Apophysomyces variabilis: draft genome sequence and comparison of predictive virulence determinants with other medically important Mucorales

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Abstract

Background: Apophysomyces species are prevalent in tropical countries and A. variabilis is the second most frequent agent causing mucormycosis in India. Among Apophysomyces species, A. elegans, A. trapeziformis and A. variabilis are commonly incriminated in human infections. The genome sequences of A. elegans and A. trapeziformis are available in public database, but not A. variabilis. We, therefore, performed the whole genome sequence of A. variabilis to explore its genomic structure and possible genes determining the virulence of the organism.

Results: The whole genome of A. variabilis NCCPF 102052 was sequenced and the genomic structure of A. variabilis was compared with already available genome structures of A. elegans, A. trapeziformis and other medically important Mucorales. The total size of genome assembly of A. variabilis was 39.38 Mb with 12,764 protein-coding genes. The transposable elements (TEs) were low in Apophysomyces genome and the retrotransposon Ty3-gypsy was the common TE. Phylogenetically, Apophysomyces species were grouped closely with Phycomyces blakesleeanus. OrthoMCL analysis revealed 3025 orthologues proteins, which were common in those three pathogenic Apophysomyces species. Expansion of multiple gene families/duplication was observed in Apophysomyces genomes. Approximately 6% of Apophysomyces genes were predicted to be associated with virulence on PHLbase analysis. The virulence determinants included the protein families of CotH proteins (invasins), proteases, iron utilisation pathways, siderophores and signal transduction pathways. Serine proteases were the major group of proteases found in all Apophysomyces genomes. The carbohydrate active enzymes (CAZymes) constitute the majority of the secretory proteins.

Conclusion: The present study is the maiden attempt to sequence and analyze the genomic structure of A. variabilis. Together with available genome sequence of A. elegans and A. trapeziformis, the study helped to indicate the possible virulence determinants of pathogenic Apophysomyces species. The presence of unique CAZymes in cell wall might be exploited in future for antifungal drug development.

Keywords: Apophysomyces variabilis, CotH proteins, Proteases, Transposons, Genome structure, Sequence analysis

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Background

*Mucoromycotina*, a subdivision of Kingdom Fungi is currently placed in *incertae sedis* and comprises three orders namely *Mucorales, Endogonales* and *Mortierellales* [1]. In the order *Mucorales* 14 families, 56 genera and around 225 species have been identified [2, 3]. The traditional taxonomy of mucoralean fungi was largely based on morphological features, sexual reproduction and ecological characters. The current taxonomical position of fungal species under the order *Mucorales* is evolving with the use of molecular techniques. Recently, Spatafora et al. proposed a phylum level phylogenetic classification of mucoralean fungi comprising of two phyla (*Mucoromycota* and *Zoopagomycota*), six subphyla, four classes, and 16 orders [4]. Of the known 225 species under *Mucorales*, approximately 20 species are known to cause human infections.

*Rhizopus arrhizus* is the predominant (~60%) agent causing mucormycosis worldwide [5, 6]. *Lichtheimia* and *Mucor* species are the next group of fungi isolated from mucormycosis cases in western world [7, 8]. In contrast, *Apophysomyces* species ranks second after *R. arrhizus* in India; around 60% of world mucormycosis cases due to *Apophysomyces* species are reported from this country [9–11]. Misra et al. isolated this fungus from the environment of a mango orchard of North India in 1979 for the first time [12]. Subsequently the fungus was reported from the soils with low nitrogen content in India [13]. *A. elegans* was considered the only species under the genus *Apophysomyces* causing human infection. However, the molecular phylogenetic studies in recent years revealed *Apophysomyces* genus encompasses 5 species: *A. elegans*, *A. mexicanus*, *A. ossiformis*, *A. trapeziformis* and *A. variabilis*. *A. elegans*, *A. trapeziformis*, and *A. variabilis* are common human pathogenic species [14, 15]. The majority of human infections are due to *A. variabilis* [16]. Though *Apophysomyces* species generally produce cutaneous and subcutaneous infection following trauma, occasional deep seated infections involving lung, paranasal sinuses and kidney have also been reported [9, 11, 17]. *A. variabilis* is the major pathogen in India and occasional cases are due to *A. elegans* [10]. *A. trapeziformis* has been reported from necrotising fasciitis cases among tornado victims from USA [18, 19]. The study on pathogenesis of mucormycosis draws attention of clinicians and scientists, as *Mucorales* are known to cause severe infection and high mortality. Recently, using transcriptome and genomic approach, various pathogenic determinants have been identified in *Mucorales* [20]. However, the genome of *A. variabilis* had not been sequenced and pathogenic determinants of *Apophysomyces* species were not clearly delineated. The present study reports the draft genome sequence of *A. variabilis* for the first time and compared the same with already available genomes of *A. elegans* and *A. trapeziformis* to identify the possible virulence determinants in pathogenic *Apophysomyces* species.

Results

Genome features and organisation

The whole genome shot-gun high quality reads of *A. variabilis* (NCCPF 102052) isolate were generated using pair-end and mate-pair libraries on Illumina platform. *A. trapeziformis* (B9324) and *A. elegans* (B7760) genomes were downloaded from NCBI genome database for the comparative analysis. The genome assembly of *A. variabilis* consisted of 1155 contigs and 431 scaffolds with N50 scaffold length of 647.5 kilobases (Kb) and total genome size of 39.38 Mb. Whereas, the genome sizes of *A. elegans* and *A. trapeziformis* were 38.46 Mb and 35.84 Mb respectively. The average G + C content of the three *Apophysomyces* genomes ranged from 41.7% to 41.9%. The genome annotation of the three *Apophysomyces* species were not clearly delineated. The present study reports the draft genome sequence of *A. variabilis* for the first time and compared the same with already available genomes of *A. elegans* and *A. trapeziformis* to identify the possible virulence determinants in pathogenic *Apophysomyces* species.

Table 1 Salient features of genome assembly of three *Apophysomyces* species

| Assembly Statistics | A. variabilis | A. elegans | A. trapeziformis |
|---------------------|--------------|------------|-----------------|
| Number of scaffolds | 431          | 1528       | 1400            |
| Number of contigs   | 1155         | 1528       | 1400            |
| Total size of scaffolds (bp) | 39,387,571 | 38,467,560 | 35,840,759 |
| Scaffold N50 (bp)   | 647,494      | 101,064    | 141,065         |
| Average Scaffold size (bp) | 91,386 | 25,175     | 25,601          |
| Max. scaffold size (bp) | 3,993,377  | 469,772    | 572,943         |
| Min. scaffold size (bp) | 500         | 301        | 205             |
| GC Content (%)      | 41.65        | 41.69      | 41.90           |

Coding sequences (CDS)

| Number of CDS | 12,764 | 12,829 | 12,108 |
| Total size of CDS (bp) | 25,752,686 | 25,262,110 | 24,132,511 |
| CDS N50 (bp) | 3141      | 3018     | 3067       |
| Average CDS size (bp) | 2018 | 1969     | 1993       |
| Max. CDS size (bp) | 34,923   | 35,019   | 35,013     |
| Min. CDS size (bp) | 57       | 44       | 43         |
| GC Content (%) | 45.49   | 45.54    | 45.69      |
| Exons (bp) | 97,463   | 98,425   | 94,508     |
| Average exon size (bases) | 263   | 255      | 254        |
| Exons/gene | 7.64      | 7.67     | 7.61        |
| Number of tRNA genes | 197   | 174      | 197         |
| Non coding sequences |
| Introns (bp) | 79,821 | 80,964 | 78,014 |
| Introns/gene | 6.25  | 6.31    | 6.44       |
| Average intron length (bp) | 77    | 76      | 76         |

bp: base pairs, CDS: coding DNA sequences
genes had multiple exons, with an average of seven exons per gene and exon length of ~255 bp (Table 1). The genomes of Apophysomyces species had six introns per gene with an average length of ~77 bp.

Protein coding gene prediction and annotation
Protein coding genes were functionally annotated using BLASTx (e-value less than 10^-5) from non-redundant protein database of NCBI. Approximately 88% of the genes in the genome were functionally annotated. The annotated genes were assigned to different Gene Ontology (GO) domains comprising of biological, cellular and molecular functions (Fig. 1a, Additional file 1: Figure S1). The PFAM annotation predicted 1877 predicted proteins for A. variabilis, 1875 for A. trapeziformis and 1884 for A. elegans (Fig. 1b). The BLASTx annotation of the predicted genes in Apophysomyces genomes revealed maximum hits with Lichtheimia ramosa (formerly Absidia idahoensis), followed by Lichtheimia corymbifera and Rhizopus microsporus.

Transposable elements (TEs)
Apophysomyces genome contained low number of transposable elements. The proportion of TEs to the whole genome of A. variabilis, A. elegans and A. trapeziformis were 5.1%, 2.8%, 1.63% respectively. The retrotransposon Ty3-gypsy was the most abundant TE found in all three Apophysomyces genomes (Table 2). The low TE content in Apophysomyces species might be attributed to the expansion of RNA interference mediated genes and heterokaryon incompatibility (HET) proteins in their genomes.

Non-coding RNA
Annotation of tRNAs was performed using infernal tool. Comparison against RFAM database revealed 197 tRNAs in A. variabilis and A. trapeziformis and 174 tRNAs in A. elegans (Table 1). Apophysomyces genomes contained single copy of nuclear ribonuclease P (RNase P), a ribozyme that cleaves the 5′ regions of precursor tRNAs to generate mature 5′ ends of tRNA. Like all eukaryotes, the genome of Apophysomyces species contained all five

![Fig. 1](image-url) Gene ontology and major protein family classification in Apophysomyces genomes. a) Functional classification of proteins by Gene Ontology in three Apophysomyces genomes, b) major protein families annotated using PFAM in Apophysomyces genomes.
small nuclear ribonucleoproteins (snRNA) essential for formation of major spliceosomes. Few components of minor spliceosome (U1, U6atac) were also present in the assembly. Other essential components like U4atac and U12 snRNA were not present.

*A. variabilis* and *A. trapeziformis* contained group I introns, a self splicing ribozyme. The histone 3′ UTR stem-loop was present only in *A. variabilis* genome and it plays a major role in transport of the histone mRNAs from nucleus to cytoplasm (Additional file 2: Table S1).

**Orthologs and paralog prediction**

OrthoMCL analysis was performed for three *Apophysomyces* genomes and compared with other *Mucorales*. A total of 3025 ortholog proteins were shared among three *Apophysomyces* genomes (Fig. 2a). Apart from the shared ortholog genes, 5078 genes were shared between *A. trapeziformis* and *A. variabilis* and 241 genes between *A. elegans* and *A. variabilis* (Fig. 2a). *Apophysomyces* genomes shared common orthologous genes with *L. corymbifera* (2078 genes), *M. circinelloides* (2248 genes) and *R. delemar* (2094 genes) (Fig. 2b, c & d). Paralogues/gene duplication events in *Apophysomyces* were predicted separately for each species using the total protein coding genes. Paralog gene prediction data showed 629 paralog/gene duplication groups for *A. variabilis* comprising 2334 genes, 733 paralog groups for *A. trapeziformis* comprising 2175 genes, and 308 paralog groups with 879 genes for *A. elegans*. Approximately 18% of the genes in *A. variabilis* and *A. trapeziformis* genomes were found to be duplicated and 6.9% in *A. elegans*. The maximum number of gene paralogs had transposons within the genes, which possibly play a role in gene duplications. Several gene expansions had been observed in cell wall components (chitin synthase, chitin deacetylase), electron transport chain (ABC membrane, E1-E2 ATPase, cytochrome p450), secreted proteases (aspartic protease), sugar transporters, CotH invasins and signal transduction pathways in all three *Apophysomyces* species.

**Predicted virulence properties**

PHIbase tool was used to determine the possible genes associated with virulence in *Apophysomyces* species. The analysis predicted diverse protein domains, families and repeats encompassing 5.7%, 6.4% and 6.1% genes of *A. variabilis*, *A. trapeziformis* and *A. elegans* respectively for virulence properties of the organisms. The protein kinase domain and Ras proteins were the predominant genes associated with virulence in *Apophysomyces* species. CotH proteins (invasins), proteases, siderophores, CBS domains, cell wall components (chitin synthase,
chitin deacetylase) and septins were the other genes determining virulence in Apophysomyces species.

Like other Mucorales, Apophysomyces genomes contain multiple copies of CotH protein (42.8 kDa protein). A. variabilis and A. trapeziformis had 17 copies of CotH genes, whereas A. elegans had 16 copies. The orthoMCL analysis revealed multiple groups of CotH paralog genes in three Apophysomyces species.

A. variabilis genome had 488 predicted protease gene families evaluated from MEROPS database, 468 in A. trapeziformis and 422 in A. elegans (Table 3). Serine proteases were the major group of proteases found in all Apophysomyces genomes ranging from 36.8 to 42.1% of all predicted proteases, followed by metallo-proteases. A. variabilis genome revealed two-fold higher expansion of aspartic protease genes (22.1%) compared to 9.2% of A. elegans and 13.6% of A. trapeziformis. Single cluster of aspartic protease paralog gene was noted in all three species. However, expansion of paralogs genes encoding aspartic protease 2 (PF13650) was observed only in A. variabilis. OrthoMCL analysis revealed the presence of transposons within the protein domain of aspartic proteases. This phenomenon suggested the possibility of aspartic protease gene insertion at multiple sites of genome. Subtilase, the major serine proteases in Apophysomyces genomes revealed multiple paralog genes; three paralog groups for A. trapeziformis, two for A. variabilis and one for A. elegans. Genes involved in iron acquisition pathways were conserved in Apophysomyces genomes (Table 4). Apophysomyces species contained high affinity iron permease (FTR1) genes (iron acquisition from host), IucA/IucC and FhuF-like transporter (regulator for siderophore biosynthesis), heme oxygenase (utilisation of heme from blood) and ferritin (iron storage) genes. However, the expansion/duplication of iron acquisition genes (siderophores, heme oxygenase, ferritin, ferrochelatase) was not found in Apophysomyces genomes. Apophysomyces genomes contained multiple copies of CBS domain (iron binding and acquisition). OrthoMCL analysis did not detect any paralog gene for CBS domain.

Carbohydrate-active enzymes (CAZymes) in Apophysomyces genomes
CAZymes were predicted and classified using dbCAN. Apophysomyces genome contained glycoside hydrolases (GHs), glycosyl transferases (GTs), carbohydrate esterases (CEs), carbohydrate binding modules (CBMs) and polysaccharide lyases (PLs) genes (Table 5, Additional file 2: Table S2). A. variabilis had 97 glycoside hydrolases (GHs) genes predominantly belonging to GH18 family (chitinases). The glycosyl transferases (GTs) were the
major CAZymes found in *A. variabilis*. This class of enzyme largely belongs to GT2 family of chitin synthase genes, an essential component for cell wall synthesis. *A. variabilis* had 75 carbohydrate esterases (CEs) genes and majority consisted of CE4 family that code for chitin deacetylase, the enzyme that helps in the conversion of chitin to chitosan. *A. variabilis* contained other CAZymes like polysaccharide lyases (PLs) and carbohydrate binding modules (CBMs) in their genome. *Apophysomyces* species had similar percentage of CAZymes in their genome (Table 5). The orthoMCL analysis revealed multiple gene duplications/paralogs in CAZymes.

**Secretory protein analysis**

A total of 435 (3.3%) genes encoding the secretory proteins were predicted for *A. elegans*, 435 (3.6%) genes for *A. trapeziformis* and 412 (3.2%) genes for *A. variabilis*. The predicted secretory proteins of *Apophysomyces* genome predominately contained carbohydrate active enzymes. The diverse group of secreted CAZymes in *Apophysomyces* genome were glycoside hydrolase (GHs) that consisted of chitinases (GH 18) and lysozymes (GH 25), followed by carbohydrate esterase (CEs) that predominantly consisted of chitin deacetylase (CE4). To predict the secretory peptidases, the secretory proteins were searched against MEROPS peptidase database using BLASTp. Aspartic protease and subtilase were the major peptidases in *Apophysomyces* genome and those enzymes might play role in tissue destruction of the host. BLASTp analysis against lipase engineering data base revealed 9 to 14 genes coding for secretory lipase in the *Apophysomyces* species (Table 6).

**Phylogenetic analysis**

We used BUSCO to identify near universal single-copy ortholog proteins from 29 species. A total of 13 single copy gene proteins, common to all 29 species were identified and used for phylogenetic tree construction (Table 7). The fungal species were classified into two major groups, Ascomycota and Basidiomycota. Fungi belonging to the subkingdom Mucoromycotina were closely related to Mortierellomycotina. The Mucorales were grouped in two major clusters. The three *Apophysomyces* species were grouped together in close association with *Phycomyces blakesleeanus* (Fig. 3). A separate phylogenetic tree was constructed by excluding *Ascomycota* and *Basidiomycota*, to increase the single copy

| Table 3 | Protease gene families in *Apophysomyces* genomes |
|---------|--------------------------------------------------|
| Proteases | *A. variabilis* N (%) | *A. elegans* N (%) | *A. trapeziformis* N (%) | *L. corymbifera* N (%) |
| Aspartic protease | 109 (22.3) | 41 (9.2) | 64 (13.6) | 60 (14.5) |
| Cysteine protease | 78 (15.9) | 85 (19.2) | 87 (18.5) | 68 (16.5) |
| Metalloprotease | 101 (20.6) | 110 (24.8) | 111 (23.7) | 146 (35.4) |
| Serine protease | 180 (36.8) | 186 (42.1) | 186 (39.7) | 124 (30) |
| Threonine protease | 18 (3.6) | 18 (4.1) | 18 (3.8) | 14 (3.4) |
| Protease of unknown type | 2 (0.04) | 2 (0.05) | 2 (0.04) | 1 (0.2) |
| Total | 488 | 442 | 468 | 413 |

*N* denotes total number of proteases predicted by MEROPS database

*The information is collected from Reference No. 22

| Table 4 | Genes responsible for iron uptake mechanism in *Apophysomyces* genomes |
|---------|--------------------------------------------------|
| Genes | *A. variabilis* N | *A. elegans* N | *A. trapeziformis* N | *L. corymbifera* N |
| Reductive pathway | | | | |
| High affinity iron permease (FTR1) | 2 | 2 | 2 | 4 |
| Multicopper oxidase | 8 | 10 | 8 | 8 |
| Ferric reductase | 3 | 3 | 3 | 3 |
| Zinc/iron permease | 7 | 7 | 8 | 6 |
| Siderophore transporter | | | | |
| IucA / IucC family and Ferric iron reductase FhuF-like transporter | 2 | 2 | 2 | 1 |
| Heme oxygenase | 2 | 2 | 2 | 2 |
| Ferrochelatase | 3 | 3 | 3 | 2 |
| Ferrochelatase | 1 | 1 | 1 | – |

*N* denotes gene copy numbers as annotated by PFAM database

*The information is collected from Reference No. 22
genes. A total of 72 single copy genes from *Mucoromycotina*, *Mortierellomycotina* and *Entomophthoromycotina* were analysed, which revealed similar findings as that of 13 single copy genes (Additional file 1: Figure S2).

**Discussion**

The current study revealed *A. variabilis* genome of 39.38 Mb. The genome was compared with available genome sequences of *A. elegans* (38.46 Mb) and *A. trapeziformis* (35.84 Mb). The genome structures of three pathogenic *Apophysomyces* species were compared with other pathogenic *Mucorales* to identify the predicted virulence properties of *Apophysomyces* species. *Apophysomyces* species have more than 12,000 protein coding genes. Among *Mucorales*, *Mucor racemosus* has the maximum number of protein coding genes (21,385 genes) with genome size of 75.32 Mbs and *Umbelopsis isabellina* has the least number of protein coding genes (7612 genes) with genome size of 21.87 Mbs [20]. The protein coding genes of common human pathogens, *Rhizopus arrhizus* (17,467 genes) and *Lichtheimia corymbifera* (12,379 genes) lie close to *Apophysomyces* species [21, 22]. The predicted protein families of *Apophysomyces* species (~1800 protein families), is also close to *Rhizomucor miehei* (1827 proteins) and *R. arrhizus* (1653 proteins) [23]. Phylogenetically, *Mucoromycotina* including *Apophysomyces* species is closely related to *Mortierellomycotina* rather than *Entomophthoromycotina*. The phylogenetic analysis of the present study confirms the findings of super tree constructed of 9748 genes using parsimony approach [22]. The present findings also confirm that pathogenic *Mucorales* are genetically different from *Entomophthorales* [20].

The transposable elements (TEs) in the fungal genomes play a major role in the genomic diversity and genome evolution [24]. The presence of TEs in fungi under *Ascomycetes* and *Basidiomycetes* leads to high degree of diversity ranging from 0.02 to 29.8% of their genomes [24]. *Mucorales* also have a varying percentage of transposable elements in their genomes ranging from 3.17% (*Rhizomucor miehei*) to 35% (*P. blakesleeanus* and *Absidia glauca*) [21, 23]. The present analysis showed that *Apophysomyces* species have lower content of TEs in their genome compared to other *Mucorales*. The lower content of TEs in *L. corymbifera* was associated with the appearance of heterokaryon incompatibility (HET) genes and RNA interference mediated RNA degradation genes like dicer, argonaute and translation initiation factor 2C [22]. Similarly, *Apophysomyces* species contain multiple copies of HET genes in their genome along with RNA interference mediated genes. In comparison, *R. arrhizus* has 35% of TEs in their genome, which may be due to the absence of HET genes [21]. The HET domain is found in all fungi under *Ascomycetes* and helps in the formation of heterokaryons in sexual reproduction. The function of HET domain in *Lichtheimia* and *Apophysomyces* species is not clearly known. The genome duplication and transposon mediated gene expansion may facilitate the pathogenic mechanism of fungi [25].

In Kingdom Fungi, the whole genome duplication has been identified in two separate group of fungi: *Saccharomyco- tin*], an ancestry of the Dikarya [26, 27] and in *R. arrhizus* [21]. In *R. arrhizus*, the whole genome

**Table 5** Comparison between carbohydrate active enzymes (CAZymes) of *Apophysomyces* species and other *Mucorales*

| Carbohydrate active enzymes               | *A. variabilis* | *A. elegans* | *A. trapeziformis* | *R. arrhizus* | *R. miehei* |
|------------------------------------------|-----------------|--------------|-------------------|--------------|------------|
| Glycoside hydrolase (GHs)                | 97 (28.2)       | 92 (27.1)    | 95 (26.9)         | 118 (35.4)   | 110 (40.7) |
| Glycosyl transferase (GTs)               | 118 (34.3)      | 113 (33.2)   | 125 (35.4)        | 143 (42.9)   | 118 (43.7) |
| Carbohydrate esterase (CEs)              | 75 (21.8)       | 81 (23.8)    | 76 (21.5)         | 39 (11.7)    | 20 (7.4)   |
| Carbohydrate-binding module (CBMs)       | 51 (14.8)       | 51 (15)      | 54 (15.3)         | 27 (8.1)     | 20 (7.4)   |
| Polysaccharide lyase (PLs)               | 3 (0.87)        | 3 (0.88)     | 3 (0.85)          | 6 (1.8)      | 2 (0.74)   |
| **Total**                                | 344             | 340          | 353               | 333          | 270        |

*N* denotes number of CAZymes as predicted by dbCAN database

* The information is collected from Reference No. 23

**Table 6** Predicted secretory proteins from *Apophysomyces* genomes

| Secretory Proteins | *A. elegans* | *A. trapeziformis* | *A. variabilis* |
|--------------------|--------------|-------------------|----------------|
| Number of CAZymes  | 68           | 66                | 66             |
| Carbohydrate-binding module (CBMs) | 11          | 14                | 10             |
| Carbohydrate esterase (CEs)         | 20           | 19                | 20             |
| Glycoside hydrolase (GHs)            | 28           | 23                | 30             |
| Glycosyl transferase (GTs)           | 8            | 9                 | 5              |
| Polysaccharide lyase (PLs)           | 1            | 1                 | 1              |
| **Total**                           | 36           | 32                | 41             |
| Aspartic protease                   | 12           | 12                | 14             |
| Cysteine protease                   | 2            | 2                 | 2              |
| Metalloprotease                     | 4            | 3                 | 5              |
| Serine protease                     | 18           | 15                | 20             |
| **Number of Lipases**               | 14           | 9                 | 10             |
duplication is an important phenomenon for the expansion of multiple gene families and plays significant role in pathogenesis and antifungal resistance of the organism [21, 28]. The presence of two copies of RNase P may attribute for whole genome duplication in R. arrhizus [21]. The analysis of L. corymbifera genome showed the presence of tandem and segmental gene duplication [22]. Single copy of RNase P was found in Apophysomyces species like L. corymbifera and indicate the possibility of tandem and segmental duplications in Apophysomyces genomes. Mucorales in general contain more duplicated genome regions (average of 4 to 13 genes) compared to other fungal genomes [29]. The duplication of genes coding for signal transduction pathways (protein kinase, ATP binding) and transport components have been reported in mucoralean genomes [29]. Multiple gene expansion were noticed in Apophysomyces genomes for ABC membrane transport, major facilitator proteins, E1-E2 ATPase and other protein families that were involved in cell wall synthesis, morphogenesis, virulence mechanism and antifungal resistance.

The pathogen-host interaction database predicted approximately 6% of genes in Apophysomyces genomes are associated with virulence of the fungi. The multiple

### Table 7

| S. No | BUSCO IDs          | Protein gene family                                                                 |
|-------|--------------------|--------------------------------------------------------------------------------------|
| 1     | BUSCOEOG7C0V786    | Orotidine 5’ phosphate decarboxylase                                                  |
| 2     | BUSCOEOG7D0FXNZ    | Nucleolar GTP-binding protein 2                                                       |
| 3     | BUSCOEOG7D28XZ     | Transcription initiation factor TFIID subunit 9                                        |
| 4     | BUSCOEOG7F168G     | Phosphoribosylaminomimidazole carboxylase                                              |
| 5     | BUSCOEOG7TJ3V6     | Proteosome core subunit beta 2 (PUP1)                                                  |
| 6     | BUSCOEOG7WDNCV     | Serine/threonine protein kinase (RO1)                                                  |
| 7     | BUSCOEOG7ZDC9      | 1-(5-phosphoribosyl)-5-((5-phosphoribosylamino)methylideneamino)imidazole-4-carboxamide isomerase |
| 8     | BUSCOEOG70KGZ3     | Proteosome regulatory particle base subunit (RPN1)                                     |
| 9     | BUSCOEOG71VTQ2     | Pre-mRNA-splicing factor (CWC22)                                                       |
| 10    | BUSCOEOG74R219     | Signal recognition particle receptor subunit alpha (SRP101)                           |
| 11    | BUSCOEOG7568SQSW   | Glycylpeptide N-tetradecanoyltransferase                                               |
| 12    | BUSCOEOG78WM1Q     | Multifunctional tryptophan biosynthesis protein (anthranilate synthase component 2) |
| 13    | BUSCOEOG779P7X     | Octanoyltransferase                                                                    |

**Fig. 3** Molecular phylogenetic analysis of Apophysomyces species. Phylogenetic tree constructed using 13 single copy genes from 29 fungal species by LG amino acid substitution model with 100 bootstrap replications.
copies of protein kinase and Ras domain in *Apophysomyces* genomes may play a role in pathogenesis. Protein kinase plays a major role in signal transduction pathways and regulates major cellular functions, cell cycle regulation and fungal morphogenesis [30]. Two-fold rise in the protein kinase gene family has been observed in *R. arrhizus*. Similarly, the present analysis showed *Apophysomyces* genomes had multiple paralog gene clusters of protein kinase. The Ras protein regulates eukaryotic growth and differentiation, and orchestrates virulence in pathogenic fungi [31].

The most common virulence trait of *Mucorales* is iron acquisition from the host. The mechanisms included reductive pathways consisting of high affinity iron permease (FTR1), multi-copper oxidase and ferric reductase, siderophore transporter and heme oxygenase [32–34]. In normal hosts, iron in blood is bound to carrier proteins like transferrin, ferrin and lactoferrin. The bound form of iron limits the toxic effects and maintains iron homeostasis [33]. During diabetic ketoacidosis (DKA), the binding affinity of iron to the carrier protein is reduced, leading to the release of free iron in the blood. Free iron helps *Mucorales* to grow in the body [35]. In *L. corymbifera*, up-regulation of FTR1 gene, multi-copper oxidase and ferric reductase is observed under iron starvation [22]. The reduction in the copy number of FTR1 gene abolishes the iron uptake from ferroxamine in *R. arrhizus* [35]. These findings suggest the role of reductase/permease pathway in iron uptake among *Mucorales*. *Apophysomyces* genome contains all the essential components of reductive pathways of high affinity iron permease (FTR1), multi-copper oxidase and ferric reductase that help the iron acquisition from the host. Like other *Mucorales*, *A. variabilis* lacks hydroxamate siderophores (NRPS genes) in their genome. However, compensatory mechanism has been observed in *L. corymbifera*, in the form of upregulation of IucA/IucC family and ferric iron reductase FhuF-like transporter (bacterial siderophores) during iron starvation [22]. *Apophysomyces* genomes also contain genes of IucA/IucC family and ferric iron reductase FhuF-like transporter. The other modes of iron acquisition genes including heme oxygenase pathway and ferritin genes are conserved in *Apophysomyces* species.

Deferoxamine therapy is a risk factor for mucormycosis. Deferoxamine acts as a xenosiderophore and helps in the iron acquisition of *R. arrhizus* [33, 34]. *R. arrhizus* contains CBS domain (Fob1/Fob2 proteins) on their cell surface, which acts as receptor to bind ferroxamine. In deferoxamine treated mice, mutants of Fob proteins have reduced iron uptake, germination and virulence [36]. *Apophysomyces* genome contains multiple copies of CBS domain, which may also play a role in the iron acquisition.

The angio-invasion and tissue necrosis are notable features in mucormycosis infections. The endothelial cell receptors play major role in the angio-invasion of *Mucorales* especially in patients with elevated serum iron concentrations (patients with diabetic ketoacidosis or on deferoxamine therapy) [33, 34]. The glucose-regulated protein-78 (GRP78) receptor on the human endothelial cell is up-regulated in the presence of iron or glucose leading to increase adherence and endocytosis of *R. arrhizus* spores [37]. CotH (invasin) protein on the surface of spore, helps in the binding of the spores to the endothelial cells [20]. The binding of CotH proteins to the GRP78 receptors mediates invasion and endothelial damage [37, 38]. Blocking of CotH receptor with anti CotH antibody abolishes their ability to adhere and invade the host cells [38]. *Apophysomyces* genome possesses multiple copies of CotH genes, which may help in angio-invasion of the fungus. Gene duplication of CotH gene families may increase the virulence properties of *Apophysomyces* species. *Entomophthorales* does not contain CotH genes and has low angio-invasive property [20].

Proteases are also considered to be important for virulence in fungi, as they are responsible for tissue destruction and necrosis in hosts. The genome of *A. variabilis* contains 488 predicted proteases (3.8%), closely similar to *R. arrhizus* (3.6%) and *L. corymbifera* (3.3%) [22]. The common secretory proteases of *A. variabilis* are serine (48.7%) and aspartic protease (34%). The number of secretory protease in *Apophysomyces* genome (8%) is similar to that of *R. delemar* (7%) [22]. However, unlike *R. arrhizus*, *Apophysomyces* species can cause severe tissue destruction in the immunocompetent host.

*Mucorales* cell wall contains unique polysaccharides molecules such as chitin and chitosan, which are not seen in mammals [39]. The *Apophysomyces* genome contains β-1,3-glucan synthase genes (FKS1), but α-1,3-glucan synthase genes are absent. Despite the presence of FKS1 gene, the inherent resistance mechanism of *Mucorales* to echinocandins is not clear [29]. CAZymes play important role in biological processes including cell wall synthesis, cell signaling and energy production [40]. The CAZymes constitute the majority of the secretory proteins in the *Apophysomyces* genomes (Table 6). However, *Apophysomyces* contains lower amount of glycosyl hydrolases and glycosyl transferase compared to *R. arrhizus* and *R. miehei* (Table 5). The expansion in the gene clusters coding for carbohydrate esterase (CEs) and carbohydrate-binding module (CBMs) possibly compensates those deficiencies in *Apophysomyces* species (Table 5). The most common CEs in *Apophysomyces* genomes are CE10 (carboxyl esterase) and CE4 (chitin deacetylase), but CE10 is absent in *R. arrhizus* and *R. miehei* [23]. The expansion of cell wall genes, chitin deacetylase and chitin synthase genes are observed in all the *Mucorales*. However, the functional consequences of these genes are not clear [20–22, 29].
Conclusion
We report the maiden draft genome sequence of A. variabilis, a common pathogen causing mucormycosis in Indian subcontinent. The genome analysis of A. variabilis together with other two pathogenic Apophysomyces species revealed multiple gene duplications in signal transduction pathways, cell signalling molecules and virulence traits. The expansion of virulence genes such as secretory protease, CotH invasins, cell wall components (chitin synthase and chitin deacetylase) is seen in Apophysomyces genomes. The unique CAZymes responsible for the biosynthesis of cell wall polysaccharides may be future antifungal target. Further exploration of Mucorales specific cell wall components may help in understanding the pathogenesis of mucormycosis in detail.

Methods
Fungal strain
Apophysomyces variabilis strain NCCPF 102052 isolated from patient with necrotising fasciitis of thigh was retrieved from the National culture collection for pathogenic fungi (NCCPF), Chandigarh, India. The isolate was grown on potato dextrose agar by incubating at 35 °C for 7 days. DNA was isolated from the fresh mycelia using phenol: chloroform: isoamyl alcohol extraction for 7 days. DNA was isolated from the fresh mycelia using phenol: chloroform: isoamyl alcohol extraction protocol as described earlier [41].

Genome sequencing, assembly and annotation
The whole genome sequence of A. variabilis NCCPF 102052 was performed using paired-end (PE)/(MP) 2*150/2*125 base pair library on Hiseq platform (Illumina, San Diego, California, USA). The paired-end sequencing library was prepared using Illumina TrueSeq Nano DNA HT library preparation kit and the mean size of the library was 645 bp. The Illumina mate- pair libraries were prepared using Cre-Lox recombination method with insert size of 3-5Kb [42]. The raw data was filtered using trimmomatic v0.30 [43] and per base sequence quality score (Q) ≥20 was considered. High quality data were assembled using Soapdenovo2 assembler as described by Li et al., to generate scaffolds using optimized parameters [44, 45]. The whole genome sequence of Apophysomyces trapeziiformis (B9324, GenBank accession ID: JNDP00000000.1) and Apophysomyces elegans (B7760, GenBank accession ID: JNDQ00000000.1) were downloaded from NCBI genome database for the comparative analysis.

Gene prediction
AUGUSTUS version–3.1.01 was used for ab initio gene prediction [46]. Rhizopus arrhizus training set was used as species for gene prediction from both strands. Prediction of incomplete genes was allowed. Using this gene model, CDS & protein sequence was extracted. Protein coding genes were annotated using BLASTx program available from NCBI [47]. Best BLAST hit annotation was transferred to the query protein when the E-value was less than or equal to 10^-5 and identity greater than or equal to 40%. Swiss-Prot database was used for homology searches [48]. Based on these annotations, Gene ontology (GO) and KEGG annotation were retrieved for individual proteins wherever applicable [49].

Protein family annotation
PFAM annotation was performed using PFAM scan tool downloaded from PFAM database [50]. This tool was used for scanning against PFAM database version 30.03 [51].

Transposable elements
TransposonPSI was used to identify diverse families of transposable elements that prevail in the genome sequences. The transposons were classified into DNA and RNA transposons [47].

Non-coding RNA
Non-coding-RNAs were identified and annotated using infernal tool of RFAM database [52].

Peptidase prediction
Peptidase analysis was conducted using BLASTp search against MEROPS database [53]. Annotations were transferred to the query proteins if the identity between query and target proteins were at-least 40% and e value = 10^-4.

Carbohydrate active enzyme (CAZymes) prediction
CAZymes annotation was done using dbCAN database [40]. Hmmscan version 3.1 was used to search against dbCAN and in-house scripts were used to identify the best hit.

Secretory protein analysis
The soluble secretory proteins satisfying the following parameters were analysed: a protein with N-terminal signal peptide, without the presence of trans-membrane domain, complete absence of GPI-anchor site and extracellular secretion of the proteins outside the cell [54, 55]. Secreted proteins were predicted using a customised bioinformatic pipeline including signalP (version 4.1) and TMHMM (version 2.0) software [56, 57]. The presence of signal peptides in the protein sequence was predicted by setting the D-cut off values to “sensitive” (option eukaryotic). The parameter setup includes that secreted proteins that did not contain any transmembrane helix or the one overlapping the signal peptide within first 40 amino acids of the protein sequences. TargetP software (version 1.1) and WolfPsort (version 0.2) were used to predict protein subcellular localization. Proteins were considered secretory if subcellular localization was assigned to extracellular secretory pathway [58, 59]. PS-SCAN (PS00014) was used to find endoplasmic reticulum localisation motif [60]. The
predicted secreted proteins were annotated using BLASTp query search (e value = 10^-5): dbCAN was used to predict CAZymes, MEROPS for peptidase and Lipase Engineering Database for lipase prediction [40, 53, 61].

Pathogen-host interaction (PHI) gene prediction
Genes responsible for possible pathogenesis and virulence mechanism were identified and annotated using PHI-base [62]. BLASTp was used to search for similar sequences from PHI-base and the annotation from best hit annotated sequence was transferred to the query proteins.

OrthoMCL analysis for ortholog and paralog gene prediction
Based on sequence similarity and graph algorithm, OrthoMCL was used to determine the orthologs, co-orthologs and paralogs. Orthologs and co-orthologs are the speciation events and paralogs describe the duplication event within a genome [63, 64]. For OrthoMCL analysis a total of six genomes including three A. elegans genomes, one genome each of R. arrhizus, L. corymbifera and R. rhizophila were used (Additional file 2: Table S3).

Phylogenetic analysis of Apophysomyces species using whole genome sequences
For phylogenetic analysis, 29 species under different genera from Ascomycota, Basidiomycota, Mucoromycota, Mortierellomycotina, and Entomophthoromycotina were included in the analysis (Additional file 2: Table S3). For phylogenetic tree construction, a single copy orthologous protein was identified using BUSCO [65]. The common proteins were used for multiple sequence alignment by MAFFT [66]. Only conserved regions were considered for further analysis by removing the non-conserved regions using Gblocks [67]. Conserved regions of different single copy proteins from each species were concatenated and then subjected to phylogenetic tree construction using PhyML and LG model [68, 69].

Additional files

Additional file 1: Figure S1. Gene ontology (GO) of Apophysomyces genomes. Figure S2. Phylogenetic analysis of Apophysomyces species with other Mucorales. (DOC 376 kb)

Additional file 2: Table S1. Non coding RNAs in genomes of three Apophysomyces species. Table S2. Carbohydrate active enzymes (CAZymes) of Apophysomyces species. Table S3. GenBank accession numbers of whole genome sequences used in phylogenetic analysis of Apophysomyces species and orthoMCL analysis. (DOC 163 kb)

Abbreviations
CAZymes: carbohydrate active enzymes; GO: Gene ontology; KEGG: Kyoto encyclopedia of genes and genomes; LG model: Le & Gascuel model; TE: Transposable elements

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Availability of data and materials
Culture information
The isolate sequenced in the study was deposited to National culture collection of pathogenic fungi (NCCPF) by Dr. Atul Patel, Sterling hospital, Ahmedabad, Gujarat, India.

Genome Accession Number
The whole genome sequence data can be accessed using BioProject accession number: PRJNA379216 and BioSample accession number: SAMN06603527. The whole genome shotgun project has been deposited at DDBJ/ENA/GenBank under the accession MZZL0000000. The version described in this paper is MZZL01000000.

Authors’ contributions
The work was designed under the guidance of AC, SMR, GSP and AKG. The experiments were performed and analysed by: HP, SMR, MK and CB. The manuscript was written and edited by: HP, AC, SMR, GSP, AKG, MK and CB. All authors read and approved the final manuscript.

Ethics approval and consent to participate
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Competing interests
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