### **RESEARCH ARTICLE**

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# Saprophytic and pathogenic fungi in the Ceratocystidaceae differ in their ability to metabolize plant-derived sucrose

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#### **Abstract**

**Background:** Proteins in the Glycoside Hydrolase family 32 (GH32) are carbohydrate-active enzymes known as invertases that hydrolyse the glycosidic bonds of complex saccharides. Fungi rely on these enzymes to gain access to and utilize plant-derived sucrose. In fungi, GH32 invertase genes are found in higher copy numbers in the genomes of pathogens when compared to closely related saprophytes, suggesting an association between invertases and ecological strategy. The aim of this study was to investigate the distribution and evolution of GH32 invertases in the Ceratocystidaceae using a comparative genomics approach. This fungal family provides an interesting model to study the evolution of these genes, because it includes economically important pathogenic species such as *Ceratocystis fimbriata*, *C. manginecans* and *C. albifundus*, as well as saprophytic species such as *Huntiella moniliformis*, *H. omanensis* and *H. savannae*.

**Results:** The publicly available Ceratocystidaceae genome sequences, as well as the *H. savannae* genome sequenced here, allowed for the identification of novel GH32-like sequences. The *de novo* assembly of the *H. savannae* draft genome consisted of 28.54 megabases that coded for 7 687 putative genes of which one represented a GH32 family member. The number of GH32 gene family members appeared to be related to the ecological adaptations of these fungi. The pathogenic *Ceratocystis* species all contained two GH32 family genes (a putative cell wall and a putative vacuolar invertase), while the saprophytic *Huntiella* species had only one of these genes (a putative cell wall invertase). Further analysis showed that the evolution of the GH32 gene family in the Ceratocystidaceae involved transposable element-based retro-transposition and translocation. As an example, the activity of a *Fot5*-like element likely facilitated the assembly of the genomic regions harbouring the GH32 family genes in *Ceratocystis*.

**Conclusions:** This study provides insight into the evolutionary history of the GH32 gene family in Ceratocystidaceae. Our findings suggest that transposable elements shaped the evolution of the GH32 gene family, which in turn determines the sucrolytic activities and related ecological strategies of the Ceratocystidaceae species that harbour them. The study also provides insights into the role of carbohydrate-active enzymes in plant-fungal interactions and adds to our understanding of the evolution of these enzymes and their role in the life style of these fungi.

**Keywords:** *Ceratocystis, Huntiella*, Vacuolar invertases, Cell wall invertases, Sucrolytic ability, Gene family evolution, Molecular dating, Transposable elements, Paralog

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#### **Background**

Glycoside hydrolases (GHs; often referred to as glycosidases or carbohydrases) that target the terminal β- $(2 \rightarrow 1)$  fructosidic bonds found in sucrose and various oligo- and polysaccharides (e.g., fructans, inulin and levan) are functionally designated as invertases [1-3]. These enzymes are classified by their pH optima into the so-called neutral/alkaline invertases that belong to GH family 100 (GH100) and the acid invertases that belong to GH family 32 (GH32; [4]). While GH100 invertases are closely related to the cyanobacterial invertases, the GH32 invertases are closely related to invertases of respiratory eukaryotes such as yeasts and aerobic bacteria such as Bacillus [5]. Like the GH100 family, proteins in the GH32 family have a range of activities [6]. Those specific to GH32 inwith β-fructofuranosidase enzymes 3.2.1.26), inulinase (EC 3.2.1.7, EC 3.2.1.64, EC 3.2.1.80), levanase (EC 3.2.1.65), fructosyltransferase (EC 2.4.1.99, EC 2.4.1.100) and fructosidase (EC 3.2.1.153, EC 3.2.1.154) activities [2, 6].

At the structural level, GH32 together with GH43, GH62 and GH68, are classified as members of the furanosidase (or β-fructosidase) superfamily [7, 8]. These four GH families have a five-blade  $\beta$ -propeller catalytic domain in common, but differ in their mechanisms for glycosidic bond hydrolysis [7]. Those in GH32 and GH68 (designated as clan GH-J) cleave glycosidic bonds in a retaining manner (i.e., retaining of the substrate anomeric configuration), while those in GH43 and GH62 (designated clan GH-F) cleave glycosidic bonds in an inverting manner (i.e., inversion of the substrate anomeric configuration) [8]. GH32 enzymes differ from GH68 in that they contain an additional C-terminal βsheet domain that probably allows for the maintenance of structural stability during protein oligomerisation [9]. In terms of their known distribution across the Tree of Life, GH32 and GH43 occur in plants, fungi and bacteria, GH68 in bacteria only and GH62 in bacteria and fungi [10].

GH32 enzymes have diverse biological roles and they are also exploited for commercial and medical purposes. In plants they influence developmental processes, supply carbohydrates to sink tissues and link intracellular and extracellular stimuli to regulate source/sink relations [11, 12]. In bacteria and fungi they allow for the utilization of plant-derived sucrose as a carbon source [2, 13]. From an industrial perspective, microbial GH32 invertases have various applications [14]. They are used in the confectionery industry to produce short-chain fructooligosaccharides (FOS), which are utilized as calorie-free and non-cariogenic sweeteners [1]. These enzymes are also associated with benefits for human health, for example as immune boosters and antioxidants [15].

Fungi utilize plant-derived sucrose through the production of different GH32 enzymes [2, 16]. In Saccharomyces cerevisiae, two forms of this protein are produced. The first is a non-glycosylated cytoplasmic form that is constitutively expressed, while the second is a glycosylated form that is secreted and repressed by the presence of glucose in the growth medium [17]. Indeed, the overall access to plant-synthesized sucrose appears to be determined by the GH32 family gene copy number [2]. It was previously shown that the number of GH32 genes in a particular species is related to its ecological strategy [2, 13]. Plant pathogens typically show GH32 family expansions, likely because these enzymes play a key role in pathogen nutrition [2, 18]. In contrast, sucrose-independent species, such as animal pathogens and some mycorrhizal fungi, generally lack the genes encoding these enzymes [2]. Such differences in gene copy number can arise from intrinsic molecular processes like unequal crossover and chromosomal duplication, or from processes linked to the activity of mobile genetic elements like transposons [19].

The potential link between GH32 protein family evolution and ecological adaptation has not been explored in the Ceratocystidaceae. This monophyletic family of fungi includes several ecologically diverse lineages that lend themselves to functional comparison [20]. The genus Huntiella, for example, includes exclusively saprophytic species that typically colonize the wounded tissues of trees and other plants [20]. In contrast, the economically important genus Ceratocystis includes mainly pathogens of woody and herbaceous plants, some of which cause devastating tree diseases [21, 22]. Notable examples include the sweet potato pathogen C. fimbriata [23], the mango pathogen C. manginecans [24], and the Acacia pathogen C. albifundus [25]. Despite the availability of whole genome sequence information for all three of the latter species, as well as for H. moniliformis [21] and H. omanensis [26, 27], very little is known regarding their GH32 genes, much less their overall sucrolytic capabilities. In this regard, only one GH32 gene and its associated product has been characterised (i.e., CmINV of H. moniliformis) and tested for its ability to produce FOS [28].

This study considered the structure and evolution of the GH32 protein family in pathogenic and non-pathogenic species in the Ceratocystidaceae. The specific research objectives were: (i) Sequence and assemble the genome of a third *Huntiella* species, *H. savannae*, to allow for meaningful genomic comparison between *Huntiella* and *Ceratocystis*; (ii) Identify and annotate putative GH32 family genes in *H. savannae* and publicly available genomes of *Ceratocystis* and *Huntiella* using an in silico approach; (iii) Infer the evolutionary history of the GH32 family in Ceratocystidaceae and other Sordariomycetes; (iv)

Identify potential genomic processes that shaped the evolution of the GH32 gene family.

#### **Methods**

#### Genome sequences

Genome sequence information for three *Huntiella* species and three *Ceratocystis* species was utilized in this study (Table 1). Genomes for *H. moniliformis* ([Gen-Bank:JMSH0000000]; [26]), *H. omanensis* ([Gen-Bank:JIRZ01 000000]; [27]), *C. manginecans* ([Gen-Bank:JIRZ01 000000]; [26]), *C. fimbriata* ([Gen-Bank:JSSU00000000]; [27]) were generated in previous studies and are available from the Gen-Bank database of the National Centre for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov/). The genome sequence for *H. savannae* (isolate CMW17300, [29]) was determined in the current study (see below).

Isolate CMW17300 of H. savannae was grown on medium containing 20 g/L malt extract agar (MEA, Biolab, Johannesburg, South Africa). Mycelia were scraped from the growth medium and genomic DNA extracted using a phenol/chloroform protocol as previously described by Barnes et al. [30]. The DNA was then sequenced using the Genome Analyzer IIx platform (Illumina) at the Genome Centre, UC Davis, California, USA. Paired-end libraries with an insert size of approximately 350 and 600 bases were used to produce reads with an average length of 100 bases. CLC Genomics Workbench 6.0.1 (CLC Bio, Aarhus, Denmark) was used to discard poor-quality reads and/or terminal nucleotides at a threshold of Q13 (P = 0.05) after which de novo assembly was done using Velvet [31], and an optimal K mer length of 67 determined using VelvetOptimiser (http://bioinformatics.net.au/software.velvetoptimiser.sht ml). The pre-assemblies were scaffolded using SSPACE v.2.0 [31] with default parameters, except -x = 0 and -k= 20. The gaps were reduced with GapFiller v.2.2.1 [32] using default parameters. Open reading frames (ORFs) were predicted using AUGUSTUS [32] based on the gene models for Fusarium graminearum (http://bioinf.uni-greifswald.de/augustus), while genome completeness was evaluated using the Core Eukaryotic Genes Mapping Approach (CEGMA) pipeline [33].

#### GH32 gene identification and characterisation

To identify putative GH32 homologs in the genomes considered in this study, we utilized representative sequences that spanned the fungal GH32 gene family phylogeny [2]. These included Aspergillus oryzae (XP001823245, Group 1), A. niger (ABB59682.1, Group 2), F. verticillioides (FVEG10082.3, Group 3), Botryotinia fuckeliana 4), Stagonospora (BCIG16010.1, Group nodorum (SNOG01192.1, Group 5), Neurospora crassa (EAA32020 Group 6), *A. niger* (ABB59678.1 Group 7), *H. moniliformis* (AGV22100.1 Group 8) [29], and *A. terreus* (XP001218601 Group 9). In the various *Huntiella* and *Ceratocystis* genomes, putative invertase homologs were identified by performing local BLAST searches (tblastn, expect (E)-values  $< 10^{-5}$ ) using BioEdit v 7.2.5 [34]. For comparative purposes, putative invertase homologs among representative Sordariomycetes were identified and obtained using BLAST searches (blastp and tblastn, E-values  $< 10^{-5}$ ) on the Joint Genome Institute (JGI) portal (www.genome.jgi.doe.gov) (Table 1).

For the identified genes, functional domains and features of the predicted proteins were annotated using Inter-ProScan (v.4.8) (http://www.ebi.ac.uk/Tools/pfa/iprscan/), NCBI's Conserved Domain (CD) (http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) and Pfam searches (http://pfam.xfam.org/search), as well as SignalP v.4.1 (www.cbs.dtu.dk/services/SignalP/) and NetNGlyc v.1.0 (www.cbs.dtu.dk/services/NetNGlyc/) analyses. cellular localization analysis was performed using SignalP. Three-dimensional (3D) models of the N-terminal and Cterminal domains were respectively generated and visualised using the Swiss-Model Web server (http:// www.expasy.org/swissmod/SWISS-MODEL.html) Swiss-PdbViewer v.4.04 (http://spdbv.vital-it.ch/). To predict the 3D structure of the identified invertases, a 3D structure of a fructosyltransferase in A. japonicus (PDB id: 3lfi.1) was used as a template.

#### GH32 orthology relationships

Several methods were employed to establish the orthology relationships among the Ceratocystidaceae GH32 homologs. This was important as the characterization of homologous proteins/genes (i.e., those derived from a common ancestry) facilitates inferences regarding their evolution and function [35]. In this study, we used the definitions proposed by Koonin [36] for the terms "paralogy" and "orthology". While orthologs (i.e., homologs that evolved from a common ancestor through speciation) are expected to encode proteins with equivalent functions, paralogs (i.e., homologs that are the product of an ancestral duplication) are thought to more readily acquire novel functional roles [36].

The orthology relationships among the Ceratocystidaceae GH32 homologs were predicted using phylogenetic criteria [34, 37]. For this purpose, a Maximum Likelihood (ML) phylogeny was constructed with the putative Ceratocystidaceae and Sordariomycetes GH32 members identified in this study, as well as the protein sequences of currently described members of family GH32 in the Carbohydrate-Active enZYmes (CAZY) database (http://afmb.cnrs-mrs.fr/CAZY/), which were obtained from the NCBI database. For this purpose, the sequences were aligned using MAFFT (Multiple sequence alignment

 Table 1 Genome information of the Huntiella, Ceratocystis and Sordariomycetes species included in this study

axon	Assembly size (Mbp)	Number of scaffolds <sup>a</sup>	Number of gene models <sup>b</sup>	References <sup>c</sup>
manginecans	31.71	980	7 494	Van der Nest et al. 2014.
fimbriata	29.40	2 641	7 266	Wilken et al. 2013
albifundus	27.15	939	6 967	Van der Nest et al. 2014.
moniliformis	25.43	365	6 832	Van der Nest et al. 2014.
savannae	28.54	361	7 361	This study
omanensis	31.50	1 638	8 395	Van der Nest et al. 2014.
cremonium alcalophilum v2.0	54.42	13	9 521	JGI
nthostoma avocetta NRRL 3190 v1.0	56.23	786	15 755	JGI
piospora montagnei NRRL 25634 v1.0	47.67	686	16 992	JGI
auveria bassiana ARSEF 2860	33.69	235	10 364	Xiao et al. 2012 Scientific Reports 2
naetomium globosum v1.0	34.90	37	11 124	JGI
lletotrichum graminicola M1.001	51.60	653	12 006	O'Connell et al. 2012 Nat Genet 2012 44:1060–5
lletotrichum higginsianum IMI 349063	49.08	10 235	16 172	O'Connell et al. 2012 Nat Genet 2012 44:1060–5
niochaeta ligniaria NRRL30616 V.1.0	42.38	135	13 657	JGI
ordyceps militaris CM01	32.27	32	9 651	Zheng et al. 2011 Genome Biol 12:R116
yphonectria parasitica EP155 v2.0	43.90	26	11 609	JGI
aldinia eschscholzii EC12 v1.0	37.55	398	11 173	JGI
typa lata UCREL1	54.01	2 334	11 685	Blanco-Ulate et al. 2013 Genome Announc 1:e00390–13
sarium fujikuroi IMI 58289	43.83	12	14 813	Wiemann et al. 2013 PLoS Pathog 9:e1003475
sarium graminearum v1.0	36.45	31	13 322	Cuomo et al. 2007 Science 317:1400-2
sarium oxysporum v1.0	61.36	114	17708	JGI
sarium verticillioides 7600 v1.0	41.78	36	14 188	JGI
ectria haematococca v2.0	51.15	72	15 707	JGI
omerella acutata v1.0	50.04	307	15 777	JGI
omerella cingulata 23 v1.0	58.84	119	18 975	JGI
osmannia clavigera kw1407	29.79	289	8 312	DiGuistini et al. 2011 PNAS 108:2504–9
poxylon sp. CI-4A v1.0	37.70	899	11 712	JGI
onectria sp. v1.0	63.66	325	22 250	JGI
etarhizium acridum CQMa 102	39.42	241	9 849	Gao et al. 2011 PLoS Genet 7:e1001264
etarhizium robertsii ARSEF 23	39.15	176	10 583	Gao et al. 2011 PLoS Genet 7:e1001264
rceliophthora thermophila v2.0	38.74	7	9 110	Berka et al. 2011 Nature Biotech 29:922–927
urospora crassa OR74A v2.0	41.04	20	10785	JGI
urospora discreta FGSC 8579 mat A	176.0	37.3	9 948	JGI
urospora tetrasperma FGSC 2508 mat A v2.0	39.10	81	10 380	Ellison et al. 2011 Genetics 189:55–69
phiostoma piceae UAMH 11346	32.84	45	8 884	Haridas et al. 2013 BMC Genomics 14:373
aeoacremonium aleophilum RPA7	47.47	624	8 834	Blanco-Ulate et al. 2013 Genome Announc 1, e00390–13
odospora anserina S mat+	35.01	7	10588	JGI
diomyces alkalinus v1.0	43.45	29	9 411	JGI
ielavia antarctica CBS 123565 v1.0	40.66	153	9 204	JGI
nielavia appendiculata CBS 731.68 v1.0	32.74	109	11 942	JGI

Table 1 Genome information of the Huntiella, Ceratocystis and Sordariomycetes species included in this study (Continued)

Thielavia arenaria CBS 508.74 v1.0	30.99	69	10 954	JGI
Thielavia hyrcaniae CBS 757.83 v1.0	31.18	251	11 338	JGI
Thielavia terrestris v2.0	36.91	6.00	9 813	Berka et al. 2011 Nature Biotech 29:922–927
Trichoderma atroviride V2.0	36.10	29.00	11 863	JGI
Trichoderma asperellum CBS 433.97 v1.0	40.87	2 282	13 932	JGI
Trichoderma harzianum CBS 226.95 v1.0	40.98	532	14 095	JGI
Trichoderma longibrachiatum ATCC 18648 v3.0	40.87	2 282	13 932	JGI
Trichoderma virens Gv29-8 v2.0	39.00	93	12 427	JGI
Trichoderma reesei v2.0	34.10	89	9 129	JGI
Verticillium dahliae v1.0	33.83	52	10 535	Klosterman et al. 2011 PLoS Pathogens 7: e1002137

<sup>&</sup>lt;sup>a</sup> Number of scaffolds/supercontigs that make up an individual genome assembly

based on fast Fourier transform) v.7.0 (http://mafft.cbrc.jp/alignment/software/) with the L-INS-i option [38]. Motifs that were not present in all of the sequences (e.g., eukaryotic signal motif for extracellular localization and transmembrane motifs for intracellular localisation) were excluded from the alignment. ML analysis was performed using PhyML v.3.0 [39] with the best-fit amino acid substitution model as indicated by ProtTest v.2.4 [40]. The GH32 family ML analysis incorporated the Le-Gascuel (LG) model [41], a proportion of invariable sites (I) and the observed amino acid frequencies (F). Branch support was estimated with PhyML using 1000 bootstrap replicates and the same best-fit models and parameters. Phylogenetic trees were viewed and edited using MEGA v.5 [42].

Gene order (i.e., synteny) and gene structure information was also used to investigate orthology relationships among the Ceratocystidaceae GH32 gene family members. According to Jun et al. [43] orthologous genes typically share homologous neighbouring genes, while non-orthologous genes are typically not flanked by homologous neighbours. Also, orthologous genes will more likely be structured similarly (i.e., share specific domains and introns) than non-orthologous genes [43]. For the gene order analyses, genes and proteins were predicted on all the scaffolds harbouring GH32 gene family members using AUGUSTUS [32]. The predicted genes were then annotated using Blast2GO [44] in the CLC Genomics Workbench 6.0.1 (CLC Bio, Aarhus, Denmark). The sequences of these predicted genes, on each side of the GH32 gene family members, were then used in local BLAST searches in BioEdit. Homology between neighbouring genes was defined as those with blastp and tblastn E-values < 10<sup>-5</sup>. Gene structure similarity was measured using the intron conservation ratio (ICR) between two intron-bearing genes [43]. The ICR between two homologous genes was calculated as the number of positionally homologous introns (i.e., introns that occur at the same position in different genes) divided by the total number of intron positions from the protein alignment [43]. Non-orthologous genes are expected to have ICR-values < 0.5 according to Jun et al. [43].

Finally, OrthoMCL v.2.0.9 [35] was used in an allagainst-all BLAST search, followed by a Markov Cluster analysis to group putative orthologs and paralogs between the *Huntiella* and *Ceratocystis* species. For this analysis, we constructed a sequence database consisting of 43 052 predicted proteins, which consisted of all the AUGUSTUS-predicted proteins for each of the *Huntiella* and *Ceratocystis* species. OrthoMCL was run according to the recommended parameters, with an E-value threshold of 10<sup>-5</sup> [35].

#### Analysis of GH32 gene family evolution

To make inferences regarding GH32 gene family expansions and contractions across the fungi examined in this study, we employed CAFE v.3.1 (Computational Analysis of gene Family Evolution) [45]. For these analyses, the birth (λ) and death (μ) rates were estimated using the lambdamu tool with '-s' option, while the number of gene gains and losses on each branch of the tree was estimated with the '-t' option. The estimated birth and death rates ( $\lambda$  and  $\mu$ ) used in the subsequent analysis were 0.003 and 0.005, respectively. CAFE was run with default parameters of a P-value cut-off of 0.01 (option -p) and the number of random samples used the default value of 1000 (option -r). A time-calibrated Sordariomycetes tree (see below) was used in this analysis where transitions over individual branches were considered significant at P<0.005.

To generate the time-calibrated Sordariomycetes tree needed for the CAFE analysis, the Bayesian Evolutionary

<sup>&</sup>lt;sup>b</sup> Number of the genes predicted for each genome assembly

<sup>&</sup>lt;sup>c</sup> JGI = Joint Genome Institute portal (www.jgi.doe.gov)

Analysis by Sampling Trees (BEAST) package v.2.2.1 [46] was used. For this purpose, we utilized five single copy genes routinely used for phylogenetic analyses [20, 47, 48]. The data (see Additional file 1: Table S1) for the analysis were extracted from the Huntiella and Ceratocystis genomes by performing local tblastn analysis (E-value < 10<sup>-5</sup>) in BioEdit using reference sequences from A. clavatus. These were elongation factor-1 alpha [EF-1a, GenBank:7000001156883129], elongation factor-3 alpha [EF3, GenBank:7000001156847434], minichromosome maintenance complex component 7 [MCM7, GenBank:7000001156824401], RNA polymerase II largest subunit [RPB1, GenBank:XP 001268791] and RNA polymerase II second largest subunit [RPB2, Gen-Bank:XP 001272355)]. These respective gene sequences were also extracted from the representative Sordariomycetes included in the JGI database. The relevant sequences for outgroup taxa in the Dothideomycetes (Alternaria brassicicola, Stagnospora nodrum and Mycosphaerella fijiensis) were also obtained using the JGI portal.

The five protein sequences were aligned with MAFFT as described above and the alignment served as input for a Bayesian tree search with BEAST. A ProtTest analysis suggested the Whelan and Goldman (WAG; [49]) model as the best-fitting evolutionary model for this data. To generate a time-calibrated tree, the analysis was run using the Markov chain Monte Carlo (MCMC) method and four calibration points, which included the Dothideomycetes crown group (mean 350 Million years ago [Mya] with a 95 % credibility interval [CI] of 273–459) [50], the last common ancestor (LCA) of the Hypocreales (181 Mya with a 95 % CI of 150-213) [51], the Clavicipitaceae crown group (117 Mya with a 95 % CI of 95-144) [51], as well as the Nectriaceae crown group (125 Mya with a 95 % CI of 98-155) [51, 52]. The program BEAUTi v.2.0 was used to prepare an xml file to create a starting tree for the BEAST analyses. Priors included the strict molecular clock model with a Yule process for the model of speciation [53]. The standard deviation of all distributions was set to 1.0. Two analyses were run with 10,000,000 generations, sampling data every 1000th generation. The first 15 % of the trees were removed (burn-in) and a consensus of the remaining trees was obtained using LogCombiner and TreeAnnotator [46] and viewed using FigTree v.1.3.1 (http://tree.bio.ed.ac.uk/software/figtree). Tracer v.1.5(http:// beast.bio.ed.ac.uk/Tracer) was used to inspect the chains for convergence, and to ensure that ESS (Effective Sample Size) values exceeded 200 [46].

#### Fot5 analysis

The genomic distribution of *pogo*-like elements, which are homologous to *F. oxysporum* transposase 5 (*Fot5*; [54]) in the Ceratocystidaceae, were investigated, as this

element was located near the GH32 family genes in the genomes of the Ceratocystis species examined. For this purpose, the F. oxysporum Fot5 protein sequence [Gen-Bank: AJ608703] was used in local BLAST searches (tblastn E-value < 10-5) with BioEdit to identify homologs in the Huntiella and Ceratocystis genomes. The conserved DDD catalytic domain of Fot5 (i.e., triad of acidic amino acids [Asp-Asp-Asp or Asp-Asp-Glx] that forms the catalytic pocket for the cleavage of DNA strands) [55] of the homologs identified here, and the previously characterised pogo-like transposons [56] were aligned with MAFFT as described above. This alignment was subjected to ML tree reconstruction using PhyML with the best-fit model parameters (WAG plus gamma to account for among site rate variation) as indicated by ProtTest. Branch support was estimated with PhyML using 1000 bootstrap replicates and the same model parameters.

Whether the Fot5 homologs identified in Ceratocystis have been subjected to repeat-induced point mutation (RIP) was also considered. In filamentous fungi, RIP is a defense mechanism against mobile genetic elements [56] and involves the transition from C:G to T:A nucleotides in pairs of duplicated sequences during meiosis [57]. Therefore, the TpA/ApT ratio across the various Ceratocystis Fot5 sequences was measured. This simple index reflects the frequency of TpA RIP products, and was used as an indication of the RIP response [58]. We also calculated the (CpA + TpG)/(ApC + GpT) index, which considers both the products (TpA) and the targets (CpA and TpG) of RIP [58]. RIPCAL (http://www.sourceforge.net/projects/ripcal) was used to calculate these indices in the aligned Fot5 nucleotide sequences of Ceratocystis.

#### Results

#### Genome sequences

Illumina sequencing of the H. savannae isolate produced a total of 2 884 747 186 bases of trimmed reads with an average length of 85.68 bases (Table 2). The draft genome of this isolate contained 28.54 megabases (Mb) and was made up of 361 scaffolds larger than 500 bases, of which the largest was 1 009 760 bases in length (Table 2). The assembly had an N50 scaffold size of 229 095 bases with a GC content of approximately 47.39 %. The H. savannae draft genome assembly was predicted to encode 7 687 putative ORFs with CEGMA completeness scores of 96.37 % (partial), which is comparable to the draft genomes of H. omanensis (31.5 Mb in size and encodes 8 395 ORFs, [27]), H. moniliformis (25 Mb in size and encodes 7000 ORFs, [26]), C. fimbriata (29.4 Mb in size and encodes 7 266 ORFs), [21], C. albifundus (27.2 Mb in size and encodes 6 967 ORFs, [27]) and

**Table 2** Statistics of the *Huntiella savannae* genome assembly and gene annotations

9	
Summary data	Huntiella savannae
Total reads before trim (bp)	33 168 540
Total reads after trim (bp)	33 055 449
Average length of reads before trim (bp)	101
Average length of reads after trim (bp)	85.68
Number of scaffolds	361
Total sequence length (Mb)	28.54
Largest scaffold (bp)	1 009 760
N50 Scaffold size (bp)	229 095
GC %	47.39
Predicted gene models	7 687
CEGMA <sup>a</sup>	96.37

<sup>&</sup>lt;sup>a</sup> Genome completeness was evaluated using the Core Eukaryotic Genes Mapping Approach (CEGMA) pipeline [38]

C. manginecans (31.7 Mb in size and encodes 7 494 ORFs, [26]).

With the exception of *C. fimbriata*, the Ceratocystidaceae GH32 gene family members were all located on single contigs (C. manginecans: scaffold JJRZ01000038; C. albifundus: scaffold JSSU01001085; H. omanensis: scaffold JSUI01006495; H. savannae: scaffold NODE\_2; H. moniliformis: scaffold JMSH01000004) (see Table 3 for gene locations and sizes). The GH32 genes of C. fimbriata were located on two scaffolds (C. fimbriata: scaffold APWK02000925 and scaffold APWK02000924). To join the two *C. fimbriata* scaffolds, as well as extend the scaffolds harbouring these genes in C. fimbriata and C. albifundus we employed the option 'Map Reads to Reference' in the CLC Genomics Workbench (mismatch cost = 2, insertion and deletion cost = 3, length fraction = 1.0, similarity fraction = 0.9) using scaffold JJRZ01000038 of C. manginecans as a reference. Scaffolds that harboured the GH32 genes in H. savannae and H. omanensis were similarly extended using scaffold JMSH01000004 of *H. moniliformis* as reference. The *C. manginecans* and *H. moniliformis* scaffolds were selected as references because these assemblies were most complete (Table 1) and also had the longest scaffolds that contained the GH32 genes.

#### GH32 gene identification and characterisation

All of the Ceratocystis and Huntiella genomes investigated in this study contained at least one putative member of the GH32 family. For the Ceratocystidaceae, the Huntiella species each contained a single copy of the gene (designated as *HaINV-CW*, *HsINV-CW* and *HmINV-CW* [previously named as CmINV; [28]), while the Ceratocystis species each contained two copies (designated as CaINV-CW, CaINV-V, CfINV-CW, CfINV-V, CmINV-CW and CmINV-V) (Fig. 1; Table 3). The distribution of the GH32 family genes among other Sordariomycetes varied greatly and some taxa lacked a GH32 gene altogether (Fig. 1). For example, 0-4 genes were identified in the Xylariales, 0-12 in the Hypocreales, and 0-3 in the Sordariales and Glomerellales. As expected [2], the plant pathogens generally contained more GH32 genes (e.g., 12 in F. oxysporum; and 6 in each of A. terreus, Talaromyces stipitatus and Nectria haematococca). These genes also appeared to be absent from insect pathogens and plant pathogens that evolved from insect pathogens (e.g., Cordycipitaceae and Clavicipitaceae) [47].

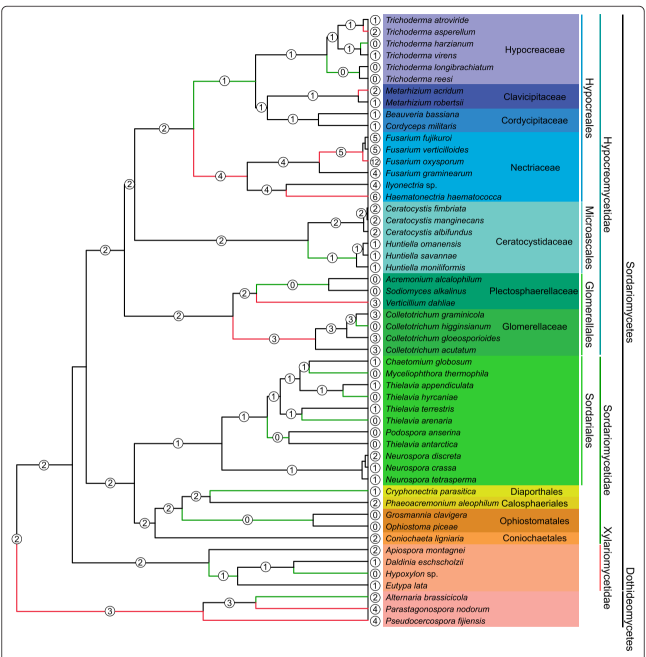
Among the examined Ceratocystidaceae GH32 family members, the InterProScan and SignalP analyses identified a conserved N-terminal (IPR013148) (Fig. 2). These analyses also identified a less conserved C-terminal (IPR013189) that is likely essential for overall protein stability ([59]; Fig. 2). These sequences, including those identified for the Ceratocystidaceae, also contain the eight well-conserved domains (A-G) and three highly conserved acidic residues characteristic of GH32 gene family members ([60]; Fig. 2). The latter include an

Table 3 GH32 family members identified in *Huntiella* and *Ceratocystis* 

	)			
Gene Name	Taxon <sup>a</sup>	Genomic Location <sup>b</sup>	Length (bp)	Length (aa)
CmINV_V	Ceratocystis manginecans	JJRZ01000038:33284-34837	1945	627
CmINV_CW	Ceratocystis manginecans	JJRZ01000038:36427-37962	1952	625
CfINV_V	Ceratocystis fimbriata	APWK02000925:4324-5817	1945	627
CfINV_CW	Ceratocystis fimbriata	APWK02000924:3-1490	1938	625
CalNV_V	Ceratocystis albifundus	JSSU01001085:20903-22444	1829	627
CalNV_CW	Ceratocystis albifundus	JSSU01001085:17498-19033	1952	625
HmINV_CW	Huntiella moniliformis	JMSH01000004:78747-80591	1848	615
HaINV_CW	Huntiella savannae	NODE_2:199823- 201667	1848	615
HoINV_CW	Huntiella omanensis	JSUI01006495:45419-47263	1848	615

<sup>&</sup>lt;sup>a</sup> The GenBank accesssion numbers for the *C. manginecans, C. fimbriata* and *C. albifundus* genomes are JJRZ01000000, APWK00000000 and JSSU00000000. Those for the *H. moniliformis, H. savannae* and *H. omanensis* genomes are JMSH00000000, LCZG00000000 and SUl00000000

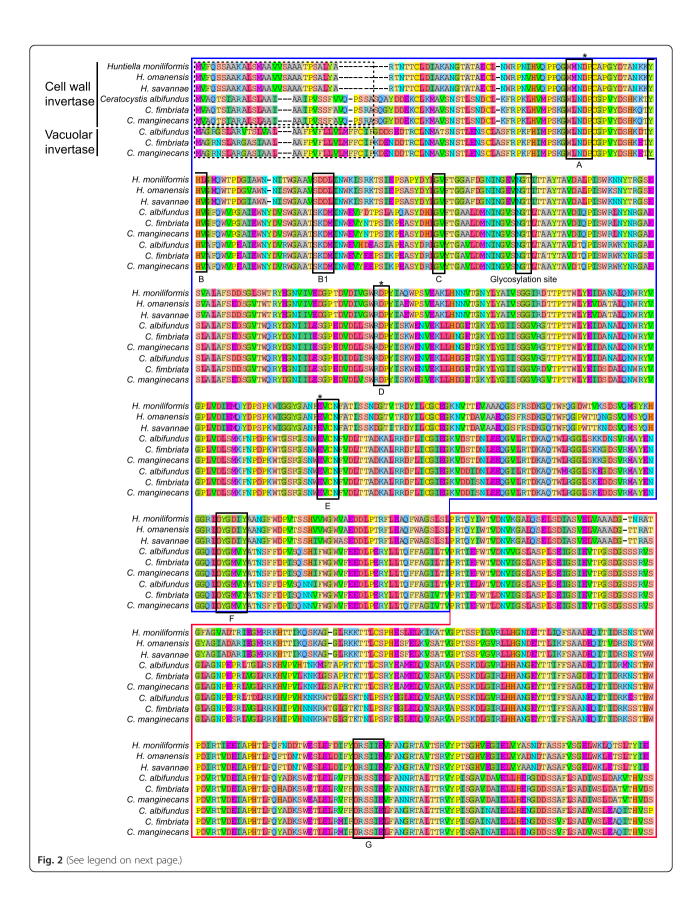
<sup>&</sup>lt;sup>b</sup> The name of the contig is followed by the nucleotide position of the gene within the contig



**Fig. 1** Glycoside hydrolase 32 (GH32) gene family expansions and contractions mapped onto the Sordariomycetes chronogram. Significant (*P*< 0.05) expansions (indicated with red lines) and contractions (indicated with green lines) were inferred using CAFE v3.1 (Computational Analysis of gene Family Evolution) [45]. The probable ancestral gene family size for each node is indicated within white circles, while the family sizes in extant species are indicated at the tips of terminal branches. Colour-coding designates the Sordariomycetes taxa to either order or family level. The chronogram was inferred in this study (see Additional file 2: Figure S1). The sequences from Dothideomycetes were used for outgroup purposes

aspartic acid located in the WMNDPNG motif (also known as  $\beta$ -fructosidase motif or sucrose-binding box) of domain A that acts as a nucleophile, and an aspartic acid located in the RDP motif of domain D that acts as a transition-state stabiliser, as well as a glutamic acid located in the EC motif of domain E that acts as the acid/

base catalyst [3]. Compared to the NG present in the WMNDPNG motif of other fungi [57], the *Huntiella* GH32 genes contain a CA, while those of the *Ceratocystis* genes contain a CG. NetNGlyc analysis also revealed that all of the Ceratocystidaceae GH32 genes contained a potential N-glycosylation site. This indicates that the

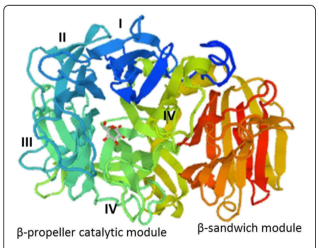


(See figure on previous page.)

**Fig. 2** Alignment of the conserved motifs of the glycoside hydrolase 32 (GH32) enzymes. These include conserved regions (labelled A-G) and various amino acids (shown with black stars). The N-terminal β-propeller module (indicated in the blue block) and the C-terminal β-sandwich module (indicated in the red block) are also highlighted. The translated sequences of one group of the *Ceratocystis* GH32 gene possess a transmembrane domain (shown with dotted lines) characteristic of vacuolar invertases [5], while the translated sequences of the other *Ceratocystis* GH32 gene and the *Huntiella* GH32 gene possess an eukaryotic secretion signal (shown with dotted lines) needed for secretion [60]

genes identified in these fungi encode a conserved asparagine residue that is predicted to attach to a glycan chain to facilitate various co- and post-translational modifications and enhance the stability, transport and secretion of proteins [61, 62]. The inferred 3D structures of the proteins encoded by Ceratocystidaceae GH32 genes further confirmed the presence of the five-bladed  $\beta$ -propeller catalytic module at the N-terminal, as well as the presence of two six-stranded  $\beta$ -sheets composed of antiparallel  $\beta$ -strands forming a sandwich-like fold at the C-terminal domain (Fig. 3).

The SignalP analyses showed that parts of the inferred amino acid sequences of the *Huntiella* genes (i.e., the first 28 residues encoded by *HaINV-CW*, *HsINV-CW* and *HmINV-CW*), as well as one of the *Ceratocystis* homologs (i.e., the first 31 residues encoded by *CaINV-CW*, *CfINV-CW* and *CmINV-CW*) are comprised of a eukaryotic secretion signal. This suggests an extracellular localisation for the proteins, which is typical of cell wall invertases [16]. These analyses also predicted possible signal peptide cleavage sites between amino acids 25 and 26 for the *Huntiella* homologs and between



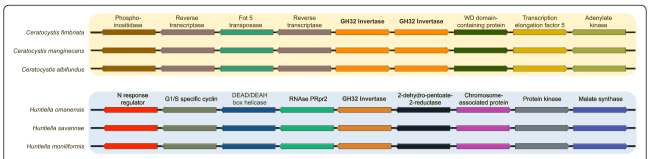
**Fig. 3** 3D structure of the *C. manginecans* invertase (CmINV-CW). Roman numerals (I–V) show the five blades of the  $\beta$ -propeller module, while the C-terminal  $\beta$ -sandwich module is indicated in dark red. These structures were inferred with the Swiss-Model Web server (http://www.expasy.org/swissmod/SWISS-MODEL.html) by making use of a fructosyltransferase from *Aspergillus japonicus* (PDB id: 3lfi.1) as template

residues 19 and 20 for the one Ceratocystis homolog (Fig. 2). However, the second homolog of the gene in Ceratocystis species lacked the N-terminal signal sequence. Instead, parts of the translated sequences of this gene (i.e., the first 32 residues encoded by CaINV-V, CfINV-V and CmINV-V) comprised a transmembrane region, which is characteristic of vacuolar invertases [5] suggesting an intracellular localisation for the protein. Our analysis also suggested that this homolog adopts the NinCout configuration that consists of a short Nterminal segment in the cytosol and a long C-terminal region in the vacuole, which is typical of MEnM of type II single-pass membrane proteins [5]. We therefore classified the Ceratocystidaceae GH32 gene family homologs as either cell wall invertases (with a CW suffix to gene and protein names; for the Huntiella homologs and one group of homologs in Ceratocystis), or as vacuolar invertases (with a V suffix to gene and protein names; for the second homolog in Ceratocystis).

The SignalP analyses of GH32 gene family members in the other Sordariomycetes showed that genes belonging to the groups designated by Parrent et al. [2] as extracellular invertases contained the eukaryotic secretion signal motif. In contrast, this motif was absent from genes that belonged to the groups they designated as intracellular invertases. Indeed, previous molecular and biochemical studies have shown that the eukaryotic secretion signal motif is present in genes encoding extracellular invertases and absent from genes encoding intracellular invertases [63, 64]. Except for the three *Ceratocystis* genes (i.e., *CaINV-V*, *CfINV-V* and *CmINV-V*), none of the other Sordariomycetes GH32 genes contained the transmembrane motif, which is characteristic of vacuolar invertase genes.

#### GH32 orthology relationships

Gene order analysis of the *Ceratocystis* scaffolds harbouring GH32 family members revealed that the cell wall and vacuolar invertase genes are located adjacent to each other in all three of the species studied. However, the *Huntiella* cell wall invertase gene is located at a different genomic region when compared to that of *Ceratocystis* (Fig. 4). This was confirmed using the gene order analysis, where homologous flanking genes (tblastn, Evalues  $< 10^{-5}$ ) were only obtained for the within-genus comparisons. Genes encoded on the examined scaffolds,



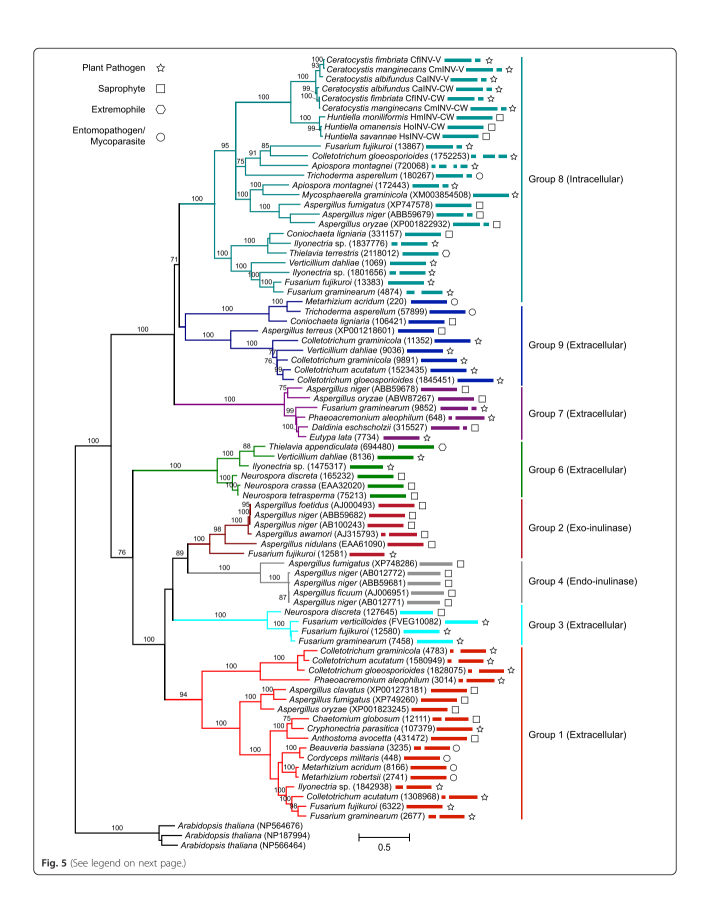
**Fig. 4** The predicted genes flanking the Glycoside hydrolase 32 (GH32) gene family members in *Huntiella* and *Ceratocystis*. Genes present on the scaffolds harbouring the putative invertases were predicted using AUGUSTUS [32] and annotated using Blast2GO [44]. Note that the genes are not drawn to scale. The *Huntiella* GH32 family gene is flanked by putative G1/S-specific cyclin Pcl5 (*Colletotrichum orbiculare*, ENH86823), RNAse P Rpr2/Rpp21 subunit domain-containing protein (*Gaeumannomyces gramini*, EPQ63823), Malate synthase-like protein (*Acremonium chrysogenum*, XP003651419), serine/threonine-protein kinase (*Metarhizium acridum*, EFY93082.1), nitrogen response regulator (*Colletotrichum gloeosporioides*, ELA29612.1), DEAD/DEAH box helicase (*Colletotrichum sublineola*, KDN64774), 2-dehydropantoate 2-reductase (*Colletotrichum gloeosporioides*, EQB48758), and structural maintenance of chromosomes 5 (*Villosiclava virens*, KDB17190) genes. The two *Ceratocystis* GH32 family genes were flanked by putative Phosphatidylinositol-specific phospholipase (*Metarhizium anisopliae*, KFG82763), putative WD domain-containing protein (*Togninia minima*, EOO00810.1), reverse transcriptases (*Sclerotinia sclerotiorum*, XP\_001588999 and *Blumeria graminis*, CCU77161), transcription elongation factor 5 (*Scedosporium apiospermum*, KEZ42236), adenylate kinase (*Magnaporthe oryzae*, XP003716198), and *Fot5* transposase (*Colletotrichum gloeosporioides*, ELA33194.1) genes

other than the GH32 family genes, had homologs elsewhere in the genomes of the two fungi (e.g. the *Ceratocystis* scaffolds harboured various putative reverse transcriptase genes, which were also present on scaffolds other than the one harbouring the GH32 gene in the *Huntiella* genomes). An exception was for the *Fot5* transposase genes that were only present in the genomes of the *Ceratocystis* species (see below). According to Jun et al. [43], such an observed lack of synteny points towards a non-orthologous relationship between the GH32 genes of *Ceratocystis* and *Huntiella*.

Analysis of gene and protein structures of the Ceratocystidaceae and Sordariomycetes GH32 family members revealed that coding sequences were interrupted by introns that vary greatly in number and distribution across all of the taxa examined in this study (Fig. 2). For example, the *Huntiella* genes (consisting of 1848 bases and encoding 615 aa) did not harbour any introns, while both the *Ceratocystis* genes (consisting of 1945–1952 bases and encoding 625–627 aa) contained a single intron at the same position (Table 3). This corresponded to an ICR of 1 for the *Ceratocystis* GH32 family members, and an ICR value of 0 for the Ceratocystidaceae GH32 family members. According to Jun et al. [43], the latter ICR value indicates non-orthology between the GH32 genes of *Ceratocystis* and *Huntiella*.

The ML phylogeny revealed that the *Huntiella* and *Ceratocystis* GH32 genes grouped with known members of this protein family (Fig. 5). The *Ceratocystis* vacuolar invertases formed part of a well-supported clade previously designated as Group 8 [2], which include invertases with intracellular localisation and that lack signal peptide cleavage sites (Fig. 5). Despite the presence of

signal peptides for extracellular localisation, however, the Huntiella and Ceratocystis cell wall invertases also formed part of Group 8. Within this clade, the Ceratocystidaceae genes grouped according to their evolutionary relationships (i.e., the two genes in Ceratocystis were more closely related to each other than to the gene in Huntiella). Within Ceratocystis, the cell wall invertases formed a sister group to the vacuolar invertases; and within each of these sister groups, the relationships among the genes matched the known relationships among species, with the sequences of C. fimbriata and C. manginecans grouping together and C. albifundus at their base. The same was also true for the Huntiella cell wall invertase genes. Therefore, CaINV-V, CfINV-V and CmINV-V are orthologs, CaINV-CW, CfINV-CW and CmINV-CW are orthologs and HaINV-CW, HsINV-CW and HmINV-CW are orthologs (Fig. 6) [34]. The Ceratocystis GH32 genes represent co-orthologs of the Huntiella genes (i.e., the two Ceratocystis GH32 genes are collectively orthologous to the Huntiella GH32 gene due to a lineage-specific duplication in the former, Fig. 6) [34]. Because the duplication that gave rise to the two Ceratocystis GH32 genes occurred in the ancestor of this genus, the cell wall and vacuolar invertase genes of these species represent outparalogs (i.e., paralogous genes derived from a gene duplication event that precedes lineage radiation [34], Fig. 6). These orthology relationships were consistent with the results of the OrthoMCL analysis. Therefore, the non-orthology of the GH32 genes in Ceratocystis and Huntiella, suggested by the results of the synteny and ICR analyses, likely reflects the involvement of retrotransposition in the evolution of these genes (see below).



(See figure on previous page.)

**Fig. 5** Maximum likelihood phylogeny of the Sordariomycetes Glycoside Hydrolase 32 (GH32) gene family. Representative sequences of the 8 groups that span the fungal GH32 gene phylogeny [2] were included in this study. GenBank accession numbers or sequence identifiers from genome projects for each of these sequences are provided in parentheses. Percentage bootstrap support (based on a 1000 repeats) is indicated at the internodes. The exon-intron structure of the genes is diagrammatically indicated next to each taxon where gaps within solid lines indicate intron positions. Colour-coding designates the groups previously identified [2]. The sequences from *Arabidobsis thaliana* were used for outgroup purposes

#### GH32 gene family evolution

BEAST and CAFE analyses were used to identify and estimate the relative ages of the losses/gains of the GH32 family genes in several orders and families in the Sordariomycetes, including Ceratocystidaceae (Fig. 1, Additional file 2: Figure S1). The ESS-values for the BEAST analysis parameters were higher than 200, which is the recommended threshold for ensuring appropriate estimation of the posterior distribution of each parameter [46]. As expected from the analysis, the root node that represents the divergence of the Sordariomycetes and Dothideomycetes was

around 362 Mya (with CI of 346–377 Mya) [51–65]. Based on these data, the estimated divergence time for the LCA of *Huntiella* and *Ceratocystis* was *ca*. 62 Mya (with CI of 50–70 Mya).

The CAFE analysis identified several gene loss and gain events in the GH32 gene family (Fig. 1). Many of these were inferred to be lineage-specific, which included significant expansions (e.g., *F. oxysporum* with 12 gene copies and *N. haematococca* with 6 gene copies) and contractions (e.g., *Hypoxylon* sp., *Thielavavia arenaria*, *Myceliophthora thermophila*, and *Colletotrichum higginsianum* all lacking GH32 family members) at the

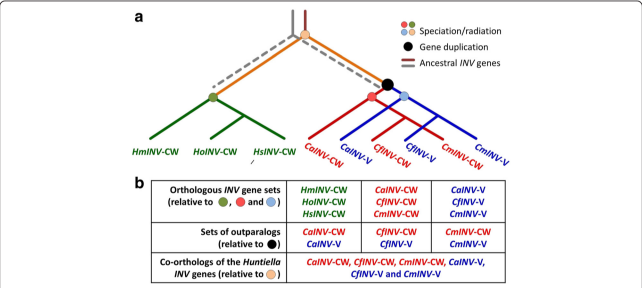


Fig. 6 The inferred evolutionary history of the Ceratocystidaceae Glycoside hydrolase 32 (GH32) gene family and the orthology relationships among these genes. a The evolutionary tree shows nine homologous genes from six species (A). The Huntiella cell wall invertase genes are depicted as HsINV-CW, HmINV-CW and HoINV-CW, while the Ceratocystis vacuolar invertase genes are depicted as CaINV-V, CmINV-V and CfINV-V and those encoding the Ceratocystis cell wall invertases as CaINV-CW, CmINV-CW and CfINV-CW. As indicated by CAFE, the genome of the Ceratocystidaceae ancestor likely encoded two invertase (INV) genes, one of which (depicted by the grey line) was subsequently lost from both the Ceratocystis and Huntiella lineages (depicted by grey broken line) before the radiation of species. However, the remaining invertase gene (depicted in orange) was duplicated in the Ceratocystis ancestor resulting in the two invertase genes encoded by the genomes of the extant species. This duplication was also reconstructed using NOTUNG 2.6 which detects duplications based on gene tree to species tree reconciliation [89] (results not shown). All of the invertase genes in the extant Ceratocystis and Huntiella species thus evolved from the same ancestral gene in the Ceratocystidaceae ancestor (depicted by the orange line). The respective Ceratocystis genes each evolved through vertical decent after their emergence (i.e., gene duplication) in the last common ancestor. **b** Following the standard nomenclature for duplicated genes (reviewed by Koonin [36]), the Huntiella cell wall invertase genes share an orthologous relationship (i.e., orthologs are related via speciation and are derived via vertical decent from the common ancestor). The same is also true for the respective cell wall and vacuolar invertase genes of Ceratocystis, where each represent a set of orthologs. Because the duplication that gave rise to the Certocystis genes occurred before radiation of this genus, the Ceratocystis cell wall and vacuolar invertase genes represent outparalogs (i.e., homologs that derive from a gene duplication event that precedes lineage radiation/speciation) [36]. However, all of the Ceratocysistis invertase genes represent co-orthologs of the gene in Huntiella. This is because the lineage-specific duplication in Ceratocystis gave rise to a set of genes that are collectively orthologous to those of Huntiella [36]

tips of branches. At deeper phylogenetic levels, significant expansions were predicted for branches leading to the Nectriaceae and the outgroup taxa in the Dothideomycetes, while significant contractions were predicted for branches leading to the Sordariales, Ophiostomatales, Xylariales, as well as the branch leading to Hypocreaceae, Clavicitpitaceae and Cordycipitaceae. Among the Ceratocystidaceae, a GH32 family contraction was predicted for the Huntiella species (ca. 62 Mya). Other GH32 family contractions and expansions in the Sordariomycetes predicted for the first time in the current study include an expansion on the branch leading to the Glomerellaceae and an expansion on the branch leading to the Nectriaceae, as well as a contraction on the branch leading to the Hypocreaceae-Clavicipitaceae-Cordycipitaceae clade.

#### Fot5 analysis

Local BLAST searches with the F. oxysporum Fot5 sequence revealed that this gene family is apparently absent from the Huntiella genomes, while the Ceratocystis genomes harbour numerous Fot5 homologs (Additional file 3: Table S2). Phylogenetic analysis of the 202 sequences (i.e., 60 from C. fimbriata, 19 from C. albifundus and 106 from C. manginecans, as well as 17 previously characterised pogo-like transposon sequences) spanning more than 75 % of the DDD catalytic domain of Fot5, confirmed that most of these sequences indeed represent putative Fot5 homologs (Fig. 7 Additional file 4: Figure S2). The identified Ceratocystis Fot5 sequences formed a monophyletic group with the known Fot5 sequence from F. oxysporum with high bootstrap support (81 %) (Fig. 7 Additional file 4: Figure S2). However, some sequences of Ceratocystis also clustered together with the other Fot family members: three copies of C. fimbriata clustered with Fot2, two copies of C. fimbriata and three copies of C. manginecans clustered with Pot3 and SCSCL. Several groups of identical and closely related Fot5 homologs were detected, where homologs belonging to the same species and homologs belonging to different species often grouped together.

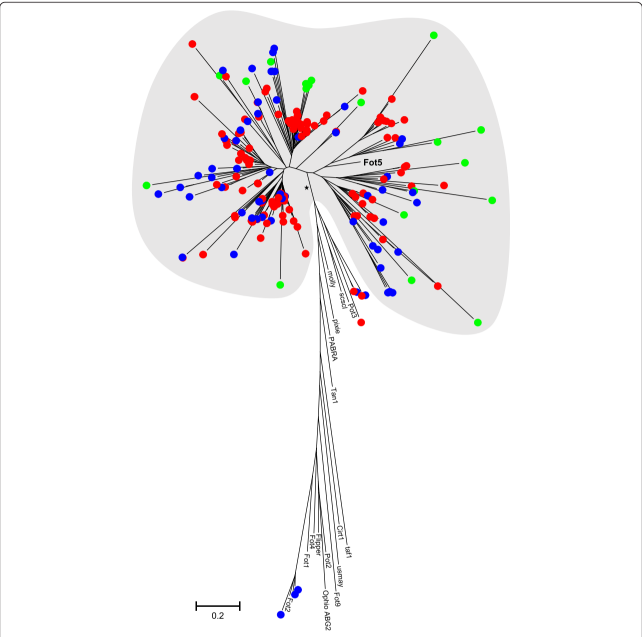
The putative *Fot5* homologs identified in the *Ceratocystis* genomes displayed the hallmarks of RIP. Overall, the *Fot5* sequences had TpA/ApT index values above 1 (1.5 for *C. albifundus*, 1.3 for *C. fimbriata* and 1.5 for *C. manginecans*), possibly due to the introduction of C:G to T:A mutations [58]. The *Fot5* sequences also had lower (CpA + TpG)/(ApC + GpT) index values (1.2 for *C. albifundus*, 1.1 for *C. fimbriata* and 1.3 for *C. manginecans*), indicating a possible RIP response [58]. Analysis of individual sequences revealed a mixture of RIPped and non-RIPped copies, with 56 % of the *C. albifundus Fot5* homologs, 35 % of the *C. fimbriata Fot5* homologs and 32 % of the *C. manginecans Fot5* homologs having TpA/ApT ratios of

>1 and A + T richness > 55 % [56]. According to Dufresne et al. [56] this is indicative of a mild RIP response, allowing the presence of potentially active *Fot5* copies.

#### **Discussion**

All of the identified Ceratocystidaceae invertase genes and inferred proteins carry hallmarks of the GH32 gene family and were considered homologs. They all have an N-terminal catalytic domain and a C-terminal βsandwich domain needed for structural stability [9]. They also contained three conserved residues (i.e., two aspartates and one glutamate) referred to as 'the catalytic triad' (see Fig. 2), which are indispensable for binding and catalysis [3, 5]. For example, it was suggested that the aspartate present in the RDP-motif provides hydrogen bonds to bind the C3 and C4 hydroxyls of fructose [3]. Although the WMNDPNG-motif present in the Ceratocystidaceae invertases is not fully conserved, they do contain the two critical amino acids (W and N) needed for transfructosylation [66]. Typical of vacuolar and cell wall invertases, all of the Ceratocystidaceae sequences also contained an N-glycosylation site where a glycan chain can potentially attach to an asparagine residue of the acceptor proteins [67]. Given these commonalities with other GH32 enzymes, it is likely that the invertases encoded by the Ceratocystidaceae represent active enzymes with sucrolytic activities. Thus far, heterologous expression of the HmINV-CW gene of H. moniliformis in S. cerevisiae yielded an active invertase that allowed the mutant yeast to utilize sucrose as sole carbohydrate source [28]. However, further studies are required to determine if both the vacuolar and cell wall invertase genes identified in this study are functional in all of the Ceratocystidaceae that harbour them.

Most functional studies of fungal cell wall invertases have focused on industrial applications [14, 68], and very little is known regarding the biological functions of these enzymes. It is possible that the cell wall and vacuolar invertases of Huntiella and Ceratocystis may enable colonization of plant tissue by facilitating uptake and transport of plant-derived sucrose [62]. Previous studies have shown that during plant-fungus interactions, both partners contribute to the overall invertase activity [69]. Plants use invertases for sugar signalling linked to stress and defence responses in addition to nutrition, whereas, fungal invertases convert extracellular and intracellular sucrose to fructose and glucose, and ensure the availability of nutrients during infection [70-72]. These enzymes may also be involved in glucose signalling that may influence fungal virulence [73]. In these fungi, vacuolar invertases may streamline sucrose utilization, especially if the sucrose-cleaving activity becomes rate-limiting for provision of sugars to the fungus during infection [71]. The functional expression of GH32 enzymes in



**Fig. 7** Maximum likelihood phylogeny of the *Fot5* DDD catalytic domain. This analysis was done using the WAG substitution model [49] and gamma correction to account for among site rate variation. The *Ceratocystis Fot5* sequences are included in the grey area and indicated according to species (green dots = *C. albifundus*, blue dots = *C. fimbriata*, red dots = *C. manginecans*). The branch labelled with an asterisk received 81 % bootstrap support based on the analysis of 1000 pseudoreplicates (see Additional file 4: Figure S2 for full information regarding bootstrap support for the tree, as well as the sequence identifiers of putative *Ceratocystis Fot5* homologs and Additional file 3: Table S2 for their genomic coordinates). GenBank accession numbers or for previously identified *Fot5* homologs are: Fot2 [Genbank:JN624854, *F. oxysporum*], *Fot5* [Genbank:CAE55867, *F. oxysporum*], Fot1 [Genbank:X64799, *F. oxysporum*], Fot4 [Genbank:AF076632, *F. oxysporum*], Fot9 [JGI:2517, *F. graminearum*], Fot9 [Genbank:CAG33729.1 *Yarrowia lipolytica*], Molly [Genbank:CAD32687, *Parastagonospora nodorum*], Ophio [Genbank:ABG26269, *Ophiostoma novo-ulmi*], PABRA [Genbank:ACY56713, *Paracoccidioides brasiliensis*], Pixie [Genbank:CAD32689, *Parastagonospora nodorum*], Pot2 [Genbank:CAA83918, *Magnaporthe grisea*], Pot3 [Genbank:AAC49418, *M. grisea*], SCSCL [Genbank:XP001592252, *Sclerotinia sclerotiorum*], Taf1 [Genbank:AAX83011, *Aspergillus fumigatus*], Tan1 [Genbank:US9946, *Aspergillus awamori*] USMA [Genbank:UM03882, *Ustilago maydis*), Flipper [Genbank:AAB63315, *Botryotinia fuckeliana*] and Cirt1 [Genbank:XP710204, *Candida albicans*]

interactions between Ceratocystidaceae and their plant hosts and substrates, should be investigated to provide insights into the potential role this gene family plays in the infection biology and pathogenesis of this group of fungi.

To the best of our knowledge, these are the first vacuolar invertases identified in fungi. It is conceivable that gene duplication followed by functional divergence of the outparalogs gave rise to the two types of invertases in the Ceratocystidaceae (see Fig. 6). In fact, gene duplication followed by functional divergence have been shown to be important drivers of the evolution of GH families [74]. For example, small changes in the primary structure of GHs can result in changes to their substrate specificities [75], while changes at their N-terminals might influence cellular localisation [8]. Such changes at the N-terminal could have allowed for the evolution of the Ceratocystidaceae cell wall invertases from ancestral Group 8 intracellular invertases. Consistent with this view, the cell wall invertases of Ceratocystis and Huntiella both contain eukaryotic signal sequences for directing proteins into the endoplasmic reticulum for secretion [5, 76]. It is also consistent with previous predictions that HmINV-CW in H. moniliformis represents an extracellular invertase [28]. In turn, the vacuolar invertase of Ceratocystis could have evolved from a cell wall invertase as has previously been suggested for plant invertases [5]. Such a process would be facilitated by the loss of the eukaryotic secretion signal sequence and acquisition of signature motifs, which in plants allow for localisation to the lytic vacuole [5]. Indeed, structural analysis suggested that the putative vacuolar invertases of Ceratocystis adopt the characteristic NinCout configuration of type II single-pass membrane proteins that are targeted to vacuoles [5]. These data, together with the results of our phylogenetic analysis, strongly suggest that the evolution of the two invertase outparalogs in Ceratocystis involved divergence from a common ancestor by the loss and gain of motifs at their N-terminals to ultimately yield a cell wall and a vacuolar invertase.

The evolutionary history of the GH32 gene family in the Ceratocystidaceae was studied in CAFE by reconstruction of ancestral states across the Sordariomycetes. This approach involves an evaluation of the probabilities of changes in family size (i.e., gene copy number expansions and contractions) from "parent to child nodes" in a time-calibrated phylogeny [77]. The CAFE analysis showed that the LCAs of most of the Sordariomycetes orders, as well as the subclass Hypocreomycetidae, likely encoded two GH32 genes (i.e., a gene family size of two represents the ancestral or plesiomorphic state for these groups) (see Fig. 1). This was also true for the Ceratocystidaceae, where the only significant transition (a contraction) in GH32 gene family size occurred approximately

62.0 Mya in the LCA of Huntiella. However, based on the GH32 gene phylogeny, the Ceratocystidaceae invertases represent a nested and monophyletic cluster within GH32 Group 8, suggesting that all of the invertases in this fungal family evolved from a single ancestral gene (i.e., the Ceratocystis genes are collectively coorthologous to the Huntiella GH32 gene). The most parsimonious explanation for these findings is therefore that the evolution of the Ceratocystidaceae GH32 gene family involved the loss of one of the two ancestral genes predicted by CAFE (i.e., one of the two GH32 genes predicted to have been encoded by the LCA of the Ceratocystidaceae was lost from both the Ceratocystis and Huntiella lineages) (Fig. 6). On the Huntiella branch, the remaining gene gave rise to the extant GH32 gene in this genus. In the LCA of Ceratocystis, a lineagespecific duplication of the remaining ancestral gene gave rise to the two GH32 genes of the extant species (Fig. 6). This duplication in the LCA of Ceratocystis also established a membership of two for its GH32 gene family. This superficially resembles the inferred ancestral state for the overall family, but the data clearly showed that the extant condition of having two GH32 genes emerged in the LCA of Ceratocystis, thus indicating that it represents the synapomorphic state for the genus.

The GH32 gene duplication in the Ceratocystis LCA likely allowed for the acquisition of novel invertase activities. A classic view popularized by Ohno [78], is that gene family expansions associated with gene duplications are the principal source of new genes that acquire new functions. This is because duplication creates a redundant gene copy that is free from selection and that can evolve a new function (i.e., neofunctionalization). It is therefore possible that following the gene duplication, relaxed selection allowed for the acquisition of novel domains by the GH32 paralogs. During this process, one of the Ceratocystis paralogs likely acquired the transmembrane region characteristic of vacuolar invertases, while the other acquired the eukaryotic signal motif characteristic of cell wall invertases. Based on the results of our ML and CAFE analyses, the evolution of the Huntiella GH32 gene followed a parallel evolutionary trajectory during which it independently acquired its eukaryotic signal motif.

As have been demonstrated for other Ascomycetes [2], the data presented here suggested a link between the ecological strategy of Ceratocystidaceae and GH32 gene family size. In fungi, changes in the repertoire of GH32 functional products are thought to influence the efficiency at which sucrolytic compounds are exploited [79]. In the Hypocreales, for example, the respective GH32 family expansions and contractions appear to be linked to the evolution of the Nectriaceae with their plant pathogenic lifestyles [80], and to that of the

Cordycipitaceae-Clavicipitaceae clade that are often insect pathogens or have undergone a host jump from insects to plants [81]. The evolution of the Glomeralles also appeared to be associated with such changes in the GH32 family, where a significant contraction was observed at the base of the Plectosphaerellaceae with its alkaliphilic representatives [82], while the Glomerellaceae clade with its plant pathogens [83] were associated with several significant expansions. Plant associated fungi likely adapted to hosts through a larger repertoire of invertases that allow these species to access plantsynthesised sucrose [2]. This might be the case for Ceratocystis species with their two GH32 invertases. On the other hand, restrictions in functional invertase repertoires (e.g., in the saprophytic Huntiella) might be important for exploiting niches with limited sucrose resources, as well as for potentially avoiding plant defence mechanisms, thus conferring the ability to colonise plant-associated niches [84]. Although the apparent link between GH32 gene family size and the ecology of the Ceratocystidaceae is consistent with the results of previous studies [2, 75], additional work is needed to fully understand the role(s) of GHs or carbohydrases available to these fungi in determining their ecological capabilities.

Similar to previous studies, results of this study suggest that transposon-like elements may have played a role in the evolution of the Ceratocystidaceae GH32 invertases. For example, retrotransposonlike elements that are part of Class I transposable elements (TEs) [5] have been used to explain why the number of introns differ between certain groups of plant invertases [5]. Local synteny information and intron conservation ratios indicated that the Huntiella invertase might represent a retrotransposed copy of the ancestral gene (i.e., the ancestral GH32 gene that gave rise to all of the Ceratocystidaceae genes examined here). Similar to what has been shown for other retrotransposed gene copies [43], the Huntiella invertase genes lack introns, and the genomic region containing them appears to be non-homologous to the invertase gene-bearing genomic region of Ceratocystis (i.e., the GH32 genes of these two genera are flanked by completely different sets of genes). Retrotransposons facilitate intron loss/gain via a copy and paste mechanism involving, first, reverse transcription of messenger RNA (mRNA) into complementary DNA (cDNA), followed by homologous recombination between the original gene (or a homolog) and cDNA [55]. Therefore, as have been suggested for Oryza and *A. thaliana* [5], the activity retrotransposon-like elements in the genomes of the Ceratocystidaceae and its ancestors could have been responsible for or involved in the initial loss of one of the two ancestral GH32 genes predicted for the Ceratocystidaceae, and the subsequent duplication in the LCA of *Ceratocystis*.

Another group of transposon-like elements that could have influenced the evolution of the Ceratocystidaceae invertases is the Fot5 or pogo-like elements (Class II of TEs; also referred to as DNA transposons). Fot5 utilizes a 'cut-and-paste' mechanism for transpositioning, during which a specific DNA region is excised and inserted into a target site elsewhere in the genome [85]. The activity of Fot5 in Ceratocystis may thus have given rise to genomic rearrangements that also affected the region harbouring the two GH32 invertase genes. In fact, the apparent abundance of Fot5 homologs in the genomes of the Ceratocystis species and the presence of short terminal branches on the Fot5 phylogeny suggests that these elements were active relatively recently [56]. Our Fot5 phylogeny further suggests that many Fot5 elements were active in the ancestral lineages of Ceratocystis (i.e., homologs from different Ceratocystis species group together in a cluster), while others were active after speciation (i.e., homologs represent unique Fot5 lineages or group according to species) [56]. Analysis of the Ceratocystis Fot5 elements also showed that their lifestyles most likely match those of other TEs and parasitic DNA elements [85]. Once inside the genome of the fungal individual, the Fot5 element likely increased in copy number and persisted until all its copies become inactive due to either vertical inactivation by the TE itself [86] or host-associated mechanisms that protect the genome from parasitic DNA elements (e.g., RIP) [55, 85]. Indeed, our analysis of the Fot5 elements suggested a possible RIP response in Ceratocystis. Over time, these inactivated copies will degenerate further through mutation and genetic drift, until no identifiable remnants of the original TE remain in the genome [85]. The fact that none of the three Huntiella genomes harboured detectable Fot5 elements thus suggests that the lineage never harboured these TEs, and if they were present they have degenerated to a point where standard in silico tools can no longer detect them.

An important hypothesis emerging from this study is that the activity of *Fot5* elements facilitated assembly of a genomic region or island key to the ecological success of *Ceratocystis* species. In addition to the two GH32 invertase genes, this genomic region encodes various other genes potentially involved in the ability of this taxon to infect and colonize health woody and herbaceous plants. In *Fusarium*, the genomic regions harbouring *Fot5* elements are commonly associated with strain- or species-specific regions that are enriched for genes involved in pathogenicity and/or adaptation [87]. Virulence genes in other pathogens are also often found in genomic regions dense with TEs where the genomic plasticity associated with

these elements is believed to contribute to the evolution of virulence and pathogenicity related genes [88]. The GH32-bearing genomic region identified in *Ceratocystis* may therefore represent a key target for future studies into the molecular basis of the ability of these fungi to cause plant disease. Also, further investigation of the diversity and evolution of *Fot5* and other TEs will undoubtedly provide valuable clues regarding gene and genome evolution in the Ceratocystidaceae with their diverse ecologies, modes of reproduction and potential biotechnological benefits.

#### **Conclusions**

In this study, we considered the capacity of Ceratocystidaceae and a selection of Sordariomycetes species to utilize sucrose by GH32 invertase enzymes. The publicly available genome sequences for these taxa, and the H. savannae genome sequenced here, were used to identify novel GH32-like sequences. The number of GH32 gene family members in a particular fungus appeared to be related to the ecological strategy employed by the fungus, which was similar to previous studies. The genomes of the plant pathogenic Ceratocystis species harboured two invertase genes. This was in contrast to their saprophytic relatives in the genus Huntiella that contained only one. Our results further showed that several processes have shaped the evolutionary trajectories of these Ceratocystidaceae genes. Based on these data, we posit that the evolution of the Ceratocystidaceae GH32 gene family involved divergence of invertase gene paralogs that presumably arose from a single Group 8 type of intracellular invertases present in the LCA of this fungal family. These paralogs acquired specific terminal motifs to give rise to genes encoding a cell wall invertase and a vacuolar invertase in extant species of Ceratocystis. A similar scenario likely also occurred in Huntiella where the ancestral invertase was remodelled into a cell wall invertase through the acquisition of relevant sequence motifs. The genes in the GH32 family of Ceratocystis and Huntiella were also located at non-homologous loci or regions in the genomes and were flanked by completely different sets of genes in the examined species, which indicated these genes are not orthologous (sensu Koonin; [36]) between the two sister genera. The genomic rearrangement that caused this was potentially linked to the activity of the putative *Fot5* element(s) found in Ceratocystis. Our results thus suggested a role for TEs in shaping the evolution of GH32 family genes, and thereby the sucrolytic activities and related ecological strategies of the Ceratocystidaceae that harbour them.

#### Availability of supporting data

This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession

LCZG00000000. The version described in this paper is version LCZG01000000.

#### **Additional files**

**Additional file 1: Table S1.** Genomic location, Protein ID and GenBank accession numbers for the sequences used in the present study. (DOC 67 kb)

Additional file 2: Figure S1. The chronogram was inferred using published calibration time points (see text for detail) and a Bayesian strict clock approach as implemented in BEAST (Bayesian evolutionary analysis by sampling trees) v.2 package v.2.2.1 [46]. Horizontal bars mark the lower and upper time boundaries of the indicated mean age (Million years ago) estimates of the nodes. The black arrows indicate the four calibration points, which include the Dothideomycetes crown group (mean 350 Million years ago [Mya] with 95 % credibility interval [CI] of 273–459) [50], the last common ancestor (LCA) of the Hypocreales (181 Mya with 95 % CI of 150–213) [51], the Clavicipitaceae crown group (117 Mya with 95 % CI of 98–144) [51], as well the Nectriaceae crown group (125 Mya with 95 % CI of 98–155) [52]. (PDF 227 kb)

**Additional file 3: Table S2.** List of the putative *Fot5* homologs identified in this study. (DOC 160 kb)

Additional file 4: Figure S2. Maximum likelihood tree of the Fot5 DDD catalytic domain. This analysis was done using the WAG substitution model [49] and gamma correction to account for among site rate variation. Percentage bootstrap support (based on a 1000 repeats, with cut-off value of 50 %) is indicated at the internodes. Genomic coordinates of putative Ceratocystis Fot5 homologs are provided in Additional file 3: Table S2. GenBank accession numbers or sequence identifiers for previously identified Fot5 homologs are: Fot2 [Genbank:JN624854, F. oxysporum), Fot5 [Genbank:CAE55867, F. oxysporum], Fot1 [Genbank:X64799, F. oxysporum], Fot4 [Genbank:AF076632, F. oxysporum], Fot9 [JGI:2517, F. araminearum], Fotvl [Genbank:CAG33729.1 Yarrowia lipolytica], Molly [Genbank:CAD32687, Parastagonospora nodorum], Ophio [Genbank:ABG26269, Ophiostoma novo-ulmi], PABRA [Genbank:-ACY56713, Paracoccidioides brasiliensis], Pixie [Genbank:CAD32689, Parastagonospora nodorum], Pot2 [Genbank:CAA83918, Magnaporthe arisea], Pot3 [Genbank:AAC49418, M. grisea], SCSCL [Genbank:XP001592252, Sclerotinia sclerotiorum], Taf1 [Genbank:AAX83011, Aspergillus fumigatus], Tan1 [Genbank:U58946, Aspergillus awamori] USMA [Genbank:UM03882, Ustilago maydis), Flipper [Genbank:AAB63315, Botryotinia fuckeliana] and Cirt1 [Genbank:XP710204, Candida albicans]. For the Ceratocystis sequences JS = Ceratocystis albifundus (green dots), J = Ceratocystis manginecans (red dots) and A = Ceratocystis fimbriata (blue dots), followed by the genomic position. (PDF 59 kb)

#### **Competing interests**

The authors declare that they have no competing interests.

#### Authors' contributions

MVDN, ETS, BDW and MJW conceived the experiment. BDW and MJW provided the resources. KN carried out DNA extraction and sequencing. CT, TG, ETS, ES, DR, ARM, PWM and MVDN performed bioinformatic analyses and all authors participated in the interpretation of findings. MVDN, ETS, PMW, ARM and MPAC produced the draft manuscript, and all the authors participated in its further development. All authors read and approved the final manuscript.

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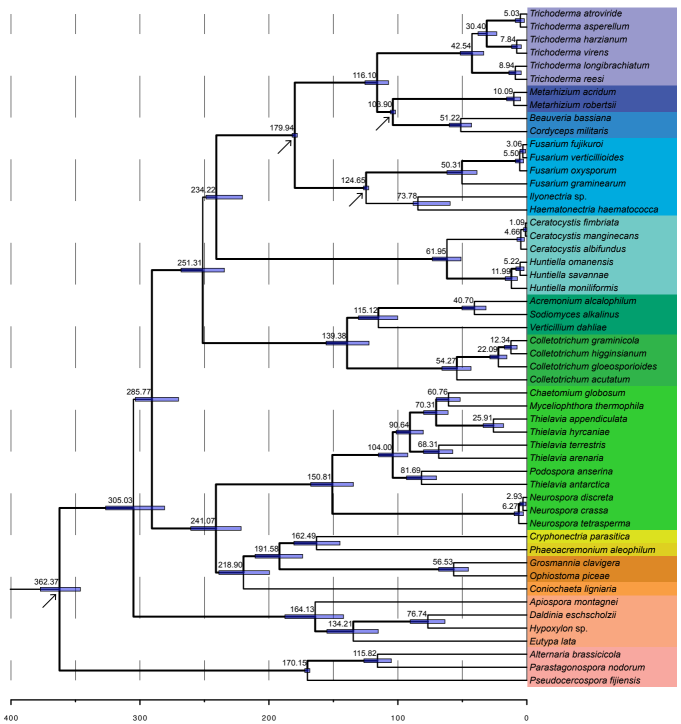


## **Supporting Material**

Additional file 1: Table S1 Genomic location, Protein ID and GenBank accession numbers for the sequences used in the present study.

Taxon	RPB2	RPB1	MCM7	EF3	EF1
Acremonium alcalophilum v2.0	1062944	1093047	2025206	2131370	1054920
Alternaria brassicicola	3555	6953	8151	6101	4028
Anthostoma avocetta NRRL 3190 v1.0	358799	486375	366025	358521	427765
Apiospora montagnei NRRL 25634 v1.0	165778	493678	198198	50284	53056
Beauveria bassiana ARSEF 2860	10291	260	1172	6276	2388
Ceratocystis albifundus	JSSU01001074:	JSSU01001372:	JSSU01000973:	JSSU01001081:	JSSU01001043:
	17650-16520	98713-99534	6705-7463	14425-15222	14465-13434
Ceratocystis fimbriata	APWK02000915:	APWK02000296:	KM495432	APWK02000274:	APWK0200014
	18679-19938	6011-5382		34298-33501	0: 31660-32691
Ceratocystis manginecans	JJRZ01000317:	JJRZ01000001:	KM495440	JJRZ01000106:	JJRZ01000044:
	13970-12711	51511-50882		53401-52604	8654-9685
Chaetomium globosum v1.0	18331	11528	10900	14798	16368
Colletotrichum graminicola M1.001	719	3226	9817	7890	8402
Colletotrichum higginsianum IMI 349063	11479	3226	4433	9852	13479
Coniochaeta ligniaria NRRL30616 V.1.0	10961	2621	5349	5900	10615
Cordyceps militaris CM01	6678	5469	5557	5648	808
Cryphonectria parasitica EP155 v2.0	348358	81184	263526	254675	355033
Daldinia eschscholzii EC12 v1.0	318358	53585	385894	386825	320720
Eutypa lata UCREL1	2978	7447	9192	4830	6820
Fusarium fujikuroi IMI 58289	7812	1359	4389	12267	8475
Fusarium graminearum v1.0	3039	1079	8215	3861	9400
Fusarium oxysporum v1.0	3867	884	7088	4679	11183
Fusarium verticillioides 7600 v1.0	3039	685	1814	4125	7139
Glomerella acutata v1.0	1356853	1444979	1457852	1571219	1333254
Glomerella cingulata 23 v1.0	1845705	1727691	1819621	1725208	1771438
Grosmannia clavigera kw1407	3040	7178	6801	2788	3764
O Company	JMSH01000100:	JMSH01000104:	JMSH01000011:	JMSH01000009:	JMSH01000070
Huntiella moniliformis	35095-33929	27732-28352	212164-211406	328536-329333	: 123062-
·					124093
II 11	JSUI01006186:	JSUI01006274:	JSUI01006482:	JSUI01006507:	JSUI01006383:
Huntiella omanensis	9662-8514	17350-16709	39381-40139	26080-25283	43877-44908
TI .: 11	NODE_6005:	NODE_3982:	KM495462	NODE_4668:	NODE_3974:
Huntiella savannae	96370-95237	106186-105545		115552-114755	9462-8431
Hypoxylon sp. CI-4A v1.0	11137	34528	1393894	123171	1390744

Ilyonectria sp. v1.0	1466688	1624773	1550485	1535645	1528939
Metarhizium acridum CQMa 102	2463	7408	1075	4249	964
Metarhizium robertsii ARSEF 23	336	658	5659	2762	3797
Myceliophthora thermophila v2.0	2307570	2294525	2295768	73461	2298136
Mycosphaerella_fijiensise_v2.0	136855	58269	56719	210369	209593
Nectria haematococca v2.0	70495	103209	92685	70096	59329
Neurospora crassa OR74A v2.0	9487	328	89	5493	436
Neurospora discreta FGSC 8579 mat A	128789	73118	91067	166279	160296
Neurospora tetrasperma FGSC 2508 mat A	124967	70543	88816	116274	118521
v2.0					
Ophiostoma piceae UAMH 11346	2689	816	8019	1044	3764
Phaeoacremonium aleophilum UCRPA7	5318	5500	7516	4179	4720
Podospora anserina S mat+	5635	297	726	4339	2076
Sodiomyces alkalinus v1.0	273095	143113	334328	354039	318117
Stagonospora_nodorum SN15 v2.0	4533	4606	10072	11732	5231
Thielavia antarctica CBS 123565 v1.0	442445	184819	85725	445493	442817
Thielavia appendiculata CBS 731.68 v1.0	581449	562737	639459	647848	350199
Thielavia arenaria CBS 508.74 v1.0	783437	700748	774767	778860	773701
Thielavia hyrcaniae CBS 757.83 v1.0	499486	128871	556260	499996	494733
Thielavia terrestris v2.0	2110960	2121340	2114018	2115984	2111611
Trichoderma asperellum CBS 433.97 v1.0	77086	68995	157112	35803	158395
Trichoderma atroviride V2.0	151043	89498	294914	156340	300828
Trichoderma harzianum CBS 226.95 v1.0	9591	508530	490345	518654	12328
Trichoderma longibrachiatum ATCC 18648	58810	67760	1427374	1398382	1397263
v3.0					
Trichoderma reesei v2.0	79225	79315	3212	71363	46958
Trichoderma virens Gv29-8 v2.0	76818	85034	74376	74376	83874
Verticillium dahliae v1.0	1143	1942	9482	3649	7717



Additional file 3: Table S2 List of the putative Fot5 homologs identified in this study.

Taxon <sup>1</sup>	Scaffold <sup>2</sup>		
Ceratocystis fimbriata	APWK02000105	2472 - 1779	
Ceratocystis fimbriata	APWK02000123	30129 - 31523	
Ceratocystis fimbriata	APWK02000123	8603 - 7812	
Ceratocystis fimbriata	APWK02000136	3179 - 4007	
Ceratocystis fimbriata	APWK02000148	74022 - 74840	
Ceratocystis fimbriata	APWK02000197	44307 - 45160	
Ceratocystis fimbriata	APWK02000198	13282 - 14288	
Ceratocystis fimbriata	APWK02000202	6057 - 4657	
Ceratocystis fimbriata	APWK02000311	11474 - 12699	
Ceratocystis fimbriata	APWK02000311	22252 - 23652	
Ceratocystis fimbriata	APWK02000312	1492 - 2416	
Ceratocystis fimbriata	APWK02000312	7208 - 6344	
Ceratocystis fimbriata	APWK02000356	5944 - 5183	
Ceratocystis fimbriata	APWK02000374	23167 - 22031	
Ceratocystis fimbriata	APWK02000374	7593 - 8337	
Ceratocystis fimbriata	APWK02000374	8539 - 9495	
Ceratocystis fimbriata	APWK02000455	36425 - 37405	
Ceratocystis fimbriata	APWK02000480	30105 - 28382	
Ceratocystis fimbriata	APWK02000506	2481 - 1694	
Ceratocystis fimbriata	APWK02000506	2519 - 3517	
Ceratocystis fimbriata	APWK02000544	1132 - 608	
Ceratocystis fimbriata	APWK02000557	11119 - 10841	
Ceratocystis fimbriata	APWK02000557	7553 - 6235	
Ceratocystis fimbriata	APWK02000559	7009 - 8409	

Ceratocystis fimbriata	APWK02000578	22299 - 22994
Ceratocystis fimbriata	APWK02000587	19846 - 18485
Ceratocystis fimbriata	APWK02000607	17088 - 17918
Ceratocystis fimbriata	APWK02000663	5126 - 3661
Ceratocystis fimbriata	APWK02000666	5454 - 4177
Ceratocystis fimbriata	APWK02000692	9449 - 9747
Ceratocystis fimbriata	APWK02000730	24422 - 23022
Ceratocystis fimbriata	APWK02000744	1927 - 1367
Ceratocystis fimbriata	APWK02000756	12574 - 13923
Ceratocystis fimbriata	APWK02000757	6330 - 5743
Ceratocystis fimbriata	APWK02000758	17279
Ceratocystis fimbriata	APWK02000758	7138 - 6403
Ceratocystis fimbriata	APWK02000814	2160 - 1530
Ceratocystis fimbriata	APWK02000839	1359 - 147
Ceratocystis fimbriata	APWK02000856	6877 - 5481
Ceratocystis fimbriata	APWK02000902	8262 - 7294
Ceratocystis fimbriata	APWK02000903	4260 - 5203
Ceratocystis fimbriata	APWK02000970	8053 - 8768
Ceratocystis fimbriata	APWK02000971	1372 - 230
Ceratocystis fimbriata	APWK02000974	18833 - 20178
Ceratocystis fimbriata	APWK02000987	6401 - 5892
Ceratocystis fimbriata	APWK02001023	18815 - 18234
Ceratocystis fimbriata	APWK02001031	18176 - 18733
Ceratocystis fimbriata	APWK02001038	5041 - 3794
Ceratocystis fimbriata	APWK02001045	10162 - 8913
Ceratocystis fimbriata	APWK02001045	12306 -

Ceratocystis fimbriata	APWK02001046	2796 - 2038
Ceratocystis fimbriata	APWK02001065	626 - 1698
Ceratocystis fimbriata	APWK02001080	2763 - 4090
Ceratocystis fimbriata	APWK02001097	4556 - 3542
Ceratocystis fimbriata	APWK02001098	10156 - 11040
Ceratocystis fimbriata	APWK02001102	11856 - 12170
Ceratocystis fimbriata	APWK02001128	2808 - 2062
Ceratocystis fimbriata	APWK02001128	517 - 1234
Ceratocystis fimbriata	APWK02000925 and APWK02000924	22604 - 23811
Ceratocystis manginecans	JJRZ01000038	36427-37962
Ceratocystis manginecans	JJRZ01000007	31904 - 32740
Ceratocystis manginecans	JJRZ01000007	90570 - 91646
Ceratocystis manginecans	JJRZ01000010	25746 - 24799
Ceratocystis manginecans	JJRZ01000012	179023 - 178172
Ceratocystis manginecans	JJRZ01000027	122476 - 123591
Ceratocystis manginecans	JJRZ01000027	149594 - 150571
Ceratocystis manginecans	JJRZ01000035	101693 - 100920
Ceratocystis manginecans	JJRZ01000035	115607 - 114546
Ceratocystis manginecans	JJRZ01000035	58804 - 59439
Ceratocystis manginecans	JJRZ01000037	39081 - 37951
Ceratocystis manginecans	JJRZ01000038	22556 - 23808
Ceratocystis manginecans	JJRZ01000040	108812 - 109645
Ceratocystis manginecans	JJRZ01000040	15282 - 16682
Ceratocystis manginecans	JJRZ01000040	93043 - 91826
Ceratocystis manginecans	JJRZ01000040	93604 - 94776

Ceratocystis manginecans	JJRZ01000040	98685 - 99658
Ceratocystis manginecans	JJRZ01000047	109605 - 110621
Ceratocystis manginecans	JJRZ01000047	42943 - 43647
Ceratocystis manginecans	JJRZ01000047	68514 - 69785
Ceratocystis manginecans	JJRZ01000053	63381 - 62626
Ceratocystis manginecans	JJRZ01000053	8471 - 7500
Ceratocystis manginecans	JJRZ01000060	43612 - 42422
Ceratocystis manginecans	JJRZ01000062	82408 - 83190
Ceratocystis manginecans	JJRZ01000063	15135 - 14359
Ceratocystis manginecans	JJRZ01000063	33842 - 34771
Ceratocystis manginecans	JJRZ01000063	727 - 68
Ceratocystis manginecans	JJRZ01000071	102960 - 103565
Ceratocystis manginecans	JJRZ01000071	17708 - 18418
Ceratocystis manginecans	JJRZ01000071	42347 - 43342
Ceratocystis manginecans	JJRZ01000071	44150 - 45061
Ceratocystis manginecans	JJRZ01000071	66866 - 67495
Ceratocystis manginecans	JJRZ01000071	67888 - 68775
Ceratocystis manginecans	JJRZ01000087	1218 - 2234
Ceratocystis manginecans	JJRZ01000087	60655 - 59912
Ceratocystis manginecans	JJRZ01000087	60797 - 62197
Ceratocystis manginecans	JJRZ01000087	69618 - 68743
Ceratocystis manginecans	JJRZ01000087	8534 - 9682
Ceratocystis manginecans	JJRZ01000092	16631 - 17467
Ceratocystis manginecans	JJRZ01000092	73101 - 73814
Ceratocystis manginecans	JJRZ01000097	28329 - 28961
Ceratocystis manginecans	JJRZ01000111	82144 - 82884

Ceratocystis manginecans	JJRZ01000125	23678 - 22878
Ceratocystis manginecans	JJRZ01000140	19610 - 18201
Ceratocystis manginecans	JJRZ01000144	39985 - 38576
Ceratocystis manginecans	JJRZ01000149	19825 - 18758
Ceratocystis manginecans	JJRZ01000149	36490 - 35312
Ceratocystis manginecans	JJRZ01000150	50435 - 51702
Ceratocystis manginecans	JJRZ01000194	18316 - 18996
Ceratocystis manginecans	JJRZ01000200	45632 - 44712
Ceratocystis manginecans	JJRZ01000211	21268 - 20648
Ceratocystis manginecans	JJRZ01000222	24694 - 25881
Ceratocystis manginecans	JJRZ01000222	28996 - 27719
Ceratocystis manginecans	JJRZ01000222	40879 - 40034
Ceratocystis manginecans	JJRZ01000222	41331 - 42740
Ceratocystis manginecans	JJRZ01000225	36626 - 35349
Ceratocystis manginecans	JJRZ01000225	9256 - 8657
Ceratocystis manginecans	JJRZ01000235	10435 - 9374
Ceratocystis manginecans	JJRZ01000235	20285 - 21352
Ceratocystis manginecans	JJRZ01000235	24590 - 23337
Ceratocystis manginecans	JJRZ01000235	30070 - 31080
Ceratocystis manginecans	JJRZ01000235	35692 - 34808
Ceratocystis manginecans	JJRZ01000247	1191 - 265
Ceratocystis manginecans	JJRZ01000260	31859 - 32488
Ceratocystis manginecans	JJRZ01000268	873 - 235
Ceratocystis manginecans	JJRZ01000287	19905 - 19222
Ceratocystis manginecans	JJRZ01000289	5055 - 3796
Ceratocystis manginecans	JJRZ01000302	3220 - 2561

Ceratocystis manginecans	JJRZ01000307	15842 - 16795
Ceratocystis manginecans	JJRZ01000307	23778 - 22906
Ceratocystis manginecans	JJRZ01000307	7642 - 8640
Ceratocystis manginecans	JJRZ01000312	22185 - 21553
Ceratocystis manginecans	JJRZ01000320	17136 - 16378
Ceratocystis manginecans	JJRZ01000352	25106 - 24237
Ceratocystis manginecans	JJRZ01000357	6336 - 7217
Ceratocystis manginecans	JJRZ01000358	10206 - 9196
Ceratocystis manginecans	JJRZ01000358	2331 - 3602
Ceratocystis manginecans	JJRZ01000362	18170 - 16683
Ceratocystis manginecans	JJRZ01000368	1225 - 1839
Ceratocystis manginecans	JJRZ01000400	8772 - 7999
Ceratocystis manginecans	JJRZ01000407	11241 - 9970
Ceratocystis manginecans	JJRZ01000412	11744 - 12691
Ceratocystis manginecans	JJRZ01000412	61213 - 61836
Ceratocystis manginecans	JJRZ01000426	1464 - 193
Ceratocystis manginecans	JJRZ01000430	16313 - 17260
Ceratocystis manginecans	JJRZ01000430	1904 - 1305
Ceratocystis manginecans	JJRZ01000430	5539 - 6321
Ceratocystis manginecans	JJRZ01000438	12839 - 11976
Ceratocystis manginecans	JJRZ01000438	15248 - 15988
Ceratocystis manginecans	JJRZ01000450	1027 - 2178
Ceratocystis manginecans	JJRZ01000462	13221 - 14288
Ceratocystis manginecans	JJRZ01000472	2690 - 3682
Ceratocystis manginecans	JJRZ01000485	13102 - 11831
Ceratocystis manginecans	JJRZ01000485	5699 - 4422

Ceratocystis manginecans	JJRZ01000485	6534 - 7811
Ceratocystis manginecans	JJRZ01000500	11942 - 12619
Ceratocystis manginecans	JJRZ01000500	4682 - 3405
Ceratocystis manginecans	JJRZ01000548	8111 - 7236
Ceratocystis manginecans	JJRZ01000553	6072 - 7223
Ceratocystis manginecans	JJRZ01000563	8127 - 7312
Ceratocystis manginecans	JJRZ01000564	7383 - 6745
Ceratocystis manginecans	JJRZ01000601	1292 - 2569
Ceratocystis manginecans	JJRZ01000603	5037 - 6227
Ceratocystis manginecans	JJRZ01000680	1868 - 597
Ceratocystis manginecans	JJRZ01000723	1324 - 71
Ceratocystis manginecans	JJRZ01000752	2156 - 2938
Ceratocystis albifundus	JSSU01000917	4384 - 3785
Ceratocystis albifundus	JSSU01000992	13389 - 13150
Ceratocystis albifundus	JSSU01001059	11392 - 11984
Ceratocystis albifundus	JSSU01001108	14800 - 14189
Ceratocystis albifundus	JSSU01001122	11103 - 10804
Ceratocystis albifundus	JSSU01001122	17389 - 17033
Ceratocystis albifundus	JSSU01001122	25755 - 25381
Ceratocystis albifundus	JSSU01001193	35553 - 35521
Ceratocystis albifundus	JSSU01001217	41643 - 42286
Ceratocystis albifundus	JSSU01001264	29167 - 28544
Ceratocystis albifundus	JSSU01001267	46882 - 46499
Ceratocystis albifundus	JSSU01001301	6742 - 7030
Ceratocystis albifundus	JSSU01001335	34450 - 34863
Ceratocystis albifundus	JSSU01001340	51154 - 50897

Ceratocystis albifundus	JSSU01001365	3963 - 3352
Ceratocystis albifundus	JSSU01001396	161193 - 161116
Ceratocystis albifundus	JSSU01001403	184780 - 184155
Ceratocystis albifundus	JSSU01001405	65665 - 65318

The GenBank accesssion numbers for the *C. manginecans*, *C. fimbriata* and *C. albifundus* genomes are JJRZ01000000, APWK00000000 and JSSU00000000.

<sup>&</sup>lt;sup>2</sup> The name of the contig is followed by the nucleotide position of the gene within the contig.

