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Formation of mushrooms and lignocellulose degradation encoded in the genome sequence of

*Schizophyllum commune*

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The wood degrading fungus *Schizophyllum commune* is a model system for mushroom development. Here, we describe the 38.5 Mb assembled genome of this basidiomycete and application of whole genome expression analysis to study the 13,210 predicted genes. Comparative analyses of the *S. commune* genome revealed unique wood degrading machinery and mating type loci with the highest number of reported genes. Gene expression analyses revealed that one third of the 471 identified transcription factor genes were differentially expressed during sexual development. Two of these transcription factor genes were deleted. Inactivation of *fst4* resulted in the inability to form mushrooms, whereas inactivation of *fst3* resulted in more but smaller mushrooms than wild-type. These data illustrate that mechanisms underlying mushroom formation can be dissected using *S. commune* as a model. This will impact commercial production of mushrooms and the industrial use of these fruiting bodies to produce enzymes and pharmaceuticals.

The fungal kingdom comprises diverse and important organisms that impact agriculture, human health, carbon cycling, and biotechnology. The mushroom fruiting body is the most conspicuous form of the fungi and is found primarily in the basidiomycete group. Mushrooms produce anti-tumor and immuno-stimulatory molecules\(^1\,^2\) and enzymes that can be used for bioconversions\(^3\). Moreover, they have been identified as promising cell factories for the production of pharmaceutical proteins\(^4\). The main economic value of mushrooms, however, is their use as food\(^1\,^2\). The world-wide production of edible mushrooms amounts approximately 2.5 million tons annually. Despite their economical interest, relatively little is known about how mushroom-forming fungi obtain nutrients and how fruiting bodies are formed. Many mushroom-forming fungi cannot be cultured in the lab nor genetically modified. The basidiomycete *Schizophyllum commune* is one of the notable exceptions. It can be cultured on defined media and it completes its life cycle in approximately 10 days. Moreover, molecular tools to study growth and development of *S. commune* have been developed. In fact, it is the only mushroom-forming fungus in which genes
have been inactivated by homologous recombination. The importance of *S. commune* as a model system is also exemplified by the fact that its recombinant DNA constructs will express in other mushroom-forming fungi.

*S. commune* is one of the most commonly found fungi and can be isolated from all continents except for the arctic regions. *S. commune* has been reported to be a pathogen of humans and trees but it mainly adopts a saprobic life style by causing white rot. It is predominantly found on fallen branches and timber of deciduous trees. At least 150 genera of woody plants are substrates for *S. commune*, but it also colonizes softwood and grass silage (see). The mushrooms of *S. commune* that are formed on these substrates are used as a food source in Africa and Asia.

In the life cycle of *S. commune* meiospores germinate to form a sterile monokaryotic mycelium, in which each hyphal compartment contains one nucleus. This mycelium grows initially submerged but after a few days aerial hyphae are formed (Fig. 1A, E). Monokaryons that are confronted with each other will fuse. A fertile dikaryon is formed when the alleles of the mating-type loci *matA* and *matB* of the partners differ. A short exposure to light is essential for fruiting, while a high concentration of carbon dioxide and high temperatures (30-37°C) are inhibitory. Mushroom formation is initiated with the aggregation of aerial dikaryotic hyphae. These aggregates (Fig. 1B, F) form fruiting body primordia (Fig. 1C, G), which further develop into mature fruiting bodies (Fig. 1D, H). Karyogamy and meiosis take place in the basidia within the mature fruiting body. The resulting basidiospores can give rise to new monokaryotic mycelia.

Here, we report the genomic sequence of the monokaryotic *S. commune* strain H4-8 and demonstrate the role of this basidiomycete as a model system to study mushroom formation.

**RESULTS**

**Genome of *S. commune***
Sequencing genomic DNA of *S. commune* strain H4-8 with 8.29x coverage (Supplementary Table 1 online) revealed 38.5 Megabase genome assembly with 11.2% repeat content (Supplementary Text 1 online). The assembly is contained on 36 scaffolds (Supplementary Table 2 online), which represent 14 chromosomes. A total of 13,210 gene models are predicted, with 42% supported by expressed sequenced tags (ESTs) and 69% being similar to proteins from other organisms (Supplementary Tables 3 and 4 online). Clustering of the proteins of *S. commune* with those of other sequenced fungi (a phylogenetic tree of the organisms used in the analysis is shown in Supplementary Fig. 1 online) resulted in 7055 groups containing at least one *S. commune* protein (Supplementary Table 5 online). Analysis of these clusters suggested that 39% of the *S. commune* proteins have orthologs in the Dikarya and are thus conserved in the Basidiomycota and Ascomycota (Supplementary Table 6 online). Intriguingly, a similar percentage of proteins (36%) are unique to *S. commune*. Of these proteins, 46% have at least one inparalog (i.e. a gene resulting from a duplication within the genome) in *S. commune*. The uniqueness of the *S. commune* proteome is also illustrated by the presence of protein family (PFAM) domains (Supplementary Text 2 online) and the fact that only 43% of the predicted genes (5,703 out of the 13,210) could be annotated with a gene ontology (GO) term.

**Global gene expression analysis**

Whole genome expression was analysed in four developmental stages (monokaryon, stage I aggregates, stage II primordia, mature fruiting bodies; see Fig. 1) using Massively Parallel Signature Sequencing (MPSS). The majority of genes are either expressed in all stages (4859 genes) or not expressed (5308 genes) (Fig. 2A,D, Supplementary Table 7 online). 59.8% of the 13,210 predicted genes are expressed in at least 1 developmental stage (Supplementary Table 7 online). Fewer of the unique *S. commune* genes meet this threshold, whereas a higher percentage is observed for genes that share orthologs with Agaricomycetes or more distant fungi (Supplementary Table 6 online). This suggests that genes specific to *S. commune* are subject to a
more stringent regulation. This is consistent with the observation that *S. commune* specific genes are over-represented in the pool of genes that are differentially expressed during the four developmental stages (Supplementary Tables 8 and 9 online).

Anti-sense transcription is a widespread phenomenon in *S. commune*. 18.7% of the tags that could be related to a gene model originate from an anti-sense transcript. 42.3% of the predicted genes have anti-sense expression in one or more developmental stages (Supplementary Tables 7 and 10 online). Northern hybridization with strand-specific probes confirmed the existence of anti-sense transcripts of *sc4* (Protein ID 73533) (data not shown). In the anti-sense direction, a relatively large number of genes are uniquely expressed in stage II (2888 genes) and relatively few genes are expressed in all stages (1195 genes) (Fig. 2B). In stage II, 4302 genes are expressed in both sense and anti-sense direction (Fig. 2C). This overlap is large compared to the other developmental stages.

**Fruiting body development**

An enrichment analysis of functional annotation was performed on the expression profiles of the four developmental stages (monokaryon, stage I aggregates, stage II primordia, mature fruiting bodies). Functional terms involved in protein production, energy production and hydrophobins are over-represented in genes that were up-regulated during formation of stage I aggregates (Fig. 1 and Supplementary Table 9 online). Genes involved in signal transduction, regulation of gene expression, cell wall biogenesis and carbohydrate metabolism are enriched in the group of down-regulated genes during formation of stage I aggregates. These functional terms are enriched in the up-regulated genes during formation of stage II primordia, whereas terms involved in protein and energy production are enriched in the down-regulated genes (Fig. 1 and Supplementary Table 9 online). During formation of mature fruiting bodies, genes encoding transcription factors, and genes involved in amino acid, glucose and alcohol metabolism are enriched in the group of down-regulated genes.
Whole genome expression analysis during mushroom formation has also been performed in *Laccaria bicolor*\(^{10}\). Regulation of orthologous gene pairs of *L. bicolor* and *S. commune* could therefore be correlated during fruiting. To this end, microarray expression profiles of free-living mycelium and mature fruiting bodies of *L. bicolor* were compared to the MPSS expression profiles of monokaryotic mycelium and mature fruiting bodies of *S. commune*. 6751 expressed genes from *S. commune* had at least 1 expressed ortholog in *L. bicolor*. The correlation of changes in expression of the functional annotation terms to which these orthologous pairs belong was determined. There were 15 GO terms, 2 KEGG terms, 4 KOG terms and 4 PFAM terms that showed a positive correlation in expression (p < 0.01) (Supplementary Table 11 online). These terms include metabolic pathways (such as valine, leucine and isoleucine biosynthesis) and regulatory mechanisms (such as transcriptional regulation by transcription factors and signal transduction by G-protein alpha subunit). This indicates that regulation of these processes during mushroom formation is conserved in *S. commune* and *L. bicolor*.

**Analysis of specific gene groups**

Formation of a fertile dikaryon is regulated by the *matA* and *matB* mating type loci. Proteins encoded in these loci activate signalling cascades (see Supplementary Text3 online) thus regulating target genes. These target genes include proteins that fulfil structural functions such as hydrophobins (see Supplementary Text4 online) and enzymes. As a result, fruiting bodies are formed.

*matA*

The *matA* locus of strain H4-8 appears to have the highest homeodomain gene number in a fungal mating type locus described so far. This locus consists of two subloci, *Aα* and *Aβ*, which are separated by 550 kb on chromosome I of strain H4-8. Annotation revealed that the *Aα* locus of H4-8 contains two divergently transcribed genes encoding Y and Z homeodomain proteins of the HD2
and HD1 class, respectively (Fig. 3; Supplementary Table 12 online). These two genes, \textit{aay4} and \textit{aaz4}, have been previously described\textsuperscript{1}. One homeodomain gene had also been identified in the \textit{A\beta} locus of H4-8\textsuperscript{11}. The genomic sequence revealed that this locus actually contains six predicted homeodomain genes \textit{abq6} (HD1), \textit{abr6} (HD2), \textit{abs6} (HD1), \textit{abt6} (HD1, but lacking the NLS), \textit{abu6} (HD1) and \textit{abv6} (HD2) (Fig. 3; Supplementary Table 12 online).

\textit{matB}

Annotation of the genomic sequence of \textit{S. commune} has revealed that the \textit{matB} system contains more genes than previously envisioned. The \textit{matB} locus comprises two linked loci \textit{B\alpha} and \textit{B\beta}, which encode pheromones and pheromone receptors (Fig. 3; see\textsuperscript{1}). Previously, one pheromone receptor gene was identified both in \textit{B\alpha3} and \textit{B\beta2} of strain H4-8 (called \textit{bar3} and \textit{bbr2}, respectively)\textsuperscript{12}. The genome sequence revealed four additional genes with high sequence similarity to these pheromone receptor genes, which we call \textit{B} receptor-like genes 1-4 (\textit{brl1-4}) (Fig. 3). Three of these genes are located near \textit{bar3} and \textit{bbr2} on scaffold 10, whereas one (\textit{brl4}) is located on scaffold 8. MPSS analysis showed that the \textit{brl} genes are expressed (Supplementary Table 13 online). In fact, of all receptor and receptor-like genes, \textit{brl3} shows the highest expression under the conditions tested.

Three and eight pheromone genes have been previously identified in the \textit{B\alpha3} and \textit{B\beta2} loci, respectively\textsuperscript{13}. One additional pheromone gene, \textit{bpl5} (\textit{B} pheromone-like), has been identified in the \textit{B\alpha3} locus. Moreover, four additional pheromone-like genes were detected in the \textit{B\beta2} locus, called \textit{bpl1-4} (Fig. 3). Based on the MPSS analysis, only \textit{bpl2} failed to show expression (Supplementary Table 13 online). The \textit{B\alpha} gene \textit{bpl5} and three of the new \textit{B\beta} pheromone-like genes show deviations from the consensus farnesylation signal with CASR for \textit{Bpl5}, CTIA for \textit{Bpl1}, CRLT for \textit{Bpl2} and CQLT for \textit{Bpl3}. Previously, one of the pheromone genes (\textit{bhp2(6)}) was shown to function with the deviating farnesylation signal CEVM\textsuperscript{12}. This suggests that in \textit{S. commune} only one amino acid residue in the consensus sequence of the farnesylation signal needs to be aliphatic.
Transcription factors

Genes for 471 putative transcription factors have been identified in the genome of *S. commune*, of which 311 are expressed in at least one developmental stage (Supplementary Table 14 online). Of these genes, 56% are expressed in all developmental stages. 268, 200, 283 and 253 of these transcription factor genes were expressed, respectively, in the monokaryon, and during formation of stage I aggregates, stage II primordia and mushrooms. Interestingly, a cluster of monokaryotic specific transcription factors and a cluster of transcription factors that are upregulated in stage II primordia and/or in mature mushrooms were identified (Fig. 4). The latter cluster includes *fst3* (Protein ID: 257422) and *fst4* (Protein ID: 66861). These genes encode transcription factors that contain a fungal specific Zn(II)2Cys6 zinc finger DNA binding domain.

Genes *fst3* and *fst4* were inactivated by targeted gene deletions. The Δ*fst3* and Δ*fst4* monokaryons showed no phenotypic differences when compared to the wild-type. In contrast, the Δ*fst4*Δ*fst4* dikaryon did not fruit but produced more aerial hyphae when compared to the wild-type (Fig. 5). Apparently, Fst4 is involved in the switch between the vegetative phase and the reproductive phase. The Δ*fst3*Δ*fst3* dikaryon did form fruiting bodies. In fact, the mutant formed more, but smaller reproductive structures than those of the wild type (Fig. 5). Spatial and temporal regulation of fruiting body formation and sporulation was not altered in the Δ*fst3*Δ*fst3* strain. From these data we conclude that Fst3 inhibits formation of clusters of mushrooms.

Wood degradation by *Schizophyllum commune*

Degradation of lignin

*S. commune* has evolved its own set of enzymes that may degrade lignin. Such enzymes are classified as FOLymes14. These oxidative enzymes consist of lignin oxidases (LO families) and lignin-degrading auxiliary enzymes that generate H₂O₂ for peroxidises (LDA families). The LO
family consists of laccases (LO1), lignin peroxidases, manganese peroxidases, and versatile peroxidases (LO2) and cellobiose dehydrogenases (CDHs; LO3). *S. commune* contains 16 FOLyme genes and 11 genes that encode enzymes that are distantly related to these enzymes (*Table 1* and *Supplementary Table 15* online). The genome lacks genes encoding peroxidases of the LO2 family. However, it contains a CDH gene (LO3), 2 laccase genes (LO1), and a total of 13 LDA genes including 4 genes encoding glucose oxidases (LDA6), and benzoquinone reductases (LDA7) (*Table 1*).

*Ustilago maydis, Cryptococcus neoformans, Aspergillus nidulans, Neurospora crassa* and *Saccharomyces cerevisiae*, which have not been reported to have ligninolytic activity, and the brown rot fungus *Postia placenta* contain a lower diversity of FOLymes compared to *S. commune* (*Table 1*). In contrast, the coprophillic fungus *Coprinopsis cinerea* and the white rot fungus *P. chrysosporium* contain many more FOLymes than *S. commune* with 40 and 27 members, respectively.

**Polysaccharide degradation**

*S. commune* has the most complete polysaccharide breakdown machinery of all basidiomycetes examined. The Carbohydrate-Active Enzyme database (CAZy) identified 240 candidate glycoside hydrolases (GH), 75 candidate glycosyl transferases (GT), 16 candidate polysaccharide lyases (PL), and 30 candidate carbohydrate esterases (CE) that are encoded in the genome of *S. commune* (*Table 1*, *Supplementary Table 16* online). Compared to the genomes of other basidiomycetes, *S. commune* has the highest number of GHs, and PLs. *S. commune* is rich in genes encoding pectin, hemicellulose and cellulose degrading enzymes (*Supplementary Table 17* online). In fact, *S. commune* has genes in each family involved in the degradation of these plant cell wall polysaccharides. The *S. commune* genome is particularly rich in members of the glycosyl hydrolase families GH93 (hemicellulose degradation), and GH43 (hemicellulose and pectin degradation), and the lyase families PL1, PL3, and PL4 (pectin degradation) (*Supplementary Table 17* online). The
pectinolytic capacity of *S. commune* is complemented with pectin hydrolases from family GH28, GH88 and GH105.

**DISCUSSION**

The phylum Basidiomycota contains roughly 30,000 described species, accounting for 37% of the true fungi\(^ {15} \). The Basidiomycota comprises of two class level taxa (Wallemiomyctes, Entorrhizomycetes) and the subphyla Pucciniomycotina (rust), Ustilaginomycotina (smuts), and Agaricomycotina\(^ {16} \). The Agaricomycotina include the mushroom and puffball forming fungi, crust fungi, and jelly fungi. Currently five genomic sequences of Agaricomycotina are available: *P. chrysosporium*\(^ {17} \), *L. bicolor*\(^ {10} \), *P. placenta*\(^ {18} \), *C. neoformans*\(^ {19} \) and *C. cinerea*\(^ {20} \). We here report the 38.5 Megabase genome assembly of *S. commune*, which represents the first genomic sequence of the family of the Schizophyllaceae. The genome of *S. commune* is predicted to have 13,210 genes. 36% of the encoded proteins have no ortholog in other fungi. The fact that much about the proteome of *S. commune* is still unknown is also illustrated by the fact that only 43% of the predicted genes could be annotated with a gene ontology (GO) term. This is a similar percentage as seen in other basidiomycetes: 30% in *L. bicolor*\(^ {10} \), 48% for *P. placenta*\(^ {18} \) and 49% for *P. chrysosporium*\(^ {17} \).

**Substrate utilization**

*S. commune* is reported to be a white rot fungus\(^ {6} \). White-rot fungi degrade all woody cell-wall components, including the recalcitrant lignin. In contrast, brown-rotters efficiently degrade cellulose but only modify lignin, leaving a polymeric residue. *S. commune* primarily invades wood by growing through the lumen of vessels, tracheids, fibres and xylem rays. Adjacent parenchymatic cells in the xylem tissue are invaded via simple and bordered pits. As a consequence of this way of invasion, cellulose, hemicellulose, and pectin can serve as the primary carbon source for *S. commune*. Indeed, the genome of *S. commune* is rich in genes that encode enzymes that are
involved in the degradation of these polysaccharides. *S. commune* has genes in each family involved in the degradation of cellulose, hemicellulose, and pectin. The high number of pectinase genes correlates with earlier studies describing *S. commune* as one of the best basidiomycete pectinase producers\(^ {21}\). *S. commune* also encodes carbohydrate active enzymes that degrade other polymeric sugars (*e.g.* those that act on starch, mannan and inulin). In fact, *S. commune* has the most complete polysaccharide breakdown machinery of all basidiomycetes examined. This complete machinery is consistent with the wide variety of substrates that support growth of *S. commune*.

Compared to plant polysaccharides, relatively little is known about how fungi degrade lignin. Fungi are assumed to degrade lignin with FOLymes\(^ {14}\). These oxidative enzymes consist of lignin oxidases (LO families) and lignin-degrading auxiliary enzymes (LDA families). Evidence has shown that at least the LO2 family is involved in lignin degradation. This family consists of the lignin peroxidases, manganese peroxidases and versatile peroxidases. Involvement of laccases (LO1) and cellobiose dehydrogenases (CDHs; LO3) is still controversial. *S. commune* contains 16 genes encoding FOLymes. Genes of the LO2 family are lacking but the genome contains one CDH gene and 2 laccase genes. CDHs may participate in the degradation of cellulose, xylan and, possibly, lignin by generating hydroxyl radicals in a Fenton-type reaction. Laccases catalyse the one-electron oxidation of phenolic, aromatic amines and other electron-rich substrates with the concomitant reduction of O\(_2\) to H\(_2\)O. They are classified as having either low or high redox potential\(^ {22}\). It is not clear whether they belong to the high or low redox potential enzyme categories.

When the genomes of the white rot fungi *S. commune* and *P. chrysosporium*\(^ {17}\) and the brown rot fungus *P. placenta*\(^ {18}\) are compared, it is clear that *S. commune* has evolved its own set of FOLymes. *P. chrysosporium* lacks genes encoding laccases (LO1). It is thought to degrade lignin with the enzymes encoded by 16 isogenes of peroxidases (LO2), one CDH gene (LO3) and 4 genes of the multicopper oxidase (MCO) super family. In contrast, *P. placenta* contains 2 laccase-encoding genes (LO1) but lacks members of the LO2 and LO3 families. Since *S. commune* and *P.*
*placenta* lack true LO2 FOLymes, one would expect a low number of LDAs that are responsible for 
H₂O₂ production for the peroxidases. This is not the case. *S. commune* contains more and a higher 
diversity of LDAs when compared to *P. chrysosporium*. For instance, *S. commune* contains four 
glucose oxidase (LDA6) genes, whereas one or none are normally present in fungi. In the absence 
of peroxidases of the LO2 family, it is expected that the glucose oxidases of *S. commune* serve 
another function. Glucose oxidases convert glucose into gluconic acid. This acid solubilises 
inorganic phosphate and thus aids in the uptake of this nutrient.

**Mushroom formation**

*The mating type loci*

Monokaryons of *S. commune* will fuse when they are confronted with each other. Formation of a 
fertile dikaryon is regulated by the *matA* and *matB* mating type loci. The genome sequence has 
revealed that *matA* and *matB* of *S. commune* represent the fungal mating type loci that contain the 
highest number of genes. The *matB* locus comprises two linked loci \( B\alpha \) and \( B\beta \), which encode 
pheromones and pheromone receptors. Nine allelic specificities have been identified for both loci, 
resulting in 81 different mating types for *matB*. It was previously described that the \( B\alpha 3 \) and \( B\beta 2 \) 
loci of H4-8 each contain one pheromone receptor gene and three and eight pheromone genes, 
respectively. 5 additional pheromone genes and 4 pheromone receptor-like genes were 
identified in the genome of H4-8. These newly identified receptor-like genes are present in a *matB* 
deletion strain, which has no pheromone response with any mate (T. Fowler, unpublished results). 
This raises the question whether the four receptor genes are functional in *matB*-regulated 
development. MPSS analysis showed that they are all expressed, which suggests that they do not 
represent pseudogenes.

The *matA* locus consists of two subloci, \( A\alpha \) and \( A\beta \), of which 9 and 32 allelic specificities 
are expected to occur in nature. These loci are separated by 550 kb on chromosome I of strain H4-
8. This large distance has not been found in other fungi that have a tetrapolar mating system. The functionally well-characterized $A\alpha$ locus showed no significant differences from the published descriptions$^1$. It is composed of two genes encoding Y and Z homeodomain proteins of the HD2 and HD1 class, respectively. The Y and Z proteins, as in other basidiomycetes, interact in non-self combinations to activate the A-pathway of sexual development$^{1,24}$. Notably, a nuclear localization signal is present in Y but not in Z. This is consistent with non-self interaction of the two proteins taking place in the cytosol, which is followed by the translocation of the active protein complex into the nucleus$^1$.

$A\beta$ of S. commune has been studied much less compared to the $A\alpha$ locus. Interestingly, $A\beta$ reflects the highest homeodomain gene complexity of a fungal mating type locus described to date. It contains 4 homeodomain genes of the HD1 class and two of the HD2 class. The $A\beta$ locus of S. commune thus resembles that of C. cinerea, which consists of two pairs of functional HD1 and HD2 homeodomain genes (b and d)$^{25}$. The large number of genes in mat$A\beta$ would explain why recombination analyses predict as many as 32 mating specificities for this locus$^{26}$. Taken together, S. commune seems ideal to identify the evolutionary pathways for creating high numbers of allelic specificities for enhancing outbreeding versus inbreeding rates.

**Whole genome expression**

Little is known about molecular processes that control formation of fruiting bodies in basidiomycetes other than the role of the mating type loci$^8$. Therefore, we performed a whole genome expression analysis in four developmental stages (i.e. in the sterile monokaryon, and in stage I aggregates, stage II primordia, and mature fruiting bodies of the dikaryon; see Fig. 1). MPSS showed that relatively few genes were specifically expressed in the monokaryon (284 genes) and in stage I aggregates and the mature mushrooms (128 genes in both cases). Interestingly, 467 genes were specifically expressed in stage II primordia. This suggests that this stage represents a major developmental switch. This is supported by the fact that genes involved in signal
transduction and regulation of gene expression are enriched in the group of up-regulated genes during formation of stage II primordia. Of interest, expression of these gene groups, among others, show a positive correlation during mushroom formation of *S. commune* and *L. bicolor*. This suggests that regulation of mushroom formation is a conserved process in the Agaricales.

Anti-sense expression was found to occur widely in *S. commune*. About 20% of all sequenced mRNA tags originated from an anti-sense transcript and more than 5600 of the predicted genes showed anti-sense expression in one or more developmental stages. Anti-sense transcription was most pronounced in stage II primordia. In this stage, more than 4300 genes were expressed in both sense and anti-sense direction and more than 800 genes were expressed in the anti-sense direction only. Previously, MPSS showed anti-sense transcripts in *Magnaporthe grisea*\(^2^7\). Little is known about the function of these transcripts in fungi. The circadian clock of *N. crassa* is entrained in part by the action of an anti-sense transcript of a clock component locus\(^2^8\), possible by RNA interference. It is tempting to speculate that anti-sense transcripts also regulate mRNA levels in *S. commune*. Natural anti-sense transcripts in eukaryotes have also been implicated in other processes such as translational regulation, alternative splicing and RNA editing\(^2^9\). The anti-sense transcripts of *S. commune* may also have such functions. In all these cases, the anti-sense transcripts could be functional in the developmental switch that would occur when stage II primordia are formed.

**Transcriptional regulators**

The fact that gene regulation seems to be conserved in the Agaricales made us decide to study the transcriptional regulators in more detail. 471 genes were identified that are predicted to encode a transcription factor. 268 of these transcription factors were expressed in the monokaryon, whereas 200, 283 and 253 were expressed during formation of stage I aggregates, stage II primordia and mushrooms, respectively. The relatively high number of transcription factors that are expressed during formation of stage II primordia again point to a major switch that takes place during this developmental stage.
A cluster of monokaryotic specific transcription factors and a cluster of transcription factors that are up-regulated in stage II primordia and/or in mature mushrooms were identified. Genes \(fst3\) and \(fst4\) that encode transcriptional regulators with a fungal specific Zn(II)2Cys6 zinc finger DNA binding domain belong to the latter cluster and were inactivated by targeted gene deletion. Growth and development of monokaryotic strains in which \(fst3\) or \(fst4\) were inactivated were not affected. Phenotypic differences, however, were observed in the dikaryon. The \(\Delta fst4\Delta fst4\) dikaryon did not fruit but produced more aerial hyphae when compared to the wild-type. On the other hand, the \(\Delta fst3\Delta fst3\) dikaryon formed more, but smaller, fruiting bodies than the wild-type. Taken together, we conclude that Fst4 is involved in the switch between the vegetative and the reproductive phase and that Fst3 inhibits formation of clusters of mushrooms. Inhibition of such clusters could be important in a natural environment to ensure sufficient energy is available for full fruiting body development. Interestingly, \(fst3\) and \(fst4\) have homologs in other mushroom forming fungi and it is tempting to speculate that they have a similar function in these organisms. This is supported by the observation that the homologs of \(fst3\) and \(fst4\) are up-regulated in young fruiting bodies of \(L. bicolor\) compared to free-living mycelium\(^{10}\). In mature fruiting bodies of \(L. bicolor\), the expression level of the homolog of \(fst3\) remains constant compared to young fruiting bodies, whereas the \(fst4\) homolog returns to the level of expression in the free-living mycelium.

**Summary**

The genomic sequence of \(S. commune\) will be an essential tool to unravel mechanisms by which mushroom forming fungi degrade their natural substrates and form their reproductive structures. The large variety of genes that encode extracellular enzymes that act on polysaccharides likely explains why \(S. commune\) is so common in nature. Moreover, the genome sequence suggests that \(S. commune\) has a unique mechanism for lignin degradation. The deep capture of gene expression via MPSS has provided us with leads on how mushroom formation is regulated. In addition to certain transcription factors, anti-sense transcription may play an important role in this process. Without
doubt, the understanding of physiology and sexual reproduction of *S. commune* will have an impact on the commercial production of edible mushrooms and the use of mushrooms as a cell factory.

**ACKNOWLEDGEMENTS**

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**CONTRIBUTION TO THE MANUSCRIPT**

JSH, HW, TF, WL, AG, EK, and SB wrote the proposal; LL, JdJ, EL and RO isolated RNA and DNA and made libraries; EL and SL coordinated sequencing of the genome; JSc assembled the genome; IG, RO, KB, JSt, SB, EK and HW coordinated the annotation process; IG, AA, AS, JSt, TF, ER, AL, FP, UK, JM, JdJ, RdV, PC, VL, BH, WL, AG, PvK, KB, JSH, EK, SE, NK, RO, MR annotated genes; RO, JdJ, FS, and LL performed experiments; RO, JdJ, FS, LL and HW interpreted and designed experiments; RO, AA, JSt, ER, JdJ, RdV, FS, KB, EK, JSH, IG, SH, and HW wrote the paper; RO and HW coordinated writing of the paper.

**LEGENDS**

**Figure 1** Development of *S. commune*. Four-day-old (A-C; E-G) and eight-day-old (D, H) colonies grown from homogenates showing typical developmental stages in the life-cycle of *S. commune*. A monokaryon forms sterile aerial hyphae that form a fluffy white layer on top of the vegetative mycelium (A, E). Aerial hyphae of a dikaryon interact with each other to form stage I aggregates.
(B, F) which, after a light stimulus, develop into stage II primordia (C, G). These primordia further differentiate into sporulating mushrooms (D, H). An enrichment analysis shows that particular functional terms are over-represented in genes that are up- or down-regulated during a developmental transition. These terms are indicted below panels A-H. A-D represent cultures grown in 9 cm Petri-dishes, whereas E-H represent magnifications thereof. Bar represents 1 mm (H), 2.5 mm (E, F) and 5 mm (G).

**Figure 2** Gene expression in four developmental stages of *S. commune* illustrated by VENN diagrams (A-C) and a heat map (D). The cut-off for expression is 4 TPM. VENN diagrams in (A) and (B) show the overlap of genes that are expressed in sense and anti-sense direction in the four developmental stages, respectively. As an example, in (A) 61 genes are expressed in sense direction in stage I and stage II, 4859 genes are expressed in all stages, 132 genes are expressed in the monokaryon and mature fruiting bodies and 5308 genes are not expressed in any of the stages. (C) VENN diagram of the overlap in genes that show sense and anti-sense expression in a particular developmental stage and with all stages combined. (D) Heat map of expression of the *S. commune* genes in the four developmental stages. Bar on top of the panel represents expression values between 0 and 300 TPM. Genes with expression values higher than 300 TPM are also indicated in red. The bar on the right indicates a cluster of 366 highly expressed and differentially regulated genes. Annotation information of the genes in this cluster is given in **Supplementary Table 18** online.

**Figure 3** Distribution of genes encoding HD1 and HD2 homeodomain proteins in the *matA* locus and genes encoding pheromone receptors and pheromones in the *matB* locus of *S. commune* strain H4-8. The *matA* and *matB* loci are positioned on scaffold 1 and 10, respectively. One additional pheromone receptor gene, *brl4*, has been identified on scaffold 8.
**Figure 4** Histogram (A) and heat map (B) of expression of the 471 transcription factors in the genome of *S. commune*. The histogram (A) shows the percentage of transcription factor genes that are differentially expressed between stages of development. The heat map (B) shows a cluster containing predominantly monokaryon specific transcription factors and a cluster containing predominantly stage II and/or mushroom specific transcription factors. These clusters are enlarged on the right part of the heat map. The latter group contains two fungal specific transcription factor genes, called *fst3* and *fst4* (protein IDs 257422 and 66861, respectively).

**Figure 5** Fruiting body formation in a wild-type dikaryon (A, D) and in dikaryons in which *fst3* (B, E) and *fst4* (C, F) have been inactivated. D-F show a magnification of part of the colonies shown in A-C. Bar represents 5 mm (D-F).
Table 1. Comparison of the number of FOLymes and CAZymes of *S. commune* with those of other fungi. LO1 represents the laccases, LO2 peroxidases, LO3 cellobiose dehydrogenases, LDA1 aryl alcohol oxidases, LDA2 vanillyl-alcohol oxidases, LDA3 glyoxal oxidases, LDA4 pyranose oxidases, LDA5 galactose oxidases, LDA6 glucose oxidases, LDA7 benzoquinone reductases, and LDA8 alcohol oxidases. GH represent the glycoside hydrolases, GT glycosyl transferases, PL polysaccharide lyases, and CE carbohydrate esterases.

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METHODS

Strains and Culture conditions

*S. commune* was routinely grown at 25 °C on minimal medium (MM) with 1% glucose and with or without 1.5% agar. Liquid cultures were shaken at 225 rpm. Glucose was replaced with 4% glycerol for cultures used in the isolation of genomic DNA. All *S. commune* strains used were isogenic to strain 1-40. Strain H4-8 (*matA43matB41*; FGSC #9210) was used for sequencing. EST libraries were generated from H4-8 and from a dikaryon that resulted from a cross between H4-8 and strain H4-8b (*matA4 matB43*). Strains 4-39 (*matA41matB41*; CBS 341.81) and 4-40 (*matA43matB43*; CBS 340.81) were used for MPSS analysis. These strains show a more synchronized fruiting compared to a cross between H4-8 and H4-8b. Partial sequencing of the haploid genome revealed that strains 4-40 and 4-39 have minor sequence differences (< 0.2%) with strain H4-8 (data not shown).

Isolation of genomic DNA, genome sequencing and assembly

Genomic DNA of *S. commune* was isolated as described and sequenced with the use of a whole-genome shotgun strategy. All data were generated by paired-end sequencing of cloned inserts with 6 different insert sizes using Sanger technology on ABI3730xl sequencers. The data were assembled using the whole-genome shotgun assembler Arachne (http://www.broad.mit.edu/wga/).

EST library construction and sequencing

Cultures were inoculated on MM plates with 1% glucose using mycelial plugs as an inoculum. Strain H4-8 was grown for 4 days in the light, whereas the dikaryon H4-8 x H4-8.3 was grown for 4 days in the dark and 8 days in the light. Mycelium of the dikaryotic stages was combined and RNA was isolated as described. The PolyA+ RNA fraction was obtained using the Absolutely mRNA Purification kit and manufacturer’s instructions (Stratagene, La Jolla, CA). cDNA synthesis and cloning followed the SuperScript plasmid system procedure with Gateway technology for
cDNA synthesis and cloning (Invitrogen, Carsbad, CA). For the monokaryon, two size ranges of cDNA were cut out of the gel to generate two cDNA libraries (JGI library codes CBXY for range 0.6k-2kb and CBXX for the range >2kb). For the dikaryon, cDNA was used in the range >2kb, resulting in library CBXZ. The cDNA inserts were directionally ligated into vector pCMVsport6 (Invitrogen) and introduced into ElectroMAX T1 DH10B cells (Invitrogen). Plasmid DNA for sequencing was produced by rolling circle amplification (Templiphi, GE Healthcare, Piscataway, NJ). Subclone inserts were sequenced from both ends using Big Dye terminator chemistry and ABI 3730 instruments (Applied Biosystems, Foster City, CA).

Annotation methods

Gene models in the genome of *S. commune* were predicted using Fgenesh\(^{33}\), Fgenesh\(^{+}\)\(^{33}\), Genewise\(^{34}\) and Augustus\(^{35}\). Fgenesh was trained for *S. commune* with a sensitivity of 72% and a specificity of 74%. Augustus *ab initio* gene predictions were generated with parameters based on *C. cinerea* gene models\(^{20}\). In addition, about 31,000 *S. commune* ESTs were clustered into nearly 9,000 groups. These groups were either directly mapped to the genomic sequence with a threshold of 80% coverage and 95% identity, included as putative full-length (FL) genes, or used to extend predicted gene models into FL genes by adding 5’ and/or 3’ UTRs. Since multiple gene models were generated for each locus, a single representative model at each locus was computationally selected based on EST support and similarity to protein sequences in the NCBI non-redundant database. This resulted in a final set of 13,210 predicted genes, of which 1314 genes have been manually curated. In 66 cases, models were created or coordinates were changed.

All predicted gene models were functionally annotated by homology to annotated genes from NCBI non-redundant set and classified according to Gene Ontology (GO)\(^{36}\), eukaryotic orthologous groups (KOGs)\(^{37}\), KEGG metabolic pathways\(^{38}\), and Protein Family (PFAM) domains\(^{39}\).
RepeatModeler 1.0.3 (http://www.repeatmasker.org/RepeatModeler.html) was used to generate \textit{de novo} repeat sequence predictions for \textit{S. commune}. Repeats were classified by comparison to the RepBase database (http://www.girinst.org/repbase/index.html). RepeatModeler produced 76 families of repeats used as a search library in RepeatMasker (http://www.repeatmasker.org).

\textbf{Data availability}

\textit{S. commune} assemblies, annotations, and analyses are available through the interactive JGI Genome Portal at http://jgi.doe.gov/Scommune. Genome assemblies together with predicted gene models and annotations were also deposited at DDBJ/EMBL/GenBank under the project accession ADMJ00000000.

\textbf{Orthologs of \textit{S. commune} proteins in the fungal kingdom}

Proteins of \textit{S. commune} were assigned to orthologous groups with OrthoMCL V2.0\textsuperscript{40} with an inflation value of 1.5. Members of such groups were assigned as orthologs (in the case of proteins from another species) or inparalogs (in the case of proteins from \textit{S. commune}). Orthologs were determined in \textit{C. cinerea}\textsuperscript{20}, \textit{L. bicolor}\textsuperscript{10}, \textit{P. placenta}\textsuperscript{18}, \textit{P. chrysosporium}\textsuperscript{17}, \textit{C. neoformans}\textsuperscript{19}, \textit{U. maydis}\textsuperscript{41}, \textit{S. cerevisiae}\textsuperscript{42}, \textit{A. nidulans}\textsuperscript{43} and \textit{N. crassa}\textsuperscript{44}. All versus all BLASTp analysis was performed using NCBI standalone BLAST v2.2.20 with an E-value of 1e-5 as a cut-off. Custom scripts were used to further analyse the orthologous groups resulting from the OrthoMCL analysis. The evolutionary conservation for each orthologous group was expressed as the taxon this orthologous group was most specifically confined to (see \textbf{Supplementary Fig. 1} online).

\textbf{Representation analysis}

FuncAssociate 2.0\textsuperscript{45} was used to study over- and under-representation of taxon-specific genes and of functional annotation terms in sets of differentially regulated genes. Default settings were used with a p-value of 0.05 or 0.01 as the cut off.
**Protein families**

The PFAM database version 24.0\(^{39}\) was used to identify PFAM protein families. Custom scripts in Python were written to group genes on basis of their PFAM domains. Differences in the number of predicted proteins belonging to a PFAM family across the fungal domain was determined using the Student’s t-test. When Agaricales were compared to the rest of the Dikarya or when S. commune was compared to the Agaricales, only groups with a minimum of 5 members in at least one of the fungi were analysed. In the case S. commune was compared to the rest of the Dikarya, only groups with a minimum of 5 members in at least four of the fungi were analysed. In all cases, a p value of 0.05 was used as a cut off. Similar results were obtained using the non-parametric Mann-Whitney-U test.

**CAZy annotation**

Annotation of carbohydrate-related enzymes was performed using the Carbohydrate-Active Enzyme database (CAZy) annotation pipeline\(^{46}\). Ambiguous family attributions were processed manually along with all identified models that presented defects (deletions, insertions, splicing problems, etc.). Each protein was also compared to a library of experimentally characterized proteins found in CAZy to provide a functional description.

**FOLy annotation**

Lignin oxidative enzymes (FOLymes)\(^{14}\) were identified by BLASTP analysis of the S. commune gene models against a library of FOLy modules using an e-value <0.1. The resulting 68 protein models were manually analysed using the BLASTP results as well as multiple sequence alignments and functional inference based on phylogeny\(^{47}\). Basically, a protein was identified as a FOLyme when it showed a similarity score above 50% with sequences of biochemically characterised
enzymes. When the similarity score was <50% the proteins were scored as a FOLyme related protein.

**MPSS expression analysis**

Total RNA was isolated from the monokaryotic strain 4-40 and from the dikaryon resulting from a cross between 4-40 and 4-39. A 7-day-old colony grown on solid MM at 30 °C in the dark was homogenized in 200 ml MM using a Waring blender for 1 min at low speed. 2 ml of the homogenized mycelium was spread out over a polycarbonate membrane that was placed on top of solidified MM. Vegetative monokaryotic mycelium was grown for 4 days in the light. The dikaryon was grown for 2 and 4 days in the light to isolate mycelium with stage I aggregates and stage II primordia, respectively. Mature mushrooms of 3 days old were picked from dikaryotic cultures that had grown for 8 days in the light. RNA was isolated as described. MPSS was performed essentially as described except that after DpnII digestion MmeI was used to generate 20 bp tags. Tags were sequenced using the Clonal Single Molecule Array technique (Illumina, Hayward, CA, US). Between 4.2 and 7.6 million tags of 20 bp were obtained for each of the stages. The data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE21265 (http://www.ncbi.nlm.nih.gov/geo/).

Programs were developed in the programming language Python to analyze the data. Tag counts were normalized to tags per million (TPM). Those with a maximum of < 4 TPM in all developmental stages were removed from the data set. This data set consisted of a total of 40,791 unique tags. Of these tags, 61.7% and 58.6% could be mapped to the genome sequence and the predicted transcripts, respectively, using a perfect match as the criterion. The mapped tags accounted for 71.4% and 70.8% of the total number of tags, respectively. For comparison, 97.4% of the ESTs from *S. commune* strain H4-8 could be mapped to the assembly. Unmapped tags can be explained by sequencing errors in either tag or genomic DNA. Moreover, RNA editing may have altered the transcript sequencing to produce tags that do not match the genome perfectly. It may
also be that the assigned UTR is incomplete or that the DpnII restriction site that defines the 5’ end of the tag is too close to the polyA-tail of the mRNA. TPM values of tags originating from the same transcript were summed to assess their expression levels. A transcript is defined as the predicted coding sequence (CDS) extended with 400 bp flanking regions at both sides.

**Comparison of gene expression in *L. bicolor* and *S. commune***

Whole genome expression analysis of *L. bicolor*\(^{10}\) and *S. commune* was done essentially as described\(^5\). For *L. bicolor*, the microarray values from replicates were averaged. Expression values of genes were increased by one and the ratio between monokaryon and mushrooms (for *S. commune*) and free-living mycelium and mature fruiting bodies (for *L. bicolor*) was log-transformed. All expressed genes from *S. commune* that had at least 1 expressed ortholog in *L. bicolor* were taken into account, resulting in a total of 6751 orthologous pairs. These pairs were classified on basis of functional annotation terms. Correlation of changes in expression of these gene classes was expressed as the Pearson correlation coefficient. Only GO-terms with 10-200 pairs were used in the analysis. In the case of PFAM domains, a minimum number of 10 ortholog pairs were used.

**Deletion of transcription factors *fst3* and *fst4***

The transcription factor genes *fst3* (proteinID: 257422) and *fst4* (proteinID: 66861) were deleted using vector pDelcas\(^3\). Transformation of *S. commune* strain H4-8 was done as described\(^3\). Regeneration medium contained no antibiotic, whereas selection plates contained 20 µg ml\(^{-1}\) nourseothricin. Deletion of the target gene was confirmed by PCR. Compatible monokaryons with a gene deletion were selected from spores originating from a cross of the mutant strains with wild-type strain H4-8.3.

**REFERENCES**


Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.