen of cucurbits and other plants isolated in Korea. We determined the functional profiles of the *C. cucurbitarum* genome and compared them with the relatives and other plant pathogenic fungal genomes to identify the genetic features responsible for the mechanisms of plant pathogenicity unique to this genome. In conclusion, exploring the pathogenicity of *C. cucurbitarum* using genomics can provide crucial insights, which may facilitate the development of disease control strategies.

**Methods and Materials**

**Strain isolation and genomic DNA/RNA preparation.** The strain *C. cucurbitarum* KUS-F28377 (Korean Agricultural Culture Collection No. 48046, http://www.genebank.go.kr/eng/about/aboutus.jsp) was isolated from *Cucurbita moschata*. Cultures were grown for 10 days on potato dextrose agar (PDA) at 25°C and the fungal mycelium was harvested. Genomic DNA was extracted by following the protocol provided by Kohler et al.43.
Genome sequencing and assembly. In total, 10 million read pairs (300 bp per read and 2.79 billion bases) were generated by Illumina MiSeq paired-end sequencing. The raw reads were trimmed and filtered on the basis of sequence quality (20 Q-score cutoff) and length (20 bp cutoff) using HTQC 1.92.344. A preliminary assembly was generated by ABySS 1.9.045 with the options of k-mer = 41 and insert size = 400 bp. The assembled contigs were used to construct a simulated jumping library with wgsim (https://github.com/lh3/wgsim), which generated 4.5 million read pairs with an insert size of 3 kbp without errors (-e 0 -d 3000 -s 300 -N 5000000 -1 150 -2 150 -r 0 -R 0 -X 0). Finally, 2,814 scaffolds were assembled by Allpaths-LG r4730046 with 86.2× sequence coverage. Scaffolds less than 1 kbp were excluded. The potential sequences of contaminants, symbionts, or parasites was checked with Blobology 47, which plots read-depth coverage calculated by read alignment using Bowtie2 48 and GC content of each scaffold.

Gene prediction. Transcriptomic data were obtained by Illumina MiSeq sequencer, which generated 1.2 million paired-end reads with 725 million bases. The raw reads were quality controlled by HTQC, as described above, and assembled with Trinity 49, which yielded 14,759 contigs (8,895 contigs >500 bp). We generated a gene prediction pool using three tools: Maker 2.31.850, Augustus 3.2.151, and Braker1 (http://exon.gatech.edu/genemark/braker1.html), which produced 32,974 preliminary gene predictions. When gene coding regions are overlapped between two genes, we only kept the more feasible predictions based on the BLASTp bit score against the NCBI nr database, which yielded 11,977 genes as the final predicted genes. The proteomes of six relatives were used to support protein homology-based gene prediction in Maker: L. corymbifera, L. ramosa, M. ambiguus, M. circinelloides, R. microsporus, and R. delemar.

Table 3. Pathogen-host interaction-related genes. Organism names corresponding to the abbreviations are listed in the methods section.
Genome comparisons. The genomes of ten Mucorales, two Glomerales, and three plant pathogenic dikarya species were used for comparisons with the C. cucurbitarum genome. The abbreviations for each organism were used in the figures: Ceu, Choanephora cucurbitarum; Agl, Absidia glauca CBS 101.48; Lco, Lichtheimia corymbifera JMR; Fsu, Fungi studio ATCC 62417; Lrr, Lichtheimia ramosa; Mmm, Mucor ambiguum NBRC 6742; Mci, Mucor circinelloides I. circinelloides 1006P; Ppa, Parasatella parasitica CBS 412.66; Rde, Rhizopus delemar 99-880; Rmi, Rhizopus microsporus ATCC 62417; UisB, Umbelopsis isabellina B7317; UisN, Umbelopsis isabellina NBRC 7884; Rir18, Rhizophagus irregularis DAOM 181602; Rir19, Rhizophagus irregularis DAOM 197198w; Ggr, Colletotrichum graminicola M1.001; Chi, Colletotrichum higginsianum IMI 349063; and Uma, Ustilago maydis 521. We predicted genes on two Umbelopsis genomes because they were not available in the NCBI database. The gene predictions were performed in the same way that we did to C. cucurbitarum genome using RNA-seq reads downloaded from NCBI sequence read archive (accession ID: SRX1003360).

Pfam domain annotation. InterProScan 5.16–55 was used to annotate the Pfam domains in the C. cucurbitarum and comparative genomes. An enrichment test was performed using Fisher's exact test embedded in the Python Scipy module. Secreted proteins were predicted using WoLF-PSORT.

Gene tree of SSI-like proteins. We used curated seed alignment of SSI-like domain sequences downloaded from the Pfam database (http://pfam.xfam.org/family/PF00720), which contained 25 sequences. We extracted domain sequences from the C. cucurbitarum SSI-like protein sequences inferred by InterProScan (64–83 aa in length). Using Mafft, we added these sequences into the seed alignment (seed option). The neighbor-joining domain sequences from the RAxML 8.2.75 were estimated using FastTree (with a local bootstrap based on 1,000 resamples) and RAxML (-f a -x 12345 -p 12345 -# 100 -m PROTGAMMAWAG) for multiple sequence alignment, eliminating poorly aligned positions, PHYLIP format conversion, and tree inference, respectively. Aspergillus nidulans (http://www.aspgd.org) was used as an outgroup for the rooted tree.

CAZy prediction. CAZymes were identified using BLASTp, dbCAN, and Pfam annotations. We assigned a CAZyme annotation when at least two methods yielded the same predictions. For BLASTp, we used the hits with a bit score of >50 and alignment coverage of >25% to remove poorly aligned sequences. An enrichment test was performed using Fisher's exact test embedded in the Python Scipy module.

PHI-related genes. Genes potentially involved in PHIs were identified using BLASTp with a bit score cutoff of 50 and a minimum coverage of 25% against the PHI-base database (http://www.phi-base.org).

Sexual communication genes. We retrieved the sexual communication protein sequences from GenBank: tsp1, Q01213 (Mucor mucedo); tsp2, AM937248.1 (Mucor mucedo); tsp3, CAL64769 (Blakeslea trispora); tsp4, XP_018293784.1 (Phycomyces blakesleeanus); tsp5, AAO46892.1 (Blakeslea trispora); and tsp6, AAO46893.1 (Blakeslea trispora). We ran BLASTp to the reference sequences and selected the true orthologs using gene families of best hit.

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Author Contributions
J.P. and H.S. isolated and identified the species; J.P., H.S., H.P., and I.C. designed and performed the experiments; B.M. and I.C. performed bioinformatics and statistical analyses; B.M., J.P., and I.C. wrote the manuscript.

Additional Information
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