A role for cell polarity in lifespan and mitochondrial quality control in the budding yeast Saccharomyces cerevisiae
A role for cell polarity in lifespan and mitochondrial quality control in the budding yeast *Saccharomyces cerevisiae*

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**SUMMARY**

Babies are born young, largely independent of the age of their mothers. Mother-daughter age asymmetry in yeast is achieved, in part, by inheritance of higher-functioning mitochondria by buds and retention of some high-functioning mitochondria in mother cells. The mitochondrial F box protein, Mfb1p, tethers mitochondria at both poles in a cell cycle-regulated manner: it localizes to and anchors mitochondria at the mother cell tip throughout the cell cycle and at the bud tip before cytokinesis. Here, we report that cell polarity and polarized localization of Mfb1p decline with age in *Saccharomyces cerevisiae*. Moreover, deletion of genes (*BUD1*, *BUD2*, and *BUD5*) that mediate symmetry breaking during establishment of cell polarity and asymmetric yeast cell division cause depolarized Mfb1p localization and defects in mitochondrial distribution and quality control. Our results support a role for the polarity machinery in lifespan through modulating Mfb1 function in asymmetric inheritance of mitochondria during yeast cell division.

**INTRODUCTION**

Mitochondria are cell fate determinants that are differentially segregated during asymmetric cell division in stem cells and budding yeast. Indeed, new and old mitochondria are segregated during division of human mammary stem-like cells. The daughter cell that preferentially inherits new mitochondria retains stem cell properties, whereas the daughter that inherits old mitochondria becomes a tissue progenitor cell (Katajisto et al., 2015). Similarly, in the budding yeast, *Saccharomyces cerevisiae*, the daughter cell or bud inherits fitter mitochondria that are more reduced, have lower reactive oxygen species levels (ROS), and higher membrane potential (Δψ). This in turn, allows daughter cells to live a full lifespan, whereas mother cells, which inherit lower-functioning mitochondria, continue to age (Higuchi et al., 2013; McFaline-Figueroa et al., 2011).

Three critical events occur during asymmetric cell division: breaking of cellular symmetry, polarity establishment and maintenance to generate anterior and posterior poles, and polarized cell division (Rose and Gόnczy, 2014). In yeast, bud site selection, determining the site for the formation of daughter cells or buds on mother cells, is one hallmark of cell symmetry breaking. Bud site selection is controlled by a system of polarity cues and culminates in the activation of Bud1p/Rsr1p, a Ras-like GTPase, at the selected bud site (Bender and Pringle, 1989). The activity of Bud1p is tightly controlled: it is activated by *BUD5*, the guanine nucleotide exchange factor (GEF), and quickly inactivated by *BUD2*, the GTPase-activating protein (GAP) (Bender, 1993; Chant et al., 1991; Chant and Herskowitz, 1991; Kang et al., 2001; Park et al., 1993, 1999; Powers et al., 1991).

These bud site selection proteins promote polarization of the cytoskeleton toward the selected bud site. Bud1p/Rsr1p that is activated at the selected bud site recruits Cdc24p to that site (Zheng et al., 1995). This, in turn, results in region-specific activation of Cdc42p, Cdc42p-dependent assembly of actin cables at that site, and actin cable-driven transport of all cellular constituents toward the selected bud site leading to bud formation and growth (Fehrenbacher et al., 2004; Moseley and Goode, 2006). Polarized actin cables also transport and further enrich Cdc42p at the bud tip (Slaughter et al., 2009). Finally, actin cables enable segregation of mitochondria in dividing yeast by mediating 1) preferential transport of higher-functioning mitochondria from the mother cell to bud (Higuchi et al., 2013) and 2) transport of one of the tethers
(Mmr1p) that anchors and retains those higher-functioning mitochondria at the bud tip (Shepard et al., 2003; Swayne et al., 2011). Indeed, promoting actin cable function in asymmetric mitochondrial inheritance during cell division promotes daughter cell fitness and extends lifespan in yeast (Higuchi et al., 2013). Interestingly, although the bud site selection machinery acts to break cell symmetry at a specific site on the yeast cell surface and to maintain a stable mother-to-bud axis between cell divisions (Bi and Park, 2012; Slaughter et al., 2009; Wu and Lew, 2013), it is not essential for polarized cell division in yeast per se (Kozubowski et al., 2008). In the absence of BUD1/RSR1, polarity factors cluster at multiple, seemingly random sites throughout the mother-cell cortex at the onset of cell division. This leads to a competition between different sites, and ultimately, to establishment of polarity at a single site followed by polarized yeast cell division (Wu et al., 2013).

Nonetheless, the bud site machinery is critical for lifespan control in yeast: loss of BUD1 and polarity cue-guided bud site selection reduces replicative lifespan (RLS) (Clay et al., 2014) and chronological lifespan (Campos et al., 2018). These studies also revealed that BUD1/RSR1 is required for establishing a diffusion barrier in the ER at the bud neck, which promotes retention of oxidatively damaged ER proteins in the mother cell (Clay et al., 2014). Because asymmetric inheritance of oxidized proteins is critical for normal RLS, the bud site selection machinery may contribute to RLS through its function in the ER diffusion barrier at the bud neck. However, deletion of BUD1 actually extends RLS in yeast exposed to ER stress (Clay et al., 2014). Thus, the reduced lifespan observed upon deletion of BUD1 is not because of effects on the diffusion barrier in the ER at the bud neck.

Moreover, although the bud tip is well established as the anterior pole in asymmetrically dividing yeast, with distinctive membrane, organelle, mRNA, and protein constituents, emerging evidence indicates that the tip of the mother cell that is distal to the bud may be the posterior pole. Specifically, a small population of mitochondria that are higher functioning compared to other mitochondria in the mother cell, are anchored at and accumulate in the mother cell tip (Pernice et al., 2016; Yang et al., 1999). This process is critical for mitochondrial quality control, yeast cell fitness and lifespan, and mother-daughter age asymmetry (Pernice et al., 2016; Yang et al., 1999).

Mfb1p, a mitochondria-associated F-box protein (Dürr et al., 2006; Kondo-Okamoto et al., 2006), serves as a tether to anchor higher-functioning mitochondria in the mother cell tip (Pernice et al., 2016). Mfb1p localization and function are polarized at two levels (Pernice et al., 2016). First, Mfb1p is asymmetrically distributed between mother and daughter cells. Through most of the cell cycle, Mfb1p is excluded from the daughter cell. That is, it localizes to mitochondria exclusively in the mother cell. However, shortly before cytokinesis, some Mfb1p localizes to mitochondria in the bud. Second, Mfb1p exhibits polarized localization within the mother cell: it localizes solely to mitochondria that accumulate at the mother cell tip. Indeed, Mfb1p is the only known protein that localizes to the mother cell tip through the entire cell cycle.

Finally, Mfb1p functions in mitochondrial distribution and quality control. Specifically, it functions to tether mitochondria in the mother cell tip throughout the cell cycle and contributes to anchorage of mitochondria in the bud tip after cytokinesis. Importantly, the mitochondria that are tethered at the mother cell tip and bud tip are more reduced and therefore higher functioning compared to other mitochondria in the yeast cell. Mfb1p function in mitochondrial distribution and quality control contributes to lifespan control in yeast: deletion of MFB1 results in defects in anchorage of mitochondria in the mother cell tip, reduced mitochondrial function throughout the cell, and premature aging (Pernice et al., 2016).

Here, we tested whether the bud site selection machinery is critical for Mfb1p localization and function in mitochondria distribution and function, and how this contributes to lifespan. We provide evidence that the fidelity of bud site selection and the polarized localization of Mfb1p decline early in the aging process in budding yeast. We also identify a role for the bud site selection machinery in lifespan and mitochondrial quality control through effects on the polarized localization of Mfb1p at the distal tip of the yeast mother cell.

RESULTS
Budding polarity declines with age
One model for the aging process in yeast is replicative lifespan (RLS), a measure of the number of times that a mother cell can divide before senescence (Longo et al., 2012). Although yeast strains can undergo up to
35–45 generations of replicative aging, age-associated phenotypes are evident at earlier replicative ages. Indeed, mitochondrial redox state begins to decline in yeast that have undergone five rounds of cell division (age 5) (McFaline-Figueroa et al., 2011). Similarly, decreases in the acidification of the yeast cell and vacuole (the lysosome of yeast) are evident at age 4 (Henderson et al., 2014; Hughes and Gottschling, 2012). Finally, an increase in DNA damage, which is detected by an increase in foci containing the DNA repair protein Rad52p in the yeast nucleus, is detected at age 7 (Alvaro et al., 2007; Novarina et al., 2017). These findings raise the possibility that changes which occur early during the yeast RLS may cause or contribute to the aging process.

Here, we studied whether bud site selection changes during aging. Yeast cells of defined replicative age were isolated using a miniature chemostat aging device (mCAD) (Hendrickson et al., 2018). Mid-log phase yeast, which are largely young cells, were immobilized and allowed to undergo replicative aging in the mCAD (Figure 1A). Daughter cells produced from immobilized cells were removed by continuous media exchange, and yeast of defined replicative age were harvested from the mCAD. We confirmed the efficacy of the mCAD for these studies: immobilized cells isolated from mCAD exhibit increasing age and aging phenotypes, including an increase in cell size and a decrease in viability, as a function of the time of propagation in mCAD (Figures 1B and S1A–S1C).

The bud scar is a chitinous ring on the cell wall of the mother cell that forms where buds separate from mother cells. It serves as a marker for the number of times a mother cell divides and as a landmark for cell symmetry breaking during establishment of cell polarity. In haploid yeast cells that undergo axial or unipolar budding, new bud sites form adjacent to the bud site from the previous round of cell division, which is marked by the bud scar (Chant and Pringle, 1995).

To assess the fidelity of axial budding, bud scars were visualized using wheat germ agglutinin (WGA), a chitin-binding agent. Inspired by TrackScar (Maxwell and Magwene, 2017), we sequentially stained yeast bud scars with WGA conjugated to different fluorescent dyes to distinguish new from old bud scars. Cells were first stained with Alexa 488-WGA to label all bud scars. They were then propagated for 2 h and stained with Alexa 594-WGA to label the two newest bud scars (Figure 1A). Here, bud site selection was scored as polarized if the two newest bud scars were adjacent to each other and random if those bud scars were not adjacent.

We found that bud site selection is polarized in >97% of the young cells (1-4 generations) examined. However, loss of budding polarity is evident early in the aging process. Random bud site selection was detected in 11.0% of aged cells at age 6–10 in RLS and increased further to 27.7 and 30.6% in cells of more advanced age, 11–20 and >20 generations, respectively (Figures 1C and 1D). Thus, polarity establishment during bud site selection declines with age in yeast and these declines are evident early in the aging process. Interestingly, the proportion of cells displaying randomized bud-site selection plateaus at ca. 30% for cells of age 11–20 and beyond.

Polarization of Mfb1p within mother cells but not between mother cells and buds declines with age

Next, we tested whether aging affects the function and polarized localization of Mfb1p. We visualized mitochondria and Mfb1p using Cit1p-mCherry (a mitochondrial matrix marker) and Mfb1p-GFPEnvy, respectively, in yeast of defined replicative age. Because Mfb1p localizes to the mother cell throughout the cell cycle and to the bud tip late in the cell division cycle, we restricted analysis to cells in early stages of the cell cycle (bud-to-mother ratios between 0.2 and 0.6) when Mfb1p localizes exclusively to the mother tip.

We detect co-localization of Mfb1p with mitochondria throughout the aging process in yeast (Figure 2A). Quantitative analyses of co-localization of Mfb1p with mitochondria using Pearson’s coefficient and Manders’ M1 coefficient indicate that there is no significant difference in the co-occurrence of Mfb1p with mitochondria as a function of replicative age (Figures S2A and S2B). Interestingly, Manders’ M2 coefficient, a measure of co-localization of mitochondria with Mfb1p, gradually increases with age.

In contrast, we find that the polarized localization of Mfb1p within mother cells declines with age (Figures 2A and 2B). Moreover, this decline is evident early in the aging process. We confirmed that Mfb1p co-localizes with mitochondria exclusively at the distal tip of young mother cells (1-4 generations), which reflects
Figure 1. Polarized bud site selection declines as yeast cells age

(A) Illustrations of the miniature chemostat aging device (mCAD) used to isolate cells of different replicative age (left) and the approach used to monitor budding polarity with age by labeling older bud scars with Alexa 488-WGA and the two newest bud scars with Alexa 594-WGA (right).

(B) Quantification of cell diameter as a function of time of propagation in the mCAD and replicative age. The diameter measured was the longest axis of the mother cell. Data are represented as mean ± SEM.

(C) Quantification of percentage of cells with random bud site selection. Data are represented as mean ± SEM, n = 3 independent trials. n > 50 cells/age group/trial. The unpaired two-tailed t test was used for statistical analysis. p values are denoted as: **p < 0.01; *p < 0.05.

(D) Representative images of the budding pattern of cells of different replicative age. Replicative age (number in right corner of each image) was determined by scoring the number of WGA-stained bud scars. Older bud scars were labeled with Alexa 488-WGA (green). Yellow arrowheads: new Alexa 594-WGA-stained bud scars (red).
Mfb1p function as a tether for mitochondria in the mother distal tip. However, we detect a decrease in the amount of Mfb1p-GFPEnvy and mitochondria in the mother distal tip and a corresponding increase in Mfb1p-GFPEnvy and mitochondrial levels in the center of the mother cell early during the aging process (age 5–9) (Figures 2B and 2C).

Figure 2. Polarized distribution of Mfb1p and mitochondria within the yeast mother cell but not between mother cells and buds declines with age

Age-associated changes early in the cell cycle
(A) The localization of Mfb1p and mitochondria in cells during early stages in the cell cycle as a function of replicative age. Mitochondria and Mfb1p were visualized by tagging the mitochondrial marker protein Cit1p with mCherry (red) and Mfb1p with GFP (green). To visualize cells early cell cycle stages, cells with mother:bud ratio of 0.2–0.6 were analyzed. The replicative age was determined by scoring calcofluor white-stained bud scars (blue). Cells were grouped according to age: 1–4, 5–9, and >9 generations (n >49, 72, and 31 cells, respectively, in three independent trials). Images shown are maximum projections. Cell outlines: white. Scale bars: 1 μm.

(B and C) Quantification of the distribution of Mfb1p and mitochondria by measurement of the integrated fluorescence in five regions of interest (ROIs) defined at the upper right panel. (B) Data are represented as mean ± SEM, p <0.0001 (one-way ANOVA with Tukey’s multiple comparisons test).

(D and E) Quantification of the integrated fluorescence of Mfb1p and mitochondria in mother cells and buds. Data are represented as mean ± SEM. (D) p <0.3312 (one-way ANOVA with Tukey’s multiple comparisons test). (E) p <0.0001 (one-way ANOVA with Tukey’s multiple comparisons test).

(F–J) Age-associated changes late in the cell cycle.
(F) The localization of Mfb1p and mitochondria in cells during late stages in the cell cycle as a function of replicative age as carried out as for panel (A). Cells with mother:bud ratios of 0.6–0.8 were grouped according to age: 1–4 and >4 generations (n >33 and 46 cells, respectively, in three independent trials).

(G–J) Quantification of the integrated fluorescence of Mfb1p and mitochondria in five ROIs (G and H) and in mother cells and buds (I and J). Data are represented as mean ± SEM, p values are denoted as: ****p <0.0001; ***p <0.001; **p <0.01; *p <0.05. (G) p <0.0024 (one-way ANOVA with Tukey’s multiple comparisons test). (I) p <0.0001 (one-way ANOVA with Tukey’s multiple comparisons test).
Interestingly, the age-associated decline in localization of Mfb1p to mitochondria in the mother cell tip results in relocalization to mitochondria at other regions within the mother cell, but not to mitochondria in the bud (Figures 2A, 2C, and 2E). Thus, asymmetric distribution of Mfb1p between mother cells and buds appears to persist with age. To test this hypothesis, we studied the localization of Mfb1p late in the cell cycle, when some Mfb1p localizes to the bud where it contributes to anchorage of mitochondria at that site (Figures 2F–2J). Age-associated declines in polarization of Mfb1p and mitochondria within mother cells are evident at late stages in the cell cycle. Mfb1p localizes exclusively to mitochondria that accumulate in the mother cell tip at age 1-4 in yeast bearing large buds. However, at age >4, some Mfb1p is detected at mitochondria throughout the mother cell (Figures 2F–2G). Interestingly, we detect accumulation of Mfb1p and mitochondria in the bud tip in young and old yeast late in the cell division cycle. Indeed, there is no significant change in the amount of Mfb1p that localizes to mitochondria in the bud tip during the aging process (Figures 2F–2I). There is a subtle but significant decrease in accumulation of mitochondria with age (Figures 2F, 2H, and 2J). Because Mfb1p contributes to but is not solely responsible for tethering mitochondria at the bud tip late in the cell cycle (Pernice et al., 2016), it is not clear whether those changes are because of effects on Mfb1p. Overall, our findings support the model that Mfb1p is polarized both within mother cells and between mother cells and buds, and that only the former, localization of Mfb1p to mitochondria in the mother cell tip, declines with replicative age, and does so early in the aging process.

Disabling bud site selection machinery disrupts the polarized localization and function of Mfb1p in the mother cell tip but not Mfb1p asymmetric localization between mother cells and buds

To determine whether the age-linked declines in Mfb1p are because of changes in the bud site selection machinery, we monitored Mfb1p distribution and function in yeast bearing a deletion in BUD1. We confirmed that deletion of BUD1 results in random bud site selection in haploid yeast cells. On the other hand, deletion of MFB1 does not affect bud site selection (Figures S3A–S3B).

Next, we quantified the distribution of Mfb1p-GFP and mitochondria in WT and bud1Δ mutant cells. Interestingly, deletion of BUD1 produces defects in Mfb1p localization and function that are similar to those produced early in the aging process. Deletion of BUD1 has no obvious effect on 1) the steady-state level of Mfb1p (data not shown), 2) association of Mfb1p with mitochondria (Figure S4), 3) inheritance of mitochondria by yeast daughter cells, or 4) the asymmetric distribution of Mfb1p between mother cells and buds (Figures 3A–3C, right panels). However, loss of axial bud site selection interferes with the polarization of Mfb1p and mitochondria within mother cells (Figures 3A–3C, left panels). Specifically, deletion of BUD1 results in a decrease in the accumulation of Mfb1p and mitochondria at the mother cell tip, and a compensatory shift of Mfb1p and mitochondrial mass to the mother cell neck in mid-log phase cultures. Thus, Mfb1p relocates to mitochondria in the mother cell but not in the bud in bud1Δ cells. Interestingly, the defects in accumulation of mitochondria in the mother cell tip in bud1Δ cells are not as severe as those observed upon deletion of MFB1. As such, our findings indicate that the accumulation of Mfb1p and its
Figure 4. Deletion of MMR1 rescues depolarized localization of Mfb1p, defects in mitochondrial distribution, and decreased mitochondrial quality in bud1Δ cells.

(A) Representative images of Mfb1p (green) and mitochondria (red) in mid-log phase WT, bud1Δ, mmr1Δ, or bud1Δ mmr1Δ cells visualized as for Figure 2A. Cell outlines: white. Scale bars: 1 µm.
function in anchorage of mitochondria in the mother cell tip, but not its binding to mitochondria or asymmetric distribution between buds and mother cells, is dependent on Bud1p/Rsr1p. Equally important, the mitochondrial phenotypes associated with deletion of BUD1 resemble those observed during early stages of yeast cell replicative aging.

To further explore the relationship of the bud site selection machinery to Mfb1p and its function in mitochondrial distribution, we studied the effect of deletion of BUD2 and BUD5, genes that regulate Bud1p, in these processes. As expected, deletion of BUD2 or BUD5 results in random bud site selection (Figure S5A). Moreover, we found that the phenotypes observed in bud2Δ and bud5Δ cells are similar to those observed in bud1Δ cells and during early stages of aging. Deletion of BUD2 or BUD5 results in defects in polarization of Mfb1p and mitochondria within mother cells (Figures S5B, S5E, and S5F) but has no effect on the asymmetric distribution of Mfb1p or mitochondria between mother cells and buds (Figures S5C and S5D).

Next, we tested whether loss of bud site selection affects mitochondrial function, using mitochondria-targeted redox-sensing GFP (mito-roGFP1) (Figures 3D and 3E). We confirmed that deletion of MFB1 results in reduced mitochondrial redox state: the ratio of reduced to oxidized mito-roGFP1 in mfb1Δ cells is 20% lower than that observed in WT cells. Beyond this, we found that mitochondrial redox state in bud1Δ, bud2Δ, and bud5Δ cells is decreased compared to WT mother cells but greater than that observed in mfb1Δ cells (Figures 3D, 3E, S5G, and S5H). Finally, in bud1Δ cells, mitochondria at the mother cell tip are not significantly higher-functioning compared to other mitochondria within the mother cell (Figure S6). Thus, mutation of bud site selection genes compromises the function of mitochondria not just in the mother cell tip but throughout the yeast mother cell.

Deletion of MMR1 rescues defects in Mfb1p and mitochondrial distribution as well as mitochondrial quality control observed in bud1Δ mutants

The premature aging phenotype of mfb1Δ has been attributed to the disproportionate loss of higher-functioning mitochondria from mother cells (Pernice et al., 2016). Deleting the cargo adaptor and bud tip mitochondrial tether, MMR1, can effectively reduce the amount of mitochondria transferred to and retained in buds, and thus rescue the overall distribution of mitochondria between mother and bud as well as the mitochondrial quality defects observed upon deletion of MFB1 (Chernyakov et al., 2013; Itoh et al., 2004; Pernice et al., 2016; Swayne et al., 2011). Therefore, if the reduced fitness and lifespan observed in bud1Δ cells are a consequence of depolarization of Mfb1p within mother cells and the associated defects in retention of higher-functioning mitochondria in the mother cell, then deletion of MMR1 should rescue the defects in mitochondrial distribution and quality observed in a bud site selection mutant.

To examine this possibility, we deleted the MMR1 gene in bud1Δ and compared the phenotypes observed in bud1Δ mmr1Δ cells to those in bud1Δ or mmr1Δ single mutants and to WT cells. Deletion of MMR1 does not change the overall distribution of Mfb1p between mother and bud cells (Figures 4A and 4D). However, it increases the abundance of mitochondria in mother cells (Figures 4A and 4E) and in the distal tip area of mother cells (Figures 4A and 4C). Intriguingly, analysis of the bud1Δ mmr1Δ mutant revealed that deletion of MMR1 partially restores the defects in accumulation of mitochondria and Mfb1p at the mother distal tip of bud1Δ cells (Figures 4A and 4B). Finally, deletion of MMR1 rescues the defect in mitochondrial quality in
Figure 5. *BUD1* and *MFB1* function in the same pathway for mitochondrial distribution and quality control and RLS

(A) Maximum projections of mitochondria visualized using Cit1p-mCherry in mid-log phase WT, *mfb1Δ*, *bud1Δ*, and *bud1Δ mfb1Δ* cells. Cell outlines: white. Scale bars: 1 μm.

(B and C) Quantification of the mitochondrial distribution of (A). ROIs are defined in Figure 2C. >21 cells/strain/trial. Data are represented as mean ± SEM. The means of three independent trials are shown in different shaped and colored...
Finally, we used epistasis analysis to determine whether mean RLS of WT, deletion of both genes will not have additive effects on RLS compared to single mutants. We find that the mother cells and buds produced by deletion of BUD1 (Figures 4F and 4G). Taken together, we find that inhibition of mitochondrial inheritance by deletion of MMR1 rescues the overall defects in mitochondrial distribution and function produced by loss of BUD1.

**BUD1 and MFB1 act in the same pathway to regulate mitochondria and lifespan**

Because mitochondria are aging determinants, it is possible that the bud site selection machinery contributes to lifespan control through effects on Mfb1p function in mitochondrial distribution and quality control. Indeed, we find that deletion of BUD1 or MFB1 results in similar phenotypes: defects in accumulation of mitochondria in the mother cell tip without affecting overall asymmetric distribution of the organelle between mother cells and buds, and defects in mitochondrial function throughout the yeast cell. We also find that bud site selection and Mfb1p polarization within mother cells both decline with age and observe striking similarities in those declines. Specifically, both processes decline at similar stages early in the yeast aging process and reach a plateau later in aging.

Here, we used analysis of epistatic interactions to determine whether the bud site selection machinery and Mfb1p function in the same pathway for control of mitochondrial distribution and quality control. If this hypothesis is correct, then accumulation of mitochondria in the mother cell tip and mitochondrial redox state observed in the bud1Δ mfb1Δ double mutant should not be greater than defects observed in either single mutant. Indeed, we obtained evidence for epistasis between BUD1 and MFB1 (Figures 5A–5E). Specifically, we found that the distribution of mitochondria, accumulation of the organelle in the mother cell tip and the redox state of mitochondria in the mother cell and bud are similar in bud1Δ mfb1Δ and mfb1Δ cells. Here too, we find that the phenotype observed in bud1Δ cells is less severe than that observed in mfb1Δ cells: mitochondrial distribution and redox state are both compromised in bud1Δ cells, but to a lesser extent than that observed in mfb1Δ cells. Moreover, we found that mitochondrial defects observed in bud1Δ mfb1Δ and mfb1Δ cells are greater than those observed in bud1Δ cells. However, the double bud1Δ mfb1Δ mutant is not significantly different from the mfb1Δ single mutant.

Finally, we used epistasis analysis to determine whether BUD1 and MFB1 function in the same pathway for RLS control (Figures 5F and 5G). If deletion of BUD1 affects lifespan through effects on Mfb1p function, then deletion of both genes will not have additive effects on RLS compared to single mutants. We find that the mean RLS of WT, bud1Δ and mfb1Δ single mutants, and bud1Δ mfb1Δ double mutants are 26.7 ± 7.51, 17.15 ± 4.94, 12.1 ± 5.01, and 15.33 ± 5.10 generations, respectively. Thus, we confirmed that deletion of BUD1 or MFB1 results in a significant decrease in yeast RLS. In addition, we find that RLS of the bud1Δ mfb1Δ double mutant is similar to that observed in the mfb1Δ single mutant and that the reduction in RLS observed in the bud1Δ mfb1Δ double mutant is not more severe than that observed in either bud1Δ or mfb1Δ single mutants. These findings provide genetic evidence that MFB1 and BUD1 function in the same pathway for control of mitochondrial distribution and quality control and for RLS in yeast.

**DISCUSSION**

Cell polarity during asymmetric cell division declines with age. For example, stem cell division becomes more symmetric in aged mice, indicating a more depolarized phenotype (Florian et al., 2018). Cell division
also becomes more symmetric with replicative age in yeast, which leads to premature aging in daughter cells (Kennedy et al., 1994). Moreover, specific components of the cell polarity machinery change with age. Cdc42 activity increases with age in hematopoietic stem cells, and inhibition of Cdc42 reduces aging phenotypes observed in those cells (Florian et al., 2012). Similarly, actin cables, the tracks for cargo movement during polarized cell division in yeast, undergo age-linked declines in stability and polarity (Sing et al., 2021).

The bud site selection machinery drives breaking of cell symmetry at the onset of polarity establishment during asymmetric cell division in yeast. Analysis of the mechanism underlying bud site selection has revealed different patterns for symmetry breaking in haploid and diploid yeast, the landmarks for bud site selection, the proteins that mediate those processes, and how bud site selection activates reorganization of the cytoskeleton to the selected polarity site. Moreover, studies in multicellular organisms indicate that symmetry breaking and defining polarity planes are critical for establishing directional cues for development. However, the benefit of bud site selection for yeast, a single-celled organism, is not well understood.

Our study also demonstrates that the decline in budding polarity in aging yeast is not linear. The bud site selection machinery is robust in younger cells (1-5 generations). This is expected because stringent bud site selection is evident in mid-log phase yeast, which are primarily young cells. However, we detect a significant increase in random bud site selection early in the aging process (>5 generations). The age-associated decline in bud site selection does not appear to be because of changes in BUD1 gene expression: two independent studies on changes in the yeast transcriptome and proteome with age indicate that BUD1 mRNA and protein levels remain constant during the yeast RLS (Hendrickson et al., 2018; Janssens et al., 2015). On the other hand, studies from our lab and others indicate that age-linked declines in established aging determinants are evident early in the yeast RLS and may therefore cause or be major contributors to the aging process (Henderson et al., 2014; Hughes and Gottschling, 2012; McFaline-Figueroa et al., 2011). Here, we find that the decline in bud site selection fidelity occurs early in replicative aging and in the same time frame as age-linked changes in other aging determinants. These findings are consistent with a causative role for declines in bud site selection in yeast cell aging.

Interestingly, this decline in axial bud site selection reaches a plateau at 10–20 generations and does not decline further as cells age. Previous studies indicate that a cell that undergoes random budding events can exhibit polarized bud site selection at subsequent rounds of cell division (Jazwinski et al., 1998). This finding raises the possibility that the polarity of bud site selection is not lost irrevocably with age. Rather, the stringency of this process may decline with age. The plateau in polarized bud site selection observed in yeast at advanced age may be a consequence of the reduced stringency of this aspect of polarity establishment in yeast.

Age-associated declines in Mfb1p localization and function in control of mitochondria

Our previous studies revealed that Mfb1p functions as a tether that anchors a small population of higher-functioning mitochondria in the mother cell tip, and that this process is required to retain some higher-functioning mitochondria in mother cells, which in turn is required for mother cell fitness and lifespan (Pernice et al., 2016). Our previous studies also revealed an age-linked decline in mitochondrial function in...
yeast (McFaline-Figueroa et al., 2011) and that defects in mitochondrial quality control can result in premature aging in yeast (Higuchi et al., 2013). Here, we find that Mfb1p function in binding to mitochondria does not decline with age. However, we detect an age-associated decline in Mfb1p’s function as a region-specific mitochondrial tether: accumulation of Mfb1p and mitochondria in the mother cell tip diminishes with age. The age-associated declines in Mfb1p localization and function in the mother cell tip do not appear to be because of changes in MFB1 gene expression: analysis of changes in the yeast transcriptome with age indicate that MFB1 mRNA levels remain constant during the yeast replicative lifespan (Hendrickson et al., 2018; Janssens et al., 2015).

Moreover, we find that the decline in Mfb1p localization at the mother cell tip occurs early in the aging process, at the same stage as age-linked declines in bud site selection. Indeed, we find that defects in Mfb1p localization also reach a plateau, similar to the plateau observed in age-linked declines in bud site selection. These findings reveal temporal links between aging of the bud site selection machinery and Mfb1p. Moreover, because declines in bud site selection and Mfb1p localization and function occur early in RLS, it is possible that these changes contribute to the aging process.

Finally, inheritance of higher-functioning mitochondria by yeast daughter cells persists during the aging process (McFaline-Figueroa et al., 2011) and is not compromised by deletion of MFB1 or BUD1. Moreover, asymmetric inheritance of Mfb1p between mother cells and buds is not affected by loss of the bud site selection machinery and does not decline with age. Therefore, Mfb1p and bud site selection function in lifespan control is not because of effects on asymmetric inheritance of mitochondria or Mfb1p. Rather, they are because of defects in polarization of Mfb1p and mitochondria within the mother cell, and the consequences of these defects on mitochondrial function.

Overall, there are multiple mechanisms to ensure that higher functioning mitochondria are inherited by yeast daughter cells. Higher functioning mitochondria exhibit higher motility activity and are preferentially transferred from mother to daughter cell (Higuchi et al., 2013). In addition, newly inherited higher functioning mitochondria are retained in daughter cells by anchorage in the bud tip (Swayne et al., 2011). Finally, any lower functioning, less motile mitochondria that enter daughter cells are transported from buds to mother cells (Higuchi et al., 2013). However, there is only one known mechanism to ensure that healthy mitochondria are retained in mother cells: Mfb1p-mediated anchorage of higher functioning mitochondria in the mother cell tip. This process affects the fitness and lifespan of mother cells. Moreover, although daughter cells preferentially inherit the fittest mitochondria from mother cells and this process persists in mfb1Δ and bud1Δ cells, a decline in mitochondrial function in mother cells impacts on all progeny produced from mother cells.

Role for the bud site selection machinery in mitochondrial quality control and lifespan

Our findings also support a role for polarized bud site selection in yeast lifespan control through effects on Mfb1p-dependent mitochondrial quality control. Specifically, we find that deletion of BUD1, BUD2, or BUD5 results in defects in localization of Mfb1p to the mother distal tip, defects in anchorage of mitochondria at that site and a decrease in mitochondrial function throughout yeast cells. Moreover, we found that deletion of MMR1, a protein implicated in movement of mitochondria from mother cells to buds and anchorage-dependent retention of newly inherited mitochondria the bud tip (Chernyakov et al., 2013; Itoh et al., 2004; Swayne et al., 2011), restores the defects in mitochondrial distribution and overall quality observed upon deletion of BUD1. Thus, our findings support a role for the bud site selection machinery in mitochondria quality control through effects on mitochondrial distribution.

Although there are clear links between the bud site selection machinery and Mfb1p functions in mitochondrial distribution and quality control, loss of bud site selection activity produces significantly less severe defects on mitochondrial distribution and function compared to deletion of MFB1. However, as described above, loss of bud site selection does not inhibit establishment of cell polarity or polarized cell division in yeast. Consistent with this, asymmetric distribution of Mfb1p between mother cells and buds is not compromised in bud1Δ, bud2Δ, or bud5Δ mutants. Moreover, deletion of bud site selection proteins does not affect association of Mfb1p with mitochondria. Thus, Mfb1p is present, active, and asymmetrically distributed in cells with no bud site selection machinery. Therefore, it is possible that the defects observed in bud1Δ, bud2Δ, or bud5Δ cells are less severe than those observed in mfb1Δ because Mfb1p can still promote some anchorage and retention of mitochondria in mother cells in the absence of bud site selection cues.
We also obtained genetic evidence that BUD1 and MFB1 function in the same pathway for mitochondrial distribution and quality control and on lifespan. First, we observed that aging has similar effects on bud site selection and Mfb1p polarization within mother cells. In both cases, declines are evident early in the aging process and reach a plateau in cells of advanced age. Second, deletion of either BUD1 or MFB1 produces similar effects on mitochondrial distribution, anchorage of mitochondria in the mother cell tip, mitochondrial redox state, and RLS. Moreover, deletion of both genes has no additive effect on those phenotypes compared to mfb1Δ or bud1Δ single mutants. Finally, we found that deletion of MMR1 and the associated decrease in mitochondrial inheritance by daughter cells rescues the polarized mitochondrial and Mfb1p distribution and defects in mitochondrial function observed in bud1Δ cells. This supports a critical role for bud site selection in mitochondrial quality control during inheritance. Thus, we conclude that the reduced lifespan produced by loss of polarized bud site selection is because of effects on Mfb1p function in mitochondrial quality control.

Is the mother distal tip the posterior pole during asymmetric cell division in haploid yeast?

During the establishment of cell polarity in model systems including frog and worm embryos, the cytoskeleton is reorganized leading to polarized localization of cortical proteins at anterior and posterior poles of the cell. Diploid yeast exhibit bipolar budding patterns (i.e., bud site selection either adjacent to or at opposite poles from the previous bud site) and asymmetric distribution of polarity factors at anterior and posterior poles of the cell (Chant and Pringle, 1991; Chiou et al., 2017; Kang et al., 2004). However, haploid yeast undergoes axial or unipolar bud site selection: each new bud is produced adjacent to the previous bud site. The bud tip is an established anterior pole and site for activation of polarity factors including Cdc42p and its effectors in haploid and diploid yeast. However, there is no obvious localization of polarity factors to a posterior pole in a haploid yeast cell. We find that the bud site selection machinery is required for 1) targeting of Mfb1p to the mother cell tip, 2) Mfb1p function in anchorage of higher-functioning mitochondria at the mother cell tip, and 3) RLS control through effects on Mfb1p localization and function. These observations support the model that a posterior pole exists in haploid yeast cells and that the mother cell tip is that pole.

Limitations of the study

Bud sites are selected by related but distinct mechanisms in haploid and diploid yeast cells: haploid cells exhibit axial budding whereas diploid cells exhibit bipolar budding. These studies were carried out in haploid yeast cells. In addition, these studies were carried out in the S288c genetic background. Previous studies indicate that the W303 yeast strain, which carries a mutation in the bud site selection gene BUD4 (Voth et al., 2005), exhibits defects in asymmetric distribution of Mfb1p (Kondo-Okamoto et al., 2006, 2007).

STAR METHODS

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Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.103957.

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AUTHOR CONTRIBUTIONS

Conceptualization, E.J.Y., W.M.P., and L.A.P.; Formal Analysis, E.J.Y. and W.M.P.; Investigation, E.J.Y. and W.M.P.; Writing – Original Draft, E.J.Y.; Writing – Review & Editing, E.J.Y., W.M.P., and L.A.P.; Funding Acquisition, L.A.P.; Supervision, L.A.P.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Chemicals, peptides, and recombinant proteins** | | |
| KAPA HiFi PCR kit | Roche Sequencing and Life Science, Kapa Biosystems, Wilmington, MA | KK1006 |
| Hygromycin B | Sigma-Aldrich, St. Louis, MO | H3274 |
| Geneticin | Sigma-Aldrich, St. Louis, MO | A1720 |
| Calcofluor® White M2R staining | Milipore Sigma, Burlington, MA | #18909 |
| Wheat Germ Agglutinin, Alexa Fluor™ 688 Conjugate | Thermo Fisher Scientific, Waltham, MA | #W11261 |
| Wheat Germ Agglutinin, Alexa Fluor™ 594 Conjugate | Thermo Fisher Scientific, Waltham, MA | #W11262 |
| Wheat Germ Agglutinin, Alexa Fluor™ 647 Conjugate | Thermo Fisher Scientific, Waltham, MA | #W32466 |
| EZ Link Sulfo-NHS-LC-LC-Biotin | Thermo Fisher Scientific, Waltham, MA | #21338 |
| Miltenyi μMACs Streptavidin MicroBeads | Miltenyi Biotec Inc., Auburn, CA | #130-048-101 |
| Miltenyi LS Columns | Miltenyi Biotec Inc., Auburn, CA | #130-042-401 |
| Miltenyi MultiStand | Miltenyi Biotec Inc., Auburn, CA | #130-042-303 |
| Miniature chemostat aging device (mCAD) | (Hendrickson et al., 2018) | |
| Dynabeads MyOne Streptavidin C1 beads | Thermo Fisher Scientific, Waltham, MA | #65001 |
| **Experimental models: Organisms/strains** | | |
| Please refer to Table S1 | | |
| **Oligonucleotides** | | |
| Please refer to Table S2 | | |
| **Recombinant DNA** | | |
| pFA6a-link-GFPEnvy-SpHis5 | (Slubowski et al., 2015) Linda Huang lab | Addgene plasmid #60782 |
| pCY3090-02 | (Young et al., 2012) Anne Robinson lab | Addgene plasmid #36231 |
| pFA6a-GFP(565T)-KanMX6 | (Bähler et al., 1998) Jurg Bähler and John Pringle lab | Addgene plasmid #39292 |
| HO-pGPD-mito-roGFP-KanMX6-HO | (Liao et al., 2020) | |
| pOM13 | (Gauss et al., 2005), Euroscarf | Euroscarf plasmid #P30388 |
| pOM12 | (Gauss et al., 2005), Euroscarf | Euroscarf plasmid #P30387 |
| pSH62 | (Gueldener et al., 2002), Euroscarf | Euroscarf plasmid #P30120 |
| **Software and algorithms** | | |
| Volocity 6.3 | Quorum Technologies | https://www.quorumtechnologies.com/volocity/volocity-downloads/legacy |
| FIJI | (Schindelin et al., 2012) | https://imagej.net/Fiji; RRID: SCR_002285 |
| Coloc2 | Daniel James White, Tom Kazimiers, Ellen Dobson | https://imagej.net/Coloc_2 |
| GraphPad Prism7 | GraphPad | https://www.graphpad.com/scientific-software/prism/; RRID: SCR_002798 |

**RESOURCE AVAILABILITY**

**Lead contact**

Further information and requests for resources, reagents and strains should be directed to and will be fulfilled by the lead contact, Liza A. Pon (lap5@cumc.columbia.edu).
Material availability
Plasmids and yeast strains generated in this study are available from the lead contact with a completed Materials Transfer Agreement.

Data and code availability
All data reported in this paper will be shared by the lead contact upon request. This paper does not report original code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Yeast growth conditions
All S. cerevisiae strains used in this study were generated in BY4741 (MATa his3Δ0, leu2Δ0, met15Δ0 and ura3Δ0) background from Open Biosystems (Huntsville, AL). The maintenance and manipulation of yeast cells were conducted as previously described (Sherman, 2002). Yeast cells were propagated in YPD medium [1% (w/v) yeast extract (BD, Franklin Lakes, NJ), 2% (w/v) Bacto-peptone (BD, Franklin Lakes, NJ), and 2% (w/v) dextrose (Sigma-Aldrich, St. Louis, MO)] at 30°C with vigorous shaking at 180–200 rotations per minutes (RPM). All experiments were carried out with liquid cultures grown to the mid-logarithmic phase (O.D.600 0.1–0.3). For aging experiments using the mCAD device, yeast cells were grown in Synthetic Complete (SC) media. For imaging experiments, all strains were washed once in SC medium [0.67% (w/v) Yeast nitrogen base without amino acids and with ammonium sulfate, amino acid mix, 2% (w/v) glucose] to reduce auto-fluorescence from YPD medium. A list of yeast strains used in this study is in Table S1.

Yeast strain construction
Knockout or tagged strains were generated essentially as described (Gardner and Jaspersen, 2014). Knockout strains were generated by using homologous recombination to replace the target genes with LEU2 selectable marker cassettes. The marker cassette was PCR-amplified from pOM13 or pOM12 plasmid (Gauss et al., 2005) with 40 bp of flanking homology upstream and downstream of the target gene. PCR reactions were performed using the KAPA HiFi PCR kit. The PCR fragments were transformed into yeast cells by the lithium acetate method described in (Gietz and Woods, 2002). In brief, yeast cells were grown to the mid-logarithmic phase, and about 2.310⁸ cells were aliquoted and harvested at 3,000 g for 1 min. Cells were washed once with 0.1 M of lithium acetate, and mixed with a transformation mixture consisting of 33.33% of PEG3350, 0.1 M lithium acetate, 0.1 mg of boiled salmon sperm carrier DNA, and 50 μL of unpurified PCR product. The transformation mixture was incubated at 30°C for 30 min, and subsequently incubated at 42°C for 45 min. Transformants were selected on agar plates containing SC medium without leucine (SC-Leu) or without uracil (SC-Ura) for 2 days. Successful transformants were confirmed by sequencing the inserted region.

To visualize Mfb1p, a bright GFP variant, GFPEnvy, was fused to the C-terminal of Mfb1p by genetic manipulation. All GFPEnvy tagged strains were generated by using the lithium acetate method described above to insert a PCR product containing the GFPEnvy gene with the SpHis5 selectable marker after the coding region of the gene of interest. The primer sets were designed to amplify PCR fragments with 40 bp of flanking homology upstream and downstream of the stop codon of the target sequence. The backbone of the PCR product was amplified from the pFA6a-link-GFPEnvy-SpHis5 plasmid (Addgene plasmid # 60782). The PCR cassette was transformed into yeast cells and selected based on the lithium acetate transformation method described above. Transformants were selected on SC medium without histidine (SC-His) plates.

The visualization of mitochondria was achieved by C-terminally fusing mCherry to α = the mitochondrial matrix protein Cit1p. The mCherry fluorophore and the hphMX4 selectable marker was amplified from pCY3090-02 with 40 bp of flanking homology upstream and downstream of the stop codon of the target sequence. The backbone of the PCR product was amplified from the pFA6a-link-GFPEnvy-SpHis5 plasmid (Addgene plasmid # 60782). The PCR cassette was transformed into yeast cells and selected based on the lithium acetate transformation method described above. Transformants were selected on YPD plates containing 200 μg/mL hygromycin B (Sigma-Aldrich, St. Louis, MO).

To perform ratiometric quantification of mitochondrial redox state, endogenously expressed GDPp-mitoGFP1 followed by the KanMX4 selectable marker was integrated into the HO locus. The PCR cassette was transformed into yeast cells and selected based on the lithium acetate transformation method described
above. Transformants were selected on YPD plates containing 200 μg/mL Geneticin (Sigma-Aldrich, St. Louis, MO).

The plasmids and primers used in strain construction are listed in key resource table and Table S2.

METHOD DETAILS

Microscopy

Fluorescence microscopy was performed with one of the following imaging systems: (1) a Zeiss Axioskop 2 Plus upright fluorescence microscope with a 100×/1.4 Numerical Aperture (NA) Zeiss Plan-Apochromat objective lens (Carl Zeiss Inc., Thornwood, NY), Light-Emitting Diode (LED) module (CoolLED pE-4000, Andover, UK) and an Orca ER cooled charge-coupled device (CCD) camera (Hamamatsu Photonics, Hamamatsu City, Japan), and (2) a Zeiss AxioObserver.Z1 inverted fluorescence microscope with a 100×/1.3 oil EC Plan-Neofluar objective lens, a metal-halide lamp and an LED Colibri system (Carl Zeiss Inc., Thornwood, NY), and Orca ER cooled CCD. The first system was controlled by NIS Elements 4.60 Lambda software (Nikon, Melville, NY), and the second system was controlled by Zen Blue (Carl Zeiss Inc., Thornwood, NY). Details about the imaging conditions are given within each experimental section below.

Visualization of bud scars

To determine the age of yeast cells, bud scars were visualized by staining with Calcofluor White M2R (Millipore Sigma, Burlington, MA) or WGA conjugated to Alexa Fluor 488, 594 or 647 (Thermo Fisher Scientific, Waltham, MA). For calcofluor staining, 25 μM Calcofluor White M2R was incubated with yeast cells for 5 min, followed by 3 washes with SC media. Calcofluor-stained cells were imaged using a standard DAPI filter set (Chroma/Zeiss filter set 49; excitation G365, dichroic FT 395, emission 445/50). For WGA staining, 1 μg/mL WGA conjugated to AlexaFluor was added to a yeast cell culture and incubated for 15 min, followed by 3 washes with SC media. Alexa Fluor 488 was excited by a 470 nm LED and Alexa Fluor 594 was excited by 561 nm LED, and emission was collected with a dual eGFP/mCherry cube (#59222, Chroma, Bellows Falls, VT), or a far-red filter set (Zeiss filter set 50 HE; excitation 640/30, emission 690/50).

Enrichment of aged cells

Aged cell enrichment by binding to magnetic beads was performed as described in (Lindstrom and Gottschling, 2009; Sinclair et al., 1997; Smeal et al., 1996) with modifications. Cultures were grown overnight in YPD at 30°C to an OD600 < 0.2 as described above. 10⁸ total cells (roughly 10 OD600) of cells were harvested by centrifuging at 1500 x g for 5 min. Cells were washed with 10 mL of ice-cold 1x phosphate-buffered saline (PBS) [137 mM sodium chloride, 2.7 mM potassium chloride, 10.14 mM sodium phosphate dibasic, and 1.77 mM potassium phosphate monobasic, and adjusted to pH 8]. Cells were labeled with 10 mM of EZ Link™ Sulfo-NHS-LC-LC-biotin (#21338, Thermo Fisher Scientific, Waltham, MA) dissolved in 1 x PBS for 30 min with gentle rotation at room temperature. The excessive biotin was quenched by washing cells three times with 1 x PBS containing 100 mM glycine. Cells were resuspended in 1 mL of YPD and used to inoculate 500 mL of YPD media at a density of 2 x 10⁸ cells/mL in a 2-L Erlenmeyer flask. Biotinylated cells were propagated at 30°C for 8 h. The OD600 of the culture was monitored and was not allowed to exceed 0.8. Cells were fixed by adding 20% of formaldehyde to a final concentration of 3.7% and incubated at 30°C with vigorous shaking for 30 min. Fixed cells were harvested by centrifuging at 3000 x g for 5 min, and washed with 10 mL of 1x PBS for 3 times. Cell density was measured and adjusted to 90 OD600 per mL, and cells were divided into 500-μL aliquots in 1.5 mL Eppendorf tubes. Miltenyi μMACs Streptavidin Micro-Beads (#130-048-101, Miltenyi Biotec Inc., Auburn, CA) were added (20 μL beads/45 OD600 cells) and rotated at room temperature for 30 min. Cells were washed twice with 1 mL of 1x PBS and then resuspended in 1 mL of 1x PBS. The cell suspension was sorted by MACS LS separation columns (#130-042-401, Miltenyi Biotec Inc., Auburn, CA). The separation columns were washed with 2 mL of 1x PBS and eluted with 2 mL of YPD by firmly pushing the plunger into the column. Enriched mother cells were stained with Calcofluor White as described above.

Enrichment of aged cells in the miniature-chemostat aging device and assessment of bud site selection

To determine the budding polarity in cells of different ages, aged cells were enriched in a miniature-Chemostat Aging Device (mCAD) and stained sequentially with WGA conjugated with Alexa Fluor 488 and Alexa Fluor 594. The assembly of the mCAD is described in (Hendrickson et al., 2018). In brief, cells were
grown overnight to mid-log phase (OD600 < 0.2). 4 OD600-mL of cells were harvested by centrifuging at 1500 × g in a Sorvall ST-16 tabletop centrifuge for 5 min, and washed twice with 5 mL of 1× PBS with 0.25% PEG3350. The cell pellet was resuspended in 0.5 mL of 1× PBS and then combined with 2 mg of EZ Link™ Sulfo-NHS-LC-LC-Biotin dissolved in 0.5 mL of 1× PBS. The biotin labeling reaction was carried out at room temperature for 30 min with gentle rotation. After the reaction, cells were washed twice with 1× PBS with 0.25% PEG3350 and then resuspended in 500 mL SC media in a 2 L Erlenmeyer flask. The biotin-labeled cells were grown at 30°C with shaking at 190–200 RPM for 8 h. The OD600 of the culture was monitored and was not allowed to exceed 0.3.

Labeled cells were counted using a hemocytometer. 0.9 μL of magnetic beads was used per 1 million labeled cells. Before the beading reaction, the Dynabeads MyOne Streptavidin C1 beads (#65001, Thermo Fisher Scientific) were equilibrated to room temperature and washed twice with 1 mL of SC media, and then resuspended in 20 mL of SC.

The post-biotinylated cell culture was harvested by centrifugation at 1500 × g for 5 min in a 50 mL conical-bottom tube, and the pellet was resuspended in 20 mL of SC. The concentrated cells were mixed with washed magnetic beads and rotated for 15 min at room temperature. The mother cells attached to magnetic beads were separated by ring magnets for 5 min. The supernatant was carefully removed and saved as young cells. The beaded mother cells were washed twice with 40 mL of SC, and then resuspended in 1 mL of SC media for mCAD loading.

For cell loading onto the mCAD, the vessel of the mCAD was removed from the magnets, and the air pump was set to the lowest airflow or none, and the effluent port was pulled above the media level or blocked by a tubing clamp. Beaded mother cells were transferred to a 1 mL Luer lock syringe and then loaded into the mCAD vessel by the Luer needle entry port. The loaded vessel was swirled to mix the cells, and placed on magnets to allow beaded mother cells to bind for 10 to 15 min. The media peristaltic pump (25 mL per hour) and air pump (~0.8-1 psi) were started after binding of beaded mother cells.

During mother cell harvesting, the media pump was off, and the air pump was set to the lowest setting. The effluent port was pulled up or blocked by a tubing clamp. The vessel was removed from the magnet and swirled to mix the cells. Mother cells were collected by using a Luer-lock syringe drawing from the loading port of the mCAD. The harvested cells were washed 4 times on magnets before the next procedure.

To distinguish the budding pattern in aged cells, cells were stained with WGA conjugated with 2 different AlexaFluor for visualizing the newest two bud scars on the cell surface. The staining process is described in the “visualization of bud scars” section. The mother cells were stained with WGA-Alexa488 for 15 min, washed 3 times, and diluted to OD600 < 0.2 in 5 mL of SC media and grown for 90 min at 30°C. Cells were then harvested and stained with WGA-Alexa594 and washed as above. Washed cells were resuspended in ~2 μL of SC and mounted on microscope slides (#3050, Thermo Fisher Scientific). Mounted cells were visualized on the microscope system 1 described above.

**Analysis of mitochondrial and Mfb1p distribution**

Analysis of mitochondrial distribution was performed as previously described (Pernice et al., 2016). In brief, yeast cells expressing endogenously tagged Cit1p-mCherry were imaged on System 1 or 2 using (1) a standard RFP filter set (Chroma) or (2) a Zeiss filter set 43 HE, excitation FT 570, dichroic FT 570, emission 605/70. Cells were imaged through the entire cell depth (6 μm total, with 0.3 μm z-steps), using 1 × 1 binning, 300 ms exposure with 561 nm LED with 55% power. For Mfb1p distribution, selected yeast strains expressing endogenously tagged Mfb1p-GFP or Mfb1p-GFPEnv were imaged on System 1 or 2, with excitation by a 470 nm LED and emission collected through either (1) a dual eGFP/mCherry cube (#59222, Chroma), or (2) a Zeiss filter set 38 HE, excitation 470/40, dichroic FT 495, emission 525/50, respectively. Acquired-wide-field images were deconvolved by an iterative restoration algorithm in Volocity (Quorum Technologies, ON, Canada) with a limit of 60 iterations. The relative distribution of mitochondrial mass was quantified by thresholding the mitochondrial voxels from 3D reconstructed microscopy images and calculating volume in 5 different regions (mother tip, mother center, mother neck, bud neck and bud tip, as defined in Figure 2B). The mother and bud diameters were measured along the longest axis of the cell based on the transmitted-light images. To quantify cells at similar cell cycle stages, cells were omitted if the bud-to-mother ratio was smaller than 0.2 or larger than 0.6.
Analysis of co-localization of Mfb1p and mitochondria

Co-localization between Mfb1p-GFPEnvy and mitochondria (Cit1p-mCherry) was calculated by using the Coloc2 plug-in in Fiji. Pearson’s and Manders’ coefficients were calculated to assess the correlation and co-occurrence, respectively, of the signals. The background of microscopic images was determined as the mean of a cell-free region near the cell of interest, and was subtracted from the image. Coloc2 was called with bisection threshold regression with a Region of Interest (ROI) flanking the cell. Coefficients and 2D intensity plots were calculated and visually inspected. Negative controls were performed by comparing a randomized channel with another channel.

Analysis of mitochondrial redox state

Analysis of mitochondrial redox state was conducted according to (Vevea et al., 2013) with modifications. The redox biosensor roGFP1 with mitochondrial targeting sequence from ATP9 was introduced into strains of interest by integration into the HO locus. Cells in mid-log phase were imaged on System 1 by excitation wavelength switching between 365 and 470 nm, and collecting emission through a modified GFP filter (Zeiss filter 46 HE without excitation filter, dichroic FT 515, emission 535/30). For the oxidized form of roGFP, a 300-ms exposure with 50% LED power was used, and for the reduced form a 200-ms exposure with 20% LED power. The wide-field images were deconvolved as described above. The background was calculated by selecting a background ROI and subtracted from the image by thresholding in the Ratio function in Volocity. The reduced-to-oxidized mito-roGFP ratio was calculated by dividing the voxel intensity of the reduced (λex = 470 nm, λem = 525 nm) channel by that of the oxidized (λex = 365 nm, λem = 525 nm) channel. The resulting ratio channel was measured with exclusion of zero values. The mother and bud diameters were measured as described above, and the same cell-size criteria were used to exclude cells in different cell cycle stages.

Analysis of replicative lifespan

Replicative lifespan (RLS) measurements were performed as described previously (Erjavec et al., 2008), without alpha-factor synchronization. Briefly, frozen glycerol stocks of select strains (stored at −80°C) were streaked out on YPD plates and grown for 2 days. Single colonies were grown overnight in liquid YPD at 30°C, diluted and grown to exponential phase for 4 h in YPD at 30°C. 2 μL of the cell suspension was streaked onto a YPD plate and small-budded cells were isolated and arranged in a matrix using a micromanipulator mounted onto a dissecting microscope (Zeiss, Thornwood, NY). Upon completion of budding, mother cells were discarded; the time and number of divisions of the corresponding daughter cells were recorded until all replication ceased.

QUANTIFICATION AND STATISTICAL ANALYSIS

All quantifications were subjected to normal distribution analysis with the D’Agostino and Pearson normality test. Statistical p values for two-group comparison were conducted by a two-tailed Student’s t test for parametric distributions and a Mann-Whitney test for non-parametric results. For more than two-group comparisons, p values were calculated by one-way ANOVA with Dunnett’s or Sidak’s test for parametric distributions and a Kruskal-Wallis test with Dunn’s post hoc test for non-parametric distributions. Results were recorded and sorted in Microsoft Excel and the statistical analyses were done in GraphPad Prism8 (GraphPad Software, San Diego, CA). Bar graphs and scatter graphs show the mean and standard error of the mean (SEM). For all statistical tests, p values are denoted as followed: ****p < 0.0001; ***p < 0.001; **p < 0.01; *p < 0.05.