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Evolution of the *MAT* locus and its Ho endonuclease in yeast species

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The genetics of the mating-type (*MAT*) locus have been studied extensively in *Saccharomyces cerevisiae*, but relatively little is known about how this complex system evolved. We compared the organization of *MAT* and mating-type-like (*MTL*) loci in nine species spanning the hemiascomycete phylogenetic tree. We inferred that the system evolved in a two-step process in which silent *HMR/HML* cassettes appeared, followed by acquisition of the Ho endonuclease from a mobile genetic element. Ho-mediated switching between an active *MAT* locus and silent cassettes exists only in the *Saccharomyces sensu stricto* group and their closest relatives: *Candida glabrata*, *Kluyveromyces delphensis*, and *Saccharomyces castellii*. We identified *C. glabrata MTL1* as the ortholog of the *MAT* locus of *K. delphensis* and show that switching between *C. glabrata MTL1a* and *MTL1 α* genotypes occurs *in vivo*. The more distantly related species *Kluyveromyces lactis* has silent cassettes but switches mating type without the aid of Ho endonuclease. Very distantly related species such as *Candida albicans* and *Yarrowia lipolytica* do not have silent cassettes. In *Pichia angusta*, a homothallic species, we found *MAT α 2*, *MAT α 1*, and *MAT α 1* genes adjacent to each other on the same chromosome. Although some continuity in the chromosomal location of the *MAT* locus can be traced throughout hemiascomycete evolution and even to *Neurospora*, the gene content of the locus has changed with the loss of an HMG domain gene (*MAT α 2*) from the *MAT α* idiomorph shortly after *HO* was recruited.

Mating in *Saccharomyces cerevisiae* is a choreographed fusion of two haploid cells with opposite mating types to form a diploid zygote. The mating type of a haploid cell is determined by its genotype at the mating-type (*MAT*) locus on chromosome III. The two variants of the *MAT* locus, *MAT α* and *MAT α* , are referred to as idiomorphs rather than alleles because they differ in sequence, size, and gene content (1). Characterization of homothallic and heterothallic strains of *S. cerevisiae* by Winge and Lindgren (2) led to the discovery of genetic loci controlling homothallism and ultimately to the cassette model of mating-type switching proposed by Herskowitz and colleagues (3). The cassette model, which has been thoroughly verified by experimentation (4, 5), states that a haploid cell can switch genotype at the *MAT* locus (from *MAT α* to *MAT α* or vice versa) by a gene-conversion process that involves replacing the genetic information at *MAT* with information copied from one of two silent cassette loci, *HML α* or *HMR α* . The gene conversion is initiated at a double-stranded DNA break made by the Ho endonuclease, encoded by the *HO* gene on chromosome IV, which cuts at a site that marks the boundary between the Y sequences unique to the *MAT α* or *MAT α* idiomorphs and the shared Z sequence flanking them. *HO* is expressed only in cells that have budded once, which means that only mother cells switch mating type and neighboring cells in a colony can mate (4). Hence, most natural isolates of *S. cerevisiae* are diploid and phenotypically homothallic.

The focus on *S. cerevisiae* as a model organism has had the consequence that relatively little investigation has been made

into the mating systems of other species in the hemiascomycetes, the group of fungi that includes *Saccharomyces*. The question of whether a mating-type system similar to that of *S. cerevisiae* is found in other hemiascomycetes has become pertinent recently because of the discovery of mating-type-like (*MTL*) loci in *Candida* species that had been regarded as asexual (6–9). The *Candida albicans* genome sequence (10) includes an *MTL* locus but not silent cassettes or a *HO* endonuclease gene. In *Candida glabrata*, three *MTL* loci (*MTL1–3*) were reported (9). Here we have used a genomics approach to study the organization and evolution of the *MAT* locus in nine hemiascomycete species. By combining our sequence data with information from previous reports and genome projects, we show that the Ho endonuclease was a relatively recent addition to a preexisting, cassette-based switching system, which in turn evolved from a simpler system without cassettes. The genomic structures are consistent with what is known about the genetics of mating-type switching and homo- and heterothallism in each species.

Methods

C. glabrata strain RND13 was generously provided by R. Ueno and N. Urano (Tokyo University of Fisheries, Tokyo) (11, 12). Although initially characterized as *Torulasporea delbrueckii* (11), strain RND13 has an 18S rDNA sequence (12) with only four differences from that of the type strain CBS 138 of *C. glabrata*, which we suspected could be sequencing errors. We amplified and sequenced the variable D1/D2 domain of 26S rDNA from RND13 using primers NL-1 and NL-4 (primer sequences are listed in Table 1, which is published as supporting information on the PNAS web site) and found only 1 nucleotide difference of 581 between it and CBS 138, which is within the normal range of intraspecies variation (13). Similarly, only 5 of 1,398 and 0 of 420 nucleotides differed between RND13 and CBS 138 in the regions we sequenced to the left and right, respectively, of *MTL1*; this level of difference is as low as that seen among clinical isolates of *C. glabrata* (9).

C. glabrata strain CBS 138 *MTL α* sequence information was obtained from a PCR product amplified from the *MTL2a* cassette by using primers X1 and Z1 (Table 1). Testing for *MTL1* genotype switching in *C. glabrata* was done by PCR amplification and sequencing across both the Y/Z1 and X/Y junctions of *MTL1* (Fig. 1). At the Y/Z1 junction we used common primers from the *EMG1* gene with either *MTL1 α* - or *MTL1 α* -specific rightward primers. At the X/Y junction we used common primers from the region upstream of *BUD5* with *MTL1 α* - or

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Abbreviations: *MAT*, mating-type; *MTL*, *MAT*-like.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession numbers AJ617300–AJ617311).

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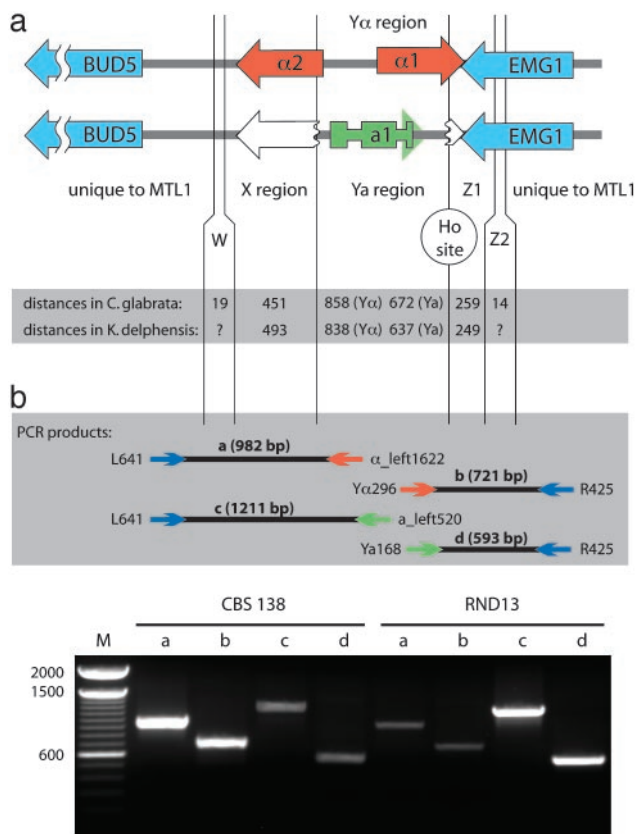


Fig. 1. (a) Organization of the α and a idiomorphs of the *C. glabrata* *MTL1* and *K. delphensis* *MAT* loci. The organization is identical in the two species. The $a1$ gene has three exons. Broken open arrows denote truncated copies of the 3' ends of the $\alpha 2$ and $\alpha 1$ genes present in the a idiomorphs. W, X, Z1, and Z2 indicate regions shared with silent cassettes. (b) PCR amplification of idiomorph-specific products from *C. glabrata* strains CBS 138 and RND13. Shown are idiomorph-specific primer pair combinations as diagrammed in Upper (lanes CBS 138 a–d and RND13 a–d); lanes 2-fold overloaded compared with the others (lanes CBS 138 c and d and RND13 a and b); and a size standard (lane M).

MTL1a-specific leftward primers. As described in Table 2 (which is published as supporting information on the PNAS web site), “switched” PCR products were detected from both RND13 and CBS 138 in amplifications across both the X/Y and Y/Z1 junctions. The sequences of switched PCR products were verified at the X/Y and Y/Z1 junctions from RND13 and at the Y/Z1 junction from CBS 138. In the sequenced region between *BUD5* and the X/Y junction, there were five nucleotide differences between strains RND13 and CBS 138 that served to show that contamination did not occur during PCR amplification.

The *C. glabrata* and *Kluyveromyces delphensis* *HO* genes were sequenced from genomic plasmid clones CG3380 and KD0425 (8). A *K. delphensis* *HMRa*-like cassette was sequenced from plasmid KD1818 (8). The *MATB* idiomorph of *Yarrowia lipolytica* was cloned by PCR using sequences from the flanking genes as primers. We sequenced loci in other species from *Génolevures* plasmids BA0AB039C05, BA0AB024C06, BB0AA018C05, BB0AA019A10, AR0AA031F11, and AZ0AA005C07 (14). Additional *Zygosaccharomyces rouxii* information was obtained from a random sequencing project similar to that described in ref. 8 (J. Gordon and K.H.W., unpublished data).

Results and Discussion

***C. glabrata* *MTL1* and *K. delphensis* *MAT* Loci and Their *Ho* Endonucleases.** We previously reported sequences of a putative *MAT* locus from *C. glabrata* and its close relative *K. delphensis* containing

homologs of the *S. cerevisiae* *MAT α 1* and *MAT α 2* genes (8). Srikantha *et al.* (9) independently identified three *MTL* loci in *C. glabrata*, one of which (*MTL1*) corresponds to the locus we described. They found both *MTL1 α* and *MTL1a* isolates of *C. glabrata*, which is haploid. In 36 of the 39 isolates that they examined (classes I and II), the *MTL* genotypes were consistent with the designation of *MTL1* as the mating-type locus and *MTL2a* and *MTL3 α* as silent cassettes similar to *HMRa* and *HML α* of *S. cerevisiae*. However, Srikantha *et al.* (9) reported that the *MTL1a1* gene has no start codon.

We verified that the *K. delphensis* *MTL1* ortholog is its *MAT* locus. From our random *K. delphensis* genomic library (8) we identified five more plasmids whose insert end sequences indicated that they span the locus. Three plasmids contained the *MAT α 1* and *MAT α 2* genes as reported (8), and the other two contained a homolog of *S. cerevisiae* *MAT α 1* (Fig. 1a). The mating-type-specific segments in *K. delphensis* are 838 and 637 bp for the *K. delphensis* *MAT α* and *MATa* idiomorphs, respectively. Given the large sizes of the plasmid inserts (7.2–11.3 kb), we can be certain that they are derived from the *MAT* locus and not from silent cassettes. This indicates that *K. delphensis* strain CBS 2170 is a *MAT α /MATa* diploid and that its *MAT* locus is orthologous to *C. glabrata* *MTL1*.

The *C. glabrata* type strain CBS 138 (ATCC no. 2001) has an *MTL1 α* genotype (8). By PCR amplification using primers from the presumed X and Z1 regions flanking *MTL1*, which are expected also to flank any silent cassettes, we obtained two products, one with *MTL α* information (from an *MTL1 α* or *MTL3 α* template) and one with *MTLa* information (from an *MTL2a* template). Comparison of the *C. glabrata* *MTLa* and *K. delphensis* *MATa* sequences showed that the $a1$ gene in each species contains two introns and has a normal start codon (Fig. 1a and see Fig. 5, which is published as supporting information on the PNAS web site).

The boundary between the Y and Z1 regions, where sequences unique to the α or a idiomorphs meet common flanking sequence, is formed by a putative cleavage site for the *Ho* endonuclease in both *K. delphensis* and *C. glabrata*. The first bases of the Z1 region have the canonical *Ho* site sequence CGCAAC in *K. delphensis* *MAT α* and *MATa* and *C. glabrata* *MTL1a*. In *C. glabrata* *MTL1 α* it is altered to CGCAGC, but this sequence also has been shown to be cleaved efficiently by the *S. cerevisiae* *Ho* endonuclease *in vivo* (15). These observations suggested that *C. glabrata* and *K. delphensis* have *HO* endonuclease genes. We found these genes by using comparative gene order information to walk from neighboring genes we had identified from genome survey sequencing (8) (Fig. 2). The predicted *C. glabrata* and *K. delphensis* *Ho* proteins have 68% sequence identity to each other and 57% identity to *S. cerevisiae* *Ho* protein and show conservation of the two LAGLIDAG endonuclease motifs and the Gly-223 residue identified as essential for mating-type switching in *S. cerevisiae* (16).

To investigate whether switching of the *MTL1* genotype occurs in *C. glabrata*, we used an environmental isolate, RND13 (with the *MTL1a* genotype), as well as the type strain CBS 138 (*MTL1 α*). RND13 was isolated recently from hot spring drainage water (40°C) and has been phenotypically characterized (11, 12). PCR amplification using one primer specific to Y α or Ya together with a primer from common flanking sequence (Fig. 1b) was used to test for *MTL1* genotype switching. The flanking primers lie outside the W and Z2 regions shared with the *MTL2* and *MTL3* cassettes (9) and so will amplify a product only from the *MTL1* locus. The RND13 isolate consistently yielded a strong PCR product with *MTL1a*-specific primers and a weaker product with *MTL1 α* -specific primers (Fig. 1b). This result was seen in amplifications with several different primer combinations and with four independent DNA preparations from RND13 (Table 2). The structures of PCR products from RND13 spanning the

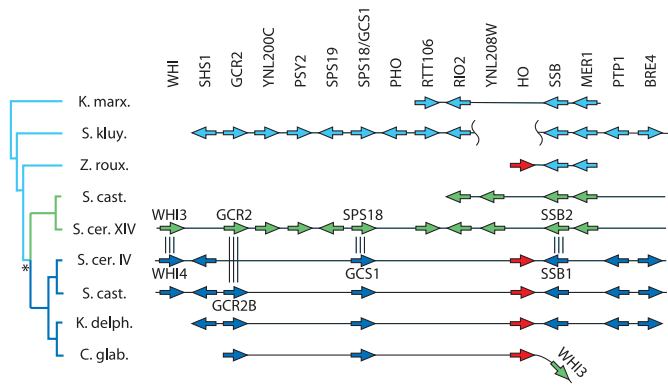


Fig. 2. Gene organization around the *HO* locus in yeast species. Homologous gene groups are aligned vertically. Triple vertical lines indicate duplicated gene pairs. The phylogenetic tree on the left shows the evolutionary relationships among sequences, with the genome duplication event (asterisk) resulting in two descendant lineages in some species. Green, lineages and genes orthologous to *S. cerevisiae* chromosome XIV; dark blue, orthologs of chromosome IV. The extent of sequence data available for each species is shown by black horizontal lines. Data for *Saccharomyces castellii* and *Saccharomyces kluyveri* are from ref. 18 (GenBank accession nos. AACE01000158, AACE01000109, AACF01000039, and AACF01000107). *K. marx.*, *Kluyveromyces marxianus*; *S. kluy.*, *S. kluyveri*; *Z. roux.*, *Z. rouxii*; *S. cast.*, *S. castellii*; *S. cer. XIV*, *S. cerevisiae* chromosome XIV; *S. cer. IV*, *S. cerevisiae* chromosome IV; *K. delph.*, *K. delphensis*; *C. glab.*, *C. glabrata*.

X/Y and Y/Z1 junctions in *MTL1 α* (switched) and *MTL1a* (unswitched) forms were confirmed by sequencing. Similarly, we obtained strong *MTL1 α* and weak *MTL1a* PCR products from CBS 138 (Fig. 1*b*), and we verified the *MTL1a* Y/Z1 junction product by sequencing. In both CBS 138 and RND13, the *MTL1 α* sequence was always found with the variant CGCAGC Ho site sequence, and *MTL1a* co-occurred with the canonical site sequence, which indicates that the Ho site was cleaved and repaired by using information from *MTL2a* (in the *MTL1 α* \rightarrow *MTL1a* switch in CBS 138) or *MTL3 α* (in the *MTL1a* \rightarrow *MTL1 α* switch in RND13). We interpret these results to indicate that a small number of RND13 cells are switching genotypes from haploid *MTL1a* to haploid *MTL1 α* , and a small number of CBS 138 cells are switching from *MTL1 α* to *MTL1a*. However, we did not find any evidence of diploid *C. glabrata* by a PCR amplification across the whole *MTL1* locus from *BUD5* to *EMG1* to look for colonies yielding equal quantities of the *MTL1 α* and *MTL1a* products, which differ slightly in size. Mating-type switching in a clinical isolate of *C. glabrata* also was reported recently (17).

Acquisition of the *HO* Endonuclease Gene in Species Close to *S. cerevisiae*. To investigate the organization and evolution of *MAT* (or *MTL*) and *HO* loci in yeasts, we combined sequences available from genome projects (10, 18) with our sequence data from several species, determined primarily from genomic clones made by the Génolevures project (14). The organization of *HO* and *MAT* loci is summarized in Figs. 2 and 3, respectively.

In *S. cerevisiae*, *C. glabrata*, and *K. delphensis*, the cleavage site for the Ho endonuclease is located within the *MAT α 1* gene, because the 3' end of this gene is located in the Z1 region (Figs. 1*a* and 3). Similarly, in *S. castellii* the Y/Z junction occurs within *MAT α 1* and resembles a Ho cleavage site, and the genome sequence includes a *HO* gene (18). Heterothallic *ho* mutants of *S. castellii* have been obtained by ethyl methanesulfonate mutagenesis (19). In *S. kluyveri*, *Kluyveromyces lactis*, *Y. lipolytica*, and *C. albicans*, *MAT α 1* is located completely within the unique Y α region of *MAT α* , and the sequences at the Y/Z junctions in these species are heterogeneous and do not resemble the Ho site (Fig. 3). Considering the phylogenetic relationships among the

species, it seems that the Y/Z boundary became situated inside *MAT α 1* relatively recently, because of the gain of the Ho site (Fig. 4).

There is no evidence that any of the yeast species that lack a Ho site in *MAT α 1* (Fig. 3) have a *HO* gene either. This observation is based on nearly complete genome sequencing of *C. albicans* and *S. kluyveri* (10, 18) and significant sequence surveying of *K. lactis*, *Pichia angusta*, and *Y. lipolytica* (14). We sequenced Génolevures plasmids containing part of a *HO* gene from *Z. rouxii* and an equivalent genomic region in *K. marxianus* from which the *HO* gene is missing (Fig. 2). From interspecies comparisons of local gene order, it seems that the *HO* gene materialized between the genes *RIO2* and *SSB* in the immediate common ancestor of *Z. rouxii*, *S. cerevisiae*, *S. castellii*, *C. glabrata*, and *K. delphensis*. Gene order around the *HO* locus is well conserved across species, but the comparisons (Fig. 2) are complicated by the occurrence of a genome duplication in the ancestor of four species after it had split off from *Z. rouxii* (20) (see Fig. 4). The gene *YNL208W* is also absent from the *K. marxianus* site but is present on a small *S. kluyveri* contig not shown in Fig. 2.

The Ho endonuclease protein is related in sequence to inteins, selfish mobile genetic elements that can insert in-frame into host genes (21–25). Their translation products contain a protein-splicing domain to autoexcise the intein from the host protein and an endonuclease domain to propagate the element into intein-lacking alleles of the host gene. The Ho protein has the same two domains but lacks a host gene and has an additional zinc finger domain at its C terminus (23). Within the phylogenetic tree of all LAGLIDADG endonucleases (26), Ho clusters with the *VMA1* intein (called VDE or PI-*Sce*I), which is the only true intein in the *S. cerevisiae* genome. The *VMA1* intein, like *HO*, is found only in hemiascomycetes, although its taxonomic distribution has been affected by horizontal transmission between species (27, 28). A hypothesis for the origin of *HO* is that an intein invaded the *VMA1* gene of a hemiascomycete ancestor from an unknown source and subsequently duplicated to give rise to *HO*. Intein endonucleases normally cut an inteinless allele of their host gene, which is then repaired by using the intein-containing allele as a template. Ho has managed to switch this specificity so that it cuts in *MAT α 1* (instead of a host gene), which is then repaired by using *HML* or *HMR* as a template (25). Because only haploid cells switch mating type, there is no second allele at *MAT* that can be used to direct repair. The switch in specificity is probably due to the extra zinc finger domain in Ho, because Ho uses this to bind its recognition site in *MAT α 1* (29), whereas the *VMA1* intein binds to its homing site in the *VMA1* gene primarily using DNA contacts in the protein splicing domain (30, 31). Many aspects of *HO* evolution, such as how it came to be asymmetrically regulated in mother and daughter cells by Ash1 or why Ho retains a protein-splicing domain, still are not understood.

Silent Cassettes. Silent cassettes of mating-type information are an essential part of the switching system. All species that have *HO* genes also have silent cassettes (Fig. 4). Cassettes often can be readily distinguished from functional *MAT* loci because they may contain truncated duplicates of the genes that flank *MAT*. In *S. cerevisiae*, part of *BUD5* is duplicated at *HML*. In the *S. castellii* genome sequence, the 5' end of *CAN1* is found beside α 1 and *a1* genes in short contigs that probably represent silent cassettes. The *C. glabrata* *MTL2* and *MTL3* loci contain truncated copies of the 3' end of *EMG1* (8, 9). In *K. delphensis* we identified a genomic clone containing a probable *HMRa*-like cassette with a 3' fragment of *EMG1* adjacent to homologs of the genes *YMR315W*–*YMR310C*, which in *S. cerevisiae* are in a subtelomeric region.

Although it apparently has no *HO* gene, *K. lactis* has a *MAT*

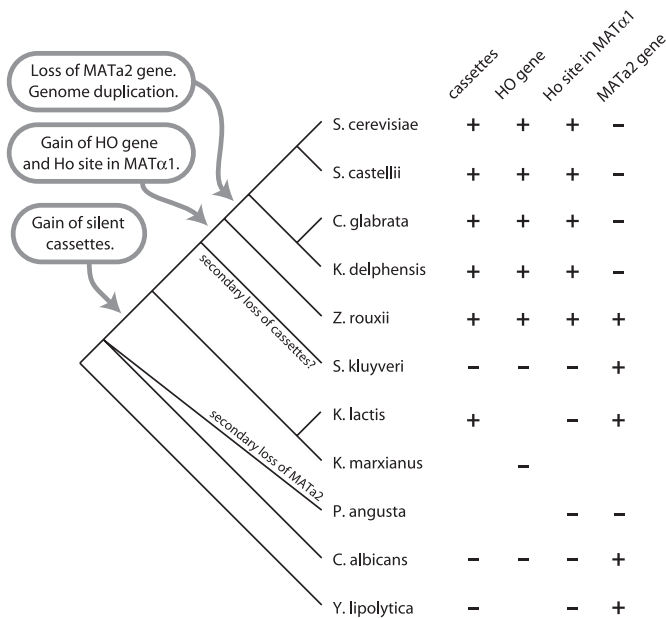


Fig. 4. Summary of events in *MAT* and *HO* locus evolution. The phylogenetic tree on the left (based on refs. 8, 50, and 51) is schematic and not drawn to scale. In the table on the right, blanks indicate missing data. *Z. rouxii* information is from Fig. 2 and low-coverage shotgun sequencing.

silent cassettes (6, 39). These observations suggest a two-step model for the evolution of mating-type loci, with the first step being the invention of cassettes and the recruitment of silencing apparatus and the second step being the acquisition of the Ho endonuclease, which may have increased the frequency and/or precision of genotype switching. In *S. cerevisiae*, the frequency of switching is $\approx 10^6$ times higher in *HO* than in *ho* cells (3). Our results support the suggestion of Keeling and Roger (24) that when Ho arose it became integrated into a preexisting passive switching system that relied on gene conversion.

Loss of an HMG Domain Gene. The *S. cerevisiae* *MAT* locus idiormorphs code for only three proteins: the homeodomain proteins $\alpha 1$ and $\alpha 2$ and the “ α -domain” protein $\alpha 1$ (40). Homologs of these three genes are found in all nine hemiascomycete species we examined (Fig. 3). It was reported previously that *Y. lipolytica* *MATA1* has no similarity to *MATA1* in other species (39), but this report resulted from overlooking a probable second exon of this gene.

An additional gene (*MATA2*) coding for an HMG domain DNA-binding protein is present in the *MATA* idiormorphs of several species (Fig. 3). This gene is called *MATA2* in *Y. lipolytica* (39) and *MATA2* in *K. lactis* (33), but it should not be confused with the unrelated ORF called *MATA2* in *S. cerevisiae*, which is unlikely to be a functional gene (40). The HMG gene is present in the *MTLa* idiormorph of *C. albicans* but was not previously recognized (6) because it is divergent in sequence and contains an intron (Fig. 6, which is published as supporting information on the PNAS web site). It is also present in *S. kluyveri* and *Z. rouxii* (Fig. 4). In *Y. lipolytica*, the function of the *MATA2* HMG domain gene is to repress conjugation in diploids (39). Very recently, Tsong *et al.* (41) identified *C. albicans* *MTLa2* as a positive regulator of α -specific genes.

We sequenced a genomic clone from *P. angusta* (also called *Hansenula polymorpha*; strain CBS 4732) that contains the genes *MAT* $\alpha 2$, *MAT* $\alpha 1$, and *MATA1*, in that order and all in the same orientation. The presence of *MAT* α and *MATA* information on the same idiormorph is surprising but consistent with

genetic evidence that *P. angusta* is a homothallic haploid in which any strain can mate with any other strain (42). *P. angusta*'s *MAT* locus organization is similar to that seen in homothallic species of *Cochliobolus* (a distantly related ascomycete classified in the Pezizomycotina), which arose from heterothallic ancestors by DNA rearrangement at the *MAT* locus (43), and in a homothallic *Gibberella* species (44). It is also surprising that the *P. angusta* *MAT* locus does not have a *MATA2* gene, although it has been suggested that *P. angusta* has a tetrapolar mating system that would necessitate a second unlinked mating-type locus (45).

Partial Conservation of the *MAT* Locus Position Among Ascomycetes.

The location of the *MAT* genes can be traced through hemiascomycete evolution and reveals a surprising degree of positional continuity despite numerous gene order changes (Fig. 3). The one exceptional species is *C. albicans*, whose gene order around *MTL* is unrelated to that in other species except for the presence of *RCY1* (*YJL204C*).

In *Y. lipolytica* the *MAT* locus lies between homologs of *S. cerevisiae* *APN2* and *SLA2* (39). Remarkably, the same two genes are beside the *MAT* loci of the filamentous ascomycetes *N. crassa* (Fig. 3) and *Gibberella zeae* (GenBank accession no. AACM01000358), so this configuration may be the ancestral one for all ascomycetes. The arrangement of *SLA2* on the right side of *MAT* (as shown in Fig. 3) is conserved in *P. angusta*, *K. lactis*, and *S. kluyveri*. In contrast, three different genes are found to the right of *MAT* in *S. cerevisiae*, *S. castellii*, and the *C. glabrata*/*K. delphensis* pair. We hypothesize that in these species the activity of Ho may have elevated the rate of evolutionary chromosome rearrangement.

On the left side, *BUD5* is immediately beside *MAT* in *S. cerevisiae*, *S. castellii*, *C. glabrata*, and *K. delphensis*. This section of *S. cerevisiae* chromosome III has a sister relationship through genome duplication with part of *S. cerevisiae* chromosome X, including the duplicated gene pair *YCR037C* (*PHO87*) and *YJL198W* (*PHO90*) (46). In *S. kluyveri*, which separated from the *S. cerevisiae* lineage before genome duplication occurred (20) (Fig. 4), the region to the left of *MAT* includes a series of genes from the same area of chromosome X, between *YJL207C* and *YJL210W* (*PEX2*). After genome duplication, a contiguous group of ≈ 10 genes must have been deleted from the left side of *MAT* in the progenitor of *S. cerevisiae* chromosome III, because in *S. kluyveri* *BUD5* is adjacent to *YJL201W* (*ECM25*). In *K. lactis*, *YJL207C* is immediately adjacent to the left side of the *MAT* locus and is truncated by ≈ 1.2 kb at its 3' end relative to other species. This suggests that a single large deletion may have removed everything between the 3' end of *YJL207C* and *MAT* $\alpha 3$ in *K. lactis*. The 3' end of the shortened gene is duplicated in the *K. lactis* *HML* and *HMR* loci.

Between *PEX2* and *MAT* in *S. kluyveri* are two genes, including one orthologous to the chromosome XII gene *DIC1*, which also appears to the left of *MAT* in *P. angusta* (Fig. 3). In summary, the chromosomal location of the *MAT* locus has been conserved throughout much of hemiascomycete evolution but has been disturbed by the effects of genome duplication and the gain of the Ho endonuclease.

Conclusion

The mating system of hemiascomycetes has evolved from an obligate heterothallic system, as seen in *Y. lipolytica*, to heterothallism with low-level switching, as seen in *K. lactis*, to Ho-catalyzed homothallic switching, as seen in *S. cerevisiae*, via a series of steps (summarized in Fig. 4). The partial conservation of gene order flanking the *MAT* loci is sufficient to indicate that the mating-type loci of hemiascomycetes and filamentous ascomycetes are orthologous despite their gene

content differences and their very different pathways of sexual development. Homeodomain and HMG domain DNA-binding proteins seem to have been interchangeable during fungal evolution, with the *MAT* loci of modern species having only homeodomains [*S. cerevisiae* and *Cryptococcus neoformans* (47)], only HMG domains (*N. crassa*), or a mixture of the

two (*K. lactis*, *C. albicans*, *Y. lipolytica*, and *Schizosaccharomyces pombe*).

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