Basidiomycete Mating Type Genes and Pheromone Signaling

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The genome sequences of the basidiomycete Agaricomycetes species Coprinopsis cinerea, Laccaria bicolor, Schizopyllum commune, Phanerochaete chrysosporium, and Postia placenta, as well as of Cryptococcus neoformans and Ustilago maydis, are now publicly available. Out of these fungi, C. cinerea, S. commune, and U. maydis, together with the budding yeast Saccharomyces cerevisiae, have been investigated for years genetically and molecularly for signaling in sexual reproduction. The comparison of the structure and organization of mating type genes in fungal genomes reveals an amazing conservation of genes regulating the sexual reproduction throughout the fungal kingdom. In agaricomycetes, two mating type loci, A, coding for homeodomain type transcription factors, and B, encoding a pheromone/receptor system, regulate the four typical mating interactions of tetrapolar species. Evidence for both A and B mating type genes can also be identified in basidiomycetes with bipolar systems, where only two mating interactions are seen. In some of these fungi, the B locus has lost its self/nonself discrimination ability and thus its specificity while retaining the other regulatory functions in development. In silico analyses now also permit the identification of putative components of the pheromone-dependent signaling pathways. Induction of these signaling cascades leads to development of dikaryotic mycelia, fruiting body formation, and meiotic spore production. In pheromone-dependent signaling, the role of heterotrimeric G proteins, components of a mitogen-activated protein kinase (MAPK) cascade, and cyclic AMP-dependent pathways can now be defined. Additionally, the pheromone-dependent signaling through monomeric, small GTPases potentially involved in creating the polarized cytoskeleton for reciprocal nuclear exchange and migration during mating is predicted.

THE MATING SYSTEMS IN BASIDIOMYCETES

The typical life cycle of the higher basidiomycetes, presently classified as Agaricomycetes (Table 1) (34), consists of haploid, monokaryotic as well as dikaryotic stages, the latter of which prevails in nature. The monokaryotic mycelium contains nuclei of only one genetic type and is thus termed homokaryon. This mycelium, grown from haploid spores, in general contains one of only one genetic type and is thus termed homokaryon. This prevails in nature. The monokaryotic mycelium contains nuclei monokaryotic as well as dikaryotic stages, the latter of which interaction occurs when both incompatible interactions occur when either of different specificities (Fig. 1), an incompatible interaction linked multiallelic loci: (i) Aα and Aβ and (ii) Bα and Bβ (80). In C. cinerea, the A complex also consists of two closely linked multiallelic loci, Aα and Aβ, while B consists of only one locus, as determined by recombination analyses (10, 12). For comparison, the plant pathogenic basidiomycetous smut fungus Ustilago maydis also contains two independent mating type loci (5), although the terminology is swapped, so that the multiallelic b locus corresponds to Aα or Aβ of the mushroom-forming agaricomycetes S. commune and C. cinerea, while the B counterpart in U. maydis is the a locus with only two specificities, a1 and a2 (5, 6, 43). On the basis of the phenotypes of the mating interactions between haploid strains with similar and different A and B loci, it is possible to determine the developmental pathway controlled by genes in each locus. In both S. commune and C. cinerea, the difference in one of the A or B subloci (α or β) is sufficient to activate the respective pathway of A- and B-controlled development. The B genes regulate reciprocal nuclear exchange and nuclear migration in both mates, while the A genes control the development of clamps involved in formation of dikaryotic hyphal compartments. This includes the initial pairing of the haploid nuclei with different A and B specificities and the synchronous division of the nuclear pair in association with the initial development of the clamp cell, the hook formation. Different B genes are then needed for the fusion of the hook cell to the subapical cell, which completes the clamp connection formation (4, 49, 80) (Fig. 1). In U. maydis, cell recognition and fusion, more reminiscent of the yeast system, is regulated by the a mating type locus, and only after fusion, the b mating type locus is called upon to regulate the formation of the dikaryotic hyphae with the ability to infect and grow in the host plant (6). Recently, the dikaryon of U. maydis also has been shown to be able to develop clamp-like structures (94).

In basidiomycetes, not all species exhibit a tetrapolar mating behavior with four potential mating interactions between spores from one fruiting body. In bipolar species (Table 1), only compatible or incompatible interactions are found, lack-
Agaricomycotina

Saccharomycetes cerevisiae
Saccharomyces pombe
Neurospora crassa

Basidiomycota

Ustilagomycotina

Ustilago maydis

Pucciniomycotina

Microbotryum violaceum

Ascomycota

Cryptococcus neoformans (teleomorph name: Filobasidiella neoformans)

Phanerochaete chrysosporium

Laccaria bicolor

Coprinellus disseminatus

Coprinopsis cinerea

Laccaria bicolor

Phanerochaete chrysosporium

Pholiota nameko

Postia placenta

Schizophyllum commune

TABLE 1. Taxonomy$^a$ and mating type systems of the fungal species discussed in the text

| Species                          | Mating system | Mating type loci                               | Reference(s) |
|---------------------------------|---------------|------------------------------------------------|--------------|
| Cryptococcus neoformans         | Bipolar       | MAT and Mata, sex chromosome                     | 35           |
| Coprinellus disseminatus        | Bipolar       | Multiallelic, multiple subloci in $A$, $B$ genes present | 41           |
| Coprinopsis cinerea             | Tetrapolar    | Multiallelic, multiple subloci in both $A$ and $B$ | 12           |
| Laccaria bicolor                | Tetrapolar    | Multiallelic $A$ and $B$                         | 70           |
| Phanerochaete chrysosporium     | Bipolar       | Multiallelic $A$, $B$ genes present              | 63, 70       |
| Pholiota nameko                 | Bipolar       | Multiallelic $A$, $B$ genes present              | 122          |
| Pleurotus djamor                | Tetrapolar    | Multiallelic $A$ and $B$                         | 40           |
| Postia placenta                 | Bipolar?      | $A$ and $B$ genes present                        | 62           |
| Schizophyllum commune           | Tetrapolar    | Multiallelic, multiple subloci in both $A$ and $B$ | 80           |
|---------------------------------|---------------|------------------------------------------------|--------------|
| $^a$ See reference 34.          |               |                                                |              |

FIG. 1. Life cycle of the tetrapolar basidiomycete Schizophyllum commune. In the fruiting bodies, basidiospores with four different combinations of $A$ and $B$ mating type genes are produced. If both $A$ and $B$ genes are different in two developing, haploid mycelia, a fully compatible mating takes place. (1) $B$ genes regulate the reciprocal nuclear migration, illustrated by movement of a nucleus via a hyphal fusion through broken septa from one haploid hypha into the other. (2) As a consequence of nuclear migration, multineucleate hyphae occur in the interaction zone of the mates. Unfused clamp connections (pseudoclamps) in the multineucleate hyphae are seen as a function of $A$-regulated development (3), which then controls the pairing of different nuclei followed by the hook cell formation and synchronous nuclear division (4). After separation of the sister nuclei, a cross wall is formed at the base of the hook and in the hypha (5). The growth of the hook toward and the subsequent fusion to the subapical cell are controlled by $B$ genes and allow the movement of the nucleus from the hook into the subapical cell to complete clamp formation (6).
connections, while the bipolar *P. placenta* (62) is known to form dikaryotic hyphae with rare clamp connections, and fruiting bodies can be found for both in nature. The mycorrhiza-forming *L. bicolor* more closely resembles *S. commune* and *C. cinerea* in that it has dikaryotic hyphae with clamp connections. It forms fruiting bodies in association with its host tree. The classical genetic studies, although performed originally by confrontation of a small number of homokaryons from the same fruiting body, suggested that the species is tetrapolar (24), which could be verified recently (61, 70).

**MOLECULAR STRUCTURE OF A MATING TYPE GENES**

**Homeodomain-encoding genes.** With the development of transformation systems, creation and screening of genomic and cDNA libraries, and expression of the isolated genomic and cDNA clones, it was possible to reveal the complex organization and to understand the function of the *A* and *B* loci of basidiomycetes. The structures of the *A* mating type loci have been well characterized in *C. cinerea*, in which the predicted “archetype” *A* locus contains at least three pairs of genes, each pair encoding one member each of the homokaryotic hyphae with rare clamp connections, and fruiting bodies can be found for both in nature. The mycorrhiza-forming *L. bicolor* more closely resembles *S. commune* and *C. cinerea* in that it has dikaryotic hyphae with clamp connections. It forms fruiting bodies in association with its host tree. The classical genetic studies, although performed originally by confrontation of a small number of homokaryons from the same fruiting body, suggested that the species is tetrapolar (24), which could be verified recently (61, 70).

**Homeodomain-encoding genes are conserved in both tetrapolar and bipolar basidiomycetes.** The screening of the *L. bicolor* genome (60) with known *HD1* and *HD2* sequences from *C. cinerea* revealed the gene pair *Lba1* and *Lba2*, encoding *HD1* and *HD2* proteins, respectively (70), without the redundancy observed in the *C. cinerea* locus (12). The *HD1* and *HD2* proteins of *L. bicolor* have the same conserved motifs identified in corresponding proteins of *C. cinerea*, suggesting functional similarity. Only one pair of homeodomain genes was also detected in the tetrapolar oyster mushroom, *Pleurotus djamor* (40), and in the *A* mating type locus of bipolar *Pholiota nameko* (122). Another bipolar species, *Coprinellus disseminatus*, contains multiallelic *Aα* and *Aβ* loci, each locus with an *HD1* and an *HD2* protein-encoding gene (41).

In *P. placenta*, the genes *mpl1* and *β-fg*, known from previous work to flank the *A* mating type loci in other basidiomycetes (39, 41), were used to identify one pair of *HD1* and *HD2*-encoding genes in both of the two putative *A* loci located on different scaffolds in the sequenced dikaryotic genome (62). The *HD* protein sequences show 40 to 50% similarity to other basidiomycete mating type proteins. This represents the level of similarity typical of the *A* mating type proteins, both between species and between allelic sequences within species. The *Aα* mating type locus with two homeodomain-encoding genes similar to those in other agaricomycetes and located next to the *mpl1* gene has also been identified in *Ph. chrysosporium* (63).

The structure and function of *HD*-protein-pairs of tetrapolar *C. cinerea*, *S. commune*, and *Pleurotus djamor*, as well as those of bipolar *Coprinellus disseminatus* and *Pholiota nameko*, are comparable to the organization and function of *be* and *bw* proteins in the tetrapolar basidiomycete *U. maydis* (44). In all these cases, a high number of allelic variants of *HD* proteins exist. The multiple pairs of genes encoding *HD* proteins in the *C. cinerea* locus are probably duplications of one original gene pair (45, 51). In *S. commune*, the evolution of multiple specificities in the *A* genes awaits detailed analyses, since so far mainly *Aα* alleles were investigated and very little information is available from genes in the *Aβ* locus (100). In contrast to the tetrapolar and bipolar species with a high number of allelic variants of *HD* proteins, the bipolar basidiomycete *Cr. neoformans* contains only two specificities, *MATα* and *MATα*, and the number of homeodomain genes is also lower. The *SXIIα* gene in *MATα* encodes an HD1 protein and *SXIIα* in *MATα* codes for an HD2 protein (5, 53).

**MOLECULAR STRUCTURE OF B MATING TYPE GENES**

**Pheromones and receptors.** The *B* mating type genes in *S. commune* and *C. cinerea* were shown to code for multiallelic
pheromone systems and for receptor systems. The first pheromone- and receptor-containing locus had been identified in the basidiomycete *U. maydis*, where the two alleles *a1* and *a2* at the *a* locus contain the pheromone genes *mfa1* and *mfa2* and the pheromone receptor genes *pra1* and *pra2* (9). Both *a1* and *a2* were shown to be comparable to the genes encoding the lipopeptide *a* factor, while both receptors share similarity with the Ste3 pheromone receptor in the yeast *S. cerevisiae*. Comparison of the structure of the cloned alleles of *Boα* and *Bβ* loci from *S. commune* (104, 112, 117), and also that of the *B* locus from *C. cinerea* (30, 74), led to detection of pheromone- and receptor-encoding genes also in these multiallelic loci. Here, a more redundant manner is seen than in *U. maydis*. The *S. commune* *Bα1* and *Bβ1* alleles contain both a receptor gene of the *S. cerevisiae* Ste3 type and several lipopeptide pheromone precursor-encoding genes, respectively. The most complex structure so far found is for the *Bβ2* allele, which codes for one receptor and eight pheromones, with three more pheromones probable from the genomic sequence (22).

The *Bβ2* allele was of special interest due to the numerous primary and secondary mutations that had been induced earlier to show that reversion of tetrapolar to bipolar mating systems is possible and indicating multiple genes responsible for *B* mating type function (76, 81, 82). The primary mutations were screened for induction of *B*-regulated development, thus rendering the mutant functionally bipolar and indicating constitutive signaling without mating (76). Secondary mutations were then induced, suppressing the constitutive phenotype and returning phenotypically to a nonconstitutive situation not exactly the same as that of the wild type but associated with loss of some specificity in tetrapolar mating interactions (81, 82). It was expected that the primary mutations should be in the receptor gene, rendering it constitutively active or responsive to (some) self-pheromones. The respective secondary mutations would then correspond to the loss of (some) pheromone genes or their respective functions (47). Both predictions were proven, since the molecular analysis of known *Bβ2* mutants could show corresponding mutations in pheromone- and receptor-encoding genes.

The pheromones contain farnesylation signals making them comparable to the *a* factor of *S. cerevisiae*. The small-lipopeptide pheromones result from the posttranslational modification of both *N* and *C* termini of the translated amino acid sequence. The pheromone-encoding amino acid sequences are deduced on the basis of the carboxyl terminus sequence CAAX, where *C* is cysteine, *A* is aliphatic, and *X* is any residue. The receptors in basidiomycetes are G-protein-coupled receptors (GPCRs) with seven membrane-passing domains (73, 112, 117, 119). The subclass of fungal, Ste3-like GPCRs is too divergent from rhodopsin-like and other serpentine receptors to allow modeling based on a known X-ray structure (7, 109). Recently, the yeast Ste3p was purified from transfected 293E cells (101), which might in the future allow structural and biochemical analyses of yeast as well as basidiomycete receptors.

**Expression of pheromones and receptors from basidiomycetes in yeast.** *S. cerevisiae* has been used as a heterologous expression system for basidiomycete receptors and pheromones (19, 31). The *α* mating type cells of *S. cerevisiae* have been used to express *S. commune* pheromones, and *α* mating type cells have been used to express compatible receptors. Culture supernatant of *S. commune* (31) or cell culture supernatant from yeast cells expressing one of the *S. commune* pheromone genes (19) has been used to activate a compatible *S. commune* receptor expressed in yeast. The strong reporter signals obtained by the pheromone receptor interaction in yeast suggested that the activation of the downstream signaling pathway took place via the yeast heterotrimeric G protein and a mitogen-activated protein kinase (MAPK) cascade. This was further confirmed by deletion of the gene *STE4* encoding the heterotrimeric G protein β subunit from the yeast host, which lowered dramatically the induction of the reporter system by *S. commune* pheromones. The deletion of the yeast *SST2* gene, involved in desensitizing of the Ga subunit, increased the reporter activity (19). The interaction with the yeast signaling cascade was also improved when the third cytoplasmic loop and proximal C terminus of the heterologous receptor protein known to be involved in heterotrimeric G-protein interaction were mutated for better homology to the yeast receptor protein (31).

The use of the heterologous expression system also made it possible to compare *S. commune* pheromone modification (19) with yeast *a* factor biosynthesis (37). The tests with yeast strains carrying either a wild-type or mutated member of the genes belonging to the *a* factor biosynthesis pathway (37) indicated that *S. commune* pheromone precursors can be farnesylated by Ram1/Ramp2, and for the removal of the C-terminal signal AAX, endoproteolysis by Rce1 or Ste24 is essential. Ste14 could also play an active role in carboxymethylation of *S. commune* pheromones. On the other hand, the yeast N-terminal proteases Ste23 and Ax1 seemed not to be required for production of an active *S. commune* pheromone. The deletion of a factor transporter Ste6p did not prevent the secretion of *S. commune* pheromones, suggesting an independent export of the heterologous pheromone from yeast (19).

In *C. cinerea*, the yeast heterologous expression of pheromones and receptors was used to analyze the activity of a mutant constitutive receptor gene by coexpressing a wild-type or a mutant receptor with a compatible pheromone (73). It was also shown that a synthetic pheromone whose structure was deduced on the basis of the *C. cinerea* pheromone sequence data (12) was able to activate the *C. cinerea* receptor expressed in yeast (73). Unfortunately, the synthetic pheromone had no effect when applied to growing *C. cinerea* hyphae, leaving the question about the site of pheromone/receptor interaction in filamentous fungi unanswered.

**S. commune** *Bmm* strain. In addition to heterologous expression in yeast, for *S. commune* an elegant system is available to screen the receptor/pheromone interactions by expression in a *Bmm* strain in which a large deletion removes all functions of the *B* complex (21, 22, 27, 28). The *S. commune* *Bmm* strain has been transformed independently with all functionally characterized *Boα* and *Bβ* pheromone- and receptor-encoding genes (21). The transformants were then mated with wild-type tester strains representing all *Boα* and *Bβ* mating types.

When a receptor is activated by a pheromone of different specificity, a dikaryon can be formed at the side of the induced receptor, providing that the transformant and tester strain had compatible *A* mating types (21, 27). This assay allowed grouping of the mature pheromones on the basis of the similarity in their 11- to 15-amino-acid sequences into five clusters, each
activating a set of \( Ba \) or \( B\beta \) receptors. Surprisingly, in three pheromone groups, two pheromones interacted with receptors from both \( Ba \) and \( B\beta \), which had not been observed before. The closer examination of one of these cases suggested that the \( Ba8 \) and \( B\beta 1 \) receptor proteins are highly similar in structure and that the mature pheromones Bbp2(4) and Bbp2(5) differ by only one amino acid. In the other two pheromone groups activating both \( Ba \) and \( B\beta \) receptors, the pheromone sequences differed by two amino acids, while the sequence information about identity/similarity of the respective \( Ba \) and \( B\beta \) receptors is still missing. These observations are extremely intriguing when the evolutionary origin of the multiallelic, multispecific \( B \) genes is considered (21, 28, 48, 87).

**Mutational analysis of pheromone receptor specificity domains.** The \( B_{null} \) strain has also been used to functionally characterize chimeric receptors. To identify specificity domains within the receptor proteins, N-terminal, central, and C-terminal parts of Bar1 and Bar2, encoded by \( Ba1 \) and \( Ba2 \), were interchanged (27). The swapping changed dramatically the properties of the receptors in recognizing different pheromones, either expressed by transformation in the \( B_{null} \) strain or from a wild-type tester strain for all \( Ba \) specificities, and led to identification of new phenotypes for receptors: constitutive, nonspecific (promiscuous), and highly selective (28). The use of one single pheromone at a time from the set of pheromones encoded in the wild-type locus showed that each pheromone has its own profile for induction of different receptors, while at the same time each receptor has a profile of pheromones accepted as ligands for activation of sexual development (22, 28). This was even true for the promiscuous receptors, which are responsive to all nine specificities if mated with a wild-type tester strains but which still excluded single pheromones when these were tested one at a time. This did show that the interaction of each single pheromone with a given receptor is dependent on recognition sites within the receptor molecule. Single point mutations at selected sites in the chimeric receptors could prove this theory, since altered specificity or reversion of a constitutive receptor to pheromone dependency was observed upon introduction of a single point mutation (28). This model is in accordance with the hypothesized multistate interaction model of GPCRs with their ligands (13, 25).

Here, a natural discriminatory system has been found that is perfectly explained by interaction of the ligand with different, distant parts of a receptor to determine mating specificity. The correct three-dimensional receptor folding pattern and the ligand-induced conformational change leading to pheromone signaling thus are dependent on multiple interactions of amino acids to allow stabilization of the induced receptor state in the case of a compatible interaction and avoid interacting with nonactivating pheromones. This also might explain why very delicate differences that needed to be balanced in \( S.\ commune \) could not be investigated in yeast, where the expression of \( S.\ commute \) receptor genes generally allowed the investigation of downstream signaling, but the differentiation between specificities was in the end different from the respective answers received when using homologous transformation assays.

**Comprehensive structure of \( B \) locus alleles.** In \( S.\ commute \), the sequencing of all pheromones and receptors from nine \( Ba \) and nine \( B\beta \) alleles is not yet complete. However, the arrangement of the entire \( B \) mating type complexes in two strains (\( Ba3 \); \( B\beta 2 \) and \( Ba3 \); \( B\beta 3 \)) has been resolved at sequence level, giving interesting material to interpret functional and evolutionary implications. Among these facts are the sequence homologies determining recombination rates between \( Ba \) and \( B\beta \) (21). The loci have been determined to be linked at 0.6 to 3.5 centimorgans (80), which already indicated a high level of diversity among natural isolates in the structure of respective \( B \) loci. This unique complexity can now be revisited using the genome sequence of \( S.\ commute \) strain H4-8.

In \( C.\ cinerea \), more extensive information on the structure of the \( B \) locus is available. The \( B \) locus arrangement in \( C.\ cinerea \) consists of three tandemly arranged groups of genes, each group consisting of one receptor-encoding gene and one to three pheromone-encoding genes. The sequencing of 13 known, allelic \( B \) mating type specificities of different geographic origin indicated that group 1 is represented by two, group 2 by five, and group 3 by seven alleles (89), allowing the generation of 70 different \( B \) mating specificities by recombination between the gene groups. The phylogenetic analysis of \( C.\ cinerea \) receptors divided the proteins into two main clusters, indicating an early duplication of an ancestral gene. Each cluster had members from the three gene groups, suggesting that the receptors in groups 1, 2, and 3 had not developed independently as separate lines but that, instead, genetic material was exchanged following by diversification. Recently, the \( C.\ cinerea \) Okayama strain 7 genome sequence revealed the receptors of all three groups of genes (89) but also a duplication of the receptor gene from group 2 and two new pheromone genes in group 1 (70). In addition, three putative nonmating type, receptor-like protein-encoding genes were identified, two of them linked to the \( B \) locus and the third gene elsewhere in the genome. The receptor genes belonging to the groups 1, 2, and 3 were named \( CcSTE3.1, CcSTE3.2a \) (and \( CcSTE3.2b \) for the duplicated one), and \( CcSTE3.3 \). The nonmating type receptor genes linked to the \( B \) locus were named \( CcSTE3-2151 \) and \( CcSTE3-2153 \) based on the protein identifications (70). It is of interest to speculate on the function of the proteins encoded by the additional receptor genes, which so far have never been detected in any functional test. The genome sequence of \( S.\ commute \) allowed the detection of a similar situation with four new pheromone receptor-like genes, indicating a common trait for mushroom-forming basidiomycetes.

**Exploring \( B \) mating type genes in newly sequenced agaricomycete genomes.** The analysis of the region containing a cluster of \( STE3 \)-like sequences in the \( L.\ bicolor \) genome led to identification of three genes encoding Ste3-like receptors, each with a pheromone-encoding gene in close proximity, and two nonmating type receptor genes with no pheromone genes in proximity. The arrangement of the \( B \) mating type gene cluster in \( L.\ bicolor \) and \( C.\ cinerea \) turned out to be highly conserved (70). A high number of pheromone-like genes with CAAX motifs at the C terminus occur throughout the \( L.\ bicolor \) genome, and several of them have multiple homologs in the \( C.\ cinerea \) genome. The function of these sequences has not yet been established. Oligoarray analysis in \( L.\ bicolor \) indicated that all the receptor-like genes were expressed but only the three with pheromone genes in close proximity were strongly expressed together with the pheromones in vegetative mycelium, ectomycorrhiza, and fruiting bodies (70).
In the bipolar *Ph. chrysosporium* and tetrapolar *Pleurotus djamor* (40), the putative *B* mating type locus seems to be represented by receptors homologous to *CcSTE3.1* and *CcSTE3.2*, respectively. A pheromone is encoded in close proximity to the *Pleurotus djamor* receptor, while no pheromone genes were linked to the receptor genes in bipolar *Ph. chrysosporium* (70).

In addition, a gene encoding a pheromone receptor-like protein was also identified in the *Ph. chrysosporium* genome (70). In *P. placenta*, the organization of the putative *B* mating type locus appears to be more complex (62). The *B* locus is represented by two allelic multimember clusters of receptor- and pheromone-encoding genes in two scaffolds of the dikaryotic genome. The phylogenetic analysis of all Ste3-type GPCRs of *P. placenta* revealed members of the three subfamilies defined for *C. cinerea* in addition to one receptor of each haploid genome, which did not group with the others, suggesting a new subtype or function other than mating. A high number of genes encoding putative pheromones occur between the receptors and in the flanking regions in each cluster. Both *Ph. chrysosporium* and *P. placenta* are assumed to represent bipolar mating type systems (62, 70), and therefore the role of the *B* mating type genes in these fungi is still unclear. Perhaps they share the reason for bipolar behavior with *Coprinellus disseminatus*, where the *B* factor receptors Ste3.1 and Ste3.2 are not linked to mating type determination but still function in clamp formation and fruiting (41). On the other hand, in the bipolar basidiomycete *Cr. neoformans*, the pheromone receptors, Ste3α and Ste3α in MATα and MATa alleles, respectively, are responsible for pheromone signaling and critical for mating specificity (35, 119). Here, the clustering of mating-associated genes led to linkage of pheromone/receptor and homeodomain transcription factor loci.

**SIGNALING PATHWAYS ASSOCIATED WITH B MATING TYPE GENES**

In agaricomycetes, the downstream components in the signaling pathways regulated by *A* and *B* mating type genes have yet to be analyzed in detail. In order to induce sexual development, mates have to respond to the ligand binding of the pheromone receptor via a downstream signaling cascade leading to dikaryotization (Fig. 1 and 2). Therefore, the last part of this review will concentrate on pheromone signaling associated with pheromone response.

The main function of the pheromone/receptor system in *S. cerevisiae* and *U. maydis* is to induce the expression of genes that encode proteins involved in attracting and directing the growth of mates toward each other. These proteins also play a role in the fusion of the conjugating cells and establishing an infectious plant pathogenic dikaryon in *U. maydis*. In agaricomycetes like *S. commune* and *C. cinerea*, hyphal fusions, or anastomoses, can occur between all vegetative hyphae and thus are thought to be independent of mating types. Therefore, the pheromone/receptor interaction is assumed to be relevant for events following fusion, specifically in the reciprocal exchange and migration of nuclei between the mates (84).

In the filamentous ascomycete *Neurospora crassa*, the differences between vegetative and sexual hyphal fusions have been investigated by screening the *N. crassa* genome for proteins known to play a central role in cell fusion during mating in *S. cerevisiae* (18, 26). No orthologs for Fus1, Fus2, Fig1, and Fig2 proteins were found (26). In *S. cerevisiae*, the expression of these proteins located at the tips of shmoos, the conjugating cells necessary for efficient mating, is regulated upon pheromone response. Instead, a pheromone-regulated membrane protein from *S. cerevisiae*, Prm1 (32), was shown to be part of the general fusion machinery in *N. crassa* (26). Deletion of the *prm1*-like gene led to 50% reduction of vegetative cell fusions and caused a complete block of fusion needed for sexual reproduction in the hook cells after fertilization, suggesting that vegetative and sexual hyphal fusions may share functions of the same proteins (17). A preliminary screening of the *S. commune* genome suggests a comparable situation in this basidiomycete; no orthologs are detected for Fus1, Fus2, Fig1, and Fig2 proteins, while a putative ortholog for Prm1 is found.

In *N. crassa*, mutations in the heterotrimeric G protein that were necessary for sexual development had no effect on vegetative fusions while effectively blocking sexual development (18). These observations point at the possibility of distinguishing two types of hyphal fusions. In agaricomycetes, this would lead to the question of whether it is possible to identify these two fusion events. One type would represent vegetative anastomoses independent of heterotrimeric G protein, whereas the other type would be hyphal fusions controlled by pheromone and receptor interaction and signaling through heterotrimeric G proteins. For sexually controlled fusion events in agaricomycetes, the fusion of a clamp cell with the peg formed at the surface of the subterminal cell also represents an example.

**FUNCTION OF G PROTEINS DURING MATING IN BASIDIOMYCETES**

Upon pheromone binding, the heterotrimeric G protein located at the plasma membrane, by virtue of the lipid tails in its Gα and Gγ subunits, will be activated by the ligand-bound GPCR that functions as a guanine exchange factor (GEF) for the Gα subunit. The conformational change associated with GPD to GTP exchange in Gα leads to release of Gα from the receptor and Gβγ subunit from Gα. Both the released Gα-GTP and Gβγ may activate downstream processes, out of which the Gβγ-activated signaling pathway of *S. cerevisiae* mating is known in detail (78, 102). From this point of view, the activation of downstream signaling in *S. cerevisiae* with heterologous expression of pheromone receptor and pheromone-encoding genes from *S. commune* (19, 31) and *C. cinerea* (73) strongly supports the idea that the respective heterotrimeric G protein is involved in downstream signaling in mating interactions of these fungi.

**Structure and function of Gα subunits in mating interactions.** In yeasts and filamentous ascomycetes, two and three Gα subunit-encoding genes are known, respectively (56). Three Gα subunit-encoding genes are also reported to be present in the basidiomycete *S. commune* (85, 120). Screening of the *S. commune* genome suggested the existence of an additional Gα gene not reported before. Previously, four Gα subunits had been reported only for *U. maydis* (88). All these proteins contain the domains typical for Gαs, the GTP-binding and GTPase domains (96). Additionally, the four *S. commune* Gα proteins share an N-terminal consensus sequence (MGX CXS/MGCXXS) with methionine, glycine, and cysteine sepa-
Note that in S. commune signaling in the agaricomycete S. commune, a hypothetical model of signaling components in the agaricomycete S. commune, has been identified in U. maydis, while an ortholog to Ste20 is detected. A homologous protein, Gpa1 (Ubc2 [46]), and in S. commune thn1 gene encodes an ortholog to regulators of G protein signaling (RGS), known as Sst2 in S. cerevisiae and FlbA in Aspergillus nidulans (20). The products of these genes regulate the rate of conversion of GDP-bound Gα to GDP-bound Gα, thus controlling the intensity and duration of G protein signaling by working as GTPase activating proteins (GAPs) on the Gα. The thin mutation, as in the constitutively active Gα and Gpc genes, reduces the growth of aerial hyphae and, when homoallelic in a dikaryon, inhibits fruiting body formation (20).

In basidiomycetes, it is not yet known which of the known Gαs interacts with pheromone receptors. In S. cerevisiae, Gpa1 is necessary for sexual development and interacts with both S. cerevisiae ligand-bound pheromone receptors, Ste2 and Ste3 (42). Deduced from positive results in heterologous pheromone and receptor expression studies of yeast, pheromone receptors Bbr1 and Bbr2 of S. commune interacted with S. cerevisiae Gpa1 (19, 31). In the heterologous expression of C. cinerea pheromone receptors, unmodified Gpa1p responded only weakly, while Gα chimeras containing the five last amino acids from mammalian Gα16 or from U. maydis Gpa3 performed better (73).

In the basidiomycete Cr. neoformans, two Gα proteins, Gpa2 and Gpa3, are involved in regulating pheromone signaling by interacting with the pheromone receptor (36). Gpa2 promotes mating, and Gpa3 inhibits the interaction. Genetic regulation is also different in that gpa2 expression is induced by pheromone signaling and gpa3 via the nutrient pathway (36). It has been suggested that involvement of two Gα subunits in sexual development could be evolutionary conserved among fungi that express several Gα proteins (36).

Screening of the S. commune genome with known Gα pro-

**FIG. 2.** Putative signaling components in fungal mating interactions. (A) The well-known components of the signaling pathway associated with mating in the ascomycete S. cerevisiae (78) and in the ustilagnomycete U. maydis (126), as well as the known and putative signaling in the agaricomycete S. commune, are shown. The arrows illustrate the direction of signaling from pheromone/receptor interaction via G protein and MAPK cascade to the transcription factor (TF). Note that in S. cerevisiae the Ste4/18 (βγ) complex interacts with Ste20 (a PAK-like kinase), which then signals further to the MAPK cascade containing the scaffolding protein Ste5. No Ste5 has been found in the S. commune genome, while an ortholog to Ste20 is detected. A homolog to Ste50, involved in mating response, invasive/filamentous growth, and osmotolerance, and acting as an adaptor that links the G protein-associated Cdc42-Ste20 complex to Ste11 in S. cerevisiae, has been identified in U. maydis (Ubc2 [46]) and in S. commune. (B) A hypothetical model of signaling components in the agaricomycete S. commune after G protein activation by pheromone/receptor interaction. The released G protein subunits activate a homologous MAPK cascade. Perceived functions include induction of enzymes necessary for hyphal fusions and septal dissolution, which might also respond to the cAMP-dependent Pka pathway. In addition to the MAPK cascade, Gβγ has been shown to activate Cdc42, thus controlling cytoskeletal rearrangement necessary for nuclear exchange and migration.
teins leads to detection of at least 11 Go-like proteins in addition to the four Gs with conserved GTP-binding and GTPase domains, further complicating the picture. Similar screening of C. cinerea and L. bicolor genomes also reveals several members of Go-like proteins. A high number of Go-like proteins has been reported for the genome of P. placenta (62). The function of these proteins is unknown, and the closer examination of the sequences for GTP binding and GTPase domains shows only weak similarity. The phylogenetic analysis of all putative P. placenta Gs separated the Go-like proteins clearly from the Gs with proper GTP binding and GTPase domains (62).

**G** protein-associated downstream pathways. The structure and function of Gβ and Gγ subunits of heterotrimeric G proteins have obtained less attention in agaricomycetes than the Gα subunits, due to the fact that the Gα subunit is thought to be involved in dissociation of the heterotrimeric G protein and in releasing Gβγ for MAPK induction upon pheromone binding (19, 73). However, experimental proof for either Gα or Gβγ signaling is still pending. In the sequenced genomes of C. cinerea, S. commune, and P. placenta, as well as in Lentinula edodes (56), there appears to be only one highly identical (90%) Gβ protein which also is conserved to Gβ subunits of ascomycetes (56). In addition, the genomes of the sequenced agaricomycetes contain several Gβ-like proteins with the typical WD-40 domains. The number of true Gγ-encoding sequences is probably two for agaricomycetes (56), and their amino acid sequence identity is close to 70%. In the S. commune genome, the two Gγs are located close to each other on the same scaffold and share 68% identity.

The mating of S. commune compatible strains leads to strong induction of the expression of pheromone- and receptor-encoding genes (112). A comparable phenomenon is seen in U. maydis, where the pheromone mfa1 transscripts increased when the cells were treated with a compatible a2 pheromone (126). This signaling could be attributed to MAPK- and cAMP-dependent signaling (16, 65, 126). While a MAPK cascade leads to activation of the transcription factor Prf1—which does not show high similarity to Ste12 of S. cerevisiae—cAMP signaling occurs via protein kinase A (Pka). In the ascomycete N. crassa, mating and fruiting body formation also involve two pathways: the MAPK cascade and cAMP signaling with Pka activation (56).

For agaricomycetes, Gα and Gβγ activation of MAPK signaling is expected to lead to phosphorylation of a transcription factor with similarity to S. cerevisiae Ste12 or U. maydis Prf1. The activated transcription factor then will repress/activate target genes necessary for changing the haploid hyphal growth pattern to B-regulated development and, ultimately, to dikaryotic growth and fruiting body development. As for the MAPK pathway, orthologs to the members of the S. cerevisiae MAPK cascade, including Ste20, Ste11, Ste7, Fus3, and Ste12, can be identified in S. commune (Fig. 2A). The complexity of the pheromone response pathway and cross talk with cAMP signaling in the basidiomycete U. maydis (44, 46, 126) prompted a search for other genes potentially involved in sexual reproduction. For cAMP signaling, the genes for Ras, adenylate cyclase, and the regulatory and catalytic subunits of Pka are also recognized in the S. commune genome.

**MONOMERIC GTPases Ras AND Cdc42**

A high number of small GTPases are suggested to be involved in regulating hyphal growth and morphogenesis (78). Among these proteins, two monomeric GTPases, Ras and Cdc42, have gained special attention. Ras is known to be involved in the regulation of cAMP/Pka signaling (52), which, in turn, has been shown to be involved in fruiting body development in C. cinerea and S. commune (99, 110, 111, 123). Cdc42 functions in pathways linking extracellular growth signals to the actin cytoskeleton, a response often initiated by binding of a signal molecule to a GPCR (95). The Ras and Cdc42 GTPases are conserved molecular switches, which are normally anchored to the plasma membrane by a C-terminal farnesyl or geranyl-geranyl tail. The ability to bind and hydrolyze GTP allows the GTPases to exist either in an active, GTP-bound form or in an inactive, GDP-bound form. Upstream regulatory proteins are involved in the cycling of Ras and Cdc42 between GTP and GDP binding forms and the GTPase-activating proteins (GAPs) facilitate efficient GTP hydrolysis, while guanine nucleotide exchange factors (GEFs) catalyze the exchange of bound GDP for GTP. Nucleotide dissociation inhibitors (GDIs) help to keep the small GTPases in soluble form (15).

**Ras proteins.** The function of Ras proteins encoded by two different genes, Ras1 and Ras2, has been extensively analyzed in S. cerevisiae (108), Cr. neoformans (68, 69, 114, 115, 118), and U. maydis (52, 64). Several aspects complicate the understanding of the function of Ras proteins, as Ras gene expression is often controlled by nutritional and environmental factors. Through activation of adenylate cyclase and increased cAMP concentrations, the Pka pathway is induced. However, this pathway is also controlled by a Gα subunit, and thus cross talk has to be considered (113). In Cr. neoformans, Ras1 is involved in regulation of early stages of mating, and it is necessary for pheromone production (115), while the cAMP production and Pka pathway are controlled by Gpap (118). In U. maydis, dominant/constitutively active Ras2 changed the growth pattern from yeast-like to filamentous, while dominant active Ras1 elevated pheromone expression, probably via increased cAMP levels (52, 64). Also, the expression of both Ras proteins varies. In Cr. neoformans, Ras1 expression is higher than Ras2 and the protein functions mainly overlap (114). Interestingly, in Cr. neoformans Ras1 is also an effector to Cdc24, the GEF of Cdc42, and in this way is involved in the regulation of actin organization at high temperatures (68, 69). In Schizosaccharomyces pombe with only one Ras gene, Ras protein is activated by two different GEFs, Efc25 and Ste6. Depending on which Ras GEF is binding, different effectors are activated: one couples Ras regulation to the MAPK pathway and the other to Cdc42 signaling (75). In U. maydis, the Ras2 GEF, sql2, with similarity to S. cerevisiae Cdc25, has been isolated as a suppressor of the morphology induced by a dominant/constitutively active gpa3 mutation, which again shows the interaction between Gα and Ras signaling (64).

In S. commune, two ras genes with 45% amino acid identity can be found. Ras1 is about 60% and Ras2 over 80% identical with C. cinerea, L. bicolor (79), and Suillus bovinus Ras1 and Ras2 proteins in GenBank, respectively. While Ras2 awaits functional analysis, constitutively active Ras1 in S. commune
(121), as well as the effect of deletion of a Ras-specific Gap gene, gap1 (97), has been investigated. These two mutations of the Ras signaling pathway would be expected to have a similar phenotype, since both mutants lead to accumulation of active, GTP-bound Ras. Indeed, the phenotypes are largely overlapping but not identical. Both mutations led to decreased hyphal growth and a disoriented, curving growth pattern. Homozygous crosses of Δgap1 strains produced a strong phenotype in dikaryons, with a failure in clamp formation. The trapped nucleus in the nonfused clamp-like structure subsequently was rescued by side branch formation originating from the mother cell (97). The growth of the branch and the outgrowing clamp cell seemed attracted toward each other, indicating involvement of pheromone recognition between the two monokaryotic cells. This view is supported by the fact that in unfused clamp cells, pheromone receptor expression is strongly upregulated (E. Kothe, unpublished data).

A third phenotype of gap1 deletions was seen in poorly developed pseudolumellae, altered hymenium formation, and lack of spore production during fruiting body development in homozygous crosses (97). Together, these results indicate that Ras1 has an effect on hyphal growth, clamp connection, and fruiting body development. The existence of the second ras gene in S. commune requires reinvestigation of the effect of Ras proteins on cAMP-, MAPK-, and Rho-dependent signaling and morphogenesis in order to reveal whether the two Ras proteins have different functions, as seen in U. maydis (64), or are more or less redundant in function, as is the case for the two Ras proteins in S. cerevisiae and Cr. neoformans. The characterization of Ras GEFs would also be informative for understanding the specificity of Ras signaling.

Cdc42 and actin. In the yeast S. cerevisiae, membrane-localized Cdc42 is required for proper bud site selection and for the development of mating projections where Cdc42 links pheromone-dependent Gβγ activation to polarized growth. Cdc42-specific GEF Cdc24 is involved in interaction with several proteins like Far1, Ste20, Bem1, and Bni1 (78). Through these interactions, GTP-bound Cdc42 activates weakly the MAPK cascade, but more importantly, it influences the proper distribution and orientation of the actin cytoskeleton for the tip growth of the conjugating cells. In yeast mating interactions, Far1p appears to play two central roles: it is necessary for cell cycle arrest but also for the polarized growth of the mates.

In S. commune, the ectopically expressed, constitutively active mutant allele of cdc42 regulates branch site selection and subsequent branch development of monokaryotic hyphae (116). In mating interactions, most of the hyphal fusions involve side branches that either grow toward each other or grow toward the hypha of the mate (84). These side branches show a strong actin cytoskeleton (M. Raudaskoski, unpublished data). The tip of the hook at clamp formation also shows actin accumulation during growth and fusion to the subapical cell (92), a process regulated by B mating type genes and Ras (80, 97). A role for Cdc42 in B-regulated mating interactions can be inferred from poor dikaryotization, along with irregular clamp connections in mating of strains both expressing a constitutively active cdc42 allele (116). Protein(s) should be identified that is needed to link the signal from pheromone/receptor interaction, and potentially also Ras, through Cdc42 to actin. Specifically, the respective GEF that can be postulated to play a central role leading to polarization of the actin cytoskeleton would be of interest.

PHEROMONE RESPONSE AND NUCLEAR MIGRATION

Mating in yeast cells and in U. maydis necessitates synchronization of the mitotic phase prior to the fusion of the conjugating cells. The cell cycle arrest is relevant to allow the nuclear fusion for becoming diploid in S. cerevisiae or to allow the establishment of a dikaryotic hypha waiting for signals to start the pathogenic growth in planta in U. maydis. In S. cerevisiae, the gene responsible for cell cycle arrest at the G1 phase of the mitotic cell cycle is Far1 (78). An ortholog to this gene is not found in the U. maydis genome (126) nor in the S. commune genomes. Instead, in U. maydis, several cell-cycle-associated genes were repressed in a microarray analysis of pheromone response, indicating cell cycle arrest in spite of the absence of Far1. In S. commune, a function in cell cycle arrest might also be envisioned from phenotypic analyses. When intercellular nuclear movements have been followed in living hyphae, or when such movements in fixed hyphae have been labeled for visualization of nuclei and microtubules, synchronous nuclear divisions were always observed in close connection with hyphal fusions (83, 84).

An interesting feature specifically well investigated in S. commune and C. cinerea is the nuclear exchange following hyphal anastomosis of cells with different B mating types (Fig. 1 and 2B). This B-specific nuclear migration was followed by looking for dikaryotization of distal hyphal parts in compatible matings. The intercellular nuclear migration in S. commune was characterized with a speed of 1 to 5 mm min⁻¹ (103), while speeds from 1 to 3 mm h⁻¹ for C. cinerea (49) and as fast as 40 mm h⁻¹ for Coprinellus congregatus (91) were reported. The reports based on cytological observations on nuclear migration indicated that the nuclei move with a speed 10 to 20 times their hyphal growth rate (71). The extensive microtubule tracks seen in migration hyphae around nuclei (84) suggest that microtubule tracks, together with microtubule-associated motors, could be used in intercellular nuclear exchange and migration. Genes for kinesins, performing mainly plus-end-directed transport, and dynein, for minus-end-directed transport, can be identified in the basidiomycete genomes. In U. maydis, 10 kinesin genes and 2 genes encoding the N-terminal and C-terminal parts of one dynein heavy chain have been identified (98). A similar situation with a bipartite dynein gene and several kinesins can also be found in S. commune (E. Kothe and M. Raudaskoski, unpublished data).

In S. cerevisiae, it is reported that a mating-specific Gø, Gpa1, interacts with kinesin-14 (Kar-3), a minus-end-directed microtubule-associated motor, in a pheromone-responsive way (125). The interaction regulates pheromone-induced nuclear migration to the shmoo tip. Gpa1 was immunoprecipitated with Kar3, showing protein-protein interaction, and Gpa1 and Kar3 were visualized at the shmoo tip after pheromone treatment. The use of Gpa1 mutants indicated that Gpa1 affects positively the association of Kar3 to the shmoo tip as well as the dynamics and orientation of microtubules. These results suggest that Gpa1 serves as an externally regulated position determinant anchoring Kar3, a role not previously known for Gø proteins (125).
In *S. commune* hyphal fusions, not only an actin cytoskeleton but also prominent microtubules extending from the nuclear spindle pole body toward the hyphal tip (84) are seen, reminiscent of the nuclear position at the yeast shmoo tip. In *S. commune*, this nuclear position might thus also be Gα dependent, separating sexual from vegetative hyphal fusions.

At the time of clamp fusion, in addition to actin cytoskeleton (92), strong microtubule tracks are involved in nuclear positioning at the fusion point (84, 86). Thus, there are two B-regulated processes in *S. commune* with similarities to the Gα-controlled processes in conjugating yeast cells that warrant further investigation of nuclear positioning during hyphal fusion and clamp formation. In animal cells, Gα proteins are suggested to link extracellular signals to microtubule cytoskeleton more dynamic (124). Thus, new functional analyses which could finally show whether G protein interacting with the pheromone receptor. It would also allow an understanding of the tetrapolar versus bipolar mating systems in fungi but will also link up to general signal transduction pathways common to all fungi and even to all eukaryotes.

**DOWNSTREAM TARGETS FOR A AND B MATING TYPE GENES**

Downstream of the major regulators of sexual differentiation, the A and B mating type genes, several target genes have been identified. In *C. cinerea*, *num1* was isolated by REMI mutagenesis and screening for altered nuclear migration. The mutation led to unilateral mating, where the mutant acts as a nuclear donor. The gene *num1* encodes a 217-amino-acid protein with leucine zippers at the N and C termini and a coiled-coil domain in the C-terminal half. Its transcription is down-regulated by activation of the B-regulated pathway. On the basis of its structure, Num1 is thought to be a part of a larger protein complex (58). In *S. commune*, differential RNA display identified *brt1*, which is repressed during B-regulated development, and *mat1*, upregulated in fully compatible matings. The gene *brt1* appears to encode a translation initiation inhibitor (54), while the product of *mat1* shows similarity to a peptide transporter (55). These genes are detected in the genome of other agaricomycetes, but no further functional characterization is available.

In *C. cinerea*, two genes have been shown to be involved in regulating clamp connection formation: *pcc1* and *clp1* (38, 66). Orthologs to the *pcc1* gene are found in other fungal and eukaryotic genomes, while *clp1* appears to be restricted to basidiomycetes (94). The gene *pcc1* encodes a transcription factor with a HMG box and is suggested to repress A-regulated genes in the homokaryon, while the A pathway activates the expression of *clp1* in the dikaryon. The *clp1* gene product is thought to repress the expression of *pcc1* in the dikaryon, which in turn would allow the transcription of genes necessary for clamp formation (45, 94).

**FUTURE PROSPECTS**

Present knowledge on the structure and function of mating type genes in fungi has already shown that homeodomain transcription factors are involved in the regulation of mating throughout the fungal kingdom (11). Although the pheromone/receptor system seems to be basically conserved, differences in its regulation, the newly *in silico* discovered receptor-like proteins, and cross talk of several intracellular signal transduction pathways make the understanding of the origin and function of the pheromone receptor system a challenge, especially in the highly complex systems of agaricomycetes.

In *S. cerevisiae* (42), *Schizosaccharomyces pombe* (72), and *N. crassa* (8), the genes of MAT loci regulate the transcription of pheromone- and receptor-encoding genes in haploid cells. In agaricomycetes, the receptor- and pheromone-encoding genes of the B mating type loci themselves regulate the specificity of the reciprocal exchange and migration of nuclei at the beginning of mating interactions (11), and they are involved in morphogenesis at the stages of clamp formation and fruiting body development. Noteworthy is that during clamp formation the nuclei with different A and B mating type genes become separated from each other, one into the forming clamp cell and the other into the new subapical cell, thus leading to a temporarily monokaryotic stage (Fig. 1), reminiscent of the situation at the beginning of a yeast mating in which pheromone/receptor signaling plays a central role. This allows future investigation of the output of pheromone signaling with a new direction, potentially even with regard to the still completely unanswered question of how the different nuclei might be recognizing each other, organizing themselves into pairs at the beginning of mating, and then maintaining the pairing in the dikaryon (Fig. 1). This leads to another interesting question about the function of both A and B mating type genes in agaricomycetes without clamp connections on dikaryotic hyphae, like *Amanita regalis*, *Suillus bovinus* (93), and *Tricholoma species* (59).

In the pheromone response pathway of agaricomycetes, the deletion or silencing (14, 33, 67) of the heterotrimeric G-protein α subunits would clarify which one is or which ones are interacting with the pheromone receptor. It would also allow functional analyses which could finally show whether G protein activation is regulating hyphal fusions. In contrast, G protein signaling limited to the time after hyphal fusions would then be responsible only for the induction of nuclear movements and breakdown of hyphal cross walls (84) necessary for nuclear exchange and migration between the mates. In addition to gene silencing, it seems necessary and possible now to identify target genes by transcriptome analyses. This will allow a more complete view of the regulatory pathways involved in sexual differentiation in basidiomycetes. The answers that can now be obtained in addition to *in silico* analyses of the fungal genomes not only will allow an understanding of the tetrapolar versus bipolar mating systems in these fungi but will also link up to general signal transduction pathways common to all fungi and even to all eukaryotes.

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