

Convergent evolution of linked mating-type loci in basidiomycetes: an ancient fusion event that has stood the test of time

Short title: Independent origins of fused mating type loci in basidiomycete yeasts

Sheng Sun¹, Marco A. Coelho¹, Joseph Heitman¹, and Minou Nowrousian^{2*}

¹Department of Molecular Genetics and Microbiology, Duke University Medical Center,
Durham, North Carolina, United States of America

²Lehrstuhl für Allgemeine und Molekulare Botanik, Ruhr-Universität Bochum, 44780
Bochum, Germany

* Corresponding author:

Minou Nowrousian

Lehrstuhl für Allgemeine und Molekulare Botanik

Ruhr-Universität Bochum

Universitätsstr. 150

44780 Bochum

Germany

phone +49 234 3224588

fax +49 234 3214184

email minou.nowrousian@rub.de

Abstract

Sexual development is a key evolutionary innovation of eukaryotes. In many species involves the interaction of compatible mating partners that can undergo cell and nuclear fusion and subsequent steps of development including meiosis. Mating compatibility in fungi is governed by mating type determinants, which are localized at mating type (*MAT*) loci. In basidiomycetes, the ancestral state is hypothesized to be tetrapolar (bifactorial), with two genetically unlinked *MAT* loci containing homeodomain transcription factor genes (*HD* locus) and pheromone and pheromone receptor genes (*P/R* locus), respectively. Alleles at both loci must differ between mating partners for completion of sexual development. However, there are also basidiomycete species with bipolar (unifactorial) mating systems, which can arise through genomic linkage of the *HD* and *P/R* loci. In the *Tremellales*, which comprise mostly yeast-like species, bipolarity is found only in the pathogenic Cryptococci, e.g. in the well-studied human pathogen *Cryptococcus neoformans*. Here, we describe the analysis of *MAT* loci from the *Trichosporonales*, a sister group to the *Tremellales*. We analyzed genome sequences from 29 strains comprising 24 species, including two new genome sequences generated in this study. Somewhat surprisingly, in all of the species analyzed, the *MAT* loci are fused and a single *HD* gene is present in each allele. This is similar to the organization in the pathogenic Cryptococci, which also have linked *MAT* loci and carry only one *HD* gene per *MAT* locus instead of the usual two *HD* genes found in the vast majority of basidiomycetes. However, almost all *Trichosporonales* strains analyzed carry either the combination of the *HD* gene *SXII* with the pheromone allele *STE3a*, or the combination of the *SXII* gene and *STE3 α* allele. This is in contrast to *MAT* alleles in *C. neoformans*, where *SXII* is linked with *STE3 α* , and *SXII* is linked with *STE3a*. The differences in allele combinations as well as the existence of tetrapolar *Tremellales* sister species to the bipolar Cryptococci suggest that the fusion of the *HD* and *P/R* loci and the loss of one *HD* gene per allele occurred independently in the *Trichosporonales* and the pathogenic Cryptococci, supporting the hypothesis of convergent evolution at the molecular level towards fused mating-type regions in fungi. A phylogenetic analysis of divergence times suggests that the *MAT* fusion in the *Trichosporonales* is the oldest fusion of *MAT* loci observed to date.

Author summary

Sexual development in fungi is governed by mating-type (*MAT*) genes, and the corresponding *MAT* loci show similarities to sex chromosomes in animals and plants. One common feature is an evolutionary trend towards combining sex-associated genes on the same chromosome, which can evolve by selection because it facilitates linkage of favorable allele combinations. Here, we show that this occurred in the *Trichosporonales*, a sister group to the *Tremellales*, similar to the expanded, fused *MAT* loci discovered previously in the human pathogen *Cryptococcus neoformans*. Our data suggest that fusion of *MAT* loci occurred independently in the *Trichosporonales* and pathogenic Cryptococci, supporting the hypothesis of convergent evolution towards fused *MAT* regions in fungi.

Introduction

Sexual reproduction is pervasive among eukaryotic organisms, but despite its rather conserved core features (syngamy/karyogamy and meiosis), many aspects of sexual development show high evolutionary flexibility [1-3]. This includes the determination of compatible mating partners that can successfully undergo mating and complete the sexual cycle. In many species, compatibility is determined by one or more genetic loci that differ in compatible mating partners. The evolution of such genes has been studied in many systems including plants, animals, algae, and fungi [1, 4-6]. In animals and plants, genes that determine sexual compatibility are often found on sex chromosomes, which have distinct evolutionary histories compared to autosomes. In fungi, genetic systems for determining mating compatibility vary widely, ranging from single genetic loci on autosomes to chromosomes that have the hallmarks of sex chromosomes, e.g. suppressed recombination. Studies in different fungal groups have revealed that transitions towards larger, sex chromosome-like regions have occurred several times in fungal evolution, with some systems having evolved only recently [5, 7-9]. Thus, fungi are excellent model systems to study the evolution of genomic regions involved in mating and mating type determination.

Mating compatibility in fungi is governed by mating-type genes, which are located in mating-type (*MAT*) loci [10, 11]. While ascomycetes and the Mucoromycotina are bipolar (unifactorial), i.e. harboring only one *MAT* locus [12-14], in basidiomycetes, the ancestral state is considered to be tetrapolar (bifactorial), with two genetically unlinked *MAT* loci controlling mating-type determination at the haploid stage [15, 16]. One *MAT* locus usually

contains tightly linked pheromone and pheromone receptor genes (*P/R* locus) involved in pre-mating recognition, and the other encodes homeodomain transcription factor genes encoding homeodomain (HD) proteins of class 1 and class 2 (*HD* locus) determining viability after syngamy. Importantly, alleles at both loci must differ between mating partners for completion of the sexual cycle [17].

However, there are also many basidiomycete species with bipolar mating systems, which can arise through genomic linkage of the *HD* and *P/R* loci, or when the *P/R* locus loses its function in determining mating specificity [15, 17, 18]. Bipolarity through *MAT* loci fusion is found in the subphylum *Ustilaginomycotina*, e.g. in *Malassezia* species and *Ustilago hordei*, several *Microbotryum* species (subphylum *Pucciniomycotina*), as well as in the pathogenic Cryptococci from the class *Tremellomycetes* (subphylum *Agaricomycotina*) [1, 7, 19-23] (Fig. 1). In *Microbotryum*, several convergent transitions to linked *MAT* loci were shown within the genus [7]. One feature that known species with fused *MAT* loci have in common is that they are associated with plant or animal host species as pathogens or commensals. It has been hypothesized that the necessity to find a mating partner while associated with a host might have favored linkage of *MAT* loci, because having one linked instead of two unlinked *MAT* loci increases the compatibility of gametes derived from a single diploid parent. This would improve mating compatibility rates on a host where other mating partners might be difficult to find, e.g. when a host is initially colonized by a single diploid genotype or meiotic progeny from a single tetrad [4, 8, 15, 17, 24].

Two additional evolutionary features can be associated with the linkage and expansion of the *MAT* loci. One is the recruitment of other development-associated genes into the fused *MAT* loci, and the second is suppression of recombination between the fused *MAT* loci that can extend along the *MAT*-containing chromosome [11, 23, 25]. Suppression of recombination is a hallmark of sex chromosomes in other eukaryotes as well, and thus might point towards convergent evolutionary transitions for the regulation of sexual development in eukaryotes [26, 27]. Suppression of recombination between the linked *MAT* loci would further increase compatibility under inbreeding conditions, and recruitment of sex-associated genes into the *MAT* locus might facilitate the inheritance of favorable allele combinations through genetic linkage [28].

A well-studied case of fused *MAT* loci is the one of *Cryptococcus neoformans*, a member of a group of closely related, pathogenic *Cryptococcus* species [1]. Fusion of the *HD* and *P/R* loci most likely occurred in the ancestor of the pathogenic Cryptococci, because other analyzed *Tremellales* species including the closely related, but non-pathogenic *Cryptococcus*

amylolentus, *Kwoniella heveanensis*, *Kwoniella mangrovensis*, *Cryptococcus wingfieldii*, and *Cryptococcus floricola* are all tetrapolar [29-33]. The fused *C. neoformans* *MAT* locus encompasses more than 20 genes over a region spanning more than 100 kb and has two alleles designated **a** and α . In the majority of basidiomycetes, each *MAT* allele at the *HD* locus carries both the *HD1* and the *HD2* transcription factor genes, whereas in *C. neoformans*, the *MAT* α locus contains only the *HD1* gene *SXII* α , and *MAT***a** contains only the *HD2* gene *SXII***a**. Except for a gene conversion hotspot, the *C. neoformans* *MAT* locus displays suppressed meiotic recombination [23, 34, 35].

Among the *Tremellomycetes*, pathogenic Cryptococci are so far the only species for which fused *MAT* loci have been described [17, 23]. In a previous study, we analyzed the genome sequence of *Cutaneotrichosporon oleaginosum* strain IBC0246 (formerly *Trichosporon oleaginosus*), which belongs to the *Trichosporonales*, a sister order to the *Tremellales* within the *Tremellomycetes* class. *Trichosporonales* species are widely distributed in the environment and have been isolated from a variety of substrates including soil, decaying plant material, and water. Many species are saprobes, but some have also been found to be associated with animals including humans either as commensals or pathogens [36-38]. Despite their common occurrence in the environment, *Trichosporonales* are an understudied fungal group, and sexual reproduction has not yet been observed for any of the known species [39, 40]. Recently, several *Trichosporonales* were studied with respect to their biotechnological properties, including the oil-accumulating *C. oleaginosum*, which was first isolated from a dairy plant, and has the ability to metabolize chitin-rich and other non-conventional substrates [41-44]. The sequenced *C. oleaginosum* strain is haploid, and similar in genome size and gene content to genomes from the sister order *Tremellales*, and this was also the case for several other *Trichosporonales* genomes that have since been sequenced [43, 45-50]. Interestingly, *C. oleaginosum* showed some similarities to *C. neoformans* in the organization of *MAT* loci. This included recruitment of genes with diverse functions during mating into the *HD* and *P/R* loci, as well as the presence of only a *SXII* homolog at the *HD* locus [43]. However, in the draft genome assembly, *HD*- and *P/R* loci were situated on different scaffolds. Furthermore, a sexual cycle has not yet been described for *C. oleaginosum*. Thus, it was not possible to conclusively distinguish whether this species carries two unlinked or fused *MAT* loci, with the seemingly more likely possibility being a tetrapolar mating system as this is present in the majority of species from the sister group of *Tremellales*.

Since the analysis of the first *C. oleaginosum* genome, several more *Trichosporonales* genomes were sequenced, although none were analyzed with respect to their *MAT* loci [45-51]. Here, we describe the sequencing of two additional *Trichosporonales* genomes for *C. oleaginosum* ATCC20508 and *Vanrija humicola* CBS4282, and the analysis of *MAT* loci organization in *Trichosporonales* genomes from 24 different species. Surprisingly, we found that all of the species analyzed have fused *MAT* loci that contain both *P/R* and *HD* genes. Furthermore, all the analyzed strains of each species contain only one of the two ancestral *HD* genes (*SXII* and *SXI2*), with almost all of the species carrying either the combination of the *HD* gene *SXII* with the pheromone receptor allele *STE3a*, or the combination of the *SXI2* gene and the *STE3 α* allele. This is in contrast to *C. neoformans*, where *SXII* is combined with *STE3 α* in the *MAT α* allele, and *SXI2* is combined with *STE3a* in the *MATa* allele [34]. The differences in allele combinations as well as the existence of tetrapolar *Tremellales* sister species to the bipolar Cryptococci suggest that the fusion of the *HD* and *P/R* loci as well as the loss of one *HD* gene per allele occurred independently in the *Trichosporonales* and the pathogenic Cryptococci. This provides further evidence that convergent evolution leading to fused *MAT* loci is evolutionary beneficial under certain circumstances, and has been selected for multiple times in basidiomycetes, with the *MAT* loci fusion in *Trichosporonales* representing the oldest such event observed to date.

Results

Trichosporonales species have fused *HD* and *P/R* mating type loci

Since 2015, genome sequences have been published for *Trichosporonales* species from the genera *Apiotrichum*, *Cutaneotrichosporon*, *Takashimella*, *Trichosporon*, and *Vanrija* [45-52] (Table 1). We analyzed the genomes of 29 isolates that belong to 24 *Trichosporonales* species for the organization of the *MAT* loci, and found that all of them contain fused *MAT* loci, with both mating-type determining genes (*STE3* and *SXI*) located ~55 kb apart from each other. Except for four strains described below, all analyzed *MAT* loci comprise either the combination of the *HD* gene *SXII* with the pheromone allele *STE3a*, or the combination of the *SXI2* gene and *STE3 α* allele (Table 1, Figs 1 and 2). Each *MAT* allele carries a single pheromone precursor gene in the vicinity of the *STE3* gene (Fig. 2, S1 Text). The *SXI/STE3* combinations found in the *Trichosporonales* are different from the *MAT* alleles in *C.*

neoformans, where the *SXII* gene is combined with *STE3 α* in the *MAT α* allele, and *SXI2* is combined with *STE3a* in the *MATa* allele [23, 34]. To avoid confusion with the *C. neoformans* nomenclature and be compatible with allele designations in other basidiomycetes, we named the *STE3 α* -containing allele A1, and the *STE3a*-containing allele A2 (Figs 1 and 2).

The fact that the genomes of 24 *Trichosporonales* species carry fused *MAT* loci with two allelic combinations distributed throughout the phylogenetic tree (Fig 1) suggests that the linkage event predates the diversification of the *Trichosporonales* clade. However, at the beginning of this study, only one *MAT* allele was known for each of the *Trichosporonales* species. To verify that both *MAT* alleles, A1 and A2, can be found in different strains within a single species, we analyzed additional *Vanrija humicola* strains (Table 2). *V. humicola* is a member of a phylogenetically early-branching lineage within the *Trichosporonales*, and the previously sequenced strain (JCM1457) carries the A2 allele (Figs 1 and 2) [50]. Several other strains obtained from culture collections were analyzed by RAPD genotyping, which revealed banding patterns similar to those from the type strain *V. humicola* JCM1457 (=CBS571, *MAT* A2) (S1 Fig). These strains were analyzed further by PCR for the mating type genes. Strains other than CBS4282 and CBS4283 gave PCR products when using oligonucleotides for *SXII* and *STE3a* derived from the JCM1457 (A2) genomic sequence. The two genes adjacent to the *SXI* gene in both *MAT* alleles are conserved in the analyzed *Trichosporonales* genomes (Fig 2), therefore we used oligonucleotide primers derived from these genes to amplify the predicted *SXI2*-containing region of CBS4282 and CBS4283 confirming the presence of an *SXI2* gene in these strains. To assess this further, we sequenced the genome of CBS4282 using Illumina sequencing. The genome was assembled into 21 scaffolds with 5089 predicted genes. At 22.63 Mb, the assembly size is similar to that of strain JCM1457 (A2) (22.65 Mb, [50]). A k-mer analysis of the Illumina reads for CBS4282 showed a single peak as expected for a haploid genome (S1 Fig). Analysis of the *MAT* region revealed fused *MAT* loci carrying the *SXI2* gene and *STE3 α* allele, confirming that this strain is indeed an A1 strain, and thus showing that alternative alleles can be found in different strains of a single *Trichosporonales* species (Figs 2 and 3, S2 and S3 Figs, Table 2). This was further confirmed by an analysis of the recently published genome of a second *Apiotrichum porosum* strain [51], with the two *A. porosum* genome sequences available now representing the A1 and A2 alleles (Figs 2 and 3).

Alternative *MAT* alleles in *Trichosporonales* show significant chromosomal rearrangements, but each displays highly conserved structure across species

The gene order within the core *MAT* region between the *SXI2* and *STE3 α /MF α* genes is very well conserved between *V. humicola* CBS4282 (A1) and the A1 alleles of the other *Trichosporonales* species analyzed, whereas it differs from the *V. humicola* A2 strain JCM1457 by two inversions, one of which encompasses most of the core *MAT* region (Fig 2). The same is true for the A2 alleles, where the core *MAT* region is also rather conserved between *V. humicola* JCM1457 (A2) and other *Trichosporonales* (Fig 2).

In addition to the key mating-type determinants, the *MAT* loci of the *Trichosporonales* contain other genes previously shown to be required for mating and filamentation in other species (e.g. *STE11* and *STE20*), and their common presence in the *MAT* loci of *Tremellales*, suggests these genes were anciently recruited to the *MAT* locus (Fig 2). In the *Trichosporonales*, *STE11* and *STE20* are found within the core *MAT* region (between the *HD* and *P/R* genes), and *STE12* is in the vicinity of the core *MAT* region in most species (Fig 2). One exception turned out to be *STE12* in the *Cutaneotrichosporon* lineage. In *C. oleaginosum* strains IBC0246 (A2) and ATCC20509 (A2), *STE12* is located on a different scaffold than the *MAT* locus. While *STE12* in IBC0246 (A2) is located at the end of its respective scaffold, its location in ATCC20509 (A2) is around position 1.1 Mb of a 2.5 Mb scaffold, indicating that even if this scaffold was linked to the *MAT* scaffold, the linkage would not be very tight.

To confirm that *STE12* is indeed not linked to the *MAT* locus in *C. oleaginosum*, we sequenced the genome of the third of the three known strains of this species, ATCC20508. The genome was sequenced with Pacific Biosciences SMRT sequencing and assembled into eight contigs with 8208 predicted genes. CHEF gel electrophoresis of the *C. oleaginosum* strains shows a karyotype comprising seven chromosomes, with the sizes of the six largest chromosomes corresponding well to the sizes of the six largest contigs of the ATCC20508 assembly (S4 Fig). Similar to the other two strains, ATCC20508 carries the *MAT* A2 allele and *STE12* on different contigs (contigs 1 and 3, respectively; S4 Fig), making it unlikely that they are located the same chromosome (S4 Fig). Additionally, analyses of the ATCC20508 (A2) and ATCC20509 (A2) genome assemblies showed that both the *MAT* locus and *STE12* are located within large co-linear scaffolds in both strains, providing further evidence that *STE12* is not linked to the *MAT* locus in this species (Fig S4). Consistent with this, in the other *Cutaneotrichosporon* species analyzed, *STE12* is either on the same scaffold as *MAT*, but at a distance of 400 to 750 kb (S5 Fig), or is present on different scaffolds than *MAT*. Thus, while this suggests that the location of *STE12* in the vicinity of *MAT* likely represents

the ancestral state in the *Trichosporonales*, its relocation to a different genomic location in the *Cutaneotrichosporon* species indicates that *STE12* might not be fully linked to *MAT* in the *Trichosporonales*.

Another difference in *MAT* organization that can be observed in the genus *Cutaneotrichosporon* is the combination of *HD* genes and *STE3* alleles. While all other strains of each species carry either A1 (*STE3 α* +*SX12*) or A2 (*STE3 α* +*SX11*) alleles, *Cutaneotrichosporon dermatis* carries an allele that has the overall genomic organization of A1, but has instead an *SX11* gene (S5 Fig). Because the A1 vs. A2 designation is based on the *STE3* variant, this allele was called A1*. Interestingly, A1* alleles can also be found in *Apiotrichum veenhuisii*, whereas a corresponding A2* allele, in which *STE3 α* is linked to the *SX12* gene, can be found in *Apiotrichum gracile* (Fig 1 and S5 Fig). Furthermore, both A1* and A2* alleles can be found in *Cutaneotrichosporon mucooides*, one of several hybrid species within the *Trichosporonales* (S6 Fig, S2 Text) [49, 50]. These *MAT* alleles seem to represent a derived state and might have been independently generated by recombination involving crossing-over between the *MAT* region adjacent to the *SX11/SX12* genes that is co-linear between the A1 and A2 alleles (S5 Fig).

The fusion of the *MAT* loci in *Trichosporonales* is ancient

The overall high degree of conservation in gene order and allele combinations of *HD* and *P/R* loci in the *Trichosporonales* suggests that the fusion of the *MAT* loci occurred in the common ancestor of the *Trichosporonales* lineage. To determine at what time this fusion might have occurred, we estimated divergence times of the basidiomycete lineages included in Fig 1 using three calibration points (see S7 Fig and Materials and Methods). Based on this analysis, the *Trichosporonales* and *Tremellales* have a common ancestor dating back approximately 179 million years ago (MYA), whereas the earliest split within the *Trichosporonales* occurred approximately 147 MYA (S7 Fig). Thus, under the assumption of a *MAT* fusion in the common ancestor of the *Trichosporonales*, this fusion would have occurred between 179 and 147 MYA, making it the oldest observed *MAT* fusion event in basidiomycetes, because the *MAT* fusions in the pathogenic Cryptococci, the *Ustilaginomycotina*, and the *Microbotryum* lineage occurred more recently (S7 Fig). While our estimates of divergence times for the *Microbotryum* and *Trichosporonales* lineages agree well with recent analyses [7, 53], it has to be noted that previous estimates for the last common ancestor of the pathogenic Cryptococci resulted in earlier divergence times ranging

from 40 to 100 MYA [54-57]. Nevertheless, even if the divergence time of the ancestor of the pathogenic Cryptococci were underestimated in our analysis, this would still place the last common ancestor of the *Trichosporonales*, and thus the likely *MAT* fusion in this group, at a much earlier time point.

Suppression of recombination is restricted to the fused *MAT* loci in *V. humicola*

Suppression of recombination is a hallmark of sex chromosomes of animals and plants and the mating-type chromosomes of algae and fungi [9, 25, 31, 34, 58-60]. One consequence of recombination cessation can be the accumulation of transposable elements as well as increased genetic differentiation between allelic sequences [26]. This was observed in *C. neoformans*, where the *MATa* and *MAT α* alleles differ significantly in gene organization, and the *MAT* locus contains more remnants of transposons and other repeat sequences than other genomic regions except for the centromeres and rDNA repeats [23, 61]. Furthermore, both *C. neoformans* *MAT* alleles are highly rearranged in comparison with their *Cryptococcus gattii* counterparts [34].

In contrast, the *MAT* A1 and A2 alleles of *V. humicola* are overall co-linear apart from two inversions, and this is generally the case throughout the *Trichosporonales* (Figs 2 and 3A). An analysis of repetitive sequences shows that there is no accumulation of repeat regions within the *MAT* alleles of *V. humicola* (S1 Table). Thus, whereas the two inversions might impair meiotic pairing and therefore recombination in this region, the conserved gene order and lack of repeats made us consider if the *V. humicola* *MAT* loci, and more generally the *Trichosporonales* *MAT* loci, are regions of suppressed recombination.

To test this, we first analyzed phylogenetic trees for several genes present within or adjacent to the *MAT* loci of *Trichosporonales* (Fig 4). For genes in regions undergoing meiotic recombination, alleles associated with alternative mating types are expected to display a species-specific topology in a phylogenetic tree, whereas genes in *MAT* regions should cluster by mating type if recombination suppression predates speciation (i.e. with the A1 alleles of the different species branching together rather than each of the alleles clustering with the A2 allele from the same species) [30, 34]. In the *Trichosporonales*, only *STE3* clearly shows a mating type-specific pattern with an ancient trans-species polymorphism in *Trichosporonales* and *Tremellales* (Fig 3B). None of the other genes tested showed a mating type-specific phylogenetic pattern for the *Trichosporonales* at such a deep phylogenetic level, indicating that recombination was not suppressed already at the base of the clade. However,

for several genes within the core *MAT* locus, sequences from two *V. humicola* A2 strains (JCM1457 and UJ1) group apart from the A1 sequence (CBS4282) (Fig 3B), and therefore, these genes might be mating type-specific at the species level, although more strains from more species would have to be investigated to exclude this finding occurring by chance (Fig 3).

This finding is further supported by BLASTN analyses comparing alleles in the three available *V. humicola* genomes, which showed that within the core *MAT* region, alleles from the A2 strains (JCM1457 and UJ1) are more similar to each other than to the A1 strain CBS4282, whereas outside of this region this is not the case (Figs 4A and 4B). This suggests that the *MAT* locus of *V. humicola* might be a region of recent recombination suppression.

To further test whether recombination is suppressed between the *MAT* alleles in *V. humicola*, we analyzed levels of synonymous divergence (d_s) between alleles on the *MAT*-containing scaffold as well as on three other scaffolds of CBS4282 (A1) and the two *V. humicola* A2 strains (S8 Fig). In the four longest scaffolds, genes within large regions of the scaffolds are more similar between CBS4282 (A1) and the JCM1457 (A2) than between the two A2 strains, where stretches of low d_s values occur much less frequently (S8 Fig). Interestingly, we also noted regions of high similarity between CBS4282 (A1) and JCM1457 (A2) that are interrupted by stretches containing more divergent alleles. This pattern is suggestive of ongoing recombination in natural *V. humicola* populations, even though sexual reproduction has not yet been observed.

In contrast to other genomic regions, including the co-linear regions outside of the core *MAT* region where overall divergence is lowest between CBS4282 (A1) and JCM1457 (A2) (Fig 4C and S8 Fig), most of the alleles in the core *MAT* region are slightly more divergent between the A1 strain and the two A2 strains than between the two A2 strains (Fig 4C). This is consistent with an absence of genetic exchange between the A1 and A2 alleles, as one would expect if recombination in this region is suppressed. These findings might be explained by a reduced recombination rate in the core *MAT* region carrying the inversions in alternate *MAT* alleles, which could lead to accumulation of mutations in the two *MAT* alleles and thus to elevated synonymous divergence between A1 and A2 alleles. However, divergence between A1 and A2 strains within the core *MAT* region is only moderately elevated for most genes, and the difference between the average d_s values is only statistically significant when comparing the analysis of CBS4282 (A1) vs. UJ1 (A2) with the analysis of JCM1457 (A2) vs. UJ1 (A2) (Fig 4D). One possible explanation could be that genetic exchange within the inverted regions may not be completely inhibited, because exchange via non-crossover gene

conversion or double crossover can still occur within the inverted regions [62]. This should result in a so-called suspension bridge pattern with divergence in the middle of an inverted region lower than towards the inversion breakpoints [63]. To test this, we performed BLASTN analyses on sliding windows of genomic sequences of the *MAT* locus and adjacent regions (S9 Fig). The results show less sequence similarity in the regions of putative inversion breakpoints between A1 and A2 strains compared to regions within the inversions and outside of the *MAT* region, consistent with the hypothesis that a certain amount of recombination is occurring within the inverted regions.

Even though recombination within the *MAT* region might not be fully suppressed possibly due to non-crossover gene conversion, the gene order within each A1 and A2 allele in different species is surprisingly well conserved (Fig 2). To test if this degree of gene order conservation extends beyond the *MAT* locus, we compared the *MAT*-containing scaffolds of *Trichosporonales*. As expected for a range of species separated by millions of years of evolution, synteny is conserved between closely related, but not between species that diverged long ago (S10 Fig), except for the *MAT* region. This suggests that the *MAT* region of *Trichosporonales* is an ancient cluster of tightly linked loci (also known as 'supergenes') that seem to segregate as a stable polymorphism within the populations of each species. Thus, the *Trichosporonales* appear to contain very stable *MAT* alleles with respect to gene order, combined with (slightly) suppressed recombination between different alleles.

The *MAT* loci in the *Trichosporonales* have a significantly lower GC content compared to the overall genomic GC content

One curious observation was found during the analysis of the *Trichosporonales* *MAT* region, namely a lower GC content in the *MAT* regions compared to the surrounding regions for the analyzed *Vanrija*, *Cutaneotrichosporon*, and *Trichosporon* strains (S10 and S11 Figs). One possible explanation for a lower GC content could be an accumulation of AT-rich transposable elements, but an analysis of repeats in the three *V. humicola* strains showed that there are only few (strain CBS4282, A1) or no (strains JCM1457 and UJ1, both A2) repeats present within the *MAT* region of these strains (S1 Table). Another explanation might be a lower density of coding regions, which tend to have a higher GC content than non-coding regions. However, an analysis of the GC content only in the coding sequences within and around the *MAT* region of strain CBS4282 showed the same pattern of lower GC content within the *MAT* region (S11B Fig). Another possible explanation for the lower GC content

might be the accumulation of mutations due to reduced recombination. Under the (simplistic) assumption that mutation frequencies for all nucleotide exchanges are similar, this would drive the GC content towards 50%. In genomes with an average GC content of more than 50%, this would appear as a region with lower GC content, and this would apply to the analyzed strains with an average genomic GC content of 58 to 63 % (S11 Fig). It has been shown that spontaneous mutations tend to be AT-biased in many species including several fungi [64-68], and thus higher mutation rates would generally lead to a lower GC content in the corresponding regions. Both explanations can be supported by an analysis of the codon usage within the *MAT* region of the three *V. humicola* strains (S2 Table). Among the codons for amino acids that can be encoded by more than one codon, there is a trend for GC-rich codons to be used less frequently in the *MAT* region compared to the genome-wide usage, consistent with (GC content-equalizing or AT-biased) mutations combined with selection for conserved protein sequences due to functional constraints. Additional studies of species with a GC content of less than 50% might be useful to test these hypotheses.

Discussion

Fused *MAT* loci evolved several times in basidiomycetes

In this study, we analyzed *MAT* loci from the *Trichosporonales*, and found that all of the analyzed species harbor fused *MAT* loci with a single *HD* gene, an arrangement that among the *Tremellomycetes* has so far only been found in the pathogenic Cryptococci [17]. Fused *MAT* loci have been identified previously in other basidiomycetes (*Malassezia*, *Microbotryum*, *Sporisorium*, *Ustilago*), but these *MAT* loci carry the ancestral arrangement of two *HD* genes [7, 17, 19-22, 69, 70]. An analysis of several strains from *V. humicola* showed that there are (at least) two *MAT* alleles, and homologs of both alleles are distributed throughout the *Trichosporonales* phylogenetic tree. This suggests that these alleles might be the most prevalent alleles in this group as the selection of sequenced strains was not based on and therefore most likely not biased towards certain *MAT* features. Indeed, if the mating system is predominantly inbreeding, rare alleles have no advantage and can be gradually lost by genetic drift. Thus, fusion of *MAT* loci, which can evolve by selection because it is beneficial in selfing mating systems, is predicted to lead to a reduction in the number of *MAT* alleles [4, 71]. In addition, theoretical modelling predicts that a combination of facultative and

rare sexual reproduction, low mutation rates, and a small effective population size should lead to a reduction in the number of mating types [72]. Population sizes and mutation rates are, however, not known for *Trichosporonales* species, and no sexual development has been observed so far making it possible that asexual reproduction is the predominant form of propagation for many *Trichosporonales*.

Convergent evolution of fused *MAT* loci in *Trichosporonales* and *Tremellales*

Overall, our finding of fused *MAT* loci with a single *HD* gene in all of the *Trichosporonales* species investigated is most consistent with the hypothesis that a fusion event of the *HD* and *P/R* loci followed by loss of one *HD* gene per *MAT* allele occurred independently in the *Trichosporonales* and the pathogenic Cryptococci lineages. The alternative hypothesis that the fusion of the *HD* and *P/R* loci occurred in the common ancestor of the *Trichosporonales* and pathogenic Cryptococci is less parsimonious as it would imply multiple independent reversions to tetrapolarity in the non-pathogenic Cryptococci. In addition, while several mating-associated genes can be found in the *MAT* loci of both groups, the majority of genes within the *MAT* locus of the *Trichosporonales* is not found in the *MAT* locus in the *Tremellales*, consistent with independent fusion events in the two lineages. This hypothesis is also supported by the combination of the *SXII/SXI2* genes and the *STE3a/α* alleles in the A1 and A2 alleles of the *Trichosporonales*, which is different from those of the *MATa* and *MATα* alleles of the pathogenic Cryptococci. A model is proposed in Fig 5 to explain the current situation in the *Tremellomycetes*, where recruitment of several mating-associated genes into the *P/R* loci of a tetrapolar ancestor occurred first as these genes can also be found within the *P/R* loci of extant *Tremellales* with unlinked *MAT* loci [29-31]. After the split of the *Tremellales* and *Trichosporonales* lineages, fusion of the *HD* and *P/R* loci as well as the loss of one *HD* gene per *MAT* allele occurred in the ancestor of the *Trichosporonales*, whereas within the *Tremellales* this happened only in the ancestor of the pathogenic Cryptococci.

An independent fusion might suggest that similar selective pressures were acting to result in similar evolutionary trajectories in the *Trichosporonales* and pathogenic Cryptococci. In the latter group, it has been hypothesized that the pathogenic lifestyle might make it difficult to find a mating partner while associated with a host. In this case, linkage between the two *MAT* loci is expected to be favored by selection, as this increases the odds of compatibility between the gametes derived from a single diploid zygote [4, 7, 8, 17, 25]. Extant members of

the *Trichosporonales* can be associated with hosts as commensals or pathogens, but this lineage also includes many soil- or water-associated saprobes [36, 37]. It is possible that the ancestor in which *MAT* fusion occurred was associated with a host, and that the fused *MAT* loci remained stable during evolution of species with saprobic lifestyles. However, a scarcity of compatible mating partners might also occur under other conditions, e.g. if a population is derived from few progenitor cells that propagated through mitotic cell divisions during favorable conditions and switched to sexual development when nutrients were depleted.

In addition to the fused *MAT* loci, the presence of only one *HD* gene per *MAT* allele in all examined *Trichosporonales* is similar to pathogenic Cryptococci. This was noted in the initial analysis of genome sequences for two *Trichosporonales* strains now designated *MAT* A2 [43], and the current study shows that this is a conserved feature throughout the *Trichosporonales*. Outside of the *Tremellomycetes*, the majority of analyzed basidiomycete species harbor at least two *HD* genes per *MAT* allele, one *HD1* and one *HD2* gene, which are not compatible with each other, irrespective of whether the mating system is tetrapolar or bipolar [15, 17]. An exception is the genus *Wallemia*, where a fused *MAT* exists with only one *HD* gene, but until now, only one *MAT* allele harboring a *SXII* homolog is known in this genus [17, 73]. Functionally, two compatible *HD1* and *HD2* genes from different *MAT* alleles are necessary and sufficient for sexual development not only in *C. neoformans*, where only two genes are present after mating [74, 75], but also in *U. maydis* and *Coprinopsis cinerea*, where two (*U. maydis*) or more (*C. cinerea*) compatible *HD1/HD2* combinations can be present after mating [76-78]. The presence of multiple *HD* gene paralogs or increased *HD* allele diversity within a species is advantageous under outcrossing, because it allows more frequent dikaryon viability after mating. However, if a species is predominantly selfing, a single *MAT* locus with two alleles will provide the highest percentage of compatibility between gametes from a single tetrad [4, 15, 17]. In such cases, the loss of one *HD* gene per *MAT* locus should not be a problem from either a functional or evolutionary point of view, and thus this genomic configuration might be observed in other species with fused *MAT* loci, unless the *HD* genes have other functions unrelated to mating.

One common feature of fungal *MAT* loci and sex chromosomes in other eukaryotes is the recruitment of sex-associated genes. The presence of the *STE11* and *STE20* within the core *MAT* loci of *Trichosporonales* and *Tremellales* suggests an ancient linkage of these genes to a *MAT* locus. In *C. neoformans*, *STE20* is required for the formation of proper heterokaryotic filaments and basidia after mating, and *STE11* α is required for mating and filament formation [79, 80]. *STE20* displays *MAT*-specific alleles [29, 30] and is also located within the *P/R* locus

of other basidiomycete species, e.g. *Leucosporidium scottii* and red yeasts in the *Pucciniomycotina* [16].

Genomic signatures that distinguish the *MAT* loci from surrounding genomic regions in *Trichosporonales*

Another common feature of fungal *MAT* loci and the sex-determining regions in other eukaryotic groups is that recombination is usually suppressed in these regions. In several fungi, suppressed recombination is observed not only within the *MAT* locus itself, but spreading out from the *MAT* locus along the *MAT*-carrying chromosome resulting in so-called evolutionary strata of stepwise recombination suppression, similar to findings in sex-chromosomes of mammals and plants. Examples are the ascomycete *Neurospora tetrasperma* and the basidiomycete genus *Microbotryum* [7, 9, 25, 28, 81-85]. In the latter, it was recently shown that the linkage of *HD* and *P/R* loci and subsequent establishment of regions of suppressed recombination extending beyond the fused *MAT* loci evolved at least five times independently within the genus [7]. In contrast, the analysis of *V. humicola* did not yield indications for suppressed recombination beyond the core *MAT* region that lies between the *SXII/SXI2* and pheromone genes. Furthermore, the gene order within the *MAT* alleles and the *HD/STE3* allele combinations in the analyzed *Trichosporonales* species are largely conserved, suggesting that the fusion of *HD* and *P/R* loci occurred once in a common ancestor. Regions outside of the *MAT* locus have in general undergone substantial rearrangements (e.g. translocations) in different species as seen by the varying genomic locations of *STE12* in the genus *Cutaneotrichosporon* and the lack of synteny outside of the *MAT* region in more distantly related *Trichosporonales*.

In *Microbotryum*, it was shown that evolutionary strata ranging from mating-type- to species-specific patterns can be distinguished along chromosomes with recently linked *MAT* loci [7, 25]. These analyses were performed based on alleles from several strains per species and of both mating types. One challenge for the phylogenetic analysis in the *Trichosporonales* is that most species are currently represented by a single strain. The analysis of additional alleles from more strains from each species will be necessary to fully evaluate the phylogenetic distribution of alleles.

In the *Trichosporonales*, the two *MAT* alleles, A1 and A2, differ in general by two inversions. Because such inversions suppress single-crossover recombination in the inverted region, and in this way keep locally adapted alleles linked, they may be under positive

selection within the different species. However, if the *MAT* fusion is relatively ancient, i.e. occurred in the last common ancestor of the *Trichosporonales*, an open question is why the genetic differences in the two known alleles are not more pronounced. One explanation could be that recombination within the inverted regions might still be possible via non-crossover gene conversion or double crossover [63]. This would be consistent with the pattern observed in *V. humicola* of lower divergence between A1 and A2 alleles within the inverted regions compared to divergence at the inversion breakpoints. However, the evolutionary consequences of double crossover and gene conversion on genetic divergence within the inverted region likely differ: the former is more likely to transfer a large segment, whereas the latter only allows small segments of DNA to be exchanged. Hence, if an inversion captures multiple locally adapted loci, gene conversion may more easily erase genetic divergence while double crossovers are more likely to result in maladaptive associations. In *C. neoformans*, gene conversion occurs in a GC-rich intergenic region within the *MAT* locus and was proposed as a mechanism for maintaining functionality of those genes within the *MAT* locus that are essential [35]. Gene conversion has also been observed in the regions with suppressed recombination of the mating-type chromosomes of *N. tetrasperma*, in the mating type locus of the green algae *Chlamydomonas reinhardtii*, and in sex chromosomes of animals [86-92].

Gene conversion might be associated with an increase in the GC content as it has been observed in a number of eukaryotes that gene conversion can be biased towards GC [93-95], which would not be compatible with the observed lower GC content in the *MAT* region of *Trichosporonales*. However, a recent analysis of two fungal species found no evidence of a GC-bias in gene conversion [96], and therefore at present gene conversion as an explanation for low divergence between *MAT* alleles in *Trichosporonales* remains possible. It is tempting to speculate that the lower GC content observed in the *MAT* region of *Trichosporonales* is connected to the suppression of recombination or the (yet unknown) mechanisms that lead to the low levels of divergence between the *MAT* alleles. In *C. neoformans*, two regions of higher GC content outside of the *MAT* locus are associated with increased recombination [97]. A correlation between higher GC content and increased recombination has been demonstrated in humans, although it is not clear in this case what is cause and what is effect, whereas in the yeast *Saccharomyces cerevisiae*, a correlation was described at the kilobase range, but no long-range correlation was found [98-100].

Another hypothesis to explain the low levels of divergence between the *MAT* alleles could be same mating-type mating ("unisexual reproduction", i.e. sexual reproduction without the

need of a partner with a different mating type), which occurs in several pathogenic Cryptococci. Similar to reproduction after fusion of *MAT_a* and *MAT_α* cells, "unisexual reproduction" also entails diploidization and meiosis including genetic exchange [101-103]. Diploidization can occur without cell fusion, e.g. through endoreplication, or through fusion of two cells carrying the same *MAT* allele [104]. In the latter case, this would allow recombination within the *MAT* region, which could prevent degeneration of the *MAT* locus despite suppressed recombination when paired with a different *MAT* allele. Recombination within the *MAT* locus during "unisexual reproduction" was shown for *C. neoformans* [105]. So far, no sexual cycle has been observed in the *Trichosporonales*, and therefore it is not known what forms of sexual reproduction occur in this group. This is an important open question that requires detailed investigation in future studies.

Conclusions

In summary, we have shown that all analyzed *Trichosporonales* species contain fused mating type loci with a single *HD* gene. The two known *MAT* alleles differ by two inversions, but each allele is relatively stable across different species, making it more likely that the fusion of the *MAT* loci occurred in the last common ancestor of the *Trichosporonales*. Thus, this would be the most ancient fusion of *MAT* loci observed to date. The apparent evolutionary stability of the alleles A1 and A2 through many speciation events (even though neighboring regions underwent significant recombination) suggests strong selective pressure operates to maintain integrity of the alleles, probably due to retaining the advantages of fused *MAT* loci with few alleles. Another possible explanation might be that the observed combinations (*STE3_α* and *SXI2* in the A1, and *STE3_a* and *SXII* in the A2 allele) have a higher fitness than the alternative combinations. However, the presence of the A1* and A2* alleles in several *Trichosporonales* species shows that these combinations also occur, making this hypothesis less likely. Mechanistically, a combination of gene conversion during meiosis after fusion of cells with different mating types to account for the limited suppression of recombination, and meiotic recombination by crossing over during "unisexual reproduction" might explain the apparent evolutionary stability of the *MAT* region while maintaining (at least) two distinct alleles.

Based on the phylogenetic distribution, gene content, and allele combinations in fused *MAT* loci in pathogenic Cryptococci of the sister order *Tremellales*, fusion of *MAT* loci as well as the loss of one *HD* gene per *MAT* locus occurred independently in the

Trichosporonales and *Tremellales*. Thus, our data support a model of convergent evolution of the *MAT* locus at the molecular level in two different *Tremellomycetes* orders, similar to patterns observed in other fungal groups [1, 7, 21, 22]. Future analyses of additional *Tremellomycetes* groups that have not yet been analyzed with respect to their *MAT* loci will allow tests of the hypothesis that this evolutionary route has been repeated independently in other cases. It will be especially interesting to analyze earlier-branching sister lineages to the *Tremellales* and *Trichosporonales* with respect to their *MAT* loci configurations and the number of *HD* genes per *MAT* locus. Other important questions are the frequency and mechanisms of recombination in the region between the fused *HD* and *P/R* loci. These questions can be addressed once *Trichosporonales* species have been identified that undergo sexual reproduction in the laboratory, so that the progeny of genetic crosses can be analyzed.

Materials and Methods

Strains and growth conditions

C. oleaginosum and *V. humicola* strains used in this study are given in Table 2. Strains were grown on YPD medium at 25°C on solid medium or in liquid culture with shaking (250 rpm) for preparation of nucleic acids.

CHEF (contour-clamped homogeneous electric field) electrophoresis analysis of the *C. oleaginosum* isolates

CHEF plugs of the *C. oleaginosum* isolates were prepared and the electrophoresis was carried out as described in previous studies with slight modification [29, 31]. Specifically, the plugs were run using two different sets of switching time to get better resolution of the larger (S4A Fig top, 20 – 30 minutes linear ramp switching time) and smaller (S4A Fig bottom, 120 – 360 seconds linear ramp switching time) chromosomes, respectively.

Genotyping and Analysis of mating type genes in *V. humicola* strains

A collection of *V. humicola* isolates were genetically screened by modified RAPD using primer 5'-CGTGCAAGGGAGCACC-3' with 48°C as annealing temperature (S1A Fig). The *V. humicola* strains showing identical genotypic profile as the type strain CBS571 (Table 2) were further analyzed for the presence of the *SXII*, *SXI2*, *STE3*, and *MYO2* genes using PCR

with oligonucleotides given in S3 Table. PCR fragments were either sequenced directly or cloned into pDrive (Qiagen, Hilden, Germany) and sequenced.

Genome sequencing, assembly, and annotation of *V. humicola* CBS4282 and *C. oleaginosum* ATCC20508

Genomic DNA samples were extracted using a modified CTAB protocol as previously reported [31, 106]. Specifically, to enrich samples with high molecular weight, after precipitation, the genomic DNA was picked out from the solution instead of spun down, and the samples were checked by CHEF for their sizes and integrity, following manufacturer's protocol (BioRad, Hercules, CA, USA). Sequencing of the ATCC20508 and CBS4282 genomes was carried out at the Sequencing and Genomic Technologies Core Facility of the Duke Center for Genomic and Computational Biology, using large insert library (15-20 kb) and PacBio Sequel (2.0 chemistry) for ATCC20508, and generating 151 nt paired-end Illumina reads for CBS4282. The PacBio sequence reads for *C. oleaginosum* ATCC20508 were assembled using the HGAP4 assembly pipeline based on the Falcon assembler [107] included in the SMRT Link v5.0.1 software by Pacific Biosciences. The Illumina sequence reads of *V. humicola* CBS4282 were trimmed using Trimmomatic (v0.36) [108] with the following parameters to remove adapter contaminations: ILLUMINACLIP:TruSeq3-PE-2.fa:2:30:10:1:TRUE LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:40. Trimmed reads were assembled with SPAdes (v3.11.1) [109], and contig sequences were improved using pilon [110] based on the Illumina reads mapped to the assembly using Bowtie2 (v2.2.6) [111]. k-mer frequencies were analyzed based on the CBS4282 Illumina reads as described previously [112, 113]. Gene models for the newly sequenced strains as well as for strains for which genome assemblies, but not annotation was available from GenBank were predicted *ab initio* using MAKER (v2.31.18) [114] with predicted proteins from *C. oleaginosum* as input [43].

Analysis of mating type regions, synonymous divergence, and repeat content

MAT regions in *Trichosporonales* genomes were identified by BLAST searches [115] against the well-annotated *MAT*-derived proteins from *C. neoformans* [34], and manually reannotated if necessary. The short and not well conserved pheromone precursor genes were not among the predicted genes, and were identified within the genome assemblies using

custom-made Perl scripts searching for the consensus sequence M-X(15-60)-C-[ILMVST]-[ILMVST]-X-Stop.

Synteny analysis of the genomes of *V. humicola* strains and *C. oleaginosum* strains was done with nucmer from the MUMmer package (v3.23) [116]. Synteny plots were drawn with Circos [117]. Synteny between *MAT* regions of different *Trichosporonales* species was based on bidirectional BLAST analyses of the corresponding predicted proteins. For the d_S plots, alleles in *V. humicola* strains were identified by two-directional BLAST analysis, and MUSCLE (v3.8.31) [118] was used to align the two alleles per gene per strain pair. Synonymous divergence and standard errors were estimated with the yn00 program of the PAML package (v4.9) [119]. Analysis of transposable elements and other repeats in *V. humicola* genomes was performed with RepeatMasker (Smit AFA, Hubley R, Green P. RepeatMasker Open-4.0. 2013-2015, <http://www.repeatmasker.org>) based on the RepbaseUpdate library [120] and a library of *de novo*-identified repeat consensus sequences for each strain that was generated by RepeatModeler (Smit AFA, Hubley R. RepeatModeler Open-1.0. 2008-2015, <http://www.repeatmasker.org/RepeatModeler/>) as described [113].

Linear synteny comparison along the *MAT*-containing scaffolds (S10 Fig) was generated with Easyfig [121] using a minimum length of 500 bp for BLASTN hits to be drawn.

Species tree and gene genealogies

Of the 24 *Trichosporonales* species for which genome assemblies were available we selected only 21 for phylogenetic analysis as the remaining three species (*Trichosporon coremiiforme*, *Trichosporon ovoides* and *Cutaneotrichosporon mucooides*) were shown to be hybrids in previous studies [49, 50]. Additionally, we selected other well-studied bipolar and tetrapolar representatives belonging to the three major Basidiomycota lineages, plus two ascomycetes as outgroup. To reconstruct the phylogenetic relationships among the selected members, the translated gene models of each species were clustered by a combination of the bidirectional best-hit (BDBH), COGtriangles (v2.1), and OrthoMCL (v1.4) algorithms implemented in the GET_HOMOLOGUES software package [122] to construct homologous gene families. The *Cryptococcus neoformans* H99 protein set was used as reference and clusters containing inparalogs (i.e. recent paralogs defined as sequences with best hits in its own genome) were excluded. A consensus set of 32 protein sequences was computed out of the intersection of the orthologous gene families obtained by the three clustering algorithms. Protein sequences were individually aligned with MAFFT v7.310 [123] using the L-INS-i

strategy and poorly aligned regions were trimmed with TrimAl (-gappyout). The resulting alignments were concatenated to obtain a final supermatrix consisting of a total of 21,690 amino acid sites. We inferred a maximum-likelihood phylogeny using the LG+F+R5 model of amino acid substitution in IQ-TREE v1.6.5 [124]. Branch support values were obtained from 10,000 replicates of both ultrafast bootstrap approximation (UFBoot) [125] and the nonparametric variant of the approximate likelihood ratio test (SH-aLRT) [126].

For phylogenetic analysis of selected genes within and outside the *MAT* region in *Trichosporonales*, protein alignments were generated and trimmed as above and subsequently used to infer maximum likelihood phylogenies in IQ-TREE. Consensus trees were graphically visualized with iTOL v4.3.3 [127].

Divergence time analysis

Divergence times were estimated in MEGA-X [128] using the RelTime approach [129]. Contrary to other methods, RelTime does not require assuming a specific model for lineage rate variation and was shown to be as accurate as other approaches using relaxed and strict molecular clock models [130]. The reconstructed species tree, with branch lengths in the units of number of substitutions per site, was used as input and transformed into an ultrametric tree with relative times. The final timetree was obtained by converting the relative node ages into absolute dates by using three calibration constraints: 0.42 million year (MY) corresponding to the divergence between *Microbotryum lychnidis-dioicae* and *Microbotryum silenes-dioicae* [131]; 41 MY for the *Ustilago* - *Sporisorium* split; and 413 MY representing the minimum age of Basidiomycota. The latter two calibration points were obtained from the Timetree website (<http://www.timetree.org/>), which should be referred to for additional information and references.

Data availability statement

The *V. humicola* CBS4282 genome sequence (BioProject PRJNA475686) has been deposited at DDBJ/ENA/GenBank under the accession QKWK00000000. The version described here is version QKWK01000000. Illumina reads have been deposited in the NCBI SRA database under accession SRP150316. The *C. oleaginosum* ATCC20508 genome sequence (BioProject PRJNA475739) has been deposited at DDBJ/ENA/GenBank under the accession QKWL00000000. The version described here is version QKWL01000000. Pacific

Biosciences reads have been deposited in the NCBI SRA database under accession SRP150334.

Acknowledgements

MN would like to thank Swenja Ellßel and Silke Nimtz for excellent technical assistance, and Ulrich Kück and Christopher Grefen for support at the Botany Department of the Ruhr-University Bochum.

Tables

Table 1. Genome assemblies that were analyzed in this study.

Genus	Species	Strain	<i>MAT</i> allele ²	GenBank acc. number	Reference
<i>Apiotrichum</i>	<i>brassicae</i>	JCM1599	A1	BCJI00000000	[50]
<i>Apiotrichum</i>	<i>domesticum</i>	JCM9580	A1	BCFW01000000	[45]
<i>Apiotrichum</i>	<i>gamsii</i>	JCM9941	A1	BCJN00000000	[50]
<i>Apiotrichum</i>	<i>gracile</i>	JCM10018	A2*	BCJO00000000	[50]
<i>Apiotrichum</i>	<i>laibachii</i>	JCM2947	A1	BCKV01000000	[50]
<i>Apiotrichum</i>	<i>porosum</i>	DSM27194	A1	RSCE00000000	[51]
<i>Apiotrichum</i>	<i>porosum</i>	JCM1458	A2	BCJG01000000	[50]
<i>Apiotrichum</i>	<i>veenhuisii</i>	JCM10691	A1*	BCKJ00000000	[50]
<i>Cutaneotrichosporon</i>	<i>arboriformis</i>	JCM14201	A2	BEDW01000001	[50]
<i>Cutaneotrichosporon</i>	<i>curvatus</i>	DSM101032	A2	LDEP01000000	[47]
<i>Cutaneotrichosporon</i>	<i>cutaneum</i>	JCM1462	A1	BCKU01000000	[50]
<i>Cutaneotrichosporon</i>	<i>cyanovorans</i>	JCM31833	A1	BEDZ01000001	[50]
<i>Cutaneotrichosporon</i>	<i>daszewskae</i>	JCM11166	A2	BEDX01000001	[50]
<i>Cutaneotrichosporon</i>	<i>dermatis</i>	JCM11170	A1*	BCKR01000001	[50]
<i>Cutaneotrichosporon</i>	<i>mucoides</i>	JCM9939 ¹	A1*, A2*	BCJT01000001	[50]
<i>Cutaneotrichosporon</i>	<i>oleaginosum</i>	ATCC20508	A2	QKWL01000000	this study
<i>Cutaneotrichosporon</i>	<i>oleaginosum</i>	ATCC20509	A2	MATS01000000	[46]
<i>Cutaneotrichosporon</i>	<i>oleaginosum</i>	IBC0246	A2	JZUH00000000	[43]
<i>Takashimella</i>	<i>koratensis</i>	JCM12878	A2	BCKT01000001	[50]
<i>Takashimella</i>	<i>tepidaria</i>	JCM11965	A1	BCKS01000001	[50]
<i>Trichosporon</i>	<i>asahii</i>	CBS8904	A2	JH925096	[132]
<i>Trichosporon</i>	<i>coremiiforme</i>	JCM2938 ¹	A1, A2	JXYL01000000	[49]
<i>Trichosporon</i>	<i>faecale</i>	JCM2941	A1	JXYK01000000	[49]
<i>Trichosporon</i>	<i>inkin</i>	JCM9195	A2	JXYM01000000	[49]
<i>Trichosporon</i>	<i>ovoides</i>	JCM9940 ¹	A1, A1	JXYN01000000	[49]
<i>Vanrija</i>	<i>fragicola</i>	JCM1530	A1	BEDY01000001	[50]
<i>Vanrija</i>	<i>humicola</i>	CBS4282	A1	QKWK01000000	this study
<i>Vanrija</i>	<i>humicola</i>	JCM1457 (CBS571)	A2	BCJF01000000	[50, 52]
<i>Vanrija</i>	<i>humicola</i>	JCM9575 (UJ1)	A2	BFAH01000000	[48]

¹Hybrid strain carrying two *MAT* alleles derived from the two different parental species that underwent hybridization (S2 Text).

²*MAT* alleles contain the following *HD* and *P/R* allele combinations: A1, *STE3α*+*SXI2*; A2, *STE3a*+*SXII*; A1*, *STE3α*+*SXII*; A2*, *STE3a*+*SXI2*.

Table 2. Strains used in this study.

Species	Strain	MAT allele	Source
<i>Cutaneotrichosporon oleaginosum</i>	ATCC20508	A2	American Type Culture Collection
<i>Vanrija humicola</i>	JCM1457 (= CBS571)	A2	Westerdijk Fungal Biodiversity Institute
<i>Vanrija humicola</i>	CBS4281	A2	Westerdijk Fungal Biodiversity Institute
<i>Vanrija humicola</i>	CBS4282	A1	Westerdijk Fungal Biodiversity Institute
<i>Vanrija humicola</i>	CBS4283	A1	Westerdijk Fungal Biodiversity Institute
<i>Vanrija humicola</i>	CBS8354	A2	Westerdijk Fungal Biodiversity Institute
<i>Vanrija humicola</i>	CBS8371	A2	Westerdijk Fungal Biodiversity Institute

Figure legends

Fig 1. Overview of the haploid *Trichosporonales* species for which *MAT* regions were analyzed. Shown here is a phylogenetic tree of basidiomycetes with a focus on groups where fusion of *MAT* loci occurred as was shown previously (*Cryptococcus*, *Ustilago*, *Malassezia*, *Microbotryum*, *Sporisorium*) [4, 7, 17]. The phylogenetic branching within the tree agrees with previous studies [4, 7, 39, 50, 133]. Next to each *Trichosporonales* species name (color coded according to their genus), the *MAT* alleles that are represented in the sequenced genomes are shown. Within one species, different strains carrying different alleles are currently known only for *V. humicola* and *A. porosum*. Abbreviations for *Trichosporonales* genus names are: *A* - *Apiotrichum*, *C* - *Cutaneotrichosporon*, *T* - *Trichosporon*, *Ta* - *Takashimella*, *V* - *Vanrija*. Abbreviations for other basidiomycete genus names are: *C* - *Cryptococcus*, *K* - *Kwoniella*, *M* - *Microbotryum*, *S* - *Sporisorium*, *Tr* - *Tremella*, *U* - *Ustilago*. Ascomycetes *Neurospora crassa* and *Saccharomyces cerevisiae* were used as outgroups. Branch lengths are given in number of substitutions per site (scale bar). Branch support was assessed by 10,000 replicates of both ultrafast bootstrap approximation (UFBoot) and the Shimodaira-Hasegawa approximate likelihood ratio test (SH-aLRT) and represented as circles colored as given in the key.

Fig 2. *MAT* loci of *Trichosporonales*. Classic mating type-defining genes (*SXII* or *SXI2*, *STE3*, and the pheromone precursor genes) are shown in red, other genes that are part of the mating type locus or flanking the mating type locus (gene *g6341*) of *C. neoformans* are shown in green, and genes not present in the *C. neoformans* mating type locus are shown in white. The *MAT* allele is indicated on the right, with A1 and A2 *MAT* alleles shown above and below the dashed line, respectively. For simplicity, the A1* and A2* alleles are not included in this Figure, and are instead depicted in S5 and S6 Figs. The proposed *MAT* locus region is enclosed in a yellow box in each strain. Orthologs are connected by grey or orange bars when in the same or opposite orientations, respectively.

Fig. 3. Phylogenetic analysis of proteins encoded by genes within and around the *MAT* loci in *Trichosporonales*. **A.** The A1 and A2 *MAT* alleles of *V. humicola* and *A. porosum* are shown for reference of the positions of the analyzed genes within the *MAT* region (depicted by black arrows). **B.** Individual genealogies of the selected genes across species. In the core

MAT region (enclosed in yellow in panel A), a deep trans-specific polymorphism is only seen for the *STE3* gene. In *V. humicola*, most of the analyzed genes within this region branch by mating type. The *MAT* alleles of the analyzed strains are given next to the phylogeny. Genus name abbreviations and other color features are given as in Figs 1 and 2.

Fig 4. *MAT* alleles of *V. humicola* strains. **A.** Two strains (JCM1457 and UJ1) are *MAT* A2, while strain CBS4282 is a *MAT* A1 strain. Genes shown in the same color (orange within the core *MAT* region, and purple outside of the *MAT* region) were the most similar pair in a pairwise BLASTN analyses among the three alleles of the different *V. humicola* strains. **B.** Schematic view of BLASTN results. For each analyzed gene, it is shown if alleles of the most similar pair came from the strains with the same (A2/A2) or different (A1/A2) *MAT* alleles. Within the core *MAT* region, higher sequence similarity was observed between the two A2 alleles than between A1 and A2 alleles. **C.** Synonymous divergence between *MAT* alleles in different *V. humicola* strains. Synonymous substitutions per synonymous site and standard errors ($d_s \pm SE$) are shown for pairwise comparisons between strains CBS4282 (A1), JCM1457 (A2), and UJ1 (A2) for genes in the region containing the mating type genes. For *STE3* genes, for which no d_s value could be calculated in comparisons of strains with different mating types, values are shown as undefined (Un) at the top of the diagram. The inset below shows d_s values in the region between the core *MAT* genes, gene order as in strain CBS4282 (A1), *STE3* is left out for clarity. **D.** Boxplots of distribution of d_s values shown in C (only genes for which values could be calculated were used in this comparison). The plots show the distribution of d_s values, with median value as a horizontal line in the box between the first and third quartiles. Outliers were left out for better visibility. Significantly different values are indicated (p-values from Student's t-test).

Fig 5. Model for the evolution of the *MAT* loci in *Tremellomycetes*. Genes at the *MAT* loci containing homeodomain transcription factor genes (*HD* locus) or pheromone and receptor genes (*P/R* locus) are shown in red/pink and blue, respectively. Genes involved in sexual development, but not originally part of a *MAT* locus are shown in green, other genes in grey. Only genes from the *C. neoformans* *MAT* locus that are also linked to the core *MAT* genes (*STE3*, *HD* genes) in the *Trichosporonales* are shown (*STE11*, *STE12*, *STE20*, *IKS1*, *MYO2*, *RPL22*). Other genes present at the *MAT* loci are left out for clarity. A trend towards integrating other developmental genes into the *MAT* loci is reflected in the recruitment of the

STE12 gene and an ancestral *STE11/STE20* cluster into the *P/R* locus. This resulted in a tetrapolar arrangement with (at least) two alleles for each *MAT* locus. In the *Tremellales*, tetrapolarity is retained in the extant *Tremella*, *Kwoniella*, and non-pathogenic *Cryptococcus* lineages, whereas in the pathogenic *Cryptococci*, the *MAT* loci fused and one *HD* gene was lost from each *MAT* allele. Fusion of the *MAT* loci occurred independently and earlier in the ancestor of the *Trichosporonales*, also with loss of one *HD* gene, resulting in two known *MAT* alleles with combinations of *SX11/SX12* and *STE3a/STE3 α* that differ from those in the known *Cryptococcus* alleles. The *STE12* gene is shown in outline only to indicate that it is not part of the *MAT* locus in all species; *STE12* may have been recruited to the *P/R* locus in the *Tremellales-Trichosporonales* common ancestor and eventually evicted from the *MAT* locus in the *Trichosporonales* possibly associated with the fusion event of *P/R* and *HD* loci, or instead *STE12* was recruited to *MAT* only in the *Tremellales*. The phylogenetic relationships and inferred dates of speciation (numbers in black on tree nodes) are depicted according to S7 Fig. The blue bars correspond to 95 % confidence intervals.

Supporting information

S1 Fig. Genome analysis of *V. humicola* strains.

S2 Fig. Analysis of Sxi proteins from *Tremellomycetes*.

S3 Fig. Multiple alignment of the N-terminus of Ste3 homologs in the *MAT* loci of several *Tremellomycetes*.

S4 Fig. Genome sequencing of *T. oleaginosus* ATCC20508 (*MAT* A2).

S5 Fig. *MAT* loci of *Apiotrichum* and *Cutaneotrichosporon* species.

S6 Fig. *MAT* loci of *Trichosporonales* lineages containing hybrid species.

S7 Fig. Analysis of divergence times of basidiomycete lineages.

S8 Fig. Synonymous divergence between alleles in different *V. humicola* strains.

S9 Fig. BLASTN scores of sliding window comparisons of *MAT* regions from *V. humicola* strains.

S10 Fig. Synteny analysis of *MAT*-containing scaffolds.

S11 Fig. GC content of the *MAT* regions and surrounding regions in several *Trichosporonales* strains.

S1 Table. Repeat analysis of *Vanrija humicola* strains.

S2 Table. Codon usage analysis of the three *V. humicola* strains CBS4282 (*MAT* A1), JCM1457 (*MAT* A2), and UJ1 (*MAT* A2).

S3 Table. Oligonucleotides used in this study.

S1 Text. Pheromone genes of *Tremellomycetes*.

S2 Text. *MAT* loci in hybrid species.

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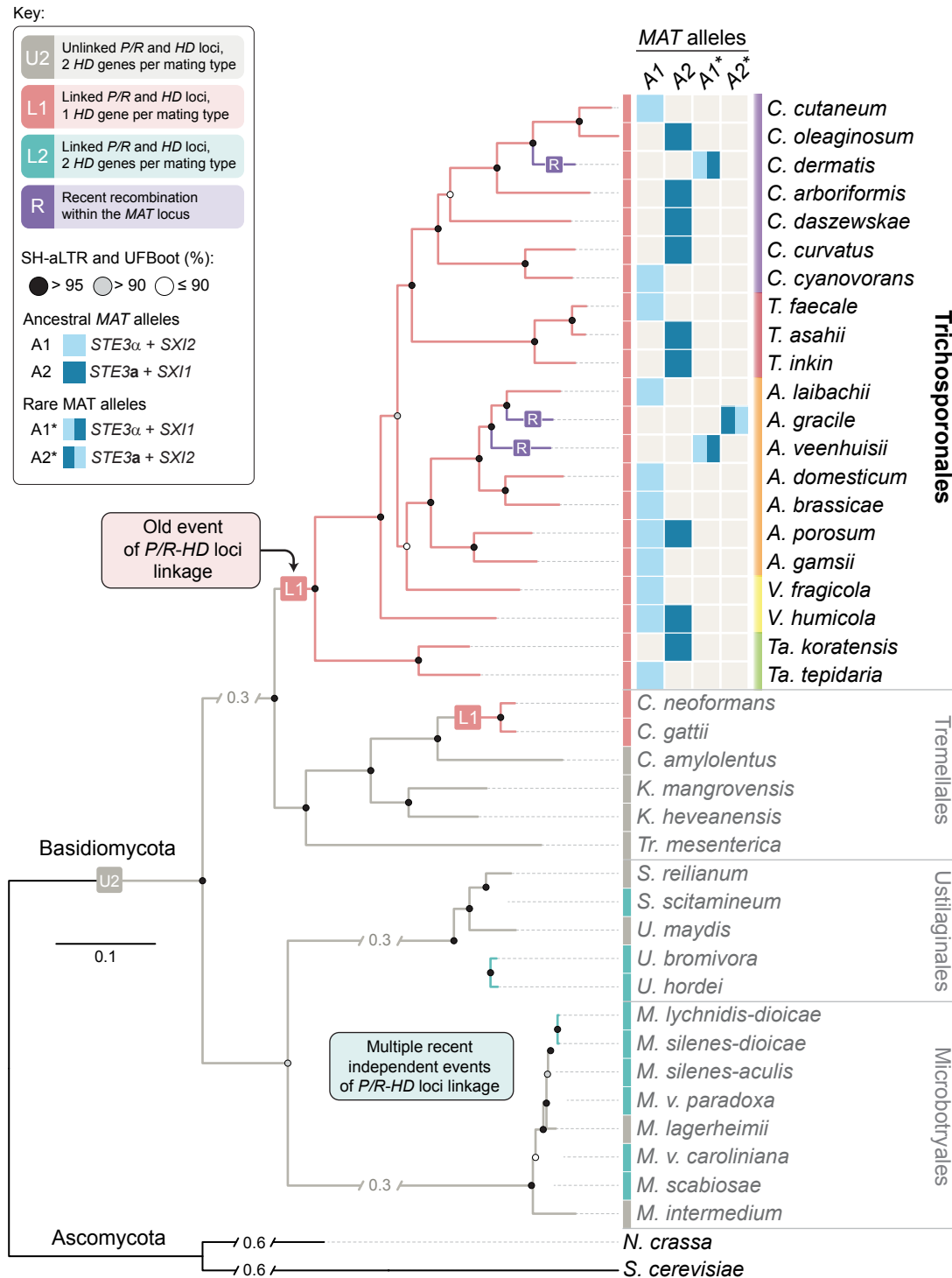


Figure 1

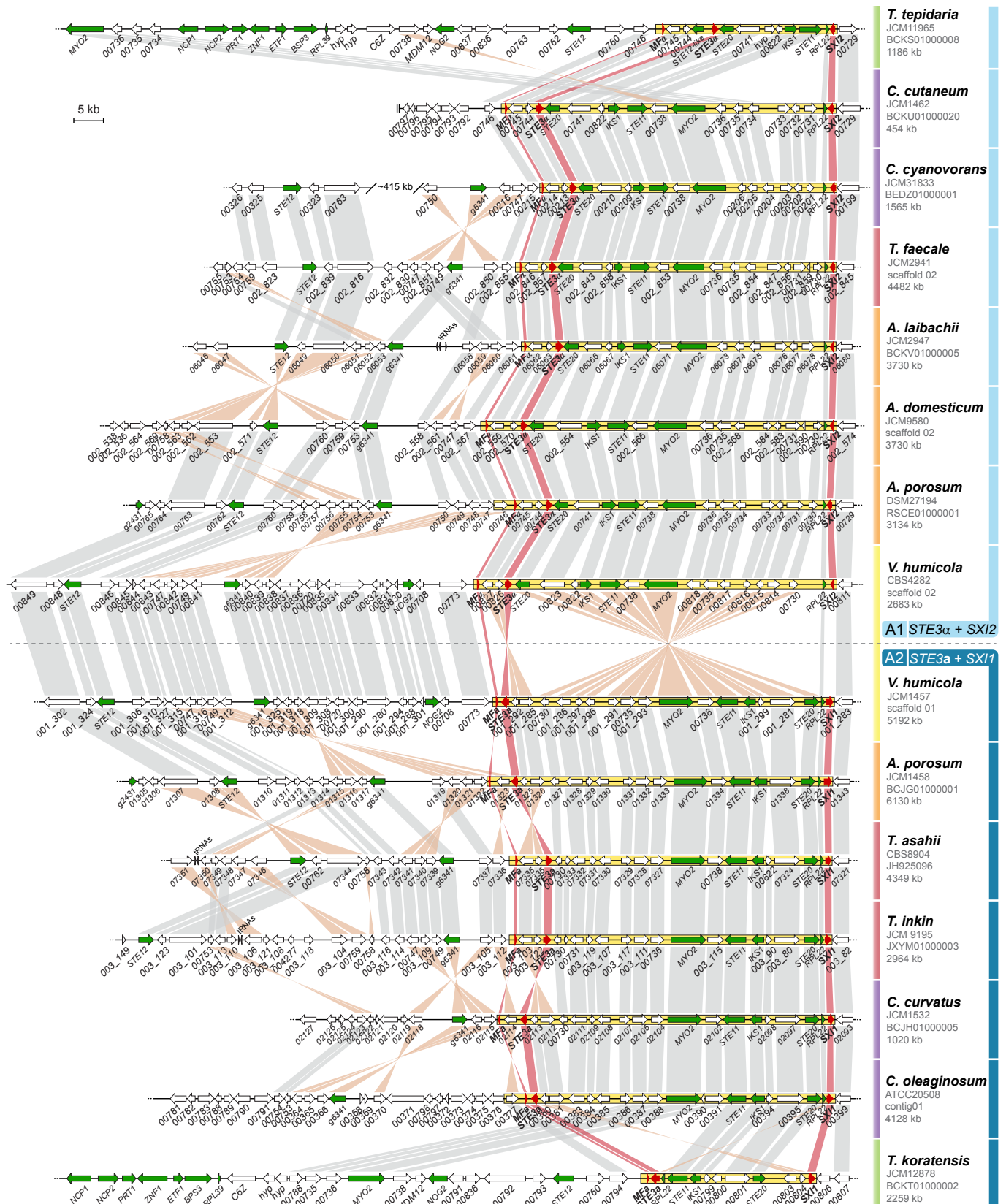


Figure 2

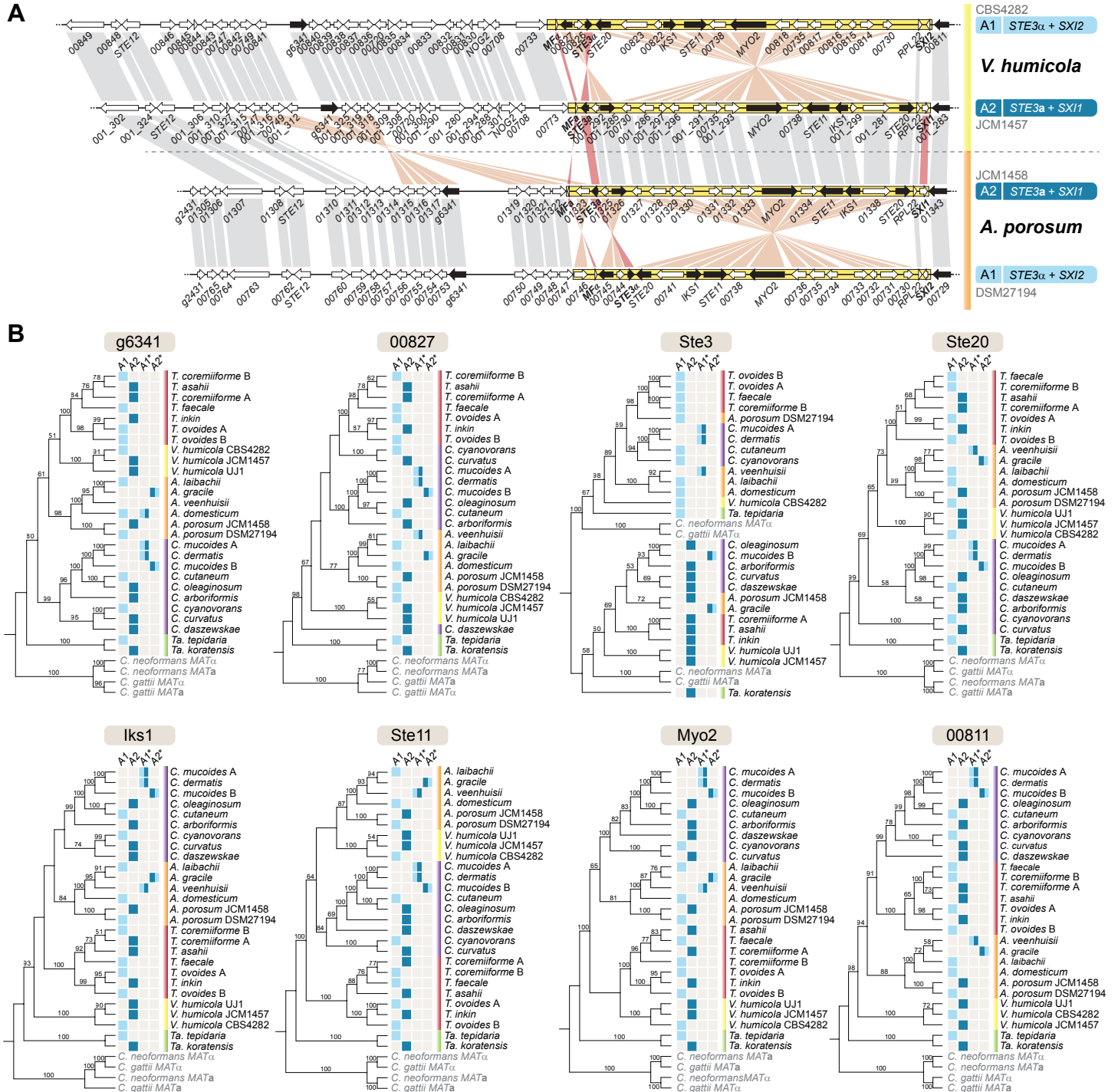
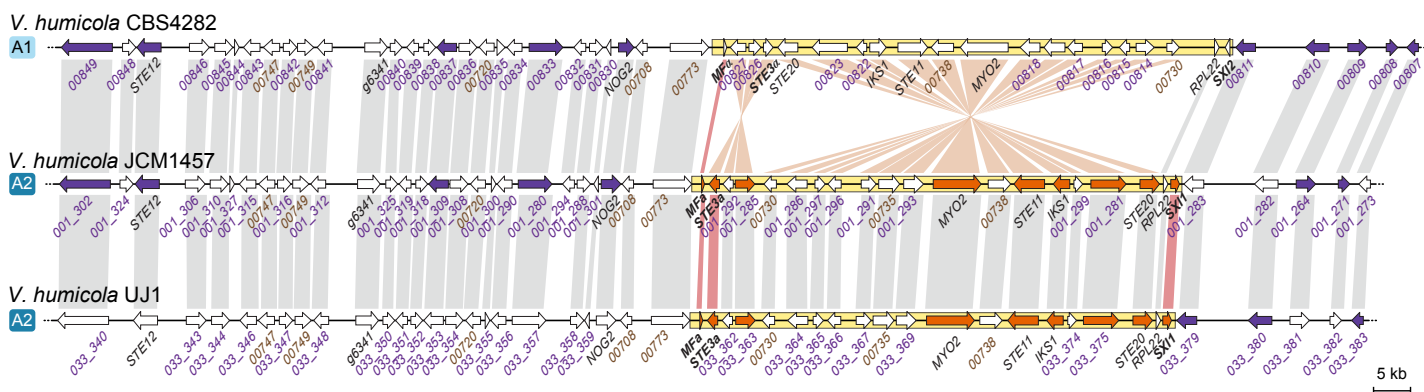
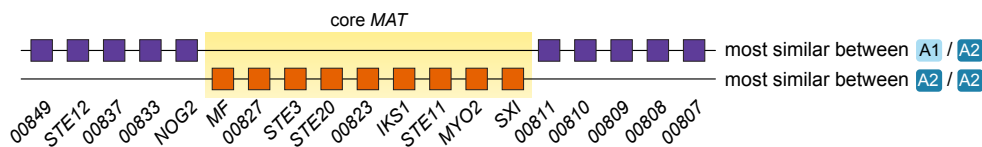


Figure 3

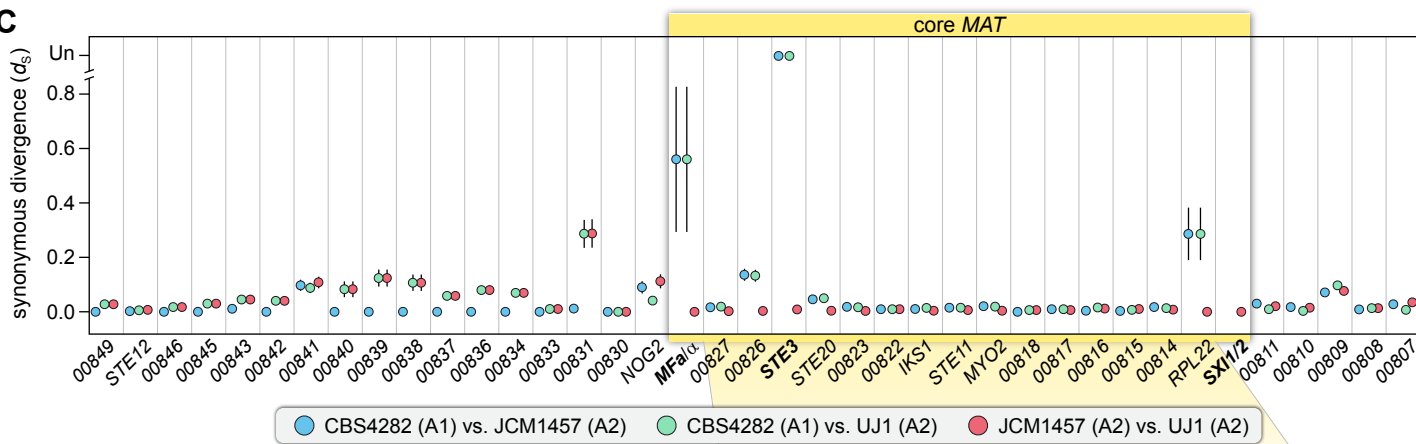
A



B



C



D

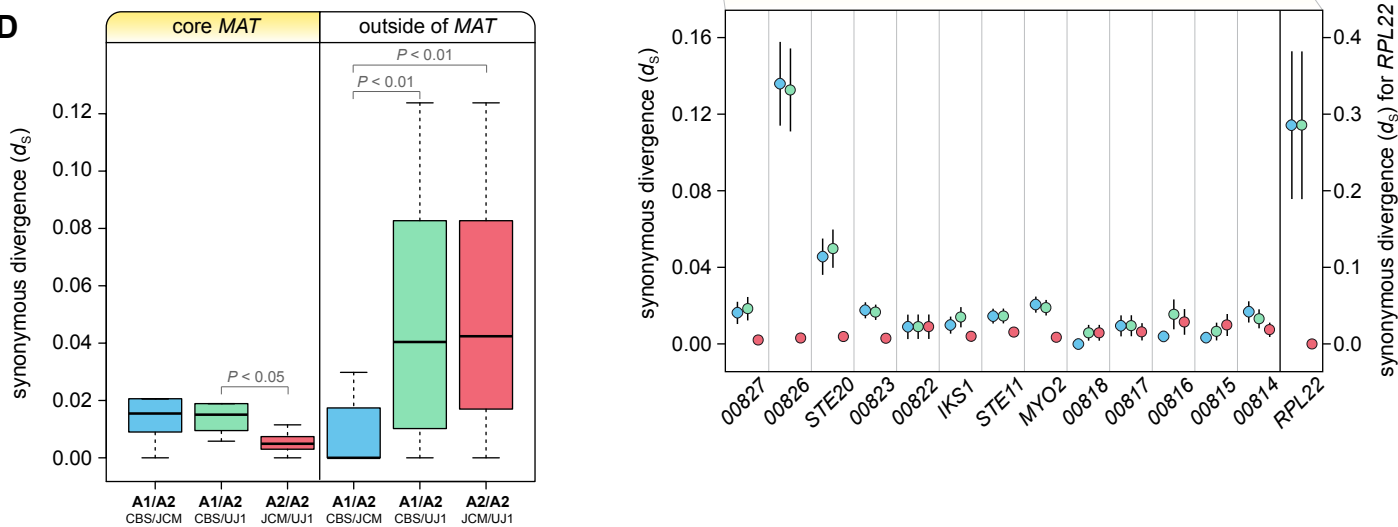


Figure 4

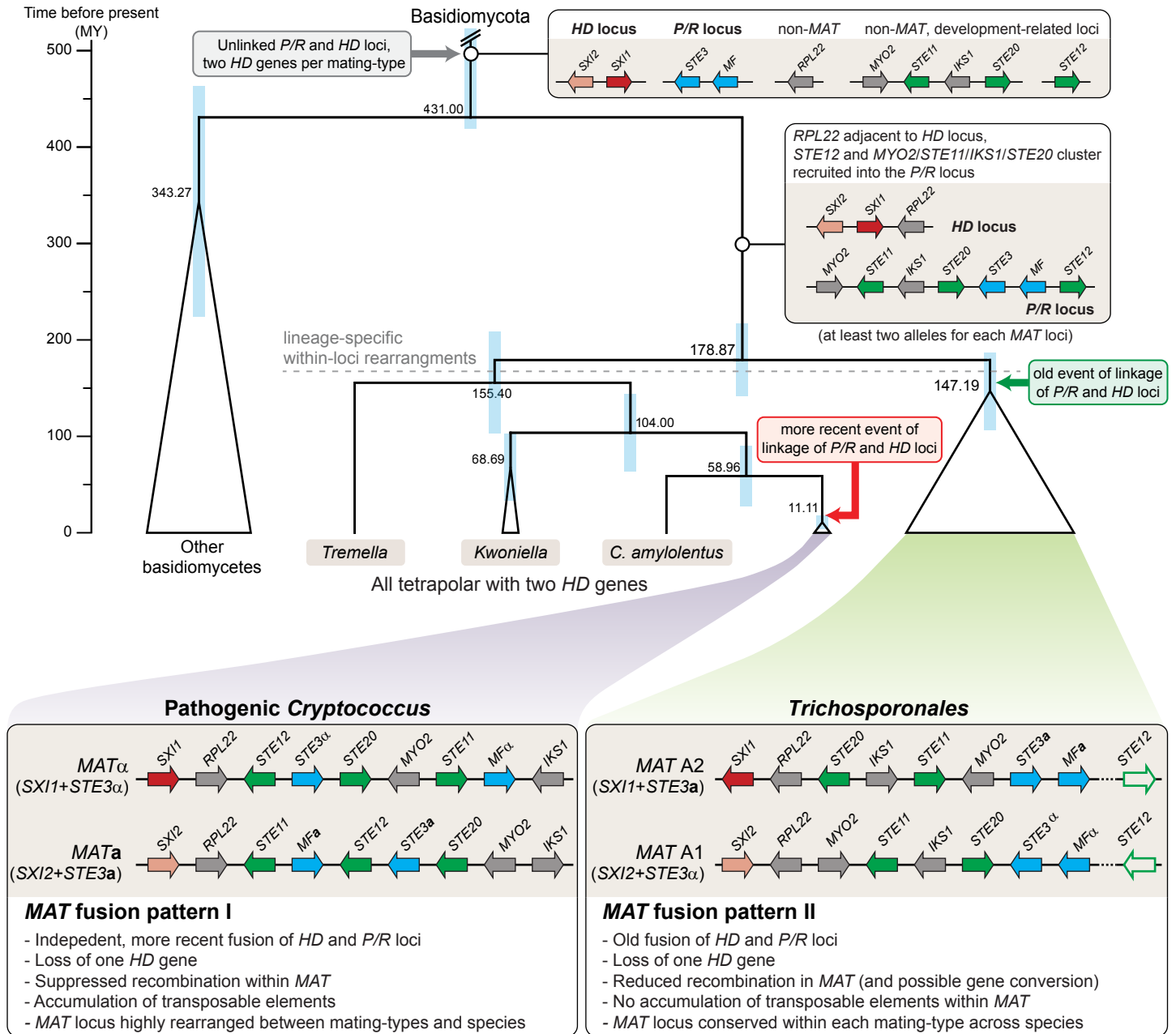


Figure 5