Evolution of the Mating Type Locus: Insights Gained from the Dimorphic Primary Fungal Pathogens Histoplasma capsulatum, Coccidioides immitis, and Coccidioides posadasii

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Sexual reproduction of fungi is governed by the mating type (MAT) locus, a specialized region of the genome encoding key transcriptional regulators that direct regulatory networks to specify cell identity and fate. Knowledge of MAT locus structure and evolution has been considerably advanced in recent years as a result of genomic analyses that enable the definition of MAT locus sequences in many species as well as provide an understanding of the evolutionary plasticity of this unique region of the genome. Here, we extend this analysis to define the mating type locus of three dimorphic primary human fungal pathogens, Histoplasma capsulatum, Coccidioides immitis, and Coccidioides posadasii, using genomic analysis, direct sequencing, and bioinformatics. These studies provide evidence that all three species possess heterothallic bipolar mating type systems, with isolates encoding either a high-mobility-group (HMG) domain or an α-box transcriptional regulator. These genes are intact in all loci examined and have not been subject to loss or decay, providing evidence that the loss of fertility upon passage in H. capsulatum is not attributable to mutations at the MAT locus. These findings also suggest that an extant sexual cycle remains to be defined in both Coccidioides species, in accord with population genetic evidence. Based on these MAT sequences, a facile PCR test was developed that allows the mating type to be rapidly ascertained. Finally, these studies highlight the evolutionary forces shaping the MAT locus, revealing examples in which flanking genes have been inverted or subsumed and incorporated into an expanding MAT locus, allowing us to propose an expanded model for the evolution of the MAT locus in the phylum Ascomycota.

In the fungal kingdom, sexual reproduction is regulated by a specialized genomic region known as the mating type (MAT) locus (12, 13). This important genomic feature has been most extensively studied in the largest of the fungal phyla, the Ascomycota. Most commonly, the sexual members of this phylum have a bipolar mating type system, where strains are one of two mating types. For mating to occur in these heterothallic species, cells of differing mating types must come together. In those species where the MAT locus has been characterized at a molecular level, this process is regulated by a bipolar system: cells normally carry one of two different alleles of the MAT locus. These alleles, known as idiomorphs, contain unrelated sequences that encode different transcription factors. In the euascomycetes, which include the medically important dimorphic pathogens and the majority of the ascomycete molds, one MAT locus allele encodes a high-mobility-group (HMG) domain-type transcription factor, and the other allele encodes an α-box domain transcription factor. When two isolates of a species with differing MAT loci come together under appropriate conditions, sexual reproduction can proceed. Importantly, for many ascomycete species, a sexual cycle has never been observed, yet analysis of the genomic sequence has revealed the presence of potentially functional mating type loci.

Over the past decade, our understanding of the structure of fungal mating type loci has blossomed, and elegant studies have begun to elucidate the evolutionary processes that fashioned these unusual genomic structures (4, 11, 18, 35). Despite the quantity of information available regarding MAT locus structures, there is a paucity of information regarding the Ongenales, one of the most medically important orders of fungi. The members of the Ongenales include a number of dimorphic primary pathogens, including Histoplasma capsulatum, Coccidioides immitis, and Coccidioides posadasii, all capable of causing life-threatening systemic mycoses (6, 38). Although the MAT locus has not been characterized in these species, there is strong evidence that functional loci exist and that they play important roles in the life cycle of these organisms.

While sexuality is uncommon among pathogenic fungi, H. capsulatum is one of the exceptions to this generalization. This primary pathogen is distributed worldwide in association with soil enriched with bird and bat guano and is thought to represent multiple cryptic species (19). In the soil, the mycelial...
phases of sexual reproduction are unknown. How mating type may regulate this dimorphic transition and increase virulence is unknown.

In contrast to H. capsulatum, which is distributed worldwide, the Coccidioides species are restricted to the Americas (32). The soilborne saprobic multicellular hyphal form of this fungus septates into single-celled arthroconidia that can become airborne (such as during dust storms) and be inhaled to produce life-threatening infections in otherwise healthy individuals (6). Although a sexual cycle has never been observed in the laboratory, molecular phylogenetic analysis revealed evidence supporting sexual reproduction (3, 21). These studies were key in revealing two extremely closely related yet independent species, C. immitis and C. posadassii, which are distinct from each other and represent discrete populations undergoing independent sexual recombination (9, 20, 21). While these studies provide evidence that C. posadassii and C. immitis undergo a sexual cycle in nature, this has never been observed in the laboratory. Furthermore, as neither mating nor the MAT locus has been identified, it is unknown whether different mating types have distinct pathogenic capacities, although different mating types have been recognized for H. capsulatum. The structure of the MAT loci of the fungi of the order Onygenales is therefore an important topic to address, both relative to the role of these fungi as pathogens and from an evolutionary perspective. Here, we describe an analysis of the MAT locus structure of the dimorphic pathogens H. capsulatum, C. posadassii, and C. immitis by combining bioinformatic analyses of genomes, direct sequence confirmation, PCR analysis of multiple isolates of each species, and comparative analyses with other fungi of the phylum Ascomycota. Our findings have revealed that all three species have canonical euascomycete bipolar MAT locus structures, with strains known or predicted to be of opposite mating types encoding either an HMG box domain and an α-box domain transcription factor at this locus. Potential mating partners have been identified for both Coccidioides species, and the + and − mating types of Histoplasma have been identified as belonging to the α-box and HMG types, respectively. Finally, this work provides critical information contributing to an extended model of MAT locus evolution in the most populated of the fungal phyla.

MATERIALS AND METHODS

Strains and media. The reference strains used in this study were H. capsulatum strains G217B, ATCC 22635, ATCC 22636, G186AR, and G217B sw8-23, C. immitis strains RS and H538.4, and C. posadassii strains Silveira, 1037, C735, and C634. C. immitis strain H538.4 is an environmental isolate, while the other isolates of Coccidioides are clinical derivatives. Strains were grown under modified biosafety level 2 (BSL2) or BSL3 conditions as appropriate. H. capsulatum strains ATCC 22635 (MAT1−/−) and ATCC 22636 (MAT1+/+) were received directly from the American Type Culture Collection as mycelial frozen stocks and were grown on solid medium at room temperature and on Histoplasma macrophage medium plates at 37°C under 5% CO2 (39).

Genomic DNA isolation. For H. capsulatum genomic DNA isolation, 100 ml of yeast culture of strains ATCC 22635 and ATCC 22636 was grown in Histoplasma macrophage medium at 37°C under 5% CO2 at 150 rpm for 3 days. Genomic DNA was isolated using a QIAEGN genomic DNA kit according to the manufacturer’s instructions. The genomic DNA was isolated from Coccidioides immitis and Coccidioides posadassii mycelia grown for 4 days in glucose-yeast extract liquid medium (Difco) as previously described (30).

Sequencing, assembly, and bioinformatics. Sequencing reactions were performed with an MJ Research thermal cycler using standard BigDye Terminator chemistry (Applied Biosystems) and analyzed on a PE3700 96-capillary sequencer. Sequence reads were assembled using Sequencher (Gene Codes Corporation, Ann Arbor, MI). Additional analysis of the data was performed using BLAST (1) and MacVector (MacVector Inc., Cary, NC). Based on the initial annotation of the end sequences, oligonucleotides were selected to close gaps in the sequence coverage by primer walking. Genes were annotated based on homology to the existing annotation in GenBank. Dot plots were generated using DOTTPUT from the EMBOS package.

PCR. Primers JOHE13616 and JOHE13619 were used to amplify the entire H. capsulatum MAT locus. Primers JOHE13950 and JOHE13951 were used to amplify the C. immitis and C. posadassii MAT1−/− idiomorph. Primers JOHE18241 to JOHE18248 were used for MAT diagnostics. PCR analysis was performed using the ExTaq polymerase blend (Miris) with primers given in the following list: JOHE13618, 5′-AGG CCA TAA CAC TGA CGA TAT-3′; JOHE13619, 5′-GCA TTA CAT CGC AGC ACA TTG-3′; JOHE13950, 5′-GTC AGC AAA CCC GGA GTA-3′; JOHE13951, 5′-AAC AAA CTT TCG CAG GCA AGG-3′; JOHE18241, 5′-TCG TCT ACT TTG GAC TTC GGA C-3′; JOHE18242, 5′-CTG CTA TTG CTT GCT CAG GTA C-3′; JOHE18243, 5′-CAG CCA ATG ACT GGT TCT AAG G-3′; JOHE18244, 5′-TTT ACG GGG ACG ATA CTG GTA G-3′; JOHE18245, 5′-AGG AAA CGA TGT CTC TGC G-3′; JOHE18246, 5′-TAC AGC AAG GAG GTA ACT TGG G-3′; JOHE18247, 5′-AAC AAG CAA CGA ACA AAG CTT C-3′ and JOHE18248, 5′-AGG TGA GAG CCG AAC AAT GAG G-3′.

H. capsulatum mating assays. ATCC 22635 and ATCC 22636 were prepared for mating tests by inoculating mycelia from soil agar plates into Sabouraud dextrose broth (Difco). The mating of each of these strains was also observed against that of two common laboratory strains, G217B (MAT1−/−) and G186AR (MAT1−/−). Two different G217B isolates were used in mating tests. The first isolate was originally a kind gift of William Goldman to the Sil laboratory and has undergone significant passage. The second G217B isolate was obtained directly from the American Type Culture Collection (ATCC 26052) and presumably has not been passaged as extensively. Mating between ATCC 22635 and ATCC 22636 and the G217B and G186AR strains was tested by streaking pairwise combinations of each strain on AlphaCol agar (MP Bioclims), sealing the plates with parafilm, and incubating the plates at 25°C. Self-mating controls for each strain were also performed. After 3 weeks, the plates were observed by eye and microscopically for the presence of perithecia or any type of mating structure. However, no mating structures were observed, even after several months of incubation.

Diagnostic amplification of the Histoplasma and Coccidioides predicted MAT regions. PCR amplification was conducted on each of six Coccidioides DNA templates, five H. capsulatum templates, and one negative control for each pair of primers. For H. capsulatum, MAT1-1 (α-box)-specific PCR employed primers JOHE18245 and JOHE18246 while MAT2-2 (HMG domain)-specific PCR employed primers JOHE18247 and JOHE18248. For Coccidioides, MAT1-1 (α-box)-specific PCR employed primers JOHE18241 and JOHE18242 while MAT2-2 (HMG domain)-specific PCR employed primers JOHE18243 and JOHE18244. Thermal cycling consisted of a 2-minute denaturation step at 95°C, followed by 30 cycles of denaturation at 94°C for 15 seconds, annealing at 55°C for 15 seconds, and extension at 72°C for 3 min and 30 seconds. After the completion of the 30 cycles, the samples were held at 72°C for 10 min. Amplicons were resolved by gel electrophoresis.
Phylogenetic analyses. For the species tree (see Fig. 5), the best reciprocal orthologs were identified via InParanoid (31) and combined with single-linkage clustering to identify gene sets with a single member per species producing 662 orthologous gene families. Protein alignments were constructed with ProbCons (8) and filtered with Gblocks (5), and phylogenetic trees were inferred with MrBayes (33) for each gene family. Gene trees were built with the topological constraint of Schizosaccharomyces pombe to be basal in the ascomycetes and with the hemiascomycetes and euascomycetes as a monophyletic group. A consensus supertree representing the putative species tree was built from the individual gene trees by using Clann (7). A species tree was also built with MrBayes by using a concatenated alignment of codons of 100 randomly selected ortholog families. The consensus supertree and the concatenated data set provided identical topologies.

MAT locus gene trees were constructed from ProbCons alignments of the predicted proteins and filtered for best-aligned blocks with Gblocks, and phylogenetic trees were inferred and bootstrapped with MrBayes and PHYML (5, 8, 16, 33). Trees were visualized with TreeViewX (29) and edited with Adobe Illustrator.

Nucleotide sequence accession numbers. The novel mating type locus sequences have been submitted to GenBank under accession numbers EF472255 and EF472256 (H. capsulatum) and EF472257, EF472258, and EF472259 (C. immitis and C. posadasi).
Employing this new sequence, we designed Coccidioides MAT1-1 contrast, H538.4, one of the two strains from which we were the control C. immitis C735. However, no product was obtained for C. posadasii HMG domain alleles, as in the strains in an effort to identify the opposite mating type.

were employed to amplify this genomic region from alternative the predicted locus (JOHE13950 and JOHE13951, respectively) tempts, these strains remained recalcitrant and no PCR prod-
correspond to

Candidate mating type locus in Coccidioides. Extending our analysis to the related primary pathogens C. immitis and C. posadasii, we searched the genomes in progress at the Broad Institute and The Institute for Genomic Research (genome strains RS and C735, respectively). In contrast to that observed with our original search of the Wu24 H. capsulatum genome, BLAST analysis using A. fumigatus SLA2 and APN2 revealed that they lie adjacent to a predicted HMG domain-type MAT1-2 allele (Fig. 1). Comparison of the two sibling species reveals that the encoded proteins are almost identical, with only three changes across the predicted 339-amino-acid protein. Again, the structure of this MAT locus is consistent with a classic ascomycete bipolar system, except in this case, the presumed second mating type encodes a MAT-1 alpha-box idiomorph. Once again, primers anchored in the APN2 and SLA2 genes flanking the predicted locus (JOHE13950 and JOHE13951, respectively) were employed to amplify this genomic region from alternative strains in an effort to identify the opposite mating type. Using this approach, ~5-kb fragments were obtained from the control C. immitis strain RS sample in addition to C. posadasii strains Silveira and 1037. Sequencing of the amplified fragments from C. posadasii strains Silveira (GenBank accession no. EF472258) and 1037 (GenBank accession no. EF472257) showed that each encodes almost identical MAT1-2 HMG domain alleles, as in the C. posadasii genome strain C735. However, no product was obtained for C. immitis strain H538.4 or C. posadasii strain C634, suggesting that they may correspond to MAT-1 alpha-box isolates. Despite repeated attempts, these strains remained recalcitrant and no PCR product could be detected.

During the course of this work, additional C. immitis genomes for strains H538.4 and RMSCC2394 were sequenced at the Broad Institute. Subsequent bioinformatic analyses of these isolates revealed that like the previous two Coccidioides genomes, RMSCC2394 is a MAT1-2 HMG domain isolate. In contrast, H538.4, one of the two strains from which we were unable to amplify MAT by using a PCR-based approach, encoded a MAT-1 alpha-box idiomorph as we had hypothesized. Employing this new sequence, we designed Coccidioides-spe-
cific diagnostic primer sets to amplify either the MAT1-1- or the MAT1-2-encoded transcription factor and facilitate mating type determination (Fig. 2). In addition to confirming the known mating types of C. immitis strains RS and H538.4 and C. posadasii strains Silveira and 1037, this approach revealed that the previously untypeable C. posadasii strain C634 corresponded to the missing MAT1-1 mating type.

Inspection of the open reading frames encoded by the alpha-box and HMG domain MAT genes revealed that both genes are apparently intact, with no evidence of loss-of-function mutations in any of the strains analyzed. This is consistent with previous population genetic studies indicative of sexual recombination and provides additional support for the hypothesis that an extant heterothallic bipolar sexual cycle remains to be defined for both species. The identification of strains of opposite mating types in both C. immitis and C. posadasii now provides a platform from which to initiate attempts to characterize the sexual cycle in these primary pathogens, an aspect of the life cycle that is relevant both as a tool in the laboratory and from an epidemiological perspective.

A step in the evolution of sex chromosomes: expansion of a MAT locus. Following the identification of Coccidioides MAT1-1 isolates, we sought an explanation for why the original PCR-based efforts were unable to identify this second mating type. Comparison of the genomic regions surrounding the C. immitis and C. posadasii MAT loci showed 100% synteny with the gene order in A. fumigatus and A. oryzae (Fig. 1). This also holds true when we compare the structures found in the genome project at the Broad Institute of the next closest sequenced relative, Uncinocarpus reesii (30), which encodes a MAT1-1 alpha-box idiomorph. As seen for C. immitis, C. posadasii, A. fumigatus, and A. oryzae, this member of the Onygenales has the COXI3-APN2-MAT-SLA2 mating type locus region arrangement.

It was not until a pairwise comparison was performed between the MAT1-1 and MAT1-2 idiomorphs of Coccidioides that the most significant changes to this locus became apparent. In contrast to the other species, in Coccidioides, the dissimilarity between the two idiomorphs expands beyond the HMG domain and alpha-box regions to encompass the adjacent APN2 and COXI3 gene pair (Fig. 3). Analysis of the amplified MAT regions revealed them to be atypically large for this type of structure in the Ascomycota, with the expanded MAT1-1 idiomorph of strain H538.4 being 8,062 bp in length while the MAT1-2 idiomorph of strain RS is 9,037 bp (Fig. 1). This is a clear example of the acquisition of genes adjacent to MAT being incorporated into the locus, expanding this important region by increasing both size and gene content. There are direct parallels between what we see here and the model we have previously proposed for the expansion of the MAT locus of the basidiomycete pathogen Cryptococcus neoformans, whose MAT locus, which is >100 kb in length and encodes more than 25 genes, is even larger than that of Coccidioides (11).

Remarkably, in this case, there is little evidence to suggest that the formation of MAT-specific alleles of captured genes is a slow and gradual process. Phylogenetic analysis of the APN2 gene showed that contrary to this gene forming closely related yet distinct MAT1-1 and MAT1-2 clades, these clades were instead dramatically different. When the C. immitis MAT1-2 alleles are not considered, this gene shows phylogeny as is
expected for any gene in these species (Fig. 4). In contrast, the \textit{C. immitis} and \textit{C. posadasii} MAT1-2 \textit{APN2} alleles form a clade basal and separate from that of the rest of the euascomycetes. How such a relationship has developed is unknown, with this example almost resembling a case of horizontal gene transfer perhaps the changes have a role in a new conformational structure. In contrast to the \textit{APN2} gene, the \textit{COX13} gene, which has also been incorporated into \textit{MAT}, has been subject to more-gradual divergence of the two \textit{MAT} alleles or more-recent gene conversion (data not shown).

The \textit{Coccidioides} MAT locus therefore represents a transitional form of the mating type locus and has evolved beyond the classical ascomycete idiomorphic form to exist instead as a type of hybrid structure. In addition to the continued presence of the idiomorphic region, since its divergence from \textit{H. capsulatum}, this structure now has a clearly delineated section that contains divergent alleles of two genes. This discovery helps to explain aspects of the unusual locus of \textit{Candida albicans}, where, in addition to the mating type transcription factors, three additional genes are now present (17, 35). These may have been enveloped by \textit{MAT} in the same way as has occurred in \textit{Coccidioides}. In the case of \textit{C. albicans}, the genes encode a phosphatidylinositol 4-kinase, a poly(A) polymerase, and an oxysterol binding protein, which are unrelated to the genes that have been incorporated into the mating type locus of \textit{Coccidioides}. Interestingly, these genes have not been incorporated into the \textit{MAT} locus of \textit{H. capsulatum} but instead have been retained as flanking sequences. Other examples of novel genes resident in the \textit{MAT} locus are also known. These include \textit{Neurospora crassa} and, most poignantly, the basidiomycetes \textit{C. neoformans} and \textit{Cryptococcus gattii}, where the \textit{MAT} locus has expanded to span more than 100 kb and more than 25 genes (at least some of which do not function directly in sexual reproduction). Whether the incorporation of originally flanking genes into the \textit{Coccidioides} MAT locus, and their subsequent divergence, contributes to cell identity or sexual reproduction in \textit{Coccidioides} or is simply a consequence of their close juxtaposition to the ancestral \textit{MAT} locus remains to be tested experimentally.

### Evolution of the \textit{MAT} locus in ascomycetes

The findings presented here provide a molecular description of the mating type locus for three thermally dimorphic fungal species, \textit{Histoplasma capsulatum}, \textit{Coccidioides immitis}, and \textit{Coccidioides posadasii}, all of which are primary pathogens of humans. A sexual cycle has been previously described only for \textit{H. capsulatum} (23, 24), but population genetic studies suggest that cryptic sexual reproduction is likely to occur in \textit{Coccidioides} as well (21). The fact that the two alleles in \textit{H. capsulatum} encode either an HMG domain protein or an \(\alpha\)-box protein, and that these alleles are present only once in the genome, is in full accord with the description of \textit{H. capsulatum} as having a heterothallic sexual cycle. That the \textit{MAT} loci of \textit{C. immitis} and \textit{C. posadasii} are arranged similarly allows the predication that the sexual cycle will also be heterothallic in these two pathogenic species. Importantly, none of the three species contain fused \textit{MAT} loci or evidence of both alleles present within the genome, and thus, all three are molecularly distinguished from homothallic species that have been defined in the fungal king-
dom, including those with silent mating type cassettes in which mating type switching can occur. Comparison of the \textit{MAT} loci of \textit{C. immitis} and \textit{C. posadasii} with that of \textit{Uncinocarpus reesii}, the closest extant species with a defined sexual cycle, further supports the assignment of \textit{MAT} in the pathogenic sibling species and buttresses the hypothesis that an extant sexual cycle remains to be defined.

Our analysis of the gene structure of all the sequenced \textit{MAT} alleles provides no evidence of loss-of-function mutations in either the HMG domain or the $\alpha$-box gene, indicating that these may still contain functional alleles and have not been subject to the pseudogene formation that would be predicted to occur if these species had evolved to be asexual. The molecular definition of the \textit{MAT} locus and the description of molecular reagents that can be used to establish the mating types of strains by rapid PCR analysis provide approaches that can now be applied to understand some of the mysteries surrounding these pathogens. These include the interesting finding that sexual fecundity is rapidly lost during laboratory passage of \textit{H. capsulatum} isolates, the link between mating type and virulence potential in \textit{H. capsulatum}, and the role that mating or mating type may have played in the speciation and geographic expansion of \textit{Coccidioides} species from North America to Central America and South America. Perhaps most significantly, these discoveries can also be applied to the generation of congenic strain pairs for genetic and virulence studies of these dangerous pathogens.

The elucidation of the molecular nature of the \textit{MAT} locus for \textit{H. capsulatum}, \textit{C. immitis}, and \textit{C. posadasii} also provides considerable insight into the evolution of mating type loci in fungi (Fig. 5). These studies reveal that the two idiomorphic \textit{MAT} alleles in all three species encode either an HMG domain or an $\alpha$-box transcription factor, similar to those of other euascomycete fungal species in which the \textit{MAT} locus has been characterized. A central question is the molecular nature of the ancestral \textit{MAT} locus in the fungal kingdom.

Ohno first hypothesized that sex-determining regions of the genome arose originally as autosomal genes which were incorporated into either a mating type locus or a sex chromosome which then expanded (28). Several types of genes have previ-
ously been found to be present in mating type loci, including those encoding key transcriptional regulators of two classes, the homeodomain proteins (as a pair of interacting proteins of HD1 and HD2 classes), and the HMG domain and α-box proteins. In a striking example in hemiascomycetes, namely, *Candida albicans*, all four of these genes are present as α1 and α2 in the MTLα locus and α1 and α2 in the MTLα locus (17, 35). Recent phylogenetic comparisons have revealed that the HMG domain protein α2 has been lost in the *Saccharomyces cerevisiae* and senso stricto lineages from a common ancestor in which all four genes were present (4).

In the euascomycete lineage, HMG domain and α-box domain genes are found in the two opposite MAT idiomorphs, but the homeodomain genes are not present (Fig. 5). In contrast, in the basidiomycete lineage, paired homeodomain genes are present in the MAT locus of all species analyzed thus far (*Ustilago maydis*, *Ustilago hordei*, *Schizosaccharomyces pombe*, *Coprinus cinereus*, *C. neoformans*, and *C. gattii*), but HMG domain or α-box genes are not. A unifying hypothesis is that there are two ancestral fungal sex determinants (one is paired HD1/HD2 homeodomain genes and the other the α-box and HMG domain pair) and that one or the other or both have been lost from, or acquired by, the MAT locus in different lineages (Fig. 5). In some cases, subsequent gene loss events have reduced the number from four to three of these genes, such as in *S. cerevisiae* and related species (4). It is possible that the additional sex-determining genes that are present in the MAT locus of some species are instead located elsewhere in the genome in other species and yet remain under the control of the MAT locus, such as has been shown for non-MAT HMG genes in *U. maydis* (2). Thus, homeodomain genes critical for cell identity and sexual reproduction may remain to be defined in euascomycetes, and HMG domain and α-box proteins may remain to be defined in members of the phylum *Basidiomycota*.

Whether the ancestral MAT locus contained all four sex determinants or whether different lineages coopted one pair or, in some cases, both pairs of determinants remains to be established. One approach to distinguish between these and other models will be to characterize the MAT locus from more-divergent fungal lineages, such as zygomycetes. The recent sequencing of the *Phycomyces* genome, with its known sexual cycle and meiotic map, will allow the nature of the MAT locus to be defined for comparison with that of ascomycete and basidiomycete fungal species.

Finally, an important consequence of our studies is to enable further insight into the evolution of the MAT locus and homothallic and heterothallic sexual cycles in other euascomycete species. In two notable species, *A. nidulans* and *A. fumigatus*, the MAT locus is organized in an unusual fashion. *A. nidulans* harbors both MAT idiomorphs in the genome of a single isolate at unlinked genomic loci, giving rise to a homothallic sexual cycle. By contrast, in *A. fumigatus*, the two idiomorphs are present at the same genomic locus, and any given isolate harbors only one or the other, but never both. This would be consistent with a heterothallic sexual cycle that remains to be described. Galagan and colleagues have hypothesized that the ancestral organization of the MAT locus in aspergilli was one where each isolate contained a locus that included both an HMG domain and an α-box-encoding gene in a homothallic ancestral species (15). Alternatively, the ancestral species may have been a heterothallic species from which both heterothallic and homothallic descendants have been derived. Based on our studies, we propose that the ancestor of the *Ozygonales* and aspergilli was heterothallic, and if the hypothesized homothallic ancestor to the aspergilli existed, it likely arose after the divergence of the lineages. Furthermore, we hypothesize that the ancestor of euascomycetes and hemiascomycetes may very well have resembled the current structures in *C. albicans* that include two homeodomain genes, paired with either an HMG domain gene or an α-box gene, but exclude the presence of the Candida-acquired PIK, OBP, and PAP genes. Further studies of fungal MAT loci will be required to fully understand this complex evolutionary system.

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