

Genetically controlled mtDNA editing prevents ROS damage by arresting oxidative phosphorylation

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Abstract: Deletion of mitochondrial DNA in eukaryotes is mainly attributed to rare accidental events associated with mitochondrial replication or repair of double-strand breaks. We report the discovery that yeast cells arrest harmful intramitochondrial superoxide production by shutting down respiration through genetically controlled deletion of mitochondrial oxidative phosphorylation genes. We show that the regulatory circuitry underlying this editing critically involves the antioxidant enzyme superoxide dismutase 2 and two-way mitochondrial-nuclear communication. While mitochondrial DNA homeostasis is rapidly restored after cessation of a short-term superoxide stress, long-term stress causes maladaptive persistence of the deletion process, leading to complete annihilation of the cellular pool of intact mitochondrial genomes and irrevocable loss of respiratory ability. Our results may therefore be of etiological as well as therapeutic importance with regard to age-related mitochondrial impairment and disease.

One-Sentence Summary: Genetically controlled editing of mitochondrial DNA is an integral part of the yeast's defenses against oxidative damage.

Main Text: Mitochondrial impairment is strongly associated with aging (1) and the pathogenesis of age-related human diseases, including Alzheimer's disease (2), Parkinson's disease (3), the deterioration of skeletal and cardiac muscle (4), and macular degeneration (5). Mitochondrial DNA (mtDNA) deletions are perceived to contribute markedly to this impairment (6). The general conception is that mtDNA deletions are caused by accidental events associated with replication or repair of double-strand breaks (7). However, considering that mtDNA deletions are prone to cripple oxidative phosphorylation (8) (OXPHOS) it is conceivable that mtDNA deletion is also under genetic control, the main rationale being that compared to a mitophagic response (9–13), it would serve as a less costly defense mechanism against an abrupt increase in intramitochondrial superoxide production inundating the primary antioxidant defenses (14–16). The disclosure of an additional genetically controlled defense layer against superoxide damage, situated between the primary antioxidant defenses and mitophagy, would bring a fresh perspective to what causes mitochondrial impairment and how it can be mitigated. This motivated us to search for the existence of such a regulatory layer in budding yeast (*Saccharomyces cerevisiae*).

Swift adaptation to enhanced superoxide production

To avoid possible confounding effects of domestication (17), we exposed wild haploid yeast cell populations expanding clonally on glucose to the mitochondrial superoxide anion ($O_2^{\bullet-}$) generator and redox cyler paraquat (N,N-dimethyl-4,4'-bipyridinium dichloride) (18). As OXPHOS complex III and mitochondrial NADPH dehydrogenases donate electrons to paraquat, which passes these on to O_2 (18, 19), this mode of $O_2^{\bullet-}$ generation is a good proxy for the *in vivo* situation (20). By titration of the paraquat dose, we ensured that the $O_2^{\bullet-}$ production was beyond

the regulatory reach of the primary antioxidant defenses, while not severely compromising cellular function (fig. S1, supplementary text 1).

We then observed how 96 clonally reproducing yeast cell populations on solid agar medium adapted to the chosen paraquat dose in terms of change in cell doubling time over approximately 240 generations (corresponding to 50 growth cycles). To provide a comparative data set, we similarly exposed a total of 672 populations to seven other stressors not explicitly challenging mitochondrial function (table S1). All 96 populations exposed to paraquat adapted much faster than every other population exposed to the other stressors (Fig. 1A). Within the first 10 generations, cell populations reduced their doubling time by on average 106 min, corresponding to 49.3% of the maximum possible reduction. Thenceforth, adaptation entered a second phase where the reduction in cell doubling time progressed much slower until it plateaued after 75 generations at 72.6% of the maximum possible reduction (mean) (Fig. 1A).

A numerical model of the adaptation process, combining population genetics and population dynamics concepts with experimental data on nuclear mutation rates and effect sizes of *de novo* point mutations and aneuploidies (21), was completely unable to reproduce the observed extraordinarily swift response to paraquat (fig. S2).

After being exposed to paraquat over a few growth cycles before being placed in a paraquat-free medium, all 96 cell populations retained their acquired tolerance to paraquat (mean reduction in cell doubling time: 106 min) for only 1-2 growth cycles before abruptly losing it (Fig. 1B). When employing the same experimental procedure to 96 cell populations from each of the two other environments to which adaptation was also fast (arsenic and glycine), we found that despite the

presence of a much stronger Darwinian counterselection (Fig. 1C), these populations lost their acquired adaptations more slowly and gradually (Fig. 1B). While a classical Darwinian mutation/selection-based adaptive process could reasonably explain the data for seven of the eight tested stressors, we concluded that the data on paraquat could hardly be reconciled with such a process.

Mitophagy is not responsible for the swift first adaptation phase

We next probed whether a mitophagic response was responsible for the swiftness of the first adaptation phase. As mitochondrial fragmentation is a well-documented prelude to canonical mitophagy (22), we assayed mitochondrial morphology before and after paraquat exposure by confocal and electron microscopy. We observed a rapid shift (<5 h) from a tubular to a fragmented mitochondrial organization (Figs. 2A, B, fig. S3A) as well as a rapid reversal (<5 h) to a tubular organization after removal of stress, consistent with the notion that mitochondrial $O_2^{\bullet-}$ generation influences the mitochondrial fission and fusion dynamics (23, 24). DNA staining indicated that the mitochondrial fragments generally contained mtDNA after 5 h of paraquat exposure (fig. S3B). The mitochondrial volume remained near pre-stress levels with at the most a marginal reduction after 77-79 h of paraquat exposure (22) (Figs. 2A, B). Moreover, cell populations ($n=16$) lacking Atg32, a key component of canonical mitophagy (25), adapted to paraquat over ~80 generations as wild type populations (Fig. 2C). These observations led us to conclude that the initial swift adaptation to paraquat was not likely to depend on canonical mitophagy.

mtDNA loss drives the first adaptation phase

We then assayed the copy numbers of mtDNA genes in nine paraquat-adapting cell populations by qPCR. In line with short read sequencing results from five populations (Fig. 3A, supplementary text 2), we found that cells lost parts of their mtDNA during the early adaptation phase. In addition, the qPCR data showed that the lost segments were unevenly distributed across the mitochondrial genome: all nine cell populations lost one or more segments within the mtDNA region spanning *COX1* to *VARI*, while a few also lost the 21S rRNA and *COX2* rapidly thereafter (Fig. 3B, fig. S4). Cells retained the mtDNA segments that were not lost at near founder levels throughout the early adaptation phase, which in all nine cases encompassed *COX3-RPM1* and 15S rRNA. Since the mtDNA coverage prior to paraquat adaptation was perfectly even (fig. S5A), the observed mtDNA loss was clearly induced by paraquat.

Anticipating that the observed mtDNA loss caused a reduction in OXPHOS activity, and thus in mitochondrial $O_2^{\bullet-}$ production, we exposed ρ^0 cell populations ($n=12$) (supplementary text 3), obtained by deletion of the sole mitochondrial DNA polymerase Mip1 (Pol γ homolog) (26), to the same concentration of paraquat as was used in the initial adaptation experiments. These *mip1 Δ* cells were highly resistant to paraquat compared to wild type cells (Fig. 3C). Moreover, exposing the pre-adapted *mip1 Δ* cell populations to paraquat for 10 growth cycles caused no or marginal additional improvement in growth (fig. S5B). To substantiate these findings, we cultivated cell populations exposed to short-term paraquat stress on the respiratory growth medium glycerol. The early adaptation correlated almost perfectly with loss of the capacity for respiratory growth (Fig. 3D), and the rapid loss of the acquired adaptations after removal of stress coincided with the restoration of the ability for respiratory growth (fig. S5C). We next repeated the stress release experiment on the five sequenced populations, and tracked by short-read sequencing the temporal change in copy numbers of the lost mtDNA segments. As

expected, the five populations quickly restored their ability for respiratory growth after removal of paraquat, and this coincided with the restoration of mtDNA and loss of the acquired adaptation (Fig. 3E, figs. S5C, D).

5 The above observations strongly suggest that the loss of OXPHOS activity was the predominant mechanism responsible for the swiftness of the first adaptation phase. Considering that 68% of the realized adaptation was achieved during this phase, the major fraction of paraquat-induced $O_2^{\bullet-}$ production is therefore arguably associated with the OXPHOS system (19).

10 **The deletion of mtDNA segments requires *SOD2***

Based on experiments that led us to conclude that hydrogen peroxide (H_2O_2) or its downstream degradation products did not play a prominent causative role in inducing the deletion of mtDNA segments (figs. S6A, B, supplementary text 4), we hypothesized that mitochondrial $O_2^{\bullet-}$ was directly responsible for this. The superoxide dismutases Sod1 (Cu/ZnSOD) and Sod2 (MnSOD),
15 located in the cytosol and mitochondrial intermembrane space and in the mitochondrial matrix, respectively, have both previously been linked to signaling responses associated with $O_2^{\bullet-}$ (20, 27). We therefore probed how *sod1Δ* and *sod2Δ* cells responded to paraquat compared to the wild type. While the *sod1Δ* populations adapted as wild type cells, the *sod2Δ* populations mostly displayed no reduction in cell doubling time over 10 growth cycles when exposed to the same
20 stress levels (Figs. 4A, B, fig. S6C, supplementary text 5).

Using qPCR, we then tracked the pattern of mtDNA deletions in eight *sod2Δ* populations exposed to paraquat. In four of the populations, the copy numbers of all mtDNA genes were retained at or near pre-stress level (Fig. 4B). Two of these failed to adapt to paraquat, while the

other two showed a weak and a much-delayed reduction in cell doubling time relative to the wild type, indicating selection for small-effect mutations not associated with mtDNA deletion. The remaining four populations showed a much-delayed reduction in cell doubling time that coincided with the acquisition of a single mtDNA deletion (fig. S6D). To clarify the importance of mtDNA deletions in relation to the Sod2-dependency, we deleted Mip1 in a *sod2Δ* background, and found that the induced loss of mtDNA increased growth as in wild type cells (Fig. 4C). This suggests that the delayed adaptation to intramitochondrial O₂^{•-} stress in cells lacking Sod2 is because the mtDNA deletions emerge at a much lower rate. Taken together, the above results strongly suggest that Sod2 is a key factor in the causal chain of events leading to the mtDNA deletions underlying the initial rapid adaptation to paraquat. As the data clearly show that the deletion of mtDNA segments causing loss of OXPHOS activity is under regulatory control, we think it is apt to denote this process as ‘mtDNA editing’ (28).

The mtDNA editing process requires anterograde mito-nuclear communication

The above results led us to suspect that the underlying regulatory circuitry might involve mitochondrial-nuclear communication. In budding yeast, deletion of mtDNA alters the expression of a multitude of genes resulting in increased glycolytic production of ATP (29), dubbed the retrograde response. Upon mitochondrial OXPHOS dysfunction, the cytosolic protein Rtg2 (30) causes the transcription factors Rtg1 and Rtg3 to translocate from the cytosol to the nucleus where they together activate retrograde transcription (31, 32). As this mechanism is the most well-documented channel in yeast for communicating mitochondrial dysfunction to the nucleus, and in particular mtDNA deletion (33), we exposed *rtg2Δ* and *rtg3Δ* cell populations (*n*=16) to paraquat for 80 generations. In both cases, all cell populations failed to adapt (Fig.

5A), and their capacity for respiratory growth was virtually unperturbed at the end of the experiment (Fig. 5B). This indicates that the two proteins induce anterograde mito-nuclear communication critical for activation of the mtDNA editing process.

5 Sustained mtDNA deletion causes irrevocable mitochondrial impairment

We found that 44 of the 96 populations had completely lost their capacity for restoring the pool of intact mtDNA genomes back to pre-stress levels after 24 generations of paraquat exposure, and after 242 generations all did (fig. S7). Moreover, in the five sequenced cell populations, the capacity to restore the copy numbers of intact mtDNA and the respiratory growth after removal
10 of stress was lost between 15 to 42 generations of paraquat exposure (Fig. 6A). This suggest that the deletion of mtDNA genes continued after the first adaptation phase in a minor fraction of cells without providing a recognizable further reduction in cell doubling time.

The sustention of mtDNA deletions led us to suspect that a long-term $O_2^{\bullet-}$ stress would finally
15 lead to a complete loss of intact mtDNA genomes. We therefore assayed the mtDNA loss in 44 sequenced endpoint (t_{50}) populations. Out of these, 25 populations had almost completely lost their entire 77 kb mtDNA (97-99%) (fig S8). 18 endpoint populations remained in a rho negative (ρ^-) state where they still possessed 6 to 34 kb mtDNA segments with copy numbers somewhat above the original founder levels (mean: 20% increase). To assess whether there was an adaptive
20 advantage associated with the sustained depletion of mtDNA, we compared the doubling times of the ρ^- populations still retaining 6 to 34 kb mtDNA segments with those that had lost almost all their mtDNA. Intriguingly, the latter group consistently grew slower on paraquat than the former (Fig. 6D). This suggests that the sustained mtDNA depletion under long-term stress, leading to irrevocable loss of intact mtDNA genomes, is a maladaptive response driven by

prolonged induction of a regulatory mechanism dimensioned by natural selection to handle $O_2^{\bullet-}$ stress it can successfully deal with before the pool of intact mtDNA genomes becomes annihilated.

5 **Chromosome duplications explain the second adaptation phase**

mip1Δ cells lacking mtDNA were not fully tolerant to paraquat (Fig. S9A), consistent with that wild type cells realized 23.3% of their adaptation potential in a second adaptation phase (Fig. 1A). The much slower doubling time reduction characterizing this second phase suggested that in this case the adaptation to on- or off-target effects of paraquat could be explained by a Darwinian mutation/selection process (fig. S2). Analysis of the sequence data of 44 random endpoint (t_{50}) populations led us to conclude that nuclear point mutations were unlikely to have played a prominent causative role, though (figs. S9B-C). However, duplications of chromosome II, III and V were common, and we found that these were able to explain the reduction in cell doubling time during the second adaptation phase (fig S10 and supplementary text 6).

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Discussion

The adaptation patterns of the *sod2Δ*, *mip1Δ*, *rtg2Δ* and *rtg3Δ* populations relative to the wild type are hard to reconcile with the operation of random $O_2^{\bullet-}$ induced mtDNA deletions (e.g. caused by genomic instability) and subsequent selection of cells possessing OXPHOS-impaired mitochondria. However, the specificity of the mtDNA deletion mechanism in terms of which mtDNA segments are lost in the first adaptation phase does seem to be moderate. Still, there was a clear preference for deletions within the *COX1-VARI* region, and the segment containing *COX3*, *RPM1* and 15S RNA were not deleted in this phase. The fact that copy number homeostasis was observed not just for these three genes, but also for genes in undeleted segments

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within the *COXI-VARI* region, indicates that partial deletion of genes within this region was sufficient for providing the marked reduction in cell doubling time during the first adaptation phase. This interpretation makes sense from an evolutionary point of view: natural selection would not be able to increase the specificity of the mtDNA editing program beyond the point where no further adaptation is achieved. However, the data suggest that even the moderate specificity characterizing the early adaptation phase is lost by sustained operation of the mtDNA deletion mechanism during long-term oxidative stress.

The restoration of intact mtDNA genomes back to pre-stress levels after release from paraquat during the first adaptation phase (Fig. 3E) is almost complete after just 2-3 growth cycles despite that there is no apparent selective advantage attached (Figs. 1B, C). This signifies that it is under strict regulatory control. A possible mechanism is selective fusion of mitochondria with intact mtDNA genomes (34, 35) during the swift return to tubular morphology after cessation of stress (Fig. 2B), and subsequent removal of mitochondria with OXPHOS deprived genomes by mitophagy. Such selective fusion may also be an integral part of the fission/fusion dynamics such that cells achieve the clearance of non-intact mtDNA genomes only after several fission/fusion cycles. In this case, one would expect the restoration to be under the control of the constituent homeostatic regulatory machinery. If the associated feedback control mechanism is oblivious to non-intact mtDNA genomes (36), and the restoration is faster than the removal of the OXPHOS-deprived genomes, one would expect a temporal overshoot of mtDNA copy numbers before a return to prestress levels. Our data are in clear agreement with this notion (fig. S5D).

Besides providing extensive support to the conception that the disclosed mtDNA-editing system defines a regulatory layer between the primary antioxidant defenses and canonical mitophagy for coping with intramitochondrial supraphysiological $O_2^{\bullet-}$ challenges, our data bring fresh perspectives to the table concerning (i) the relationship between stress-induced mitochondrial fragmentation and canonical mitophagy (37), (ii) under which conditions do mitochondria deprived of OXPHOS genes produce more $O_2^{\bullet-}$ due to increased electron leakage (38, 39), (iii) whether clonal expansion is the main mechanism underlying the propagation of mitochondria containing deletions (40), and (iv) under which conditions, and to which degree, does the main retrograde response to mtDNA deletion in yeast mediate two-way mito-nuclear communication (33). They also support the emerging notion that selective mitophagy is an important mechanism for local mitochondrial repair (13), and that selective mitophagy may be deliberately repressed while the cell experiences $O_2^{\bullet-}$ stress.

Available data on the cancer therapeutics doxorubicin and cisplatin suggest that our main results may extend to post-mitotic cells. They both accumulate in mitochondria (41, 42), where they induce $O_2^{\bullet-}$ production through redox cycling (43, 44). Their long-term administration cause oxidative injury to a variety of cells and tissues (44–47), and they both increase the frequency of mtDNA deletions (42, 48). Arrestment of OXPHOS activity by mtDNA editing, followed by homeostatic restoration of the pool of intact mitochondrial genomes, may thus be a generic eukaryotic adaptation to obviate a more costly mitophagic response in a variety of situations. If so, the biomedical implications in terms of the etiology of age-related disease and therapeutic opportunities appear to be noteworthy.

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Data and materials availability: Sequence data that support the findings of this study have been
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e0478912a209](https://data.mendeley.com/datasets/mvx7t7rw2d/draft?a=95381e47-dc80-47af-85ab-e0478912a209). All unique strains and stored populations generated in this study are available
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20 **Supplementary Materials**

Materials and Methods

Supplementary Text S1 to S6

Figs. S1 to S10

Table S1

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Data S1 to S30

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Figures and figure captions

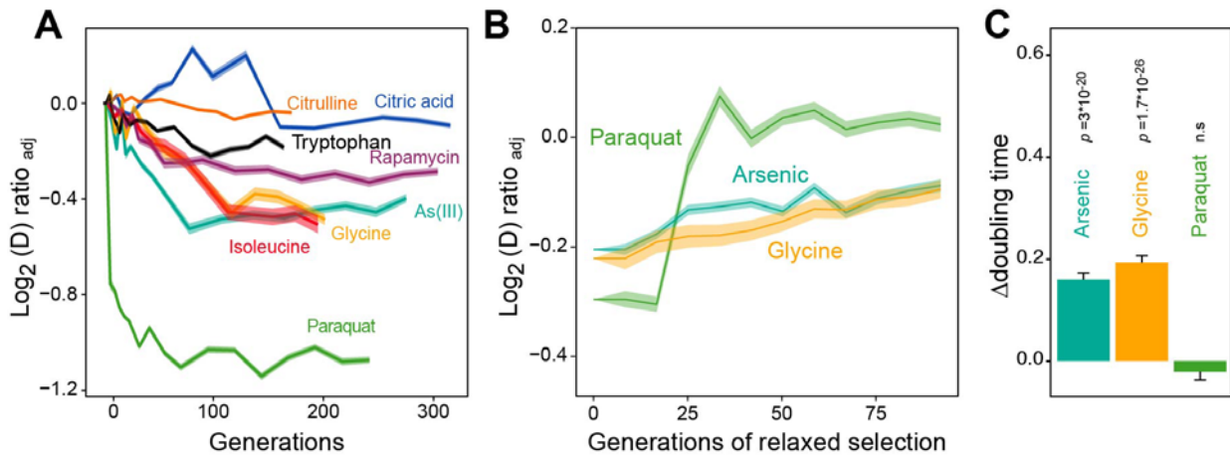


Figure 1. Distinct adaptation to paraquat.

5 (A) Mean temporal adaptive response to paraquat and seven other stressors. y-axis shows \log_2 fold reduction in cell doubling time (h) from pre-stress, adjusting for plate, position and pre-culture effects. 96 populations for each stressor ($n=6$). Shade: S.E.M. (B) Loss of the acquired adaptation as a function of number of cell generations after release from the selection pressure. Colored lines: mean of 96 populations (each measured at $n=5$). Shade: S.E.M. The populations were released from stress after reaching 70-90% of their endpoint (t_{50}) adaptation. (C) The difference in cell doubling time (h) in a no-stress environment between 96 populations (each measured at $n=5$) having achieved 70-90% of their endpoint adaptation to paraquat, arsenic and glycine, respectively, and the founder population. The difference reflects the selective advantage of losing the acquired adaptations when the populations are no longer exposed to stress. Error

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15 bars: S.E.M.

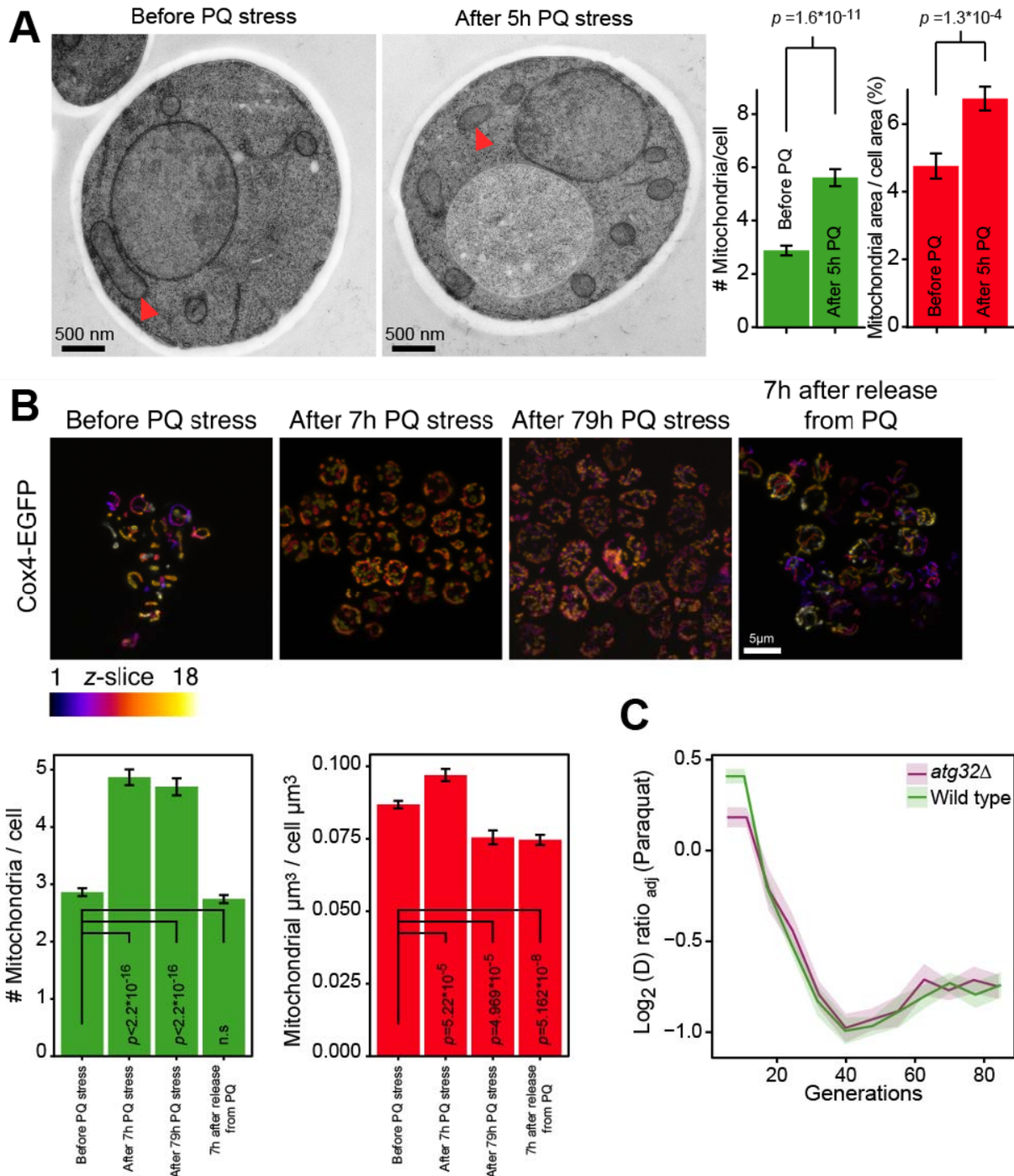


Figure 2. Mitochondrial fragmentation precedes the swift adaptation to paraquat (PQ).

(A) EM microscopy of cells before (panel 1) and after short-term stress (panel 2, red arrowheads mark representative mitochondria). Panel 3 shows the number of mitochondria per imaged cell (left) and the imaged cell area occupied by mitochondrial area (%) (right), used as proxy for

mitochondrial volume. Error bars: S.E.M. ($n=100$ cells). p -values: Welch two-sided t-test. **(B)**

Confocal microscopy of cells with a Cox4-GFP mitochondrial tag. Color: z -dimension

(yellow=front, purple=back; 18 slices). *Lower left diagram*: Number of mitochondria per imaged

cell. *Lower right diagram*: Mean sum of mitochondrial volume as a fraction of cell volume.

5 Error bars: S.E.M. ($n=473-910$ cells). p -values: Welch two-sided t-test **(C)** Adaptation of *atg32Δ*

and wild type populations to paraquat. Shade: S.E.M. ($n=16$ populations, each measured at $n=6$).

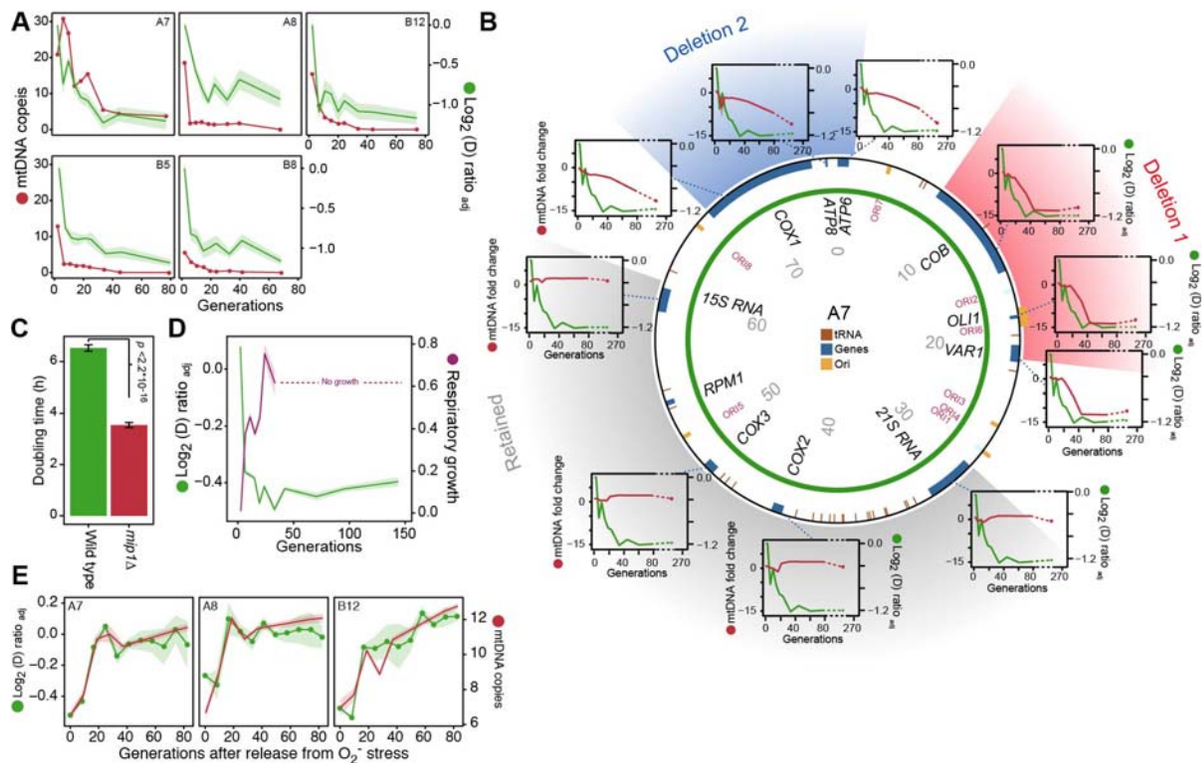


Figure 3. mtDNA editing causes the early adaptation to paraquat.

(A) mtDNA copy number change (left y-axis, red line, median coverage relative to the haploid nuclear genome) during paraquat adaptation (right y-axis, green line, $n=6$) for 5 populations (panels). Shade: S.E.M. (B) mtDNA deletions associate with the paraquat adaptation. *Circle*: mtDNA (77 kb) before exposure to paraquat. Genes, origins of replication and position (kb) are indicated. *Coloured fields*: mtDNA deletions with concerted copy number change. *Diagrams*: mtDNA copy number change (left y-axis, purple line) of individual mtDNA genes during adaptation (right y-axis, green line) in population A7. Shade: S.E.M. ($n=3$). (C) Doubling time (h) of wild type and *mip1Δ* cell populations in paraquat. Error bars: S.E.M. ($n=191$). p -values: Welch two-sided t-test. (D) The early-phase paraquat adaptation coincides with the loss of respiratory (glycerol) growth. Shade: S.E.M. 96 populations, each measured at $n=5$. Broken line indicates no growth (cell doubling time $>24h$). (E) Recovery of the copy number of deleted

mtDNA (right y-axis, red line) after release from 6 generations of paraquat exposure coincides with loss of the early-phase paraquat adaptation (left y-axis, green line, shade = S.E.M ($n=15$)) in populations A7, A8 and B12.

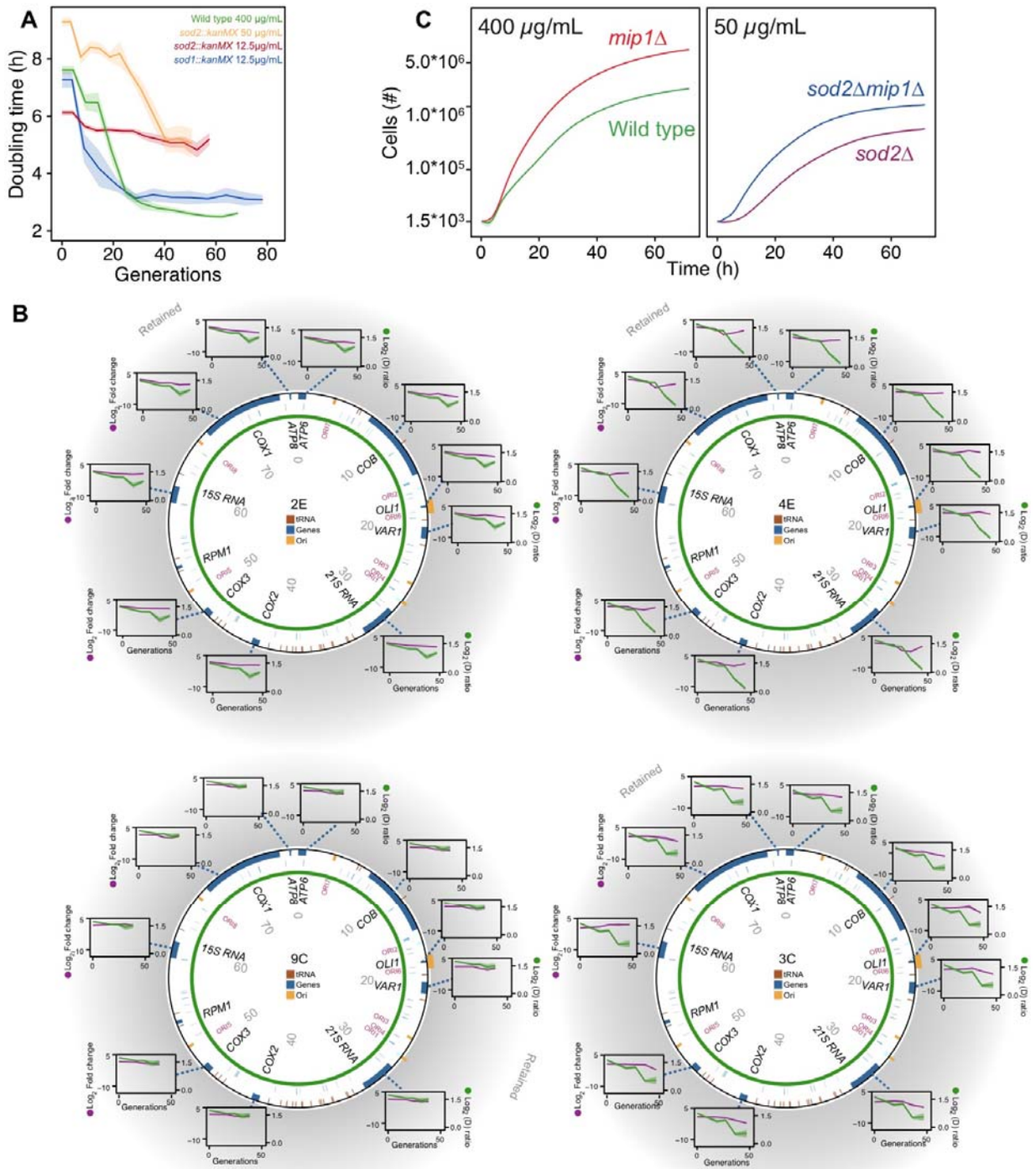


Figure 4. The mtDNA editing critically involves Sod2.

(A) Doubling time (*h*) in wild type (400 $\mu\text{g/mL}$ paraquat: green), *sod2* Δ (12.5 $\mu\text{g/mL}$ paraquat:

5

red; 50 $\mu\text{g/mL}$ paraquat: yellow) and *sod1* Δ (12.5 $\mu\text{g/mL}$ paraquat: blue) cell populations

adapting to equivalent stress levels. **(B)** mtDNA change in *sod2Δ* cell populations 2E, 4E, 9C and 3C adapting to 12.5 μg/mL paraquat. *Circle*: mtDNA (77 kb) before exposure to paraquat. Genes, origins of replication and nucleotide positions (kb) are indicated. *Colored fields*: mtDNA deletions with concerted copy number change. *Diagrams*: mtDNA copy number change (left y-axis, purple line, ($n=2$) of individual mtDNA genes during the adaptation (right y-axis, green line). Shade: S.E.M. **(C)** Mean growth of wild type ($n=480$; green), *mip1Δ* ($n=288$; red), *sod2Δ* ($n=96$; purple) and *sod2Δmip1Δ* ($n=384$; blue) cell populations in the presence of 400 μg/mL (left) and 50 μg/mL (right) paraquat.

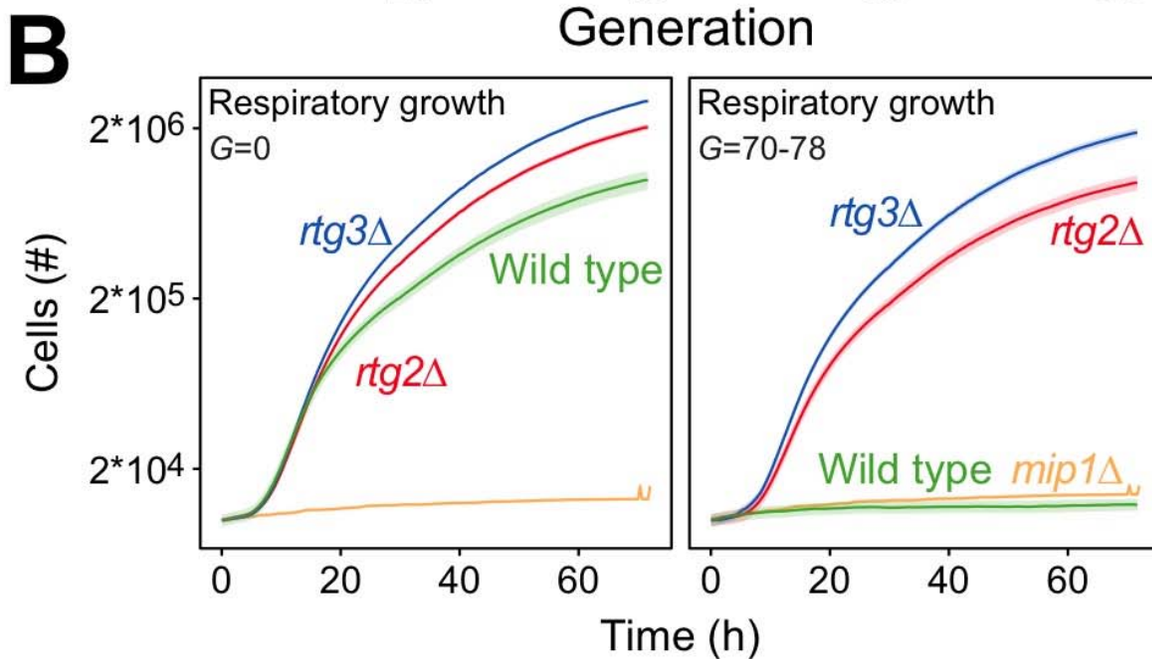
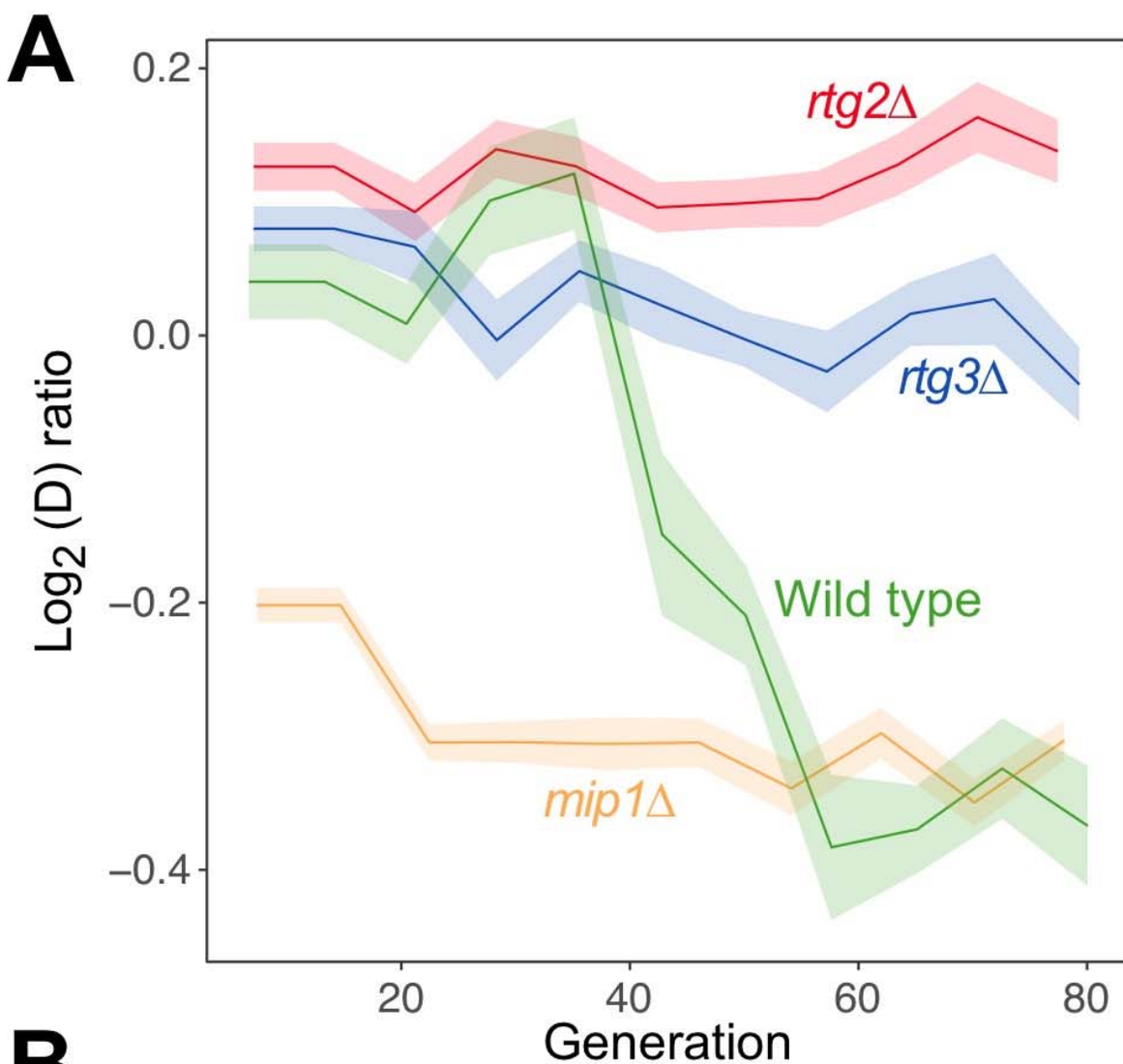


Figure 5. The mtDNA editing critically involves anterograde mito-nuclear communication.

(A) Doubling time adaptation of 18 wild type, *rtg2Δ*, *rtg3Δ* and *mip1Δ* cell populations to 400 $\mu\text{g/mL}$ of paraquat. Shade: S.E.M. Each population type measured at $n=4$. (B) Respiratory (glycerol) growth of wild type, *rtg2Δ*, *rtg3Δ* and *mip1Δ* cell populations, before (left) and after (right) 70-78 generations of paraquat adaptation. Shade: S.E.M ($n=72-144$ populations, each measured at $n=1$).

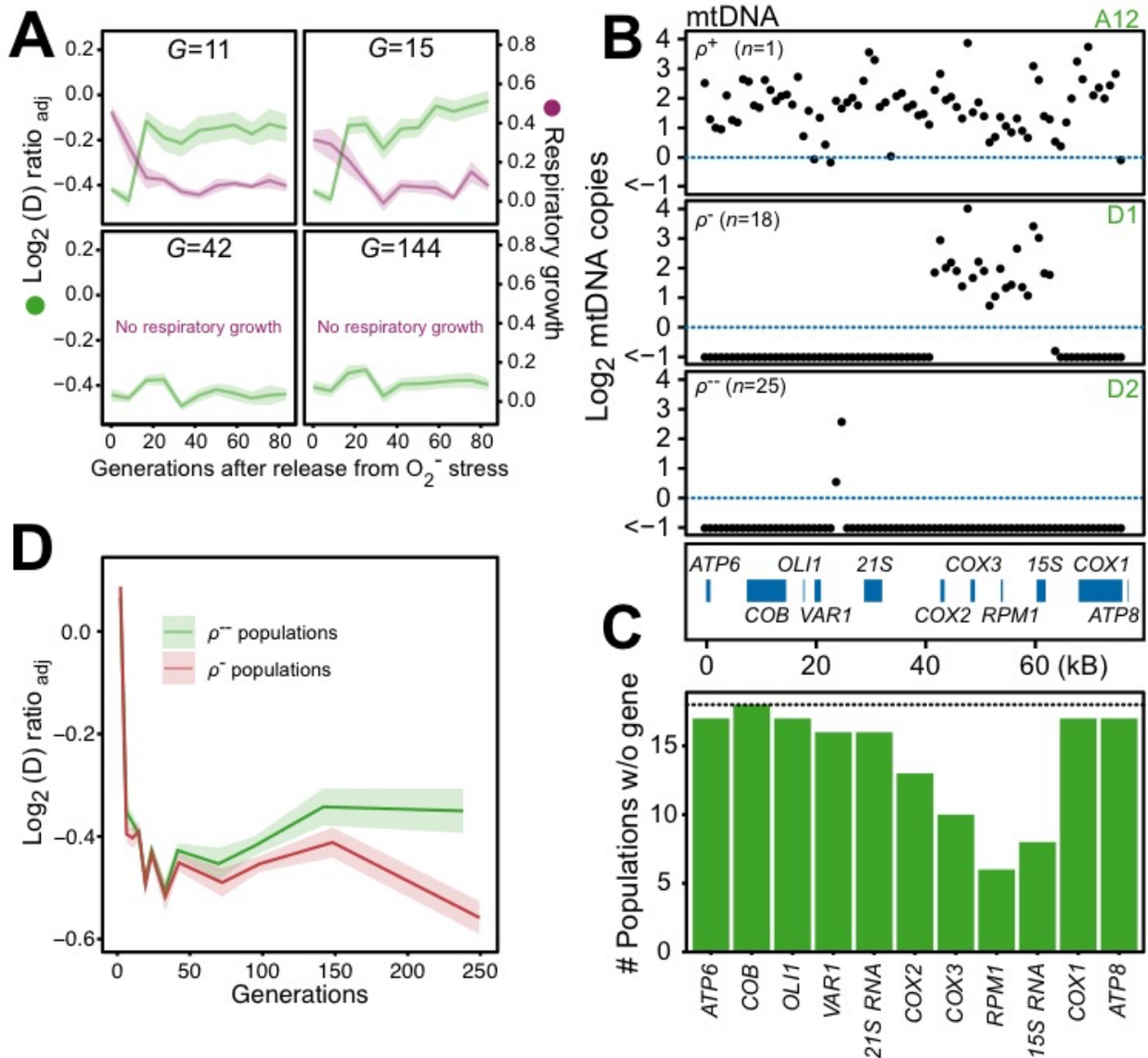


Figure 6. Chronic exposure to paraquat causes irreversible mitochondrial impairment by sustained mtDNA editing.

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(A) Paraquat adapting cell populations (G = generations of exposure to paraquat) ultimately lose their capacity to recover respiratory (glycerol) growth (right y-axis, purple line, log₂ doubling time relative to founder) and the loss coincides with the genetic fixation of the paraquat adaptation (left y-axis, green line). Shade: S.E.M. of 5 populations, each measured at $n=5$. (B)

All but one (ρ^+) sequenced cell population adapted to long-term paraquat stress (t_{50}) retain only small (6-30 kb; ρ^-) or very small (<2 kb, ρ^{--}) mtDNA segments. *Panels:* Representative populations. *y-axis:* mtDNA copy number (median coverage in 1 kb windows relative to haploid nuclear genome). Gene positions are indicated. **(C)** Number of ρ^- populations after long-term paraquat exposure (t_{50}) in which the specified mtDNA gene was lost. **(D)** The ρ^{--} populations became less fit than the ρ^- populations during a long-term exposure to paraquat.