Ploidy variation in multinucleate cells changes under stress

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ABSTRACT Ploidy variation is found in contexts as diverse as solid tumors, drug resistance in fungal infection, and normal development. Altering chromosome or genome copy number supports adaptation to fluctuating environments but is also associated with fitness defects attributed to protein imbalances. Both aneuploidy and polyploidy can arise from multinucleate states after failed cytokinesis or cell fusion. The consequences of ploidy variation in syncytia are difficult to predict because protein imbalances are theoretically buffered by a common cytoplasm. We examined ploidy in a naturally multinucleate fungus, *Ashbya gossypii*. Using integrated lac operator arrays, we found that chromosome number varies substantially among nuclei sharing a common cytoplasm. Populations of nuclei range from 1N to >4N, with different polyploidies in the same cell and low levels of aneuploidy. The degree of ploidy variation increases as cells age. In response to cellular stress, polyploid nuclei diminish and haploid nuclei predominate. These data suggest that mixed ploidy is tolerated in these syncytia; however, there may be costs associated with variation as stress homogenizes the genome content of nuclei. Furthermore, the results suggest that sharing of gene products is limited, and thus there is incomplete buffering of ploidy variation despite a common cytosol.

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INTRODUCTION

Variation in ploidy within an organism can be a defining feature of either pathologies or normal developmental programs. Understanding both the utility and the deleterious consequences of varying DNA copy number relates to problems in fields as diverse as cancer biology, microbial pathogenesis and ecology, and plant development. Copy-number variation can be considered from the scale of small insertions or deletions that affect a single gene to amplifications of whole chromosomes (aneuploidy) or the entire genome (polyploidy). DNA copy-number changes can lead to changes in the levels of the mRNA and proteins encoded in the amplified regions (Torres *et al.*, 2007; Pavelka *et al.*, 2010). Expression of altered copy-number regions thus has the potential to affect dramatically the physiology of cells.

The consequences of aneuploidy on cell function are highly variable and context dependent. The majority of human solid tumors display aneuploid karyotypes and are also composed of rapidly proliferating cells (Weaver and Cleveland, 2006). Robust tumor growth is paradoxical in light of certain model aneuploid cells, which can be observed to grow slowly (Torres et al., 2007). The mechanisms by which tumors tolerate aneuploidy are still under study. However, one known function of key tumor suppressors is to monitor and arrest growth when cells display chromosome instability, a context in which aneuploidies arise (Li et al., 2010; Thompson and Compton, 2010). These programs are dismantled with inactivation of the tumor suppressors coincident with tumor progression. Work in model aneuploid yeast and fibroblast cells with engineered extra chromosomes also indicates that one major cause of growth defects is proteotoxic stress. This stress emerges from the attempt to degrade excess proteins, and presumably tumor cells adapt to this stress (Torres et al., 2007; Pavelka et al., 2010; Oromendia et al., 2012).

As in tumors, aneuploidy is also commonly observed in diverse fungal species in natural, clinical, and laboratory environments

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(Morrow and Fraser, 2013; Bennett et al., 2014). For example, in the fungal pathogen Candida albicans, aneuploid cells readily emerge through transient tetraploidization and in response to antifungals (Forche et al., 2008; Selmecki et al., 2010; Harrison et al., 2014). This capacity to remodel the genome is beneficial to the pathogen during infection and enables the development of drug resistance. Variation in ploidy is also seen during the course of sexual development and infection with the human pathogen Cryptococcus neoformans (Idnurm, 2010; Semighini et al., 2011). In addition, under some stress conditions, aneuploid Saccharomyces cerevisiae cells are more fit than euploid strains, indicating that ploidy variation may be highly adaptive, depending on the environmental conditions (Pavelka et al., 2010; Yona et al., 2012; Zhu et al., 2012). Furthermore, industrial Saccharomyces yeast strains developed for brewing and baking are largely polyploid and/or aneuploid (Querol and Bond, 2009). Thus the capacity to induce and tolerate large genome changes can provide an adaptive advantage for diverse fungi in the context of pathogenesis and environmental stress.

Aneuploidy can arise through multiple, distinct molecular mechanisms. Current known routes to aneuploidy include failure in the spindle assembly checkpoint (SAC), altered error correction of misattached chromosomes, and transient polyploidization through cytokinesis failure or cell fusion (Burds et al., 2005; Lu and Kang, 2009; Krajcovic et al., 2011; Bakhoum and Compton, 2012). There is evidence that polyploidy can increase the probability of many cells, including those of budding yeast, of becoming aneuploid (Storchova et al., 2006). Similarly, aneuploid liver cells can arise from a polyploid starting population (Duncan et al., 2010). Whereas polyploidy has long been appreciated for functional relevance in plants and specific animal tissues, recent work in fungi has started to focus attention on polyploidy as an engine of evolution (Albertin and Marullo, 2012). Thus polyploidy could serve as a reservoir to generate functional aneuploidies under specific stresses or be useful unto itself for specific metabolic needs in a given tissue or environment.

Most filamentous fungi exist in syncytial states, in which many nuclei share a common cytoplasm during mycelial growth. Furthermore, it is clear that in many fungi, different genomes can coexist, cooperating and/or competing in a single cell (Roper et al., 2011). However, it is less clear how genomes that differ due to chromosome imbalances may function in fungal syncytia. Syncytia theoretically may be a highly permissive environment for aneuploidy, as imbalances may be complemented across multiple nuclei in the same cytoplasm. This would require that gene products are well mixed so that the cytosol buffers protein imbalances. In this case, syncytial cells may harbor high frequencies of aneuploidy and suffer fewer fitness consequences than uninucleate cells with a chromosome imbalance. This idea is supported by the multinucleate states seen in tumors that are also aneuploid (Lu and Kang, 2009). Alternatively, if the cytosol is compartmentalized in large multinucleate cells such that gene products are not shared between nuclei, then ploidy may be more tightly controlled due to functionally heterogeneous cytosol.

We monitored ploidy in the multinucleate fungus Ashbya gossypii to examine the variability of chromosome content of individual nuclei within a single cell. Previous work in Ashbya suggested that the euploid state of the system is haploid in multinucleate mycelia and uninucleate haploid spores are produced by asexual sporulation (Dietrich et al., 2004). In Ashbya, nuclei have highly variable division cycle durations, so that nuclei divide out of sync with their neighbors (Gladfelter et al., 2006). One possible source of timing variation between nuclei in a single cell is variable DNA content. In this study, we tested the hypotheses that there is ploidy variation between nuclei in the same cell and that aneuploidy is tolerated in syncytia.

RESULTS

DNA content varies among nuclei in a single cell

Ashbya cells expressing green fluorescent protein-labeled histone-4 (Hhf1-GFP) were filmed, and the fluorescence intensity of the histone signal in single nuclei was measured at birth (pre-DNA replication) and moments before division (post-DNA replication). If all nuclei are haploid, we would predict two discrete peaks of histone intensities corresponding to 1N ploidy at birth and 2N ploidy before mitosis (Figure 1A). Surprisingly, nuclei showed a wide range of intensities at birth and immediately preceding mitosis (Figure 1B, N = 47 nuclei). The two distributions of intensities show substantial overlap, although the mean intensity at mitosis is modestly but significantly higher than at birth (birth mean, $12,720 \pm 3970$ a.u.; mitosis mean, $15,030 \pm 5623$ a.u.; two-sample t test, p <0.03). The signal variation is not due to differential photobleaching, as the Hhf1-GFP intensity of nuclei is not correlated with the time point in the image acquisition (Supplemental Figure S1). This suggests that nuclei are born and divide with variable amounts of DNA in the same cell.

To ensure that histone intensities increase with cell cycle progression and DNA replication, we plotted the histone intensity for individual nuclei and found that for all nuclei the signal increases with progression toward mitosis. The time from minimum to maximum signal intensity ranged from 30 to 134 min (Figure 1C; mean, 83 min, SD = 21 min, N = 47), with an average 1.5-fold change in the histone intensity (Figure 1D). This change may not be twofold due to a soluble pool of histones that is present in the nucleus at birth and used to build nucleosomes during replication or because not all the DNA is replicated. These data suggest that the duration of S phase is likely highly variable, and this could be due to variable DNA content, heterogeneous import of Hhf1-GFP, or other factors controlling DNA replication being limiting. Therefore, as a second measure of DNA content through the cell cycle, we measured the incorporation of the DNA dye 4',6-diamidino-2-phenylindole (DAPI) into nuclei that were scored for cell cycle stage based on the state of the spindle pole body (SPB). As seen in live cells, nuclei in the same SPB state, and presumably the same cell cycle stage, show a broad distribution of DAPI intensities, supporting the idea that there may be substantial variation in the DNA content among nuclei in the same cell (Figure 1E).

Nuclei vary in the number of copies of individual chromosomes

Given the heterogeneity of histone and DAPI signals, we next investigated the numbers of individual chromosomes by integrating 32 lac operator repeats into intergenic regions of either chromosome I (smallest in size) or chromosome VI (largest in size) in a strain also expressing GFP-LacI-NLS (Figure 2, A and B). Nuclei had from zero to greater than four spots of GFP-Lacl signals for each of the chromosomes; the largest number of spots detected was six (Figure 2, A-C; >300 nuclei for each chromosome, cells scored at a stage of >100 nuclei). At very low frequency, nuclei could be seen without a Lacl signal, suggesting rare loss of the chromosome. Work using similarly marked chromosomes in S. cerevisiae showed that Lacl signals on sister chromosomes are not resolvable until anaphase as long as the repeat sequences are far from centromeres (Straight et al., 1997; Pearson et al., 2001). This suggests that the Ashbya nuclei with ≥2 spots in fact have extra chromosomes rather than having finished S phase. Nevertheless, we assessed the SPB



FIGURE 1: DNA content varies among Ashbya nuclei. (A) Schematic of predicted relative Hhf1-GFP intensity changes before replication and after replication. (B) Hhf1-GFP intensity at birth and mitosis. Background-corrected GFP intensity was measured by thresholding around the central nuclear coordinate, which was manually recorded immediately before and after mitosis (47 nuclei; Anderson *et al.*, 2013). (C) Background-corrected Hhf1-GFP intensity summed for each nucleus over time. Each line represents one nucleus (47 nuclei). (D) Fold change in Hhf1-GFP intensity normalized to the starting Hhf1-GFP intensity for each nucleus (47 nuclei). (E) Histogram of DAPI intensity for all cell cycle phases. Cell cycle stage was scored by SPB state, and GFP intensity was normalized to the one-SPB population mean. Black bars represent one SPB (G1; N = 126), green bars represent two SPBd (S/G2; N = 117), and blue bars represent two SPBs that are bioriented (M; N = 19).



FIGURE 2: Individual chromosomes are variable between nuclei. (A, B) Schematic of Ashbya chromosomes. All seven chromosomes and the mitochondrial DNA are shown to scale. Red ovals indicate centromeres; green bars indicate the location of the 32lacO::Gen3 integration onto chromosomes I (A) and VI (B). The chromosome I and IV lacO integrations are 126 and 56 kb away from the centromere, respectively. Still images of *Ashbya* nuclei containing zero to four copies of both chromosomes I and VI are shown beside each schematic. (C) Quantification of chromosome counts in *Ashbya* nuclei. Chromosome I counts are indicated in black and chromosome VI counts are in gray (chromosome I, N = 350; VI, N = 364). (D) Chromosome I counts are independent of cell cycle phase, as indicated by SPB state. Dark blue bars represent chromosome I distribution for one SPB (G1), and light blue bars represent the chromosome I distribution for two SPBs (S/G2/M). One SPB, N = 112; two SPBs, N = 66. (E) RNA FISH of individual nuclei using *CLN1/2*-TAMRA probes that hybridize to mRNA. Signals in nucleus represent sites of gene expression, and >90% nuclei express *CLN1/2* transcript. Images of DNA (Hoechst) and chromosomes and a merge image show that individual nuclei containing one or two signals indicate sites where *CLN1/2* is expressed in a single nucleus.



FIGURE 3: Ashbya chromosome variation is independent of location in the cell and increases as cells age. (A) Chromosome I counts and cell age. Black bars represent the chromosome I distribution for young cells (<40 nuclei), and gray bars represent the chromosome I distribution for older cells (>40 nuclei). Young cells, N = 178; old cells, N = 350. (B) Cumulative distribution plot of chromosome I counts and distance from cell tip (ANOVA, F = 1.56, p > 0.20).

state of nuclei along with Lacl signals and found that 40% of nuclei with a single SPB have >1 Lacl spot, suggesting that the presence of additional Lacl spots is not simply due to replication (Figure 2D). As an alternative measure of chromosome number, we used fluorescence in situ hybridization (FISH) to visualize chromosome VI at the *CLN1/2* locus. Specifically, we localized *CLN1/2* mRNA to identify sites of active transcription and determine chromosome number within nuclei. Based on this approach, 32% of nuclei have two or more copies of chromosome VI (Figure 2E; N = 436). Thus these two different approaches for detecting individual chromosomes support the idea that individual nuclei vary in their chromosome content and that a substantial fraction of the population of nuclei is not haploid.

Chromosome number increases as cells age

We next looked at how the frequency of altered chromosome number changes as cells age. Ashbya spores do not germinate synchronously, and so we used the number of nuclei within a cell as a measure of age. This is because the number of nuclei scales with growth, which is generally similar among individuals of a given age. We defined "young" cells as those with <40 nuclei and "old" cells as those with <40 nuclei. Of interest, chromosome I counts per nucleus increased with the age of mycelia, with a shift from a mode of 1 in very young cells and a mode of 2 in older cells. This suggests that deviations from a haploid complement of chromosomes accumulate with cell age (Figure 3A).

Nuclei with increased chromosome content are positioned throughout the cell

The increase in chromosome copy number with cell age prompted us to examine the position of the nuclei with extra DNA. Nuclei in *Ashbya* cells have been show to move and intermix within the mycelium (Anderson et al., 2013). Therefore we hypothesized that nuclei with many copies of individual chromosomes may be positioned further away from the actively growing tips of *Ashbya* cells. This would ensure that haploid nuclei would predominantly populate the most active portions of the cell, whereas those nuclei with altered chromosome numbers reside in the more interior portions of the cell. Surprisingly, however, there is no apparent bias in position for specific ploidies, as nuclei with each chromosome count were found both close to and distant from growing cell tips (Figure 3B; analysis of variance [ANOVA], F = 1.56, p > 0.20). Thus nuclei of variable ploidies can be found throughout the cell, indicating that altered-ploidy nuclei are functional or at least tolerated even in actively growing areas.

Chromosomes are faithfully segregated at mitosis

Given the variation in the copy number of chromosomes I and VI and the increase as cells age, we hypothesized that Ashbya nuclei were missegregating chromosomes during mitosis. We filmed cells with labeled chromosome I and, remarkably, found no evidence of missegregation as determined by the equal inheritance of LacI-GFP spots (Figure 4A and Supplemental Movies S1 and S2, N = 36 mitoses). Regardless of whether the observed nucleus started with one or two LacI spots, all the marked chromosomes were accurately segregated. Consistent with faithful segregation, most sister nuclei were born with comparable levels of Hhf1-GFP intensities, which would be unlikely if there were large chromosome imbalances (Figure 4, B and C, 15 sister pairs). The similarities between sisters were seen regardless of the intensity of the mother nucleus. These data support the idea that chromosome copy number variation does not arise from sloppy chromosome segregation in mitosis. Thus, even in a syncytial cell, where more genome copy number variability theoretically could be tolerated, there are controls of chromosome instability.

Populations of nuclei are polyploid, with limited aneuploidy

The faithful segregation of chromosomes in *Ashbya* suggests that many nuclei may be polyploid rather than aneuploid. Unfortunately, due to limited selectable markers available in *Ashbya*, we are unable to use two different marked chromosome reporters such lacO and tetO together. Therefore, to assess aneuploidy and polyploidy frequencies, we generated a strain with both chromosomes



FIGURE 4: Ashbya faithfully segregates chromosomes at mitosis. (A) Still images of chromosome segregation at mitosis. Top, frames from Supplemental Movie S1, showing one copy of chromosome I being faithfully segregated. Arrowheads point to single chromosome spots. Bottom, frames from Supplemental Movie S2, showing two copies of chromosome I being segregated. The asterisk is centered between the two copies of chromosome I in each nucleus. (B) Sister Hhf1-GFP intensity at birth. The sum Hhf1-GFP intensity of the brighter sister is plotted in black, with the dimmer sister overlaid in gray (15 pairs of sisters). (C) Kolmogorov–Smirnov (K-S) test plot of observed sister Hhf1-GFP intensities. The observed difference in sister Hhf1-GFP intensity at birth (immediately after mitosis) is displayed as a cumulative distribution plot in black. A randomized difference was calculated for two different populations of observed nuclei; the lowest 10 observed Hhf1-GFP intensities were used as a distribution for 1N, and the highest 10 observed Hhf1-GFP intensities were used as a distribution (15 sister pairs; compared with 1N [red line], D = 0.26, p = 0.26; compared with 2N [blue line], D = 0.19, p = 0.65).

I and VI marked with lacO repeats in the same nuclei (Figure 5A). If nuclei are polyploid, nuclei should have LacI spots in multiples of two, such as two, four, or eight. However, if nuclei are aneuploid, then odd numbers should be observed at some frequency, as by chance chromosome I or VI would be in excess or missing (Figure 5B). When scored in very young cells, the majority of nuclei in



FIGURE 5: Ashbya nuclei are predominantly polyploid. (A) Schematic of Ashbya chromosomes. All seven chromosomes and the mitochondrial DNA are shown to scale. Red ovals indicate centromeres, and green bars indicate the location of the 32lacO::Gen3 integration onto chromosomes I and VI. (B) Schematic of ChI 32lacO::Gen3/ChVI 32-lacO::Nat1 results. Nuclei with two, four, or eight resolvable Lacl spots are polyploid. All other Lacl spot counts would be evidence for aneuploidy. (C) Quantification of chromosome counts in ChI 32-lacO::Gen3/ChVI 32-lacO::Nat1 Ashbya nuclei. Black bars represent the distribution for young cells (<40 nuclei, N = 118), and gray bars represent the distribution for older cells (>40 nuclei, N = 253).

the strain with two marked chromosomes contain two Lacl spots, consistent with the idea that nuclei are haploid at a young age (<40 nuclei). As cells grow (≥40 nuclei), the distribution shifts to a modal number of four spots, indicative of two copies of each chromosome (Figure 5C, N = 118 [≥40 nuclei], N = 253[<40 nuclei]). Coincident with aging, there is also an increase in nuclei with odd numbers of chromosomes, which reflects modest levels (~25%) of aneuploidy. These data are consistent with the observed faithful segregation of chromosomes at mitosis in young cells (Figure 4A and Supplemental Movies S1 and S2) and supports the idea that nuclei within the syncytium can exist with chromosomal aneuploidies but are predominantly polyploid. We speculate that as the volume of cytosol increases with age, low levels of aneuploidy become tolerable. Thus nuclei vary in ploidy in the same cell, but the majority of the variation likely involves amplification of whole genomes rather than individual chromosomes.

Spindle checkpoint protein Mad2 is required for normal growth

Given that the frequency of an euploidy is low in young cells, we hypothesized that the SAC is critical for keeping an euploidy in check. Activity of the checkpoint may then decrease as the cells age when an euploid nuclei appear. We generated mutants lacking Mad2—part of the conserved spindle assembly checkpoint machinery—and found that $mad2\Delta$ cells grow slowly and had highly irregular borders, which reflects unequal growth across multiple hyphae (Figure 6). Such variable growth edges would be expected if there is stochastic loss of genome stability and therefore localized cell death or senescence in certain branches, leading to their inability to grow while others maintain growth. This observed growth phenotype is considered "haploinsufficient," as these are heterokaryons and there is still wild-type Mad2 expressed from a subset of nontransformed nuclei. Therefore, in areas where there are many transformed $mad2\Delta$ nuclei, presumably growth cannot occur, whereas



FIGURE 6: $mad2\Delta$ cells have decreased viability in Ashbya. Colony growth of wild-type (-ade2) and $mad2\Delta$::NAT1 heterokaryon strains. Bars, SD; three plates/strain. Images represent colony growth after 5 d.

those branches with enough Mad2-expressing nuclei continue to grow. Alternatively, the overall dose of Mad2 is insufficient for each nucleus to trigger a checkpoint when needed, or nuclei residing in the same cell do not generally share checkpoint components. We hypothesize that the loss of Mad2 in branches causes higher levels of aneuploidy than can be tolerated and localized growth arrest. Heterokaryon cells lacking Mad2 also fail to produce asexual spores, making it impossible to generate homokaryon, null mutants. This requirement for the spindle checkpoint suggests that aneuploidy in young cells is not well-tolerated and is likely kept in check by the SAC. It also suggests that chromosome imbalances are not compensated for even though there are multiple neighboring nuclei capable of supplying missing or imbalanced gene products. The SAC is likely active and required even in nonperturbed conditions, indicating that there may be generally high levels of errors in chromosome attachment, perhaps due to the presence of extra chromosomes in polyploid nuclei and insufficient spindle microtubules.

Stress promotes a redistribution of nuclei to a haploid state

In previous studies, it was found or postulated that variation in ploidy—primarily aneuploidy— might be adaptive under stressful or fluctuating conditions. In fact, experimental evolution experiments show that aneuploidy can be one of the fastest ways in which yeast cells adapt to fluctuating conditions (Yona et al., 2012). If so, we predict that growing cells under stress may promote even more extreme ploidy differences and potentially increase the level of chromosome variation among nuclei. We subjected Ashbya strains with a lacO-marked chromosome I to a panel of stress conditions, including cell wall stress (2 mM caffeine), osmotic stress (200 mM NaCl), excess zinc (10 mM ZnSO₄), the ergosterol synthesis inhibitor and clinical antifungal fluconazole (325 nM), and temperature (37°C). Remarkably, in all stress conditions except caffeine, the ploidy status shifted significantly to a euploid state rather than a more polyploid state, such that the majority of nuclei now contained a single detectable Lacl-GFP spot (Figure 7; Z test, p <0.05; >85 nuclei each condition). This supports the idea that Ashbya cells can vary ploidy in response to external signals but that, remarkably, stress drives the cells to a more euploid state rather than more variable ploidy.

DISCUSSION

Ploidy variation has been shown to be advantageous or deleterious, depending on the biological context. For example, aneuploidy is a



FIGURE 7: Chromosome number variation decreases in response to various cellular stresses. Quantification of chromosome I counts in *Ashbya* nuclei under normal growth and in the presence of various cellular stresses (>85 nuclei/condition).

common feature of solid tumors and is associated with infection and drug resistance in human fungal pathogens, yet when artificially engineered into cells, aneuploidy can be a fitness disadvantage. Although many aneuploidies arise from cells that undergo a multinucleate stage, the functional consequences of having variable ploidies in one cytosol have not been examined. Given that in multinucleate cells there is the potential for mixing and sharing of gene products between nuclei with different genotypes, it is unclear whether aneuploidy would be deleterious, beneficial, or neutral. Here we analyzed natural ploidy variation in the multinucleate fungus Ashbya, in which nuclei transit the division cycle asynchronously (Gladfelter et al., 2006; Nair et al., 2010). Thus the goal of this work was both to understand how ploidy might vary in syncytia and to see whether ploidy could be a source of differences among nuclei. Recent work has implicated aneuploidy in delaying cell cycle progression (Thorburn et al., 2013), and we speculated that ploidy variation may be tolerated in a shared cytoplasm and be a source of division-timing variation among nuclei.

We observed in Ashbya that nuclei have variable numbers of chromosomes, ranging from one to greater than four copies of individually tagged chromosomes. Nuclei are able to faithfully segregate chromosomes and appear to be frequently polyploid and less frequently aneuploid. Deletion of MAD2 is deleterious, suggesting that a functional spindle assembly checkpoint is necessary for normal growth in Ashbya. Of note, growth under stress causes ploidy to decrease to near euploid, suggesting that chromosome count variability is not maintained under stress. Thus polyploidy and to a certain degree aneuploidy are tolerated in Ashbya under optimal growth conditions. Electron tomography analysis of a small number of nuclei in young Ashbya cells showed the mitotic spindle contains a similar number of kinetochore microtubules as budding yeast, and yet haploid A. gossypii has less than half the number of chromosomes (seven chromosomes in Ashbya vs. 16 in S. cerevisiae; Gibeaux et al., 2012). Therefore it is probable that in diploid nuclei, sufficient kinetochore microtubules are made for correct chromosome attachments at mitosis but that, as ploidies increase with cell age, kinetochore microtubules become limiting and/or improper attachments are more common.

In addition to polyploidy, there is also a degree of aneuploidy among Ashbya nuclei, suggesting some level of chromosome loss. In the pathogenic fungus C. albicans, parasexual cycles result in tetraploid nuclei that undergo concerted chromosome loss to reach near diploidy (Hull et al., 2000; Bennett, 2003; Forche et al., 2008). Chromosome loss after polyploidization was also observed in S. cerevisiae, in which the frequency of loss increased with increased ploidy (Mayer and Aguilera, 1990; Gerstein et al., 2006, 2008). Similarly, in Drosophila rectal cells, polyploid cells produce high levels of aneuploidy (Schoenfelder et al., 2014). It is possible that predominantly polyploid nuclei infrequently undergo chromosome loss, resulting in the low levels of aneuploidy we observe in Ashbya. Consistent with this hypothesis, we see increased levels of aneuploidy in older cells, suggesting that the increased ploidy variation in more mature cells may be leading to increased chromosome loss.

Although limited aneuploidies are observed, they are likely not well tolerated in Ashbya. Deletion of MAD2 results in decreased colony growth and produces phenotypes even when some nuclei express Mad2. This indicates both that aneuploidy is likely not tolerated and that checkpoints may be nuclear intrinsic in action, which has been seen in the context of binucleate yeast and marsupial cells (Rieder et al., 1997; Demeter et al., 2000). Why is aneuploidy not tolerated when gene products can theoretically be shared between nuclei in a syncytium? In fact, the cytosol may not be as functionally homogeneous as might be expected. These results are consistent with recent work in Ashbya showing that nuclei can create functional compartments, and neighboring nuclei do not completely intermix their surrounding cytosol. These nuclear territories require cytoplasmic microtubules and functional protein aggregates that localize mRNAs (Anderson et al., 2013; Lee et al., 2013). The regionalization of cytosol around individual nuclei may decrease any hypothetical buffering capacity of the syncytium. In a context in which the cytosol is regionalized rather than homogenized, nuclei with polyploid complements of chromosomes create functional territories. Aneuploid nuclei may not be functionally complemented by neighboring nuclei and therefore may not be maintained.

The existence of functionally distinct regions of cytosol with limited exchange might help explain why external cell stress reduces ploidy variation. It is well established that for proteins that form larger complexes to function (e.g., tubulin heterodimers), an imbalance in the expression of one of the components is deleterious (Weinstein and Solomon, 1990). In addition, it has been shown that excess proteins that arise from a chromosome amplification create a proteotoxic burden on cells as they degrade excess proteins to achieve optimum stoichiometries in protein complexes (Oromendia et al., 2012). In Ashbya, in which a haploid, diploid, and tetraploid nucleus may all be neighbors, there is the possibility for some local imbalances in levels of proteins in complexes at the boundaries between nuclear territories, and this may present some low baseline of proteotoxic stress. This stress may be outweighed in normal conditions by having some nuclei at enhanced biosynthetic capacity by housing more DNA. Under stress, perhaps the burden of proteotoxic stress is too great, and haploid nuclei take over, as seen in the majority of stress conditions analyzed. Of interest, caffeine did not lead to a ploidy reduction, potentially due to a different response to this specific condition or because the level of stress was insufficient to generate proteotoxic strain. In any case, this work reveals another example in which ploidy changes under stress and shows that the number of nuclei in the cell might be an important factor in how ploidy state might change in stress. This is important in light of the observation of multinucleate cells in tumors.

How are ploidy differences established in *Ashbya*? In the filming of individual nuclei with marked chromosomes, which revealed no missegregation, there were unfortunately no hints of how nuclei become polyploid. We found no evidence of nuclear fusion, failed anaphases, or clear endoreduplication, but we predict that there must be some frequency of these events that potentially are suppressed during live-cell imaging and evade our detection. Although the direct mechanism of establishing ploidy variation remains elusive, the striking shift we observed from an apparently haploid state in very young cells to polyploid states with age suggests that germlings emerge from haploid spores. This is also supported by extensive application of reverse genetics in the system in which clean gene replacement strains are usually able to be generated from single spores of a heterokaryon transformant (Wendland *et al.*, 2000).

How do haploid spores arise out of mature polyploid/aneuploid mycelia? Ashbya cells form asexual spores only after colonies have matured for multiple days on an agar plate. The specific mechanisms of spore formation in Ashbya are poorly understood, and it is not known whether there is a meiotic component to sporulation. There are many conserved components for meiosis, and it has been speculated that nuclei fuse and then undergo meiotic-like reductions to make spores (Wasserstrom et al., 2013). However, the lack of mechanistic evidence for meiotic-like reductions suggests that all spores examined in the lab are asexual. To generate haploid spores from the variable ploidy nuclei of mature mycelia means that there is a likely a reduction in ploidy and correction of the low levels of aneuploidy during spore formation. Alternatively, the population of asexual spores formed may have variable ploidy and be a bet-hedging strategy for germination in different environments, which would be especially useful source of genetic diversity in an asexually reproducing population. We saw that only half of wild-type Ashbya spores germinate (N > 200; unpublished observations). This low germination frequency suggests that it is likely that spores of many ploidies are produced, but only haploids are able to germinate under lab conditions. Furthermore, spores from $mad2\Delta$ strains show an even lower germination rate of 33%, suggesting that altering the SAC and presumably increasing ploidy variation further decreased spore viability in lab conditions. In S. cerevisiae, >90% of spores undergo germination and are viable (Klapholz et al., 1985; Diaz et al., 2002). However, many fungi that under certain conditions have been shown to be aneuploid, including some strains of S. cerevisiae, Neurospora crassa, Sordaria macrospora, and the basidiomycetes C. neoformans and Coprinus cinereus, have decreased spore viability (Klapholz et al., 1985; Celerin et al., 2000; Diaz et al., 2002;

Storlazzi *et al.*, 2003; Bowring *et al.*, 2006). It is possible that differences in genome copy number, even if balanced as in polyploidy, are more detrimental in spores with just a single nucleus, whereas ploidy variation is more tolerated in mature cells.

The high degree of ploidy variation in *Ashbya* is striking and may be a mechanism by which nuclei cycle asynchronously through the cell cycle. The connection between ploidy, cytoplasmic compartmentalization, and adaptive fitness is fascinating, and further study will ideally link these traits to cell cycle timing variation within a common cytoplasm.

MATERIALS AND METHODS

Growth conditions and strain construction

A. gossypii media, culturing, and transformation conditions were performed as described previously (Ayad-Durieux *et al.*, 2000; Wendland *et al.*, 2000). The strains in this study are described in Supplemental Table S1, the plasmids used are listed in Supplemental Table S2, and the oligonucleotide primers used are listed in Supplemental Table S3. All restriction enzymes were from New England BioLabs, Ipswich, MA, and all oligos were from Integrated DNA Technologies, Coralville, IA.

lacO plasmid construction

To generate plasmids AGB245/246, pAKH37 and AGB21 were digested with *Kpn*I and *Nde*I (NEB). The ~1600-base pair band (pAKH37) and the 4214-base pair band (AGB21) were gel extracted and ligated using T4 Ligase (NEB). Resulting plasmids were verified by digesting with *Kpn*I and *Nde*I and sequencing. To generate plasmids AGB264/265, AGB246 and pRS416 were digested with *Nde*I and *Sbf*I. The 3343-base pair band from AGB246 was gel extracted and ligated with digested pRS416 using T4 Ligase (NEB). Resulting plasmids were verified with *Sac*I digests and sequencing with AGO37.

Chromosome I 32-lacO::Gen3

Chromosome I homology was engineered using gap repair. Approximately 250 base pairs of 5' homology to chromosome I were amplified from $\Delta l\Delta t$ genomic DNA using AGO638/639. AGB264/265 were digested with Stul and cotransformed into yeast with 5' chromosome I homology to generate AGB266/267, which were verified by Bgll digestion and sequencing with AGO640. Approximately 250 base pairs of 3' homology to chromosome I were amplified from $\Delta l\Delta t$ genomic DNA using AGO641/642. AGB266/267 were digested with Ndel and cotransformed into yeast with 3' chromosome I homology to generate AGB268/269, which were verified by digestion with Spel/Stul and sequencing with AGO640 and AGO678. A 4260-base pair fragment was gel purified from AGB268/269 after digestion with Spel/Stul and transformed into AG302 to generate AG459.1-4. Primary transformants were picked onto G418 plates (200 µg/ml), and single spores were picked to generate the homokaryon AG460. Strains were verified using oligo pairs AGO721/ AGO761, AGO723/AGO759, and AGO721/AGO723.

Chromosome VI 32-lacO::Gen3

Approximately 200 base pairs of 5' homology to chromosome VI were amplified from $\Delta l\Delta t$ genomic DNA using AGO753/754. AGB264/265 were digested with *Stul* and cotransformed into yeast with 5' chromosome VI homology to generate AGB309/310, which were verified by digestion with *Xhol* and sequencing with AGO640. Approximately 250 base pairs of 3' homology to chromosome VI were amplified from $\Delta l\Delta t$ genomic DNA using AGO765/766. AGB309/310 were digested with *Ndel* and cotransformed into yeast with 3' chromosome I homology to generate AGB322/323, which were verified by digestion with *Xnml/Xhol* and sequencing with AGO678. AGB322/323 were digested using *Spel/Stul* and transformed into AG302 to generate AG479.1-3. Primary transformants were picked onto G418 plates (200 µg/ml), and single spores were picked to generate the homokaryon AG480. Strains were verified using oligo pairs AGO757/AGO758, AGO98/AGO760, and AGO757/AGO760.

Chromosome I 32-lacO::Gen3/chromosome VI 32-lacO::Nat1

To make strain AG514/515, AGB268/269 were digested using *Spel/Stul* and transformed into AKH26.2. Primary transformants were picked onto G418 plates (200 µg/ml), and single spores were picked to generate the homokaryon AG517/518. Strains were verified using oligo pairs AGO721/AGO761, AGO723/AGO759, and AGO721/AGO723. To make a chromosome VI homology Nat1 plasmid, a 1817–base pair fragment was gel extracted from AGB09 digested with *Sapl* and *Ndel*. Using this band, Nat1 was PCR amplified using AGO1050/968 and cotransformed with AGB322/323 to generate plasmid AGB403.1/403.2. Plasmids were verified using *Kpnl* digests and sequencing with AGO234/235. AGB403.1 was transformed into AG517 to generate strain AG720. Strains were verified using oligo pairs AGO757/AGO758, AGO760/AGO234, and AGO757/AGO760.

mad2∆::Nat1

To generate strain AG600/601, AGB9 was cut with *Pvull*-HiFi. NAT1 was amplified off of this cut plasmid using AGO1101/AGO1102. PCR products were pooled and directly integrated into *Ashbya*. Primary transformants were picked onto CloNat plates (Werner Bioagents, Jena, Germany). Strains were verified using oligo pairs AGO1103/AGO235, AGO234/AGO1104, and AGO1103/AGO1105.

Microscope setup and imaging conditions

A Zeiss Axioimage-M1 upright light microscope (Carl Zeiss, Jena, Germany) equipped with a Plan-Apochromat 63×/1.4 numerical aperture oil objective was used. To visualize the fluorescence signals, an Exfo X-Cite 120 lamp was used in conjunction with Zeiss 38HE (GFP), Chroma 41002B (TAMRA), Zeiss 49 (Hoechst/DAPI), and Chroma 41043 (mCherry; Chroma Technology, Brattleboro, VT). Images were acquired on an Orca-AG charge-coupled device (CCD) camera (C4742-80-12AG; Hamamatsu, Bridgewater, NJ) driven by Volocity (Perkin-Elmer, Waltham, MA).

To image Lacl chromosome spots, cells were grown for 16 h and imaged on thin gel pads containing 2% agarose and 100% 2x lowfluorescence minimal medium. Z-stacks were acquired with 0.5-µm slices spanning the hyphae. The images were exposed for 50– 100 ms at 100% transmission. All images were processed by iterative deconvolution (100 iterations) using calculated point spread functions in Volocity, and nuclei were visually scored for number of Lacl-GFP–labeled chromosomes.

To image chromosome segregations, cells were imaged on an OMX microscope equipped with a Hamamatsu electron-multiplying CCD. A 488-nm solid-state laser was used for GFP illumination, using 1.08% laser power. Cells were imaged every minute through a 10- μ m z-depth (21 slices at 0.5 μ m).

Hhf1-GFP intensity quantification

Time-lapse movies and previous nuclear tracking were used to quantify Hhf1-GFP intensity over time (Anderson *et al.*, 2013).

Time-lapse images and nuclear coordinates were imported into MATLAB, and the nuclei around the central coordinates was found using thresholding. Photobleaching correction was applied to individual nuclei over time. Histone intensity was measured as the sum GFP intensity for each nucleus. Fold change for each nucleus was calculated using the maximum and minimum Hhf1-GFP signal throughout the tracking. S-phase duration was the time between maximum and minimum GFP intensity.

Spindle pole body scoring

Cell cycle stages were determined using SPBs as described previously (Nair *et al.*, 2010). Briefly, nuclei with a single SPB were assigned to G1, nuclei with either a single SPB that was twice the intensity and size as a G1 SPB or with two adjacent SPBs that were <0.5 μ m apart were assigned to S/G2, and nuclei with two SPBs that were >0.5 μ m apart on opposite sides of the nucleus were assigned to M.

FISH

A single-molecule RNA FISH protocol was used to visualize the chromosome expressing the CLN1/2 gene (Lee et al., 2013). This transcript is expressed in >90% of nuclei, and the site of expression of the gene is readily detected as spots that are generally brighter than the intensity of single mature transcripts in the cytosol. Because all attempts at DNA FISH were unsuccessful and a DNA FISH protocol has never been established for Ashbya, we used RNA FISH of transcription as a proxy for chromosome number. Cells were grown for 12-16 h at 30°C while shaking and fixed in 3.7% formaldehyde for 1 h. Cells were washed into buffer B (1.2 M sorbitol, 0.1 M potassium phosphate, pH 7.5), spheroplasted using Zymolyase (15 µg/ml), and incubated at 37°C until phase dark. Cells were washed twice with wash buffer and mixed with hybridization solution (100 μ l of hybridization buffer, 1.5 μ l of probe). Cells were incubated at 92°C for 3 min and then overnight at 37°C. Cells were washed twice with wash buffer and incubated for 10 min at room temperature in 500 µl of wash buffer plus 1 µl Hoechst. Cells were washed twice more, mounted in Prolong Gold mounting media (Life Technologies), and imaged.

Germination and colony growth assays

For germination assays, spores were spread onto Ashbya Full Media (AFM) plates with appropriate selection and allowed to germinate at 30°C for 8 h. Plates were imaged and scored in ImageJ (National Institutes of Health, Bethesda, MD) for spore germination. For colony growth assays, 10 μ l of spores was plated in the center of AFM plates with appropriate selection. Colonies grew at 30°C, and plates were imaged daily for up to 10 d using a Bio-Rad Chemi-Doc XR5 molecular imager with ImageLab software. Colony area and perimeter were measured using a macro written in ImageJ, which "autothresholded" images and determined colony area and perimeter.

Cellular stress tests

A. gossypii spores were germinated in liquid AFM in the presence of ampicillin (100 μ g/ml), G418 (200 μ g/ml), and Clonat (50 μ g/ml) for 10 h while shaking in baffled flasks at 30°C. Cells were transferred to fresh media with selection containing 2 mM caffeine (Sigma-Aldrich), 200 mM NaCl (Fisher Scientific), 10 mM ZnSO₄ (Sigma-Aldrich), or 325 nM fluconazole (Sigma-Aldrich) and returned to 30°C with shaking for an additional 6 h. The appropriate fluconazole concentration was determined by comparing radial growth rates on AFM plates with ampicillin (100 μ g/ml) and

1.5% agarose, supplemented with varying concentrations of fluconazole (Sigma-Aldrich). Control and heat-stressed cells were also transferred to fresh media after 10 h and returned to 30 or 37°C, respectively, for an additional 6 h. Cells were spun down at 300 rpm for 5 min, washed with 5 ml of $2\times$ low-fluorescence minimal media, and spun as before. A 10-µl amount of cells was taken directly from the pellet and plated onto glass slides for imaging.

Statistical analysis of cellular stress populations

All pairwise comparisons were performed between unstressed cells and each stress condition (0 chromosomes [control] vs. 0 chromosomes [experimental], 1 chromosome [control] vs. 1 chromosome [experimental]). p_1 = proportion of nuclei with a given number of chromosome copies (control) and p_2 = proportion of nuclei with the same number of chromosome copies (experimental), and n_1 and n_2 = the number of nuclei scored (control and experimental, respectively). A two-sample Z test for proportions was applied to each population compared with the control population. For p < 0.05, the Z value must be >1.96 or <-1.96. Strains were considered significantly different from the control if two or more categories were p < 0.05.

$$\frac{(\overline{p}_1 - \overline{p}_2) - 0}{\sqrt{\overline{p}(1 - \overline{p})} \left(\frac{1}{n_1} + \frac{1}{n_2}\right)}$$

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