Uniparental nuclear inheritance following bisexual mating in fungi

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Abstract

Some remarkable animal species require an opposite-sex partner for their sexual development but discard the partner’s genome before gamete formation, generating hemi-clonal progeny in a process called hybridogenesis. Here, we discovered a similar phenomenon, termed pseudosexual reproduction, in a basidiomycete human fungal pathogen, *Cryptococcus neoformans*, where exclusive uniparental inheritance of nuclear genetic material was observed during bisexual reproduction. Analysis of strains expressing fluorescent reporter proteins revealed instances where only one of the parental nuclei was present in the terminal sporulating basidium. Whole-genome sequencing revealed the nuclear genome of the progeny was identical with one or the other parental genome. Pseudosexual reproduction was also detected in natural isolate crosses where it resulted in mainly *MATα* progeny, a bias observed in *Cryptococcus* ecological distribution as well. The mitochondria in these progeny were inherited from the *MATα* parent, resulting in nuclear-mitochondrial genome exchange. The meiotic recombinase Dmc1 was found to be critical for pseudosexual reproduction. These findings reveal a novel, and potentially ecologically significant, mode of eukaryotic microbial reproduction that shares features with hybridogenesis in animals.
Introduction

Most multicellular organisms in nature undergo (bi)sexual reproduction involving two partners of the opposite sex to produce progeny. In most cases, following the fusion of the two haploid gametes, the diploid zygote receives one copy of the genetic material from each parent. To produce these haploid gametes, a diploid germ cell of the organism undergoes meiosis, which involves recombination between the two parental genomes, generating recombinant progeny. Recombination confers benefits by bringing together beneficial mutations and segregating away deleterious ones (Dimijian, 2005; Meirmans, 2009). In contrast, some organisms undergo variant forms of sexual reproduction, including parthenogenesis, gynogenesis, androgenesis, and hybridogenesis, and in doing so, produce clonal or hemi-clonal progeny (Avise, 2015; Neaves & Baumann, 2011).

In parthenogenesis, a female produces clonal progeny from its eggs without any contribution from a male partner (Avise, 2015; Hörandl, 2009). Gynogenesis and androgenesis occur when the fusion of an egg with a sperm induces cell division to produce clonal female or male zygotes, respectively (Lehtonen, Schmidt, Heubel, & Kokko, 2013). During hybridogenesis, an egg from one species fuses with the sperm from another species to generate a hybrid diploid zygote (Lavanchy & Schwander, 2019). However, one of the parental genomes is excluded during development, in a process termed genome exclusion that occurs before gametogenesis. The remaining parental genome undergoes replication followed by meiosis to produce an egg or a sperm. The sperm or egg then fuses with an opposite-sex gamete to generate a hemiclonal progeny. Because only one parent contributes genetic material to the progeny, but both parents are physically required, this phenomenon has been termed sexual parasitism (Lehtonen et al., 2013; Umphrey, 2006). While most of the reported cases of hybridogenesis are from female populations, recent reports suggest that it may also occur in male populations of some species (Dolezalkova et al., 2016; Schwander & Oldroyd, 2016). Currently, hybridogenesis has only been observed in the animal kingdom in some species of frogs, fishes, and snakes. Plants also exhibit parthenogenesis (aka apomixis), along with gynogenesis and androgenesis (Lehtonen et al., 2013; Mirzaghaderi & Horandl, 2016).

Unlike animals, most fungi do not have sex chromosomes; instead, cell-type identity is defined by the mating-type (MAT) locus (Heitman, 2015; Heitman, Sun, & James, 2013). While many fungi are heterothallic, with opposite mating-types in different individuals, and undergo
sexual reproduction involving two partners of compatible mating-types, other fungi are homothallic, with opposite mating-types residing within the same organism, and can undergo sexual production during solo culture in the absence of a mating partner. One class of homothallic fungi undergoes unisexual reproduction, during which cells of a single mating type undergo sexual reproduction to produce clonal progeny, similar to parthenogenesis (Heitman, 2015; Lee, Ni, Li, Shertz, & Heitman, 2010). Gynogenesis and hybridogenesis have not been identified in the fungal kingdom thus far.

_Cryptococcus neoformans_ is a basidiomycete human fungal pathogen that exists as either one of two mating types, _MATa_ or _MATα_ (Sun, Coelho, David-Palma, Priest, & Heitman, 2019). During sexual reproduction, two haploid yeast cells of opposite mating type interact and undergo cell-cell fusion (Kwon-Chung, 1975, 1976; Sun, Priest, & Heitman, 2019). The resulting dikaryotic zygote then undergoes a morphological transition and develops into hyphae whose termini mature to form basidia. In the basidium, the two parental nuclei fuse (karyogamy), and the resulting diploid nucleus undergoes meiosis to produce four daughter nuclei (Idnurm, 2010; Kwon-Chung, 1976; Sun, Priest, et al., 2019; Zhao, Lin, Fan, & Lin, 2019). These four haploid nuclei repeatedly divide via mitosis and bud from the surface of the basidium to produce four long spore chains. Interestingly, in addition to this canonical heterothallic sexual reproduction, a closely related species, _C. deneoformans_ can undergo unisexual reproduction (Lin, Hull, & Heitman, 2005; Roth, Sun, Billmyre, Heitman, & Magwene, 2018; Sun, Billmyre, Mieczkowski, & Heitman, 2014).

In a previous study, we generated a genome-shuffled strain of _C. neoformans_, VYD135α, by using the CRISPR-Cas9 system targeting centromeric transposons in the lab strain H99α. This led to multiple centromere-mediated chromosome arm exchanges in strain VYD135α when compared to the parental strain H99α, without any detectable changes in gene content between the two genomes (Yadav, Sun, Coelho, & Heitman, 2020). Additionally, strain VYD135α exhibits severe sporulation defects when mated with strain KN99a (which is congenic with strain H99α but has the opposite mating type), likely due to the extensive chromosomal rearrangements introduced into the VYD135α strain. In this study, we show that the genome-shuffled strain VYD135α can in fact produce spores in crosses with _MATa_ _C. neoformans_ strains after prolonged incubation. Analysis of these spores reveals that the products from each individual basidium contain genetic material derived from only one of the two parents. Whole-
genome sequencing of the progeny revealed an absence of recombination between the two parental genomes. The mitochondria in these progeny were found to always be inherited from the \textit{MATa} parent, consistent with known mitochondrial uniparental inheritance (UPI) patterns in \textit{C. neoformans} (Sun, Fu, Ianiri, & Heitman, 2020). Using strains with differentially fluorescently labeled nuclei, we discovered that in a few hyphal branches as well as in basidia, only one of the two parental nuclei was present and produced spores, leading to uniparental nuclear inheritance. We also observed the occurrence of such uniparental nuclear inheritance in wild-type and natural isolate crosses. Furthermore, we found that the meiotic recombinase Dmc1 plays a central role during this unusual mode of reproduction of \textit{C. neoformans}. Overall, this mode of sexual reproduction of \textit{C. neoformans} exhibits striking parallels with hybridogenesis in animals.

\textbf{Results}

\textbf{Chromosomal translocation strain exhibits unusual sexual reproduction}

Previously, we generated a strain (VYD135\(\alpha\)) with eight centromere-mediated chromosome translocations compared to the wild-type parental isolate H99\(\alpha\) (Yadav et al., 2020). Co-incubation of the wild-type strain KN99\(a\) with the genome-shuffled strain VYD135\(\alpha\) resulted in hyphal development and basidia production, but no spores were observed during a standard two-week incubation. However, when sporulation was assessed at later time points in the VYD135\(\alpha\) x KN99\(a\) cross, we observed a limited number of sporulating basidia (16/1201 = 1.3\%) after five weeks compared to a much greater level of sporulation in the wild-type H99\(\alpha\) x KN99\(a\) cross (524/599 = 88\%) (Figure 1A-D). None of these strains exhibited any filamentation on their own even after 5-weeks of incubation, indicating that the sporulation events were not a result of unisexual reproduction (Figure 1A-B). To analyze this delayed sporulation process in detail, spores from individual basidia were dissected and germinated to yield viable F1 progeny. As expected, genotyping of the mating-type locus in the H99\(\alpha\) x KN99\(a\) progeny revealed the presence of both mating types in spores derived from each basidium (Figure 1E and G, Table 1). In contrast, the same analysis for VYD135\(\alpha\) x KN99\(a\) revealed that all germinating progeny from each individual basidium possessed either only the \textit{MATa} or the \textit{MATa} allele (Figure 1E and G, Table 1). PCR assays also revealed that the mitochondria in all of these progeny were inherited from the \textit{MATa} parent, in accord with known UPI (Figure 1F-G).
These results suggest the inheritance of only one of the parental nuclei in the VYD135α x KN99α F1 progeny. The presence of mitochondria from only the MATα parent in MATα progeny further confirmed that these progeny were the products of fusion between the parent strains and were not the products of unisexual reproduction.

Fluorescence microscopy reveals uniparental nuclear inheritance after mating

Next, we tested whether the uniparental inheritance detected at the MAT locus also applied to the entire nuclear genome. To address this, we established a fluorescence-based assay in which the nuclei of strains H99α and VYD135α were labeled with GFP-H4, whereas the KN99a nucleus was marked with mCherry-H4. In a wild-type cross (H99α x KN99a), the nuclei in the hyphae as well as in the spores were yellow to orange because both nuclei were in a common cytoplasm and thus incorporated both the GFP- and the mCherry-tagged histone H4 proteins (Figure 2-figure supplement 1A and B). We hypothesized that in the cases of uniparental nuclear inheritance, only one of the nuclei would reach the terminal basidium and would thus harbor only one fluorescent nuclear color signal (Figure 2-figure supplement 1A).

After establishing this fluorescent tagging system using the wild-type strains H99α x KN99a, shuffled-strain VYD135α x KN99a crosses with fluorescently labeled strains were examined. In the wild-type cross, most of the basidia formed robust spore chains with both fluorescent colors observed in them, while a small population (~1%) of basidia exhibited spore chains with only one color, representing uniparental nuclear inheritance (Figure 2A and Figure 2-figure supplement 2A). In contrast, the majority of the basidium population in the shuffled-strain VYD135α x KN99a cross did not exhibit sporulation, and the two parental nuclei appeared fused but undivided (Figure 2B and Figure 2-figure supplement 2B). A few basidia (~1%) bore spore chains with only one fluorescent color, marking uniparental nuclear inheritance events. While the basidia with uniparental nuclear inheritance in the H99α x KN99a cross were a small fraction (~1%) of sporulating basidia, the uniparental basidia accounted for all of the sporulating basidia in the VYD135α x KN99a cross. Taken together, these results show that the uniparental nuclear inheritance leads to the generation of clonal progeny but requires mating, cell-cell fusion between parents of two opposite mating types. Thus, this process defies the main purpose of sexual reproduction, which is to produce recombinant progeny from two parents. Based on these observations, we define the process of uniparental nuclear inheritance during sporulation in C.
*neofor mans* as pseudosexual reproduction (and it is referred to as such hereafter). The progeny obtained via this process will be referred to as the uniparental progeny because they inherit a nuclear genome derived from only one of the two parents.

**Pseudosexual reproduction also occurs in natural isolates**

After establishing the pseudosexual reproduction of lab strains, we sought to determine whether such events also occur with natural isolates. For this purpose, we selected two wild-type natural isolates, Bt63a and IUM96-2828a (referred to as IUM96a hereafter) (Desjardins et al., 2017; Keller, Viviani, Esposto, Cogliati, & Wickes, 2003; Litvintseva et al., 2003). IUM96a belongs to the same lineage as H99α/KN99a (VNI) and exhibits approximately 0.1% genome divergence from the H99α reference genome. Bt63a belongs to a different lineage of the *C. neoformans* species (VNBI) and exhibits ~0.5% genetic divergence from the H99α/KN99a genome. Both the Bt63a and the IUM96a genomes exhibit one reciprocal chromosome translocation with H99α, and as a result, share a total of ten chromosome-level changes with the genome-shuffled strain VYD135α (Figure 3A). None of these strains are self-filamentous even after prolonged incubation on mating media but both cross efficiently with H99α and VYD135α (Figure 3-figure supplement 1A).

The H99α x Bt63a strains crossed rapidly (within a week) producing robust sporulation from most of the basidia observed. The VYD135α x Bt63a cross underwent a low frequency of sporulation (12 spore-producing basidia/840 basidia=1.4%) in 2 to 3 weeks (Figure 3-figure supplement 1B). Dissection of spores from the H99α x Bt63a cross revealed a low germination frequency (average of 25%) with two of the basidia showing no spore germination at all (Supplementary file 1a). This result is consistent with previous results, and the low germination frequency could be explained by the genetic divergence between the two strains (Morrow et al., 2012). Genotyping of germinated spores from the H99α x Bt63a cross revealed both MATα and MATα progeny from individual basidia, with almost 75% of the meiotic events generating progeny that were heterozygous for the MAT locus (Figure 3-figure supplement 1C and Supplementary file 1a). For the VYD135α x Bt63a cross, spores from 15/20 basidia germinated and displayed a higher germination frequency than the H99α x Bt63a cross (Supplementary file 1a). Interestingly, all germinated progeny harbored only the MATα mating-type, whereas the
mitochondria were in all cases inherited from the \textit{MATa} parent (Figure 3-figure supplement 1C). These results suggest that pseudosexual reproduction also occurs with Bt63a and accounts for the high germination frequency of progeny from the VYD135\(\alpha\) x Bt63a cross. The occurrence of pseudosexual reproduction was also identified using the fluorescence-based assay with crosses between the GFP-H4 tagged VDY135\(\alpha\) and mCherry-H4 tagged Bt63a strains (Figure 3-figure supplement 2).

Crosses with strain IUM96a also revealed a low level of sporulation (19/842=2.3\%) with VYD135\(\alpha\) but a high sporulation frequency with H99\(\alpha\) (91\%) (Figure 3-figure supplement 1D). Analysis of progeny from crosses involving IUM96a revealed a similar pattern to what was observed with crosses involving KN99\(\alpha\). The progeny from H99\(\alpha\) x IUM96a exhibited variable basidium-specific germination frequencies and inherited both \textit{MATa} and \textit{MAT\(\alpha\)} in each basidium, whereas VYD135\(\alpha\) x IUM96a progeny from each basidium inherited exclusively either \textit{MATa} or \textit{MAT\(\alpha\)} (Figure 3-figure supplement 1E, and Supplementary file 1b).

Interestingly, we observed co-incident uniparental \textit{MAT} inheritance and a high germination frequency in progeny of basidia 7, 8, and 9 from the H99\(\alpha\) x IUM96a cross as well (Figure 3-figure supplement 1E, and Supplementary file 1b). Taken together, these results suggest that this unusual mode of sexual reproduction occurs with multiple natural isolates. We further propose that pseudosexual reproduction occurs in nature in parallel with canonical sexual reproduction.

\textbf{Uniparental progeny completely lack signs of nuclear recombination between the two parents}

As mentioned previously, H99\(\alpha\) (as well as the H99\(\alpha\)-derived strain VYD135\(\alpha\)) and Bt63a have approximately 0.5\% genetic divergence. The occurrence of pseudosexual reproduction in the VYD135\(\alpha\) x Bt63a cross allowed us to test if the two parental genomes recombine with each other during development. We subjected progeny from crosses VYD135\(\alpha\) x Bt63a and H99\(\alpha\) x Bt63a to whole-genome sequencing. As expected, for the H99\(\alpha\) x Bt63a cross, both parents contributed to the nuclear composition of their progeny, and there was clear evidence of meiotic recombination as determined by variant analysis (Figure 3B). However, when the VYD135\(\alpha\) x Bt63a progeny were similarly analyzed, the nuclear genome in each of the progeny was found to be inherited exclusively from only the VYD135\(\alpha\) parent (Figure 3C and
Figure 3-figure supplement 3), and the progeny exhibited sequence differences across the entire Bt63a genome. In contrast, the mitochondrial genome was inherited exclusively from the Bt63a parent (Figure 3D and Figure 3-figure supplement 4), in accord with the PCR assay results discussed above. Additionally, the whole-genome sequencing data also revealed that while most of the H99α x Bt63a progeny exhibited aneuploidy, the genome-shuffled strain VYD135α x Bt63a progeny were euploid (Figure 3-figure supplement 5A and B), and based on flow cytometry analysis, these uniparental progeny were haploid (Figure 3-figure supplement 5C).

The progeny from crosses involving IUM96a as the MATa partner were also sequenced. Similar to the Bt63a analysis, the H99α x IUM96a progeny exhibited signs of meiotic recombination, whereas the VYD135α x IUM96a progeny did not (Figure 3-figure supplement 6). Congruent with the mating-type analysis, the progeny in each of the basidia exclusively inherited nuclear genetic material from only one of the two parents. Furthermore, the H99α x IUM96a progeny were found to be aneuploid for some chromosomes, while the progeny of VYD135α x IUM96a were completely euploid (Figure 3-figure supplement 7). We also sequenced four progeny from basidium 7 from the H99α x IUM96a cross, which were suspected to be uniparental progeny based on mating-type PCRs. This analysis showed that all four progeny harbored only H99α nuclear DNA and had no contribution from the IUM96a nuclear genome, further supporting the conclusion that pseudosexual reproduction occurs in wild-type crosses (Figure 3-figure supplement 6A). Similar to other progeny, the mitochondria in these progeny were inherited from the MATa parent (Figure 3-figure supplement 1E, and Supplementary file 1b). Combined, these results affirm the occurrence of a novel mode of sexual reproduction in C. neoformans, which is initiated by the fusion of two strains of opposite mating types, but whose progeny inherit DNA exclusively from one parent.

**Pseudosexual reproduction stems from nuclear loss via hyphal branches**

Fluorescence microscopy revealed that only one of the two parental nuclei undergoes meiosis and produces spores in approximately 1% of the total basidia population. Based on this finding, we hypothesized that the basidia with only one parental nucleus might arise due to nuclear segregation events during hyphal branching. To gain further insight into this process, the nuclear distribution pattern along the sporulating hyphae was studied. As expected, imaging of long hyphae in the wild-type cross revealed the presence of pairs of nuclei with both fluorescent...
markers along the length of the majority of hyphae (Figure 4A). In contrast, tracking of hyphae from basidia with spore chains in the genome-shuffled strain VYD135α x KN99a cross revealed hyphal branches with only one parental nucleus, which were preceded by a hyphum with both parental nuclei (Figure 4B, Figure 4-figure supplement 1A and B). Unfortunately, a majority of the hyphae (>30 independent hyphae) we tracked were embedded into the agar, and most of these could not be tracked to the point of branching. For some others, we were able to image the hyphal branching point where two nuclei separate from each other but were then either broken or did not have mature basidia on them (Figure 4-figure supplement 1B). In total, we observed seven events of nuclear loss at hyphal branching in independent experiments and were able to track two of them to observe sporulation or basidia formation at the tip. We also observed long hyphae with only one parental nucleus in the VYD135α x Bt63a cross as well, suggesting the mechanism might be similar between strains.

These results suggest that hyphal branching may facilitate the separation of one parental nucleus from the main hyphae harboring both parental nuclei. While this is the most plausible explanation based on our results, we cannot rule out other possible mechanisms, such as a role for clamp cells, leading to nuclear separation during hyphal growth. As a result, one of the parental genomes is excluded at a step before diploidization and meiosis, similar to the process of genome exclusion observed in hybridogenesis. We hypothesize that nuclear segregation can be followed by endoreplication occurring in these hyphal branches or in the basidium to produce a diploid nucleus that then ultimately undergoes meiosis and produces uniparental progeny, which will be explored in future studies.

**Meiotic recombinase Dmc1 is important for pseudosexual reproduction**

Because the genomes of the uniparental progeny did not show evidence of meiotic recombination between the two parents, we tested whether pseudosexual reproduction involves meiosis. Additionally, we sought to test our hypothesis that pseudosexual reproduction involves endoreplication that is followed by meiosis. We therefore tested whether Dmc1, a key component of the meiotic machinery, is required for pseudosexual reproduction. The meiotic recombinase gene *DMC1* was deleted in congeneric strains H99α, VYD135α, and KN99a, and the resulting mutants were subjected to crossing. A previous report documented that *dmc1Δ* bilateral crosses (both the parents are mutant for *DMC1*) display significantly reduced, but not completely
abolished, sporulation in *Cryptococcus* (Lin et al., 2005). We observed a similar phenotype with the H99α *dmc1Δ* x KN99α *dmc1Δ* cross. While most of the basidia were devoid of spore chains, a small percentage (21/760=2.7%) of the population bypassed the requirement for Dmc1 and produced spores (Figure 5A and Figure 5-figure supplement 1A). When dissected, the germination frequency for these spores was found to be very low (~22% on average) with spores from many basidia not germinating at all (Supplementary File 1c). Furthermore, *MAT*-specific PCRs revealed that some of the progeny were aneuploid or diploid. For VYD135α *dmc1Δ* x KN99α *dmc1Δ*, many fewer basidia (~0.1%) produced spore chains as compared to ~1% sporulation in VYD135α x KN99α (Figure 5A, B and Figure 5-figure supplement 1B). *dmc1* mutant unilateral crosses (one of the two parents is mutant and the other one is wild-type) sporulated at a frequency of 0.4% suggesting that only one of the parental strains was producing spores (Figure 5B). When a few sporulating basidia from the VYD135α *dmc1Δ* x KN99α *dmc1Δ* bilateral cross were dissected, two different populations of basidia emerged, one with no spore germination, and the other with a high spore germination frequency and uniparental *MAT* inheritance (Supplementary File 1c). We hypothesize that the basidia with a high spore germination frequency represent those that have escaped the normal requirement for Dmc1.

Overall, the *DMC1* deletion led to a 20-fold reduction in viable sporulation in the VYD135α x KN99α cross, observed as a 10-fold decrease from the number of sporulation events in the bilateral cross and a further 2-fold reduction in the number of basidia producing viable spores.

To further support these findings, *DMC1* was deleted in mCherry-H4 tagged KN99α and crossed with GFP-H4 tagged VYD135α. We hypothesized that GFP-H4 tagged VYD135α would produce spore chains in this cross because it harbors *DMC1*, whereas mCherry-H4 tagged KN99α *dmc1Δ* would fail to do so. Indeed, all 11 observed basidia with only the GFP-H4 fluorescence signal were found to produce spores, but only 2 out of 19 mCherry-H4 containing basidia exhibited sporulation (Figure 5-figure supplement 2). These results combined with the spore dissection findings show that Dmc1 is critical for pseudosexual reproduction. While these results provide concrete evidence for meiosis as a part of pseudosexual reproduction, they also suggest the occurrence of a preceding endoreplication event. However, further studies will need to be conducted to validate and confirm endoreplication or alternate mechanisms to achieve the ploidy necessary for a classical meiosis event.
**Discussion**

Hybridogenesis and parthenogenesis are mechanisms that allow some organisms to overcome some hurdles of sexual reproduction and produce hemiclonal or clonal progeny (Avise, 2015; Hörandl, 2009; Lavanchy & Schwander, 2019). However, harmful mutations are not filtered in these processes, making them disadvantageous during evolution and thus restricting the occurrence of these processes to a limited number of animal species (Lavanchy & Schwander, 2019). In this study, we discovered and characterized the occurrence of a phenomenon in fungi that resembles hybridogenesis and termed it pseudosexual reproduction (Figure 6- figure supplement 1). Fungi are known to exhibit asexual, (bi)sexual, unisexual, and parasexual reproduction and can switch between these reproductive modes depending on environmental conditions (Heitman, 2015; Heitman et al., 2013). The discovery of pseudosexual reproduction further diversifies known reproductive modes in fungi, suggesting the presence of sexual parasitism in this kingdom.

Hybridogenesis in animals occurs between two different species. The result of hybridogenesis is the production of gametes that are clones of one of the parents, which then fuse with an opposite-sex gamete of the second species, generating hemiclonal offspring. In our study, we observed a similar phenomenon where only one parent contributes to spores, the counterpart of mammalian gametes. However, we observed this phenomenon occurring between different strains of the same species, *C. neoformans*. It is important to note that these strains vary significantly from each other in terms of genetic divergence and in one case by chromosome rearrangements to the extent that they could be considered different species. This suggests that hybridogenesis in animals and pseudosexual reproduction in fungi are similar to each other.

Hybridogenesis requires the exclusion of one of the parents, which is followed by endoreplication of the other parent's genome and meiosis. The whole-genome sequence of the progeny in our study revealed the complete absence of one parent's genome, suggesting manifestations of genome exclusion during hyphal growth. The mechanism by which the retained parental genome increases its ploidy before meiosis remains to be further investigated in *C. neoformans*. Endoreplication is known to occur in the sister species *C. deneoformans* during unisexual reproduction, and we think that this is the most likely route via which ploidy is increased during pseudosexual reproduction.
The mechanism and time of genome exclusion during hybridogenesis in animals are not entirely understood, except for a few insights from triploid fishes of the genus *Poeciliopsis* and water frogs, *Pelophylax esculentus*. Studies using *Poeciliopsis* fishes showed that haploid paternal genome exclusion takes place during the onset of meiosis via the formation of a unipolar spindle, and thus, only the diploid set of maternal chromosomes is retained (Cimino, 1972a, 1972b). On the other hand, studies involving *P. esculentus* revealed that genome exclusion occurs during mitotic division, before meiosis, which is followed by endoreplication of the other parental genome (Heppich, Tunner, & Greilhuber, 1982; Tunner & Heppich-Tunner, 1991; Tunner & Heppich, 1981). A recent study, however, proposed that genome exclusion in *P. esculentus* could also take place during early meiotic phases (Dolezalkova et al., 2016). Using fluorescence microscopy, we examined the steps of nuclear exclusion in *C. neoformans* and found that it occurs during mitotic hyphal growth and not during meiosis. We also observed that genome exclusion could happen with either of the two parents in *C. neoformans*, similar to what has also been reported for water frogs. However, for most other species, genome exclusion was found to occur with the male genome only, leaving behind the female genome for meiosis (Cimino, 1972a; Holsbeek & Jooris, 2009; Lavanchy & Schwander, 2019; Umphrey, 2006; Uzzell, Günther, & Berger, 1976; Vinogradov, Borkin, Gunther, & Rosanov, 1991). Multiple studies have shown the formation of meiotic synaptonemal complexes during hybridogenesis, clearly establishing the presence of meiosis during this process (Dedukh et al., 2019; Dedukh et al., 2020; Nabais, Pereira, Cunado, & Collares-Pereira, 2012). Our results showed that the meiotic recombinase Dmc1 is required for pseudosexual reproduction, suggesting the presence of meiosis, whereas there is no direct evidence for the role of a meiotic recombinase in hybridogenetic animals. Taken together, these results indicate that the mechanism might be at least partially conserved across distantly related species. Future studies will shed more light on this, and if established, the amenability of *C. neoformans* to genetic manipulation will aid in deciphering some of the unanswered questions related to hybridogenesis in animals.

The occurrence of pseudosexual reproduction might also have significant implications for *C. neoformans* biology. Most (>95%) of *Cryptococcus* natural isolates belong to only one mating type, α (Zhao et al., 2019). While the reason behind this distribution is unknown, one explanation could be the presence of unisexual reproduction in the sister species *C. deneoformans* and *C. gattii* (Fraser et al., 2005; Lin et al., 2005; Phadke, Feretzaki, Clancey,
The presence of pseudosexual reproduction in *C. neoformans* might help explain the mating-type distribution pattern for this species. In this report, one of the *MATa* natural isolates, Bt63a, did not contribute to pseudosexual reproduction and the other isolate, IUM96a, produced uniparental progeny in only one basidium, while the rest of the basidia produced *MATα* progeny. We hypothesize that *MATa* isolates may be defective in this process due to either a variation in their genomes or some other as yet undefined sporulation factor. As a result, pseudosexual reproduction could lead to the generation of predominantly α progeny in nature, reducing the *MATα* population and thus favoring the expansion of the α mating-type population. However, it is still possible that the preferential inheritance of the nuclear genome from one of the two parents is decided by genetic elements located in regions other than *MAT*, and whether the uniparental nuclear inheritance is mating-type specific remains to be elucidated.

Furthermore, the occurrence of pseudosexual reproduction in other pathogenic species such as *C. deneoformans* and non-pathogenic species such as *C. amylolentus* will be investigated in future studies. Attempts to identify the occurrence of pseudosexual reproduction between species where hybrids are known to occur, *C. neoformans* and *C. deneoformans* hybrids, will also be made. These studies will help establish the scope of pseudosexual reproduction in *Cryptococcus* species and could be extended to other basidiomycetes.

We propose that pseudosexual reproduction can occur between any two opposite mating-type strains as long as each of them is capable of undergoing cell-cell fusion and at least one of them can sporulate. We speculate that pseudosexual reproduction might play a key role in *C. neoformans* survival during unfavorable conditions. In conditions where two mating partners are fully compatible, pseudosexual reproduction will be mostly hidden and might not be important (Figure 6, top panel). However, when the two mating partners are partially incompatible or completely incompatible due to high genetic divergence or karyotypic variation, pseudosexual reproduction will be important (Figure 6, left, right, and bottom panels). For example, most of the basidia in H99α and Bt63a cross largely produce aneuploid and/or inviable progeny leading to unsuccessful sexual reproduction. However, a small yet significant proportion of the basidia generate clonal yet viable and fit progeny via pseudosexual reproduction. We hypothesize that these progeny will have a better chance of survival and find a suitable mating partner in the environment whereas, the unfit recombinant progeny might fail to do so. In nature, this might allow a new genotype/karyotype to not only survive but also expand and will prove
advantageous. If a new genotype/karyotype had only the option of undergoing sexual reproduction, it might not survive, restricting the evolution of a new strain. Overall, this mode of pseudosexual reproduction might act as an escape path from genomic incompatibilities between two related isolates and allow them to produce spores for dispersal.

One of the key differences between pseudosexual reproduction and unisexual reproduction observed in the Cryptococcus species complex is the inheritance of mitochondrial DNA. While both unisexual and pseudosexual reproduction result in clonal progeny with respect to the nuclear genome, the mitochondria in pseudosexual reproduction are almost exclusively inherited from the MATa parent (Figure 6- figure supplement 1). This results in the exchange of mitochondrial DNA in the progeny that inherit the MATa nuclear genome, resembling the nuclear-mitochondrial exchange observed during cytoduction in Saccharomyces cerevisiae. During cytoduction, mutants defective in nuclear fusion produce haploid progeny with nuclear genome from one parent, but a mixture of both parents cytoplasm resulting in the inheritance of one parental mitochondrial genome with the other parent’s nuclear genome (Conde & Fink, 1976; Lancashire & Mattoon, 1979; Zakharov & Yarovoy, 1977). This process was used to study mitochondrial genetics with respect to the transfer of drug-resistance genes and other mitochondrial mutations. Similar to cytoduction, pseudosexual reproduction could be employed to study mitochondrial genetics, such as functional analysis of mitochondrial encoded drug resistance, and cytoplasmic inheritance of factors such as prions in C. neoformans.

The fungal kingdom is one of the more diverse kingdoms with approximately 3 million species (Sun, Hoy, & Heitman, 2020). The finding of hybridogenesis-like pseudosexual reproduction hints towards unexplored biology in this kingdom that might provide crucial clues for understanding the evolution of sex. Fungi have also been the basis of studies focused on understanding the evolution of meiosis, and the presence of genome reduction, as well as the parasexual cycle in fungi, have led to the proposal that meiosis evolved from mitosis (Hurst & Nurse, 1991; Wilkins & Holliday, 2009). Pseudosexual reproduction may be a part of an evolutionary process wherein genome exclusion followed by endoreplication and meiosis was an ancestral form of reproduction that preceded the evolution of sexual reproduction. Evidence supporting such a hypothesis can be observed in organisms undergoing facultative sex or facultative parthenogenesis (Booth et al., 2012; Fields, Feldheim, Poulakis, & Chapman, 2015; Hodač, Klatt, Hojsgaard, Sharbel, & Hörandl, 2019; Hojsgaard & Harandl, 2015). The presence
of these organisms also suggests that a combination of both sexual and clonal modes of
reproduction might prove to be evolutionarily advantageous.


Materials and Methods

Strains and media

C. neoformans wild-type strains H99α and KN99α served as the wild-type isogenic parental lineages for the experiments (Nielsen et al., 2003; Perfect, Ketabchi, Cox, Ingram, & Beiser, 1993), in addition to MATα strains Bt63α and IUM96-2828α (Keller et al., 2003; Litvintseva et al., 2003). Strains were grown in YPD media for all experiments at 30°C unless stated otherwise. G418 and/or NAT were added at a final concentration of 200 and 100 µg/ml, respectively, for the selection of transformants. MS media was used for all the mating assays, which were performed as described previously (Sun, Priest, et al., 2019). Basidia-specific spore dissections were performed after two-five weeks of mating, and the spore germination frequency was scored after five days of dissection. All strains and primers used in this study are listed in Supplementary File 1d and Supplementary File 1e, respectively.

Genotyping for mating-type locus and mitochondria

Mating-type (MAT) and mitochondrial genotyping for all the progeny were conducted using PCR assays. Genomic DNA was prepared using the MasterPure™ Yeast DNA purification kit from Lucigen. To determine the MAT, the STE20 allele present within the MAT locus was detected because it differs in length between the two different mating types. Primers specific to both MATα and MATα (JOHE50979-50982 in Supplementary File 1e) were mixed in the same PCR mix, and the identification was made based on the length of the amplicon (Figure 1E-G). For the mitochondrial genotyping, the COX1 allele present in the mitochondrial DNA was probed to distinguish between H99α/VYD135α and KN99α/IUM96α. For the differentiation between Bt63α and H99α/VYD135α, the COB1 allele was used because COX1 in Bt63α is identical to H99α/VYD135α. The difference for both COX1 and COB1 is the presence or absence of an intron and results in significantly different size products between MATα and MATα parents (Figure 1 and Figure 3-figure supplement 1). The primers used for these assays (JOHE51004-51007) are mentioned in Supplementary File 1e.

Genomic DNA isolation for sequencing

Genomic DNA for whole-genome sequencing was prepared using the CTAB-based lysis method, as described previously (Yadav et al., 2020). Briefly, 50 ml of an overnight culture was pelleted, frozen at -80°C, and subjected to lyophilization. The lyophilized cell pellet was broken into a fine powder, mixed with lysis buffer, and the mix was incubated at 65°C for an hour with
intermittent shaking. The mix was then cooled on ice, and the supernatant was transferred into a fresh tube, and an equal volume of chloroform (~15 ml) was added and mixed. The mix was centrifuged at 3200 rpm for 10 min, and the supernatant was transferred to a fresh tube. An equal volume of isopropanol (~18 to 20 ml) was added into the supernatant and mixed gently. This mix was incubated at -20°C for an hour and centrifuged at 3200 rpm for 10 min. The supernatant was discarded, and the DNA pellet was washed with 70% ethanol. The pellet was air-dried and dissolved in 1ml of RNase containing 1X TE buffer and incubated at 37°C for 45 min. The DNA was again chloroform purified and precipitated using isopropanol, followed by ethanol washing, air drying, and finally dissolved in 200 µl 1X TE buffer. The DNA quality was estimated with NanoDrop, whereas DNA quantity was estimated with Qubit.

**Whole-genome Illumina sequencing, ploidy, and SNP analysis**

Illumina sequencing of the strains was performed at the Duke sequencing facility core (https://genome.duke.edu/), using Novaseq 6000 as 150 paired-end sequencing. The Illumina reads, thus obtained, were mapped to the respective genome assembly (H99, VYD135, Bt63, or IUM96) using Geneious (RRID:SCR_010519) default mapper to estimate ploidy. The resulting BAM file was converted to a .tdf file, which was then visualized through IGV to estimate the ploidy based on read coverage for each chromosome.

For SNP calling and score for recombination in the progeny, Illumina sequencing data for each progeny was mapped to parental strain genome assemblies individually using the Geneious default mapper with three iterations. The mapped BAM files were used to perform variant calling using Geneious with 0.8 variant frequency parameter and at least 90x coverage for each variant. The variants thus called were exported as VCF files and imported into IGV for visualization purposes. H99, Bt63, IUM96-2828, and VYD135 Illumina reads were used as controls for SNP calling analysis.

**PacBio/Nanopore genome assembly and synteny comparison**

To obtain high-molecular-weight DNA for Bt63 genome PacBio and IUM96-2828 genome Nanopore sequencing, DNA was prepared as described above. The size estimation of DNA was carried out by electrophoresis of DNA samples using PFGE. For this purpose, the PFGE was carried out at 6V/cm at a switching frequency of 1 to 6 sec for 16 h at 14°C. Samples with most of the DNA ≥100 kb or larger were selected for sequencing. For PacBio sequencing, the DNA sample was submitted to the Duke sequencing facility core. Nanopore sequencing was
performed in our lab using a MinION device on an R9.4.1 flow cell. After sequencing, reads were assembled to obtain a Bt63 genome assembly via Canu (RRID:SCR_015880) using PacBio reads > 2 kb followed by five rounds of pilon polishing (RRID:SCR_014731). For IUM96-2828, one round of nanopolish was also performed before pilon polishing. Once completed, the chromosomes were numbered based on their synteny with the H99 genome. For chromosomes involved in translocation (Chr 3 and Chr 11), the chromosome numbering was defined by the presence of the respective syntenic centromere from H99. Centromere locations were mapped based on BLASTn analysis with H99 centromere flanking genes.

Synteny comparisons between the genomes were performed with SyMAP v4.2 using default parameters (Soderlund, Bomhoff, & Nelson, 2011) (http://www.agcol.arizona.edu/software/symap/). The comparison block maps were exported as .svg files and were then processed using Adobe® Illustrator® (RRID:SCR_010279) and Adobe® Photoshop® (RRID:SCR_014199) for representation purposes. The H99 genome was used as the reference for comparison purposes for plotting VYD135, Bt63, and IUM96-2828 genomes. The centromere and telomere locations were manually added during the figure processing.

**Fluorescent tagging and microscopy**

GFP and mCherry tagging of histone H4 were performed by integrating respective constructs at the safe haven locus (Arras, Chitty, Blake, Schulz, & Fraser, 2015). GFP-H4 tagging was done using the previously described construct, pVY3 (Yadav & Sanyal, 2018). For mCherry-H4 tagging, the GFP-containing fragment in pVY3 was excised using SacI and BamHI and was replaced with mCherry sequence PCR amplified from the plasmid pLKB25 (Kozubowski & Heitman, 2010). The constructs were then linearized using XmnI and transformed into desired strains using CRISPR transformation, as described previously (Fan & Lin, 2018). The transformants were screened by PCR, and correct integrants were obtained and verified using fluorescent microscopy.

To observe the fluorescence signals in the hyphae and basidia, a 2-3 week old mating patch was cut out of the plate and directly inverted onto a coverslip in a glass-bottom dish. The dish was then used to observe filaments under a DeltaVision microscope available at the Duke University Light Microscopy Core Facility (https://microscopy.duke.edu/dv). The images were captured at 60X magnification with 2x2 bin size and z-sections of either 1 or 0.4 µm each. GFP and mCherry signals were captured using the GFP and mCherry filters in the Live-Cell filter set.
The images were processed using Fiji-ImageJ (https://imagej.net/Fiji) (RRID:SCR_002285) and exported as tiff files as individual maximum projected images. The final figure was then assembled using Adobe® Photoshop® software for quality purposes.

**Sporulation frequency counting**

To visualize hyphal growth and sporulation defects during mating assays, the mating plates were directly observed under a Nikon Eclipse E400 microscope. Hyphal growth and basidia images were captured using the top-mounted Nikon DXM1200F camera on the microscope. The images were processed using Fiji-ImageJ and assembled in Adobe® Photoshop® software.

For crosses involving wild-type H99α, VYD135α, KN99a, Bt63a, IUM96a, approximately 1000 total basidia were counted after 4 weeks of mating, and the sporulation frequency was calculated. For crosses involving VYD135 dmc1Δ strain, three mating spots were setup independently. From each mating spot periphery, 6 images were captured after 3-4 weeks of mating. Basidia (both sporulating and non-sporulating) in each of these spots were counted manually after some processing of images using ImageJ. The sporulation frequency was determined by dividing the sporulating basidia by the total number of basidia for each spot. Each mating spot was considered as an independent experiment and at least 3000 basidia were counted from each mating spot.

**Flow cytometry**

Flow cytometry analysis was performed as described previously (Fu & Heitman, 2017). Cells were grown on YPD medium for two days at 30°C, harvested, and washed with 1X PBS buffer followed by fixation in 70% ethanol at 4°C overnight. Next, cells were washed once with 1 ml of NS buffer (10 mM Tris-HCl, pH = 7.2, 250 mM sucrose, 1 mM EDTA, pH = 8.0, 1 mM MgCl2, 0.1 mM CaCl2, 0.1 mM ZnCl2, 0.4 mM phenylmethylsulfonyl fluoride, and 7 mM β-mercaptoethanol), and finally resuspended in 180 μl NS buffer containing 20 μl 10 mg/ml RNase and 5 μl 0.5 mg/ml propidium iodide (PI) at 37°C for 3-4 hours. Then, 50 μl stained cells were diluted in 2 ml of 50 mM Tris-HCl, pH = 8.0, transferred to FACS compatible tube, and submitted for analysis at the Duke Cancer Institute Flow Cytometry Shared Resource. For each sample, 10,000 cells were analyzed on the FL1 channel on the Becton-Dickinson FACScan. Wild-type H99 and previously generated AI187 were used as haploid and diploid controls,
respectively, in these experiments. Data analysis was performed using the FlowJo software (RRID:SCR_008520).

**Data Availability**

The sequence data generated in this study were submitted to NCBI with the BioProject accession number PRJNA682203.

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**Competing interests**

The authors declare no competing interests.

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Table 1. Genotype analysis of basidia-specific spores germinated from H99α x KN99a and VYD135α x KN99a crosses.

| Basidia # | H99α x KN99a cross | VYD135α x KN99a cross |
|-----------|---------------------|------------------------|
|           | Spores germinated/ dissected | % germinated | MAT | Mito | Spores germinated/ dissected | % germinated | MAT | Mito |
| 1         | 5/14                | 36          | 4α + 1a | a    | 12/24               | 50          | All α | a    |
| 2         | 14/14               | 100         | 7α + 7a | a    | 6/10                | 60          | All α | a    |
| 3         | 12/14               | 86          | 2α + 7a | a    | 15/15               | 100         | All a  | a    |
|           | 4                   | 10/14       | 4α +6a  | a    | 22/27               | 81          | All a  | a    |
| 5         | 7/13                | 54          | 6α + 1a/α | a | 3/12               | 25          | All α  | a    |
| 6         | 13/14               | 93          | 6α + 7a | a    | 25/27               | 93          | All α  | a    |
| 7         | 11/14               | 79          | 6α + 5a | a    | 4/4                 | 100         | All α  | a    |
| 8         | 14/14               | 100         | 12α + 2a | a | 10/13               | 77          | All α  | a    |
| 9         | 10/14               | 71          | 4α +6a  | a    | 13/15               | 87          | All α  | a    |
| 10        | 14/14               | 100         | 7α + 7a | a    | 31/61               | 51          | All α  | a    |
| 11        | 14/14               | 100         | 10α + 4a | a | 10/10               | 100         | All a  | a    |
| 12        | 12/14               | 86          | 8α + 4a | a    | 4/5                 | 80          | All a  | a    |
| 13        | 4/11                | 36          | All a   | a    | 24/28               | 86          | All a  | a    |
| 14        | 13/13               | 100         | 8α + 5a | a    | 16/28               | 57          | All a  | a    |
| 15        | 14/14               | 100         | 7α + 7a | a    | 11/11               | 100         | All a  | a    |
| 16        | 14/14               | 100         | 6α + 8a | a    | 10/22               | 45          | All α  | a    |

Mito refers to Mitochondria.
Figures and Figure Legends

**Figure 1. Chromosome shuffled strain exhibits unusual sexual reproduction.** (A-B) Images of cultures for the individual strains H99α, KN99a, and VYD135α, showing no self-filamentation on mating medium. Magnification=10X. (C-D) Light microscopy images showing robust sporulation in the H99α x KN99a cross, whereas the VYD135α x KN99a cross exhibited robust hyphal development but infrequent sporulation events. The inset images in colored boxes show examples of basidia observed in each of the crosses. Bar, 100 µm. (E-F) A scheme showing the MATα (H99α and VYD135α) and MATa (KN99a) alleles at the STE20 (E) and COX1 (F) loci. Primers used for PCR analysis are marked by blue triangles. (G) Gel images showing PCR amplification of STE20 and COX1 alleles in the progeny obtained from four different basidia for both H99α x KN99a and VYD135α x KN99a crosses. PCR analysis for the parental strains is also shown, and key bands for DNA marker are labeled.

**Figure 2. Fluorescence microscopy reveals uniparental nuclear inheritance in the wild-type crosses.** (A) Crosses of GFP-H4 tagged H99α and mCherry-H4 tagged KN99a revealed the presence of both fluorescent markers in most spore chains along with uniparental nuclear inheritance in rare cases (~1%). In these few sporulating basidia, only one of the fluorescent signals was observed in the spore chains, reflecting the presence of only one parental nucleus in these basidia. (B) Crosses involving GFP-H4 tagged VYD135α and mCherry-H4 tagged KN99a revealed the presence of spore chains with only one fluorescent color. In the majority of basidia that have both parental nuclei, marked by both GFP and mCherry signals, spore chains are not produced, consistent with a failure of meiosis in these basidia. Bars, 10 µm.

**Figure 3. VYD135α progeny exhibit strict uniparental nuclear inheritance and lack the signature of meiotic recombination.** (A) Chromosome maps for H99α/KN99a, VYD135α, Bt63a, and IUM96a showing the karyotype variation. The genome of the wild-type strain H99α served as the reference. Black arrowheads represent chromosome translocations between VYD135α and H99α whereas red arrowheads mark chromosomes with a translocation between H99α and Bt63a or IUM96a. (B) Whole-genome sequencing, followed by SNP identification, of H99α x Bt63a progeny revealed evidence of meiotic recombination in all of the progeny. The
left panel shows SNPs with respect to the Bt63a genome whereas the right panel depicts SNPs against the H99α genome. H99α and Bt63a Illumina sequencing data served as controls for SNP calling. (C) SNP analysis of VYD135α x Bt63a progeny revealed no contribution of the Bt63a parental genome in the progeny as evidenced by the presence of SNPs only against Bt63a (left panel) but not against the VYD135α genome (right panel). The presence of a few SNPs observed in VYD135α, as well as all VYD135α x Bt63a progeny, are within nucleotide repeat regions. GF stands for germination frequency and P stands for progeny. (D) SNP analysis of H99α x Bt63a and VYD135α x Bt63a progeny using mitochondrial DNA as the reference revealed that mitochondrial DNA is inherited from Bt63a in all of the progeny. Progeny obtained from VYD135α x Bt63a basidium 18 also revealed recombination between the two parental mitochondrial genomes as marked by the absence or presence of two SNPs when mapped against VYD135α and Bt63 mitochondrial genomes, respectively. The green bar in each panel depicts the locus used for PCR analysis of the mitochondrial genotype in the progeny.

**Figure 4.** Pan-hyphal microscopy reveals the loss of one parental nucleus during pseudosexual reproduction. Spore-producing long hyphae were visualized in both (A) wild-type H99α x KN99a and (B) VYD135α x KN99a crosses to study the dynamics of nuclei in hyphae. Both nuclei were present across the hyphal length in the wild-type and resulted in the production of recombinant spores. On the other hand, one of the nuclei was lost during hyphal branching in the VYD135α x KN99a cross and resulted in uniparental nuclear inheritance in the spores that were produced. The arrow in B marks the hyphal branching point after which only one of the parental nuclei is present (also see Figure 4-figure supplement 1A). The images were captured as independent sections and assembled to obtain the final presented image. Bars, 10 μm.

**Figure 5.** Meiotic recombinase Dmc1 is required for pseudosexual reproduction. (A) Light microscopy images showing the impact of dmc1 mutation on sexual and pseudosexual reproduction in C. neoformans. Bar, 100 μm. (B) A graph showing quantification (n=3) of sporulation events in multiple crosses with dmc1Δ mutants. At least 3000 basidia were counted in each experiment.
Figure 6. Model for the role of pseudosexual reproduction in *C. neoformans* ecology.
Scenarios showing possible roles for pseudosexual reproduction under various hypothetical mating conditions. Except for one condition where the two parents are completely compatible with each other, pseudosexual reproduction could play a significant role in survival and dissemination despite its occurrence at a low frequency.
**Supplementary Figure Legends**

**Figure 2-figure supplement 1. Dynamics of sexual reproduction and sporulation analyzed with C. neoforms strains expressing nuclear-localized fluorescent reporter proteins. (A)** A cartoon depicting various stages of sexual reproduction in *C. neoforms*, from the formation of conjugation tube to sporulation, and possible dynamics of the nuclei at these different stages. After cell-cell fusion, tagged proteins assort into both nuclei and yield a yellow/orange fluorescence color as a result of the mixing of the green and red signals. Cartoons in the box show hypothetical scenarios where uniparental nuclear inheritance occurs after the loss of one parental nucleus. **(B)** Direct fluorescence microscopy images showing the status of GFP-H4 and mCherry-H4 tagged nuclei in post-mating hyphae as well as in spores. Both GFP and mCherry fluorescent colors were observed in hyphae and spores as hypothesized in A. Bar, 10 µm.

**Figure 2-figure supplement 2. Nuclear dynamics during sporulation in the wild-type and VYD135α crosses.** GFP-H4 and mCherry-H4 tagging revealed different localization patterns in the (A) wild-type H99α x KN99α and (B) VYD135α x KN99α crosses. Wild-type spore chains mostly harbored both the nuclear stains as a result of bisexual meiosis. On the other hand, basidia with only one of the parental nuclei produced spores in VYD135α x KN99α crosses; basidia with both nuclei failed to produce spore chains and, as a result, remained as bald basidia. Bars, 10 µm.

**Figure 3-figure supplement 1. Pseudosexual reproduction occurs in natural isolates, Bt63a and IUM96a. (A)** Images of the mating spots showing filamentation when two strains of opposite mating-type are crossed. No filamentation is observed without the presence of a mating partner. **(B and D)** Light microscopy images showing sporulation frequency in crosses involving Bt63a (B) and IUM96a (D). Bars, 100 µm. **(C and E)** Schemes depicting the *STE20* alleles used for *MAT* locus and *COB1* (for Bt63a) and *COX1* (for IUM96a) alleles for mitochondrial genotyping, respectively. Gel images show the PCR analysis on progeny from four basidia and the parental strains for all crosses as mentioned.
**Figure 3-figure supplement 2.** Bt63a fluorescence microscopy revealed pseudosexual reproduction events. GFP-H4 tagged VYD135α crossed with mCherry-H4 tagged Bt63a showed only VYD135α sporulation events as also observed in spore dissection analysis. Bars, 10 μm.

**Figure 3-figure supplement 3.** VYD135α x Bt63a progeny lack signatures of meiotic recombination. SNP analysis on VYD135α x Bt63a progeny revealed no contribution of the Bt63a parental genome in the progeny as evidenced by the presence of SNPs only against Bt63a (left panel) but not against VYD135α genome (right panel). The few SNPs observed in VYD135 as well as all VYD135α x Bt63a progeny lie within nucleotide repeat regions. GF stands for germination frequency and P stands for progeny.

**Figure 3-figure supplement 4.** Mitochondria are inherited from MATα parent in all of the progeny. (A) A map of SNPs detected in H99α x Bt63a progeny when using H99α mitochondrial DNA (upper panel) and Bt63a mitochondrial DNA (lower panel) as the reference. (B) SNP analysis revealed variants in all the progeny when using VYD135α mitochondrial DNA as the reference but not when using Bt63a mitochondrial DNA. The two SNPs detected against Bt63a DNA in progeny P19-24 (Basidium 18) suggest recombination of two parental mitochondrial DNA during mating. The green bar in each panel depicts the fragment used for PCR analysis in Figure 3-figure supplement 1. P stands for progeny.

**Figure 3-figure supplement 5.** VYD135α x Bt63a progeny are haploid. (A) Whole-genome sequencing of the H99α x Bt63a progeny revealed extensive aneuploidy in the progeny. Each progeny seemed to exhibit aneuploidy for at least one chromosome. (B) Whole-genome sequencing data revealed that the progeny obtained from VYD135α x Bt63a 5-week old crosses are euploid in nature as they show a uniform level of genomic content when mapped to the Bt63 genome. VYD135α and Bt63a whole-genome sequencing data were also mapped as controls. Each lane represents one strain, and the difference in intensity correlates with the number of reads obtained per sample. (C) Flow-cytometry analysis on progeny obtained from three basidia confirmed that all the germinating progeny are haploid. While progeny from B12 and B14 are
the same as used for the whole-genome sequencing, progeny from B3 were subjected to only
flow cytometry analysis. Bt63a and VYD135α were also analyzed as controls for this
experiment. P stands for progeny.

**Figure 3-figure supplement 6. IUM96a exhibits meiotic recombination in progeny with
H99α but not with the genome shuffle strain VYD135α.** (A) The left panel depicts SNPs with
respect to the IUM96a genome whereas the right panel shows SNPs against the H99α genome.
Whole-genome sequencing, followed by SNP analysis, for the H99α x IUM96a progeny (basidia
3 and 4) revealed evidence of meiotic recombination in the progeny. Basidium 7 from the H99α
x IUM96a cross produced uniparental progeny, which was confirmed by SNP analysis on a
subset of these progeny. The progeny exhibited SNPs only against the IUM96a genome but not
against the H99α genome. (B) SNP analysis from two different basidia revealed inheritance of
only one set of parental nuclear DNA in the progeny from VYD135α x IUM96a cross. Basidium
3 progeny possessed DNA from only the VYD135α parent, while basidium 5 progeny inherited
nuclear DNA from IUM96a alone. The results obtained from this analysis are congruent with
mating-type PCR results shown in Supplementary file 1b. GF stands for germination frequency
and P stands for progeny.

**Figure 3-figure supplement 7. Ploidy analysis of IUM96a progeny reveals haploid
uniparental progeny.** Whole-genome sequencing analysis revealed the presence of multiple
aneuploidies in the (A) H99α x IUM96a progeny, but a completely euploid genome for the (B)
VYD135α x IUM96a progeny. P stands for progeny.

**Figure 4-figure supplement 1. Hyphal branches act as a gateway for nuclear separation
during pseudosexual reproduction.** (A) Individual z-sections showing the hyphal branching
(marked by arrow) where the two parental nuclei segregate in the figure 4B. (B) Images showing
hyphal branching points where one of the parental nuclei separates from the main hyphae with
two parental nuclei (Top two panels). The branch point is marked with the arrow. The lower two
panels show the long hyphae with only one of the parental nuclei in them. The third panel shows
other hyphae with both parental nuclei suggesting that separation occurred at an early stage. The
fourth panel exhibits the same between VYD135α x Bt63a but also has a sporulating basidium on it. Bar, 10 µm.

Figure 5-figure supplement 1. Dmc1 deletion leads to severe sporulation defects in both sexual and pseudosexual reproduction. Light microscopy images showing the phenotype of DMC1 deletion in (A) H99α x KN99a unilateral crosses as well as bilateral mutant crosses and (B) VYD135α x KN99a dmc1Δ unilateral and bilateral crosses. The deletion of DMC1 led to a reduction in sporulating basidia in bilateral mutant crosses.

Figure 5-figure supplement 2. Meiotic regulator Dmc1 is required for pseudosexual reproduction. A cross between a GFP-H4 tagged VYD135α strain and an mCherry-H4 tagged KN99a dmc1Δ mutant revealed that Dmc1 is required for pseudosexual reproduction events. The majority of the KN99a dmc1Δ nucleus-containing basidia failed to produce spore chains (top two rows and bottom rows). While all 11 observed basidia with VYD135α nuclei produced spores, only 2 out of 19 observed basidia with KN99a dmc1Δ nuclei produced spores. One of these two is represented in the third row. Bars, 10 µm.

Figure 6-figure supplement 1. Unisexual, bisexual and pseudosexual reproduction in C. neoformans. A diagram depicting various types of sexual reproduction in Cryptococcus species. C. deneoformans exhibits unisexual reproduction in which two cells of the same mating-type fuse or a single cell undergoes endoreplication followed by the production of clonal progeny. Both C. neoformans and C. deneoformans show bisexual reproduction in which two cells of opposite mating-types fuse with each other and produce recombinant progeny. Pseudosexual reproduction, as proposed in this study, arises from bisexual mating but generates clonal progeny of one of the parents after the other parental nucleus is lost during development. While both unisexual and pseudosexual reproduction produce clonal progeny, they differ with respect to the inheritance of mitochondrial DNA (marked by grey color cell background in the illustration).
Supplementary File 1a. The genotype of basidia-specific spores dissected from H99α x Bt63a and VYD135α x Bt63a crosses.

Supplementary File 1b. The genotype of basidia-specific spores dissected from H99α x IUM96-2828a and VYD135α x IUM96-2828a crosses.

Supplementary File 1c. Genotype analysis of basidia-specific progeny from H99α dmc1Δ x KN99a dmc1Δ and VYD135α dmc1Δ x KN99a dmc1Δ crosses.

Supplementary File 1d. Strains used in this study.

Supplementary File 1e. Primers used in this study.
A

H99α x KN99α

H99α dmc1Δ x KN99α dmc1Δ

VYD135α x KN99α

VYD135α dmc1Δ x KN99α dmc1Δ

~88% sporulation
~2.7% sporulation
~1% sporulation
~0.1% sporulation

B

Sporulation frequency (%)

VYD135α x KN99α crosses
Successful bisexual reproduction (~99%)
No SNPs detected in the H99 x Bt63 progeny when using Bt63 mitochondrial genome as reference.

No SNPs detected in rest of the VYD135 x Bt63 progeny when using Bt63 mitochondrial genome as reference.
A

IUM96a genome

H99a genome

Basidium 7 Basidium 4 Basidium 3
(GF=64%)

Basidium 7 Basidium 4 Basidium 3
(GF=64%)

Basidium 5
(GF=99%)

P1

P2

P3

P4

P5

P6

P7

P8

P9

P10

P11

P12

P13

P14

SNPs representing H99a DNA

SNPs representing IUM96a DNA

SNPs representing VYD135a DNA

B

IUM96a genome

VYD135a genome

Basidium 3
(GF=79%)

Basidium 5
(GF=99%)

P1

P2

P3

P4

P5

P6

P7

P8

P9

P10

P11

P12

P13

P14
Endoreplication

Nuclear fusion

Filamentation

MATα

Yeast cell

Conjugation

Cell-cell fusion

Filamentation

Maintain dikaryon

Loss of one parental nucleus

Nuclear fusion

Endoreplication?

Bisexual reproduction

Meiosis

Sporulation

Unisexual reproduction

Basidium formation

Meiosis

Sporulation

MATTα (marked by grey background) from MATα parent

Pseudosexual reproduction