Evidence that mutation accumulation does not cause aging in Saccharomyces cerevisiae

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Evidence that mutation accumulation does not cause aging in *Saccharomyces cerevisiae*

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Summary

The concept that mutations cause aging phenotypes could not be directly tested previously due to inability to identify age-related mutations in somatic cells and determine their impact on organismal aging. Here, we subjected *Saccharomyces cerevisiae* to multiple rounds of replicative aging and assessed de novo mutations in daughters of mothers of different age. Mutations did increase with age, but their low numbers, < 1 per lifespan, excluded their causal role in aging. Structural genome changes also had no role. A mutant lacking thiol peroxidases had the mutation rate well above that of wild-type cells, but this did not correspond to the aging pattern, as old wild-type cells with few or no mutations were dying, whereas young mutant cells with many more mutations continued dividing. In addition, wild-type cells lost mitochondrial DNA during aging, whereas shorter-lived mutant cells preserved it, excluding a causal role of mitochondrial mutations in aging. Thus, DNA mutations do not cause aging in yeast. These findings may apply to other damage types, suggesting a causal role of cumulative damage, as opposed to individual damage types, in organismal aging.

Key words: aging; DNA damage; lifespan; mitochondria; mutations; thiol peroxidase; yeast.

Introduction

The nature of aging remains one of the grand mysteries of biology (Kirkwood & Austad, 2000). In the 1950s, Failla and Szilard proposed that random somatic mutations cause cell and tissue dysfunction, leading to aging (Failla, 1958; Szilard, 1959). Maynard Smith questioned this hypothesis (Maynard, 1959), initiating the debate that is not resolved to this day. Previous studies revealed accumulation of age-related mutations in human cells (oller et al., 1989; Jones et al., 1995), mice (Dolle et al., 2000), and flies (Garcia et al., 2010). However, the molecular mechanisms involved remain unknown. One possibility is that decreased fidelity (e.g., due to protein damage) of enzymes responsible for preventing or repairing DNA damage leads to mutation accumulation and aging (Andressoo et al., 2006; Brosh & Bohr, 2007; Gorbunova et al., 2007), but this could also be due to critical mutations in these genes accumulated through the lifespan. Deficiency in these enzymes can lead to premature aging, but whether they limit lifespan under conditions of their sufficiency is unclear. Another possibility is the accumulation of oxidative damage, which decreases or abolishes the activities of repair enzymes. Oxidative stress is considered as one of the main factors in the development and progression of cancer and other age-related diseases (Ames et al., 1993).

While previous data are consistent with increased mutation load as a function of age, it remains unclear whether mutations in the DNA, the only theoretically nonrenewable molecular species within cells (the species that can be modified irreversibly), cause aging. This is because the concept that mutations cause functional decline and other aging phenotypes could not be directly tested due to inability to fully assess age-related mutations in individual somatic cells and determine their impact on organismal aging. Recently, single-cell sequencing technology has been applied to individual normal human clonal crypt, showing that chromosomal rearrangements occur in the last decades of normal human lifespan (Hsieh et al., 2013). However, it has been hard to determine if these mutations contribute to the aging process.

In this regard, the unicellular yeast, *Saccharomyces cerevisiae*, could be very useful in addressing the half-century-old debate. Yeast cells produce a certain number of daughter cells before they die, and the total number of daughter cells produced by the mother defines her replicative lifespan (Kaeblerlein et al., 2007). Significant conservation of proteins in the DNA repair pathway between yeast and higher eukaryotes further elevates the yeast as a model to study age-related genomic instability (Bitterman et al., 2003). Most importantly, the yeast genome is well characterized, and individual cells can give rise to colonies that can be sequenced, thereby establishing the genotype of the daughter. Moreover, the yeast genome is small, so the impact of its mutations can be quantified, which is more difficult to do in the case of model organisms with large genomes. Here, we subjected *S. cerevisiae* to multiple rounds of replicative aging and assessed de novo mutations in daughters of mothers of different age. The data show that DNA mutations do not cause aging in yeast.

Results and discussion

To address the role of DNA mutations in aging, we subjected *Saccharomyces cerevisiae* to replicative lifespan analyses and sequenced the genomes of clones derived from individual daughter cells of mothers of different age. First, we collected 5th (designated young) and last alive (~30th, designated old) daughter cells from each of the 4 wild-type (WT) mother cells and generated colonies from them (Figs 1 and 2A). We then used virgin cells from these colonies as mothers and again collected 5th (from the young lineage) and last (from the old lineage) daughters and made colonies from them. This procedure was repeated six times by selecting young cells from young lineages and old cells from old lineages (Figs 1 and S1), followed by high-coverage sequencing of the genomes (>99.99% completeness) of single cell-derived colonies after 2, 4, and 6 cycles of the procedure (a total of 24 genomes).

Analyses of these nuclear genomes revealed a steady increase in de novo mutations (base substitutions and indels) (Table 1) with each cycle of the procedure (Fig. 2B). After six cycles, young cells accumulated ~0.5 mutations/cell and mutations increased threefold in old cells (P< 0.03),
consistent with the previous data in flies and mice, representing both dividing (mouse) and postmitotic (fruit fly) cells (Dolle et al., 2000; Garcia et al., 2010). However, the low number of mutations observed in old yeast cells, ~1.5 mutations per six cycles or ~0.4 mutations per lifespan of an average cell, was inconsistent with the causal role of these mutations in aging. This finding also agrees with observations of yeast lifespan. For example, although daughters of very old mother cells have a shorter lifespan, the granddaughters restored lifespan to that observed in the daughters of young mothers (Kennedy et al., 1994). We also found that the lifespan of progeny of young and old cells at the end of six cycles was similar and matched that of original cells (Fig. S2). In addition, a recent report demonstrated that spores generated by aging diploid cells had the same replicative potential as those from young cells (Unal et al., 2011). Thus, while nuclear mutations do accumulate with age, they do not cause aging.

Our further analysis of the observed mutations revealed that many of them occurred on chromosome XII, where the ribosomal DNA (rDNA) repeats are located. rDNA has been identified as a factor leading to ERCs, which accumulate during replicative aging (Sinclair & Guarente, 1997). It was also shown that rDNA instability is a contributing factor to the aging process (Kobayashi, 2008) and that aging is accompanied by a significant increase in loss of heterozygosity (Lindstrom et al., 2011). In this regard, our observation of a higher mutation rate in this region is consistent with previous findings. An increased number of rDNA reads was recently shown in aged mother cells (Hu et al., 2014). We also analyzed the reads aligned to the rDNA locus in both young and old cells and found no large deletions/insertions or an increase in the number of rDNA reads during replicative aging (Fig. S3).

To assess the contribution of oxidative stress to mutation accumulation and compare it with age-dependent mutation load, we similarly analyzed a mutant strain (designated Δ8) lacking eight thioli peroxidases (Fomenko et al., 2011; Kaya et al., 2014), which are the enzymes (peroxiredoxins and glutathione peroxidases) that detoxify hydroperoxides. We further sequenced the 24 genomes corresponding to young and old Δ8 cells after 2, 4, and 6 cycles, similarly to the procedure described above for WT cells (Table 2; Figs S1 and S4). This mutant showed a 12-fold higher mutation rate than WT cells, ~6 mutations per six cycles (Fig. 2C). However, even young Δ8 cells had twice as many mutations as the old WT cells, and this situation did not correspond to the aging patterns of these cells, that is, old WT cells had few mutations, yet stopped dividing and died, whereas young Δ8 cells had many more mutations, yet they continued dividing. The overall substitution mutation rate of WT cells was in line with the known mutation rate in yeast (Lynch et al., 2001).
Do mutations in the yeast mitochondrial genome cause aging? As mitochondria are the major source of reactive oxygen species, mtDNA may be vulnerable to oxidative damage (Harman, 1956). The mitochondrial genome has a higher mutation rate than the nuclear genome (Lynch et al., 2008), and the biology of yeast mitochondria has been implicated in aging (Breitenbach et al., 2012). We found that mtDNA was consistently lost in WT cells after they generated 10–20 daughter cells, and this effect was also seen in young cells after several aging cycles, making many of these cells respiration deficient (Fig. 3; Figs S5 and S7). The observation that cells lose mtDNA during the replicative aging process leading to nonrespiring cells was also reported previously (Veatch et al., 2009; Lindstrom et al., 2011). An interesting observation was that some of these cells showed mitochondrial mutations just prior to completely losing the mitochondrial genome, explaining the loss of the mitochondrial genome during the aging progress. Although yeast cells grown on glucose do not rely on respiration and do not utilize mtDNA, mutations may disrupt integrity of mtDNA, so it may be lost. Interestingly, among the cells that we analyzed, the two old WT lines that still had functional mitochondria possessed 10 and 6 distinct noncoding substitution mutations.

In contrast, Δ8 cells had a higher mtDNA content in both old and young cells, did not lose mtDNA during aging and progression through aging cycles, and did not accumulate mutations in the mitochondrial genome (Fig. 3; Figs S6 and S7). Although in some samples we observed heterogeneous mutations (i.e., both original and mutant sequences were present) none of them persisted, that is, mutated mtDNA was lost in subsequent cycles, whereas the original mtDNA remained. Throughout the procedure, Δ8 cells were fully dependent on respiration, yet their lifespan was shorter than that of WT cells. Thus, not only mtDNA

drial genome has a higher mutation rate than the nuclear genome

### Table 1 Mutations observed in young and old WT clones following pedigree analysis

<table>
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<th>Y2</th>
<th>Y3</th>
<th>Y4</th>
<th>O1</th>
<th>O2</th>
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Mutation distributions for each pedigree lines are shown after completion of six cycles of aging. Mean mutation rates are calculated based on the total number of substitutions and small indels divided by the total number of generations for four lines. The whole list of mutations together with positions and nucleotide changes is shown in Table S1. Y is young, and O is old.

### Table 2 Mutations observed in young and old Δ8 donees following pedigree analysis

<table>
<thead>
<tr>
<th>Substitutions Position</th>
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<th>Y3</th>
<th>Y4</th>
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Mutation distributions for each pedigree lines are shown after completion of six cycles of aging. Mean mutation rates are calculated based on the total number of substitutions and small indels divided by the total number of generations for four lines. The whole list of mutations together with positions and nucleotide changes is shown in Table S2. Y is young, and O is old.

et al., 2008) (Fig. 2D). Although larger sample sizes would have allowed measuring the mutation rates more precisely, our goal was to determine the actual number of mutations in individual yeast cells per lifespan. It was clear from our analysis that both young and old WT cells had a much lower mutation rate than either young or old Δ8 cells (Fig. 2D). We also examined mutations in repetitive regions, which are thought to exhibit higher mutation rates (Lynch et al., 2008) and impact protein quality control (Nyström & Liu, 2013), but detected no such mutations during aging. Together, these findings, and the low number of mutations in old cells (<1 per lifespan), exclude a causal role of mutation accumulation in aging in yeast.

Do mutations in the yeast mitochondrial genome cause aging? As mitochondria are the major source of reactive oxygen species, mtDNA may be vulnerable to oxidative damage (Harman, 1956). The mitochondrial genome has a higher mutation rate than the nuclear genome (Lynch et al., 2008), and the biology of yeast mitochondria has been implicated in aging (Breitenbach et al., 2012). We found that mtDNA was consistently lost in WT cells after they generated 10–20 daughter cells, and this effect was also seen in young cells after several aging cycles, making many of these cells respiration deficient (Fig. 3; Figs S5 and S7). The observation that cells lose mtDNA during the replicative aging process leading to nonrespiring cells was also reported previously (Veatch et al., 2009; Lindstrom et al., 2011). An interesting observation was that some of these cells showed mitochondrial mutations just prior to completely losing the mitochondrial genome, explaining the loss of the mitochondrial genome during the aging progress. Although yeast cells grown on glucose do not rely on respiration and do not utilize mtDNA, mutations may disrupt integrity of mtDNA, so it may be lost. Interestingly, among the cells that we analyzed, the two old WT lines that still had functional mitochondria possessed 10 and 6 distinct noncoding substitution mutations.

In contrast, Δ8 cells had a higher mtDNA content in both old and young cells, did not lose mtDNA during aging and progression through aging cycles, and did not accumulate mutations in the mitochondrial genome (Fig. 3; Figs S6 and S7). Although in some samples we observed heterogeneous mutations (i.e., both original and mutant sequences were present) none of them persisted, that is, mutated mtDNA was lost in subsequent cycles, whereas the original mtDNA remained. Throughout the procedure, Δ8 cells were fully dependent on respiration, yet their lifespan was shorter than that of WT cells. Thus, not only mtDNA

![Fig. 3](image-url) Mitochondrial content is higher in Δ8 cells and does not decrease during aging. Genomes of individual clones (four young and four old at each cycle) were sequenced following 2, 4, and 6 cycles of aging. X_{nuc} coverage (mitochondrial genome) divided by X_{nuc} coverage (nuclear genome) is shown. (A) Mitochondrial genome abundance of young WT and Δ8 cells following 2 and 6 cycles. (B) Mitochondrial genome abundance of old WT and Δ8 cells following 2 and 6 cycles.
mutations, but even the presence of the functional mitochondrial genome, did not correlate with the aging process. A possibility remains that transient heterogeneous mitochondrial mutations in Δ8 cells contribute to the aging process at the level of individual mitochondria, but deleterious mutations are selected against during colony expansion. It should also be noted that although an increase in oxidative stress levels leads to increased mutation load, oxidative stress also targets numerous other molecules. It is unlikely that these conditions affect only DNA in Δ8 cells. In future studies, it would be useful to use a strain, which has a mutated proofreading domain of a nuclear DNA polymerase, thus leading specifically to increased mutation load without affecting other cellular systems.

It would also be interesting to understand why Δ8 cells rely on respiration. It is known that age asymmetry between the mother and daughter cells is affected by the end of life of mother cells (McMurray & Gottschling, 2004). Old cells pass some of the damaged molecules including aggregated proteins and dysfunctional mitochondria to their daughter cells (Henderson & Gottschling, 2008; Steinbraus et al., 2008; Kaeberlein, 2010), and daughters of old mothers live a shorter life (Kennedy et al., 1994). However, granddaughters of old mother cells are apparently able to clear the damage (McMurray & Gottschling, 2004; Kaeberlein, 2010). Previously, age-dependent deterioration in mitochondrial genome integrity was shown in mice; however, mitochondrial mutations did not limit the lifespan of these animals (Vermulst et al., 2007). In yeast, it was also shown that the absence of the mitochondrial genome does not affect the replicative lifespan in both BY4742 and YDW2 backgrounds (Kaeberlein et al., 2005; Woo & Poyton, 2009), and the replicative lifespan is increased in J4M3 and D273-108 backgrounds (Woo & Poyton, 2009).

Aging is often viewed in terms of gradual degeneration and accumulation of damage, leading to decline in fitness. We agree with this notion of aging as a gradual process driven by damage (in the broad sense of it, that is, damage not limited to by-products and errors) (Gladyshev, 2013). Universality of the aging process and the conservation of aging pathways and genetic and dietary interventions that affect lifespan point to a common basis for aging in diverse organisms. Point mutations represent an important form of cellular damage, but not the only one. For example, genome rearrangements have been shown to increase during replicative (McMurray & Gottschling, 2004) and chronological (Longo & Fabrizio, 2012) aging in yeast. However, we found no structural genome rearrangements during aging in the cells that we examined. A possibility remains that point mutations and structural rearrangements are eliminated during cell growth, that is, they accumulate in large numbers during aging, but selection acts during colony expansion of daughters of old cells eliminating them. However, we think it is highly unlikely that all mutations and rearrangements can be eliminated, returning the genome to the original state, especially considering that most age-related mutations are expected to be neutral.

A recent study analyzed populations of yeast old mother cells, in which more than ~85% of cells ceased to divide, and observed many genomic rearrangements accompanying aging, including mitochondrial DNA transfer to the nuclear genome, translocations, and retrotranspositions (Hu et al., 2014). This study cannot be directly related to our conclusion that DNA mutations do not cause aging (but together the two studies hold significant information about progression of replicative aging and the processes after terminally old mother cells stop dividing). This is because that other study analyzed enriched, purified aged mother cells at the population level, whereas we did this at the individual cell level, which was required to assess the causal role of mutations. It is possible that a significant fraction of the genomic rearrangements observed (Hu et al., 2014) occurred after the last daughter was formed, or these mutations occurred in a fraction of dying cells in the population. Thus, we cannot exclude a possibility that mutations contribute to demise of the mother cell (which is synonymous with her inability to bud off one more daughter). But, this would be equivalent to the contribution of mutations to death of an organism, as opposed to having a causal role in the aging process.

Another recent study analyzed Ty1 retrotransposon expression and mobility during chronological aging in S. paradoxus (VanHoute & Maxwell, 2014). Increased expression and mobility of retrotransposons with age have been found in nearly every aging model and thought to promote genome instability during aging (Moskalev et al., 2013). However, this study showed that, while the strain expressing Ty1 element had 40-fold increased mobility to the rDNA region, neither Ty1 mobility nor increased mutation rate of a representative Ty1 gene led to chronological lifespan decrease. Furthermore, increased expression of a Ty1 could extend chronological lifespan under certain conditions (VanHoute & Maxwell, 2014).

Replicative aging in yeast can be defined as the progression of mitotic division until an ultimate phenotype that prevents further daughter cell formation. During progression of aging, accumulation of different types of damage has been reported. If a single damage form limits lifespan, selection on other damage forms should be relaxed, leading to a certain degree of synchronization with regard to their deleterious impact. Thus, we speculate that application of our findings to other individual damage types and to other organisms suggests that there is no individual damage form that represents, when considered in isolation, a main causal factor in aging. If a damage form decreases fitness, organism can develop protective strategies to downregulate it to a milder form or modify its metabolism to decrease production of this damage. But the mild damage forms are too numerous to be protected against (Gladyshev, 2013). Thus, even if mutations and other individual damage forms, when taken in isolation, do not cause aging, aging may still result from cumulative damage, to which these damage forms contribute. From this perspective, mutations in nuclear and mitochondrial genomes, together with the myriad of other damage forms, contribute to normal aging, but can cause aging only at the level of cumulative damage.

**Experimental procedures**

**Yeast strains**

All yeast strains were in BY4741 background. A mutant strain, lacking all eight thiol peroxidases, was reported previously (Fomenko et al., 2011). Cells were grown on yeast extract-peptone-dextrose (YPD) medium.

**Pedigree analyses and mutation detection**

WT and Δ8 cells were grown at 30°C for 2 days. Several virgin mothers were placed in predetermined positions with a micromanipulator. We isolated 5th and last daughters from these mothers and allowed them to form colonies. For the 2nd round, single virgin cells from these colonies were subjected to replicative lifespan analysis, and 5th (from each young line) and last (from each old line) daughters were collected again, to obtain additional colonies. This procedure was repeated six times. At the end of the 6th cycle, DNA was isolated from each colony and paired-end sequenced on an Illumina platform (400 bp insert size, 75 bp nucleotide reads, paired end). The depth of coverage for each line was between 50X and 150X. Burrows–Wheeler Aligner (BWA) was used for alignment.
DNA mutations do not cause aging. A. Kaya et al.

Sensitivity was tested on YPD plates containing 2% ethanol and glycerol to assess growth on media containing Antimycin A. The data were further analyzed using the SHORE toolkit (Schneeberger et al., 2009), the nucleotide variants were cross-checked with the GATK output, and mutations were manually verified with Tablet (Milde et al., 2010). We also adjusted parameters to include cases in which the fraction of reads supporting an alternative allele was lowered to 0.5; however, this did not result in an increase in the number of candidate mutations. In addition, variable tandem repeats were extracted from Vinces et al., 2009 (intragenic repeats) and Verstrepen et al., 2005 (intragenic repeats) and further subjected to our analysis.

Calculation of mutation rates

Mutation rate was calculated as the number of mutations per line divided by the number of cell divisions and the length of the genome. Calculation of mutation rates was performed using the Genome Analysis Toolkit (GATK) (McKenna et al., 2010) to identify mutations covered by at least three reads and with the GATK quality score higher than 10 (P = 0.001 for misidentification). The data was further analyzed using the SHORE toolkit (Schneeberger et al., 2009), the nucleotide variants were cross-checked with the GATK output, and mutations were manually verified with Tablet (Milde et al., 2010). We also adjusted parameters to include cases in which the fraction of reads supporting an alternative allele was lowered to 0.5; however, this did not result in an increase in the number of candidate mutations. In addition, variable tandem repeats were extracted from Vinces et al., 2009 (intragenic repeats) and Verstrepen et al., 2005 (intragenic repeats) and further subjected to our analysis.

Analyses of mitochondrial mutations and the rDNA locus

For normalization, X_nuc coverage (mitochondrial genome) was divided by X_nuc coverage (nuclear genome). The same procedure was also applied to analyze the rDNA locus (X_rDNA). The total number of reads mapped to the rDNA locus was divided by X_nuc coverage and plotted. YPEG plates containing ethanol and glycerol were used to assess growth on respiratory substrates. Petite colony formation was performed as previously described (Ferguson & Von Borstel, 1992). Antimycin A sensitivity was tested on YPD plates containing 2 μg/mL of antimycin A.

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We thank Amnon Koren for the help with mutation rate analyses, and Daniel Promislow and Matt Kaeberlein for suggestions.

Author contributions

Conceived and designed the experiments: AK, VNG. Performed the experiments: AK. Analyzed the data: AK, AVL, VNG. Wrote the paper: AK, VNG.

Funding

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Conflicts of interest

The authors declare no conflict of interest.

References


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### Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.

**Fig. S1** Age at which daughter cells of young and old WT mothers were taken for further analysis.

**Fig. S2** Lifespan of clones derived from young and old mother cells is not affected by the aging process.

**Fig. S3** Comparison of the reads mapped to the rDNA region of WT young and old lines.

**Fig. S4** Age at which daughter cells of young and old Δ8 mothers were taken for further analysis.

**Fig. S5** Depletion of mitochondrial DNA and respiratory function in WT lines during aging.

**Fig. S6** No depletion of mitochondrial DNA in Δ8 lines during aging.

**Fig. S7** Induction of petite colony formation.

**Table S1** Mutations identified in WT cells during aging.

**Table S2** Mutations identified in Δ8 cells during aging.