Relationship between oxidative stress and lifespan in *Daphnia pulex*

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Macromolecular damage leading to cell, tissue and ultimately organ dysfunction is a major contributor to aging. Intracellular reactive oxygen species (ROS) resulting from normal metabolism cause most damage to macromolecules and the mitochondria play a central role in this process as they are the principle source of ROS. The relationship between naturally occurring variations in the mitochondrial (MT) genomes leading to correspondingly less or more ROS and macromolecular damage that changes the rate of aging associated organisal decline remains relatively unexplored. MT complex I, a component of the electron transport chain (ETC), is a key source of ROS and the NADH dehydrogenase subunit 5 (ND5) is a highly conserved core protein of the subunits that constitute the backbone of complex I. Using *Daphnia* as a model organism, we explored if the naturally occurring sequence variations in ND5 correlate with a short or long lifespan. Our results indicate that the short-lived clones have ND5 variants that correlate with reduced complex I activity, increased oxidative damage, and heightened expression of ROS scavenger enzymes. *Daphnia* offers a unique opportunity to investigate the association between inherited variations in components of complex I and ROS generation which affects the rate of aging and lifespan.

**Abbreviations**

| Term        | Definition                                         |
|-------------|----------------------------------------------------|
| ROS         | Reactive oxygen species                            |
| MT          | Mitochondrial                                     |
| ETC         | Electron transport chain                           |
| ND5         | NADH dehydrogenase subunit 5                       |
| mtDNA       | Mitochondrial DNA                                  |
| LHON        | Leber’s hereditary optic neuropathy                |
| MELAS       | Mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes |
| ORF         | Open reading frame                                |
| BN          | Blue Native                                       |
| CN          | Clear Native                                       |
| SOD         | Superoxide dismutase                               |
| CAT         | Catalase                                           |

Understanding the molecular basis of aging is a priority as age is the single most important risk factor for human pathologies such as cancer, diabetes, cardiovascular disorders, and neurodegenerative diseases¹. Accumulating evidence supports that aging starts with macromolecular damage leading to cell, tissue and ultimately organ dysfunction². This intrinsic process can be viewed as an inevitable outcome of the cellular biochemistry, and forms a basis from which the diseases of older age originate. The cellular product that causes the most damage is the intracellular reactive oxygen species (ROS)³. Mitochondria play a central role in regulating the rate of macromolecular damage, as they are the principle source of ROS⁴–⁶. So far, the contribution of mitochondria to aging has been studied with a view that mitochondrial DNA (mtDNA) is the primary target of the ROS generated by mitochondria. Thus, somatic mtDNA mutations are reported in normal aging, particularly in post-mitotic tissues such as skeletal muscle and neurons, leading to a corresponding decline in the mitochondrial function with age⁷–⁹. Although this decline leads to further ROS production, cellular damage, and aging associated cellular dysfunction¹⁰; such observations do not show a causal relationship between inherited variations in mitochondrial efficiencies and variations of aging. The causal relationship between naturally occurring variations in the mitochondrial genomes leading to differences in the mitochondrial electron transport chain (ETC) efficiencies

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that could result in correspondingly less or more ROS and macromolecular damage to change the rate of aging associated organismal decline has remained relatively unexplored.

The mitochondrial ETC is comprised of 5 multi-subunit complexes (complex I- complex V), the last one being the ATP synthase. Of the complexes 1–5 of the ETC, complex I (NADH:quinone oxidoreductase) is the first and largest enzyme, the major electron entry point for the respiratory chain, and is a key source of ROS in mitochondria. Mutations in complex I subunits are implicated in human diseases and aging. Of the 44 subunits (human) of the complex I, 38 are encoded by the nuclear genome and 7 by the mtDNA. The 7 mtDNA-encoded hydrophobic subunits, NADH dehydrogenase (ND) 1–6 and 4L (ND1-ND6 and ND4L), are the core subunits that form the major membrane arm of complex I. Mutations in ND subunits disrupt either complex I assembly or enzyme activity and cause ~20% of cases of isolated complex I deficiency. In humans, variations in ND5 sequence are associated with mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (MELAS), as well as some symptoms of Leigh’s syndrome, and Leber’s hereditary optic neuropathy (LHON). Many fatal pathological conditions such as Leigh syndrome, leukoencephalopathy, LHON, mitochondrial myopathy, encephalopathy, MELAS, and Parkinson’s disease, and cancer are associated with mutations in the ND genes including the ND5 gene. ND5 mutations cause mitochondrial dysfunction due to a decrease in complex I efficiency and increased ROS generation. However, very few studies have been done to relate naturally occurring, non-pathogenic ND5 sequence variations to ETC efficiencies, and ROS generation that can affect rate of aging, senescence, and life span.

In this study, we identified naturally occurring ND5 sequence variations in ecologically distinct populations of the freshwater microcrustaceans Daphnia pulex and D. puleicaria and examined its association with complex I enzyme activity and level of oxidative damage to total and mitochondrial proteins. D. pulex and D. puleicaria are short- and long-lived ecotypes of the Daphnia pulex species complex and interbreed frequently. Although zooplankton taxonomists use two different species names for the ecotypes, they are not genetically distinct species in the traditional sense of reproductive isolation. In fact, there is substantial gene flow between the ecotypes, and thus persistent differences are likely to be associated with adaptive divergence between ecotypes, rather than true species differences that have accumulated due to non-adaptive evolutionary processes. For simplicity and to emphasize the lack of reproductive isolation, we refer to them as D. pulex throughout the manuscript. We also investigated the activities of ROS scavenging antioxidant mechanisms in the short- and long-lived Daphnia. Our results indicate that lower activity of complex I possibly leading to higher levels of ROS leads to correspondingly high levels of oxidative damage to proteins. Although there is a corresponding increase in activities of ROS scavenging antioxidant mechanisms, the short-lived clones show an imbalance between ROS production and removal thus accumulating damaged proteins and possibly contributing to their shorter lifespan. The results are important because they demonstrate that oxidative stress, which has been shown to be a proximate driver of lifespan and aging in other model organisms, also correlates similarly in Daphnia, a newer model system for research on aging. Our study demonstrates that Daphnia is an excellent model system that offers significant advantages with its unique clonal reproduction, availability of genome manipulation techniques, and naturally existing ecotypes with varied lifespans to understand natural determinants of lifespan.

**Results**

**Genetic variation of lifespan and ND5 sequence variations in D. pulex clones.** The median lifespans in 11 clones of D. pulex were determined by life tables under identical conditions. As seen in Fig. 1A, the median lifespans differ greatly (~15 to 85 days) among the clones. Once we had established the survivorship curves, we narrowed our selection of clones for all the future experiments. We chose RW20 and WF6 to represent our short-lived Daphnia clones and XVI-11 to represent our long-lived clone. We next investigated whether specific inherited variations of ND5 could be causative of differences in the ETC efficiencies that can then lead to differences in ROS generation and consequently affect the rate of aging and lifespan as observed in other established model organisms for research on aging. ND5 was chosen for initial analysis, as it is used extensively for understanding evolutionary relationships among Daphnia populations. We sequenced a 1140 bp region of the ND5 gene from the 11 clones to identify specific amino acid variations associated with characteristic short or long lifespans. When we aligned the 11 ND5 protein sequences in the order of increasing life spans (Fig. 1B), a striking pattern emerged. The most significant observations are: (1) the two shortest-lived clones carry a proline at position 307 whereas all other clones have a leucine, (2) the two longest-lived clones carry a leucine at position 320 whereas all other clones carry a phenylalanine, and (3) the three longest lived clones carry a leucine at position 354 whereas all other clones carry a phenylalanine, and (4) these variations are in the conserved C-terminal domain important for ND5 enzymatic activity.

**The MT complex I activity is diminished in the short-lived clones.** In order to test if the differences seen in the ND5 sequences correlate with the enzymatic activity of complex I, we compared the complex I enzymatic activity in two representative clones relative to total amount of complex I. We prepared MT protein extracts from mixed-age populations of WF6 short-lived clone and lake XVI-11 long-lived clone. We separated the individual ETC complexes by Blue Native polyacrylamide gel and measured the enzymatic activity of complex I using in-gel activity assay using NADH as the complex I substrate. If complex one activity is intact, it shows as a purplish dark blue band indicating active complex I (Fig. 2 and Supplementary Fig. 1). Such in-gel assay is used to measure complex I enzymatic activity as the complex remains catalytically active in native gels. As seen in Fig. 2A, (lanes 4–6), the short-lived WF6 clone showed significantly less complex I enzymatic activity when compared to the long-lived XVI-11 clone (lanes 1–3). The band intensities were measured and normalized to the band intensities of total amount of complex I (blue native panels) and are shown in Fig. 2B. The short-lived WF6 showed about 59% reduction in complex I activity compared to XVI-11. Although additional
sequence variations in other components of complex I that we have not investigated here also may contribute to the reduced enzymatic activity, these results show a similar association of complex I activity to the corresponding lifespans based on studies in other model systems.

Differences in the level of oxidative damage to proteins in the short-lived and long-lived clones. To test if the short-lived clone generated more oxidative stress in cells, we compared the levels of oxidative protein damage at different ages in two representative clones. We performed the analysis with extracts from 1, 2, and 3-week-old organisms of the short-lived clone RW20 and 1, 4, and 8-week-old organisms of the long-lived clone lake XVI-11, which were previously determined to correspond to equivalent time points in their respective life spans.

As shown in Fig. 3 and Supplementary Fig. 2, the short-lived RW20 clone showed high levels of oxidative damage at a young (Y) age of 1 wk (lane 1) that remained high throughout life (lanes 3 and 5, M and O). In contrast, the long-lived clone lake XVI-11 showed dramatically less oxidative damage at a young (Y) age (lane 2) as well...
as during middle (M) age (lane 4) and the old (O) population showed oxidized protein levels (lane 6) that were comparable to the 1 wk-old RW20 clone (lane 1). Similarly, it can be seen in Fig. 4 and Supplementary Fig. 3 that MT extract from a mixed population of short-lived WF6 clone showed more oxidative damage to proteins, when compared to the MT extract from a mixed population of long-lived XVI-11 clone. These results demonstrate that not only does the short-lived clone and long-lived clone have variation in their MTDNA encoded ND5 protein, these differences could be playing a role in why there is a big difference in their lifespans. As the short-lived clone WF6 had less complex I enzymatic activity (Fig. 2) which is known to generate higher ROS accumulation, it would result in increased oxidative damage in both the cells as well as in the mitochondria.

**Differences in the ROS scavenger mechanisms in short- and long-lived clones.** In order to investigate additional factors that could be contributing to the difference in oxidative damage to proteins, we...
compared the activity of the ROS scavenger enzymes superoxide dismutase (SOD) and catalase (CAT) in the short- and long-lived clones. SOD and CAT are ROS-scavenging enzymes known to play a central role in the antioxidant defense systems in all organisms. SOD enzymes convert the toxic superoxide radical into hydrogen peroxide (another ROS) and molecular oxygen depending on the cellular context. As seen in Fig. 5, the short-lived RW-20 displays significantly higher levels of SOD activity at young and middle ages, 10.9 and 14.5 units per microgram of protein respectively, whereas the long-lived clone XVI-11 at equivalent ages shows only 1.9 and 0.9 units per microgram of protein respectively. At old age in RW20, the SOD activity declines drastically to levels similar to old XVI-11 with 1.3 and 0.9 units per microgram of protein, respectively. Of particular note, XVI-11 exhibits relatively low amounts of SOD activity at all ages examined with very little change through the lifespan indicating lower amounts of ROS accumulation as ROS production is known to induce expression of SOD.

As the short-lived RW20 showed high level of oxidative damage to proteins at all ages during lifespan (Figs. 3 and 4) and also had elevated levels SOD activity (Fig. 5), we next examined the relative levels of hydrogen peroxide, the SOD scavenging reaction product that is also a ROS, in both short- and long-lived clones. Hydrogen peroxide causes oxidative damage in macromolecules including proteins and eleven distinct mitochondrial sites have been identified that can produce it in addition to the SOD enzyme reaction. As is shown in Fig. 6, the short-lived RW20 contains relatively high levels of hydrogen peroxide with little change during the lifespan. The young, middle and old RW20 organisms showing 153.6, 139.7, and 147.9 µM hydrogen peroxide per microgram.
protein respectively. In contrast, the long-lived XVI-11 clone contains high levels of hydrogen peroxide only in young organisms with 139.3 µM hydrogen peroxide per microgram protein, which declines drastically by mid-
middle age and is kept relatively low even in old organisms at 46.5 and 41.1 µM hydrogen peroxide per microgram protein respectively.

We then investigated the activity of catalase, another ROS scavenger enzyme that enzymatically converts the toxic hydrogen peroxide to water and molecular oxygen. To analyze the activity of catalase, we used a catalase activity assay as previously described. We initially standardized the assay using commercially available catalase (Sigma) and also optimized the assay parameters for Daphnia samples. As is shown in Fig. 7, young RW20 exhibit the highest catalase activity with an average of 63.4 units of CAT per microgram of protein with a decline to 30.9 units per microgram of protein at middle age and thereafter staying at 30.4 units of CAT per microgram of protein in older age. In XVI-11 the same overall trend is observed with higher amounts of catalase being in young organisms before declining in middle age and staying relatively unchanged in old organisms (44.6 to 10.4 to 7.7 units of CAT per microgram of protein, respectively. However, the amount of catalase activity is significantly lower in XVI-11 at equivalent time points compared to RW20 and as ROS is known to induce catalase expression, this possibly means lower overall levels of ROS in XVI-11.

Figure 6. Comparison of hydrogen peroxide levels in short-lived RW20 and long-lived XVI-11. Hydrogen peroxide levels were determined at the indicated ages for short-lived RW20 and using a kit available through Pierce. Before the kit, which uses a colorimetric assay, was used with samples from aged Daphnia, the kit was standardized for use with Daphnia samples (Data not shown). All reactions were performed in a 96 well plate that was read on a BioTek Plate reader at a wavelength of 560 nm using the Gen5 software that was included as part of the Bio-Tek plate reader. Blue corresponds to short-lived RW20 and orange corresponds to the long-lived XVI-11. Error bars indicate the standard error of the mean from 3 experiments. Student T-tests were performed and the p values are as indicated, ns means not significant.

Figure 7. Comparison of catalase activity in RW20 and XVI-11. Catalase activity was determined using a previously published protocol based upon the extinction of hydrogen peroxide over a period of time. All reactions were performed in a 96 well plate that was read on a BioTek Plate reader at a wavelength of 240 nm using the Gen5 software that was included as part of the Bio-Tek plate reader. Plates were read every 47 s for a duration of 5 min and the extinction of hydrogen peroxide was calculated. The activity of catalase was then calculated based upon the slope of the line representing the disappearance of hydrogen peroxide. Blue corresponds to short-lived RW20 and orange corresponds to the long-lived XVI-11. Error bars indicate the standard error of the mean from 3 experiments and the p-values are as indicated.
Discussion

Multiple studies from several model organisms reveal a complex relationship between the harmful ROS and aging. The role of oxidative damage and the effects of ROS on the aging process is not as straightforward as originally envisaged. Initial studies performed in *D. melanogaster* depicted ROS as being quite detrimental to organisms and was cited as being a major factor for the aging process \(^{43-45}\). Later on, other studies indicated that high ROS levels do not necessarily cause shorter lifespan and that overexpression of certain scavenger enzymes known to mitigate effects of ROS showed no effect on longevity \(^{46-47}\). One theory poses the presence of ROS as necessary for survival and can be thought of as molecules similar to AMP and NAD+, molecules that are necessary in a strict physiological range and are essential for proper homeostasis \(^5\). It is plausible that when ROS levels are elevated beyond this physiological range, they then become detrimental to the organism by causing damage to macromolecules, with the most crippling damage being to the proteins. ROS acts as a stress response signaling cascade that is not initially causative of aging and several ROS, particularly hydrogen peroxide, have been shown to be essential signaling molecules \(^1\).

In our present study we sought to examine the overall amount of oxidative damage to proteins, the activity levels of several ROS scavenger enzymes, and an associated ROS concentration level in young, middle, and old organisms of closely related *Daphnia* with vastly different lifespans, RW20 or WF6 for short-lived and XVI-11 for long-lived. Some of the previous studies have demonstrated DNA and lipid damage from ROS aren't likely involved as large causative agents for the aging process \(^{45,47}\). Therefore, we decided to examine oxidative damage in proteins and based on our data, WF6 and RW20 exhibit more oxidative damage to proteins than XVI-11 at young and middle-ages but the amount of damage is comparable in old organisms of both clones. What is causing the difference in oxidative damage to proteins? Based on our results, the naturally occurring ND5 sequence variations observed in short-lived RW20 and WF6 correspond to reduced complex I activity, which is known to lead to higher ROS generation. Thus, the short-lived clones generate more ROS and that corresponds well with higher amounts of oxidative damage to both cytoplasmic and mitochondrial proteins. Thus, we are able to associate the natural variations in mitochondrial ND5 sequence to lower complex I activity and increased oxidative damage to proteins in *Daphnia* clones with shorter lifespans.

The ROS scavenger enzyme SOD activity levels were much higher in the short-lived RW20 than long-lived XVI-11. This enzyme is known to mitigate the extremely toxic free radical superoxide anion; however, the product of the enzymatic reaction that rids the cells of these toxic species is hydrogen peroxide which is another toxic ROS. Therefore, the high amount of SOD activity in RW20 may not be beneficial for the organism as the enzyme produces a toxic product that could further damage cellular macromolecules, especially proteins. The hydrogen peroxide species is much more stable than superoxide anion and could have a lasting detrimental effect on various macromolecules and can be converted into another toxic ROS, the hydroxyl radical \(^{48}\). We also analyzed the levels of hydrogen peroxide in short- and long-lived clones at young middle and old ages. The short-lived RW20 did in fact have higher levels of hydrogen peroxide levels than XVI-11 and these higher levels are maintained throughout life. Note that SOD activity diminishes in old age of RW20 but the hydrogen peroxide levels remain high. As mitochondrial respiration could be less efficient with age and more ROS is known to be produced from aberrant ETC reactions, that could be the cause of the high level of hydrogen peroxide in old RW20 \(^{49}\). The long-lived XVI-11 also displayed higher amounts of hydrogen peroxide in young organisms which diminished by middle age and was maintained low through old age. To follow this up, we examined CAT activity, the enzyme which would convert hydrogen peroxide into water and molecular oxygen and thus can potentially alleviate some of the ROS induced stress in the cells. The results indicated that although RW20 showed higher amounts of catalase activity there was much higher amounts of hydrogen peroxide as well. This explains why, the short-lived RW20 contains higher amounts of oxidative damage to proteins, being unable to cope with the ROS produced during normal metabolism throughout the lifespan.

It’s also important to note that there are other types of ROS and ROS scavenging enzymes that we did not assay in this study, including hydroxyl anions and glutathione peroxidases \(^{51-54}\). Thus, whether free radicals and ROS are causative of aging, or act as a signaling molecules in response to other aging related insults, we definitely can detect differences in MT complex I activity, ROS levels, ROS scavenging enzymes, and oxidative damage to proteins between the short- and long-lived clones of *D. pulex*. In order to evaluate the direct causal relationship between lifespan and naturally occurring sequence variations in ND5 and possibly other MT genes as well as nuclear genomes, various mitochondrial genomes would need to be manipulated on different nuclear backgrounds and can be explored in future studies.

Crosstalk between the ROS network and the heat shock response has been observed and is important for aging process and longevity \(^{55-57}\). In a previous study, we had examined the proteotoxic response known as the heat shock response in short- and long-lived clones and found that the short-lived clone stops responding to heat stress by middle age whereas the long-lived clone can still respond to heat shock in middle age by inducing the expression of heat shock protein 70, a molecular chaperone \(^5\). Thus, the ability to respond to proteotoxic heat stress varies between the short- and long-lived clones of *Daphnia* and may also contribute to the observed differences in the longevity. Other ROS not analyzed in this work, such as the hydroxyl ions and their associated scavenging enzymes can be analyzed in future to determine their role in oxidative damage and lifespan. More importantly, we could further analyze the effects of exogenous treatment with antioxidants on lifespan extension, rate of aging, or sex-dependent aging \(^{58-60}\) using *Daphnia* model system.

Our work provides an entry point for the joint consideration of proximate and ultimate causes of differences in lifespan and aging. Ultimate causes are evolutionary, and in our system genetic differences have been shown to be associated with ecological differences in selection pressures on aging \(^{48,61,62}\). Our ecotypes, whether they are considered sister species that continue to exchange genes, incipient species in the throes of speciation, or simply divergent populations of one larger species, have strong differences of both life history and mitochondrial
function. While the differences in mitochondrial function almost certainly are proximate contributors to differences of life span and aging, they are unlikely to be the only proximate causes. We cannot tell whether the mitochondrial functions we examined were the initial molecular mechanisms underlying the phenotypic differentiation of the ecotypes, because as evolutionary change in aging occurs, feedback loops are expected to change selection pressures on other proximate mechanisms that contribute to aging (e.g., differences in heat shock response). Whether the differences we found are limited to our specific clones or are broadly representative of the adaptive divergence of the ecotypes is uncertain without broader sampling. However, it is worth noting that sequencing ND5 is often used as a shortcut for taxonomic identification because the ecotypes are morphologically indistinguishable, and thus we expect that functional differences driven by ND5 will be linked to taxonomic identifications made with ND5.

It is also important to keep in mind the limitations of our study. One of the limitations of our study is that any trait or gene that differs between the chosen short- and long-lived ecotypes will possibly score as associated with differences in lifetime and aging. Nevertheless, it is interesting that genes and molecular processes that have previously been identified as possible proximate drivers of lifespan and aging in other model organisms differ between these Daphnia ecotypes as well. In order to establish a direct link between naturally occurring mitochondrial genome variations with aging and lifespan, various mitochondrial genomes will need to be combined with different nuclear backgrounds of short- and long-lived ecotypes. This is certainly possible by setting up genetic crosses, as Daphnia can reproduce sexually and parthenogenetically allowing isogenic organisms to be easily produced. Our studies indicate that Daphnia would be the best suited model organism for examining and establishing the role of natural variations in mitochondrial genomes, oxidative damage to proteins, in rates of aging and relative lifespans.

Methods

Daphnia cultures. Daphnia clones used in this study were isolated from ponds and lakes in southwest Michigan in 2004–2008 and have since been cultured in the lab. No specific permissions are required to collect zooplankton from these public-access waterbodies in Michigan. D. pulex clones are neither endangered nor protected. For further details on the source populations, see Dudycha. Daphnia are maintained at a temperature of 20 °C with a photoperiod of 12:12 LD (12 h of light followed by 12 h of dark) within a Percival growth chamber. Daphnia were maintained at a concentration of 3 to 5 adult animals per 250 ml beaker in 200 ml of COMBO, an artificial lake water. Daphnia were cleared of young and transferred to a new beaker with fresh water on thrice weekly. They were fed every day with vitamin-supplemented algae Ankistrodesmus falcatus at a concentration of 20,000 cells/ml. To generate experimental animals, even-aged cohorts were begun by placing neonates individually in 100 ml of COMBO. Experimental animals were otherwise maintained as in the source cultures.

Survivorship curves. We compared survivorship of the 11 different clones using previously described methods for Daphnia in. Experimental conditions were 20 °C, a 12:12 LD photoperiod, with animals kept in individual 150-ml Pyrex beakers in 100 ml COMBO hardwater artificial lake water. Two generations were maintained under experimental conditions prior to initiating the life table to minimize maternal effects variation. Experimental individuals were fed 20,000 cells/ml Ankistrodesmus falcatus daily, a food level that produces normal aging processes in Daphnia. Individuals were transferred to fresh beakers and COMBO thrice weekly, and survivorship was recorded at each transfer until all experimental animals died. For each clone, n = 60 females, divided into two blocks of 30 individuals per clone per block. For survival curves, data were pooled across blocks. Individuals that died as juveniles (before age = 9 days, 47 individuals total across all clones and females, divided into two blocks of 30 individuals per clone per block. For survival curves, data were pooled across blocks. Individuals that died as juveniles (before age = 9 days, 47 individuals total across all clones and blocks) were excluded from analysis.

ND5 sequence analysis. Cloning of Daphnia ND5 ORF. Primers were designed to PCR amplify the Daphnia ND5 open reading frame (ORF) sequences of all our long- and short-lived Daphnia clones using MacVector based on the published D. pulex genome on the wleadata.org. For each PCR reaction, 10 ng genomic DNA was used with 50 pmol each of the forward and reverse primers designed to amplify the ND5 PCR product using the Promega GoTaq PCR kit. The following conditions were used for PCR: Lid 95 °C, 95 °C for 2 min (initial denaturation), denaturation at 95 °C for 15 s, annealing at 55 °C for 15 s, extension at 72 °C for 15 s for 24 cycles on the Eppendorf Scientific gradient Mastercycler. PCR products were separated on a 1% agarose gel. The PCR products were sent out for sequencing (Eton Bio). Both DNA sequences and the deduced protein sequences were aligned using MacVector and compared to the reference Daphnia mitochondrial genome. The protein sequences were checked for any amino acid sequence changes.

Primer sequences used were as follows:

Primer: ND5 Forward: 5’-GAGGTGTGGCACATTTCTTTA-3’.
Primer:ND5 Reverse: 5’-AAAGTCAATGAGCCGGGTA-3.

Daphnia cell lysate isolation preparation for Blue Native gels. Daphnia were collected from ongoing cultures into 1.5 ml microcentrifuge tubes. All water was removed from the tubes containing the Daphnia, the organisms were washed once with 1 ml cold PBS and the organisms were homogenized using a tight-fitting pestle. We homogenized 100–200 individuals in 100 µl of NativePAGE Sample buffer, 5% Digitonin containing protease inhibitor cocktails (Sigma and Calbiochem) following the manufactures protocol for the NativePAGE Sample Prep Kit (BN206, BN 2008, Life Technologies) and then centrifuged at 20,000×g for 30 min at 4 °C. The supernatant was aliquoted into sterile microcentrifuge tubes and stored at −80 °C until use. The protein concentration was...
Hydrogen peroxide assay. Hydrogen peroxide amounts were determined for each aged set of *Daphnia* using the Pierce Quantitative Peroxide Assay Kit (Lipid). This assay is based upon the oxidation of ferrous iron to ferric iron which is performed directly by the hydrogen peroxide contained within the sample. Once the iron is in the ferric state, it can bind directly to a xylenol orange dye which produces a violet color that can be measured spectrophotometrically at a wavelength of 560 nm. We first standardized the assay using various amounts of hydrogen peroxide (30% w/v) and then performed the assay with an equivalent amount of hydrogen peroxide contained within the sample. Once the iron was oxidized to ferric iron which is performed directly by the hydrogen peroxide contained within the sample, we calculated the hydrogen peroxide concentration using the formula: 

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\text{Hydrogen peroxide (U/mL)} = \frac{\text{Absorbance at 560 nm}}{\text{Absorbance of standard}} \times \text{Concentration of standard}
\]

Detection of protein carbonyls via Oxyblot. The oxidation status of proteins from *Daphnia* clones was determined using the Oxyblot protocol from EMD Millipore. This assay is based upon the fact that reactive oxygen species induce carbonyl group formation in proteins at specific residues. These carbonyl groups can then be derivatized into 2,4-dinitrophenylhydrazone (DNPH) by 2,4-dinitrophenylhydrazine (DNPH). The altered groups are detected following SDS PAGE and Western Blot Analysis with an antibody that detects the DNPH-hydrazone group of oxidized proteins. *Daphnia* protein samples were harvested using RIPA buffer and protein concentration was determined using a BCA protein quantification kit (thirty individual *Daphnia* were used for each aged set of both ecotypes). To derivatize the carbonyl group into DNPH-hydrazone, 20 µg of each protein sample was placed in a fresh tube (the samples were diluted such that 5 µl of each sample was used). Then 5 µl of 12% SDS was added to the tube to denature the proteins and 10 µl of 1X DNPH solution was added to the tube. The tubes were then incubated for 15 min at room temperature. The derivation reaction was stopped by adding 7.5 µl of neutralization solution to each tube. Note that we performed each Oxyblot with duplicate samples, one that had been derivatized and a control that was not derivatized into DNPH-hydrazone. The 20 µg samples were loaded onto a 12% polyacrylamide gel and Western blot analysis was performed using chemiluminescence ECL plus for detection of protein on a GE Typhoon LAS 4000 (antibody used was provided with the Oxyblot kit, recognizes DNPH-hydrazone, dilution 1:2000). The images were analyzed and quantified using the program ImageQuant.
Catalase activity assay. Catalase activity was detected using a modified method first developed by Beer and Sizer\(^3,4\). This spectrophotometric method utilizes the fact that the absorbance of hydrogen peroxide can be detected at 420 nm. The higher the amount of catalase present, the faster the absorbance by hydrogen peroxide will be diminished\(^4\). Seven individual Daphnia from each ecotype at the 3 different ages were homogenized in 30 µl of 67 mM Phosphate Buffer Saline (PBS) with protease inhibitor cocktail (Sigma-Aldrich, P8340). Note for each experiment 3 biological replicates and 3 technical replicates were performed. The homogenate was centrifuged at 13,200 rpms for 5 min and the cleared supernatant was kept and used in the Catalase Activity Assay. The assay was initiated by adding 300 µl of PBS-hydrogen peroxide solution (comprised of 160 µl of 30% hydrogen peroxide and 99.84 ml of 67 mM PBS) into a clear bottom well 96 well black plate (UV-star plate from Greiner Bio-one). Then 5 µl of Daphnia homogenate was added to the PBS-hydrogen peroxide solution in the well, mixed and read on a 96 well plate reader (Bio-Tek) using Gen5 software that was included as part of the Bio-Tek plate reader. Numerous reads were taken over a duration of 5 min with reads occurring every 47 s. We then calculated the slope of the diminishing hydrogen peroxide amounts and calculated the activity of catalase based upon the following equation where S is equal to the slope describing the diminishing hydrogen peroxide absorbance, V is volume of PBS-hydrogen peroxide solution added per well, and ε is the molar extinction rate of hydrogen peroxide at 240 nm (0.0436 ml µmol\(^{-1}\) cm\(^{-1}\) at 240 nm)\(^5\).

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\text{Catalase Activity} = S \left( \frac{V}{\varepsilon} \right)
\]

For each aged Daphnia set, we also determined the protein content of the samples using BCA kit from Pierce and present the data as units of catalase per microgram of protein. Note we also performed standard calculations based upon commercial catalase isolated from bovine liver (Sigma). For the standard calculation, we serially diluted the catalase and performed the assay at various concentrations to standardize the assay.

Superoxide dismutase (SOD) activity assay. To determine the activity of SOD in the two ecotypes of Daphnia at different ages, we utilized a kit from Sigma (Sigma SOD Assay kit 19160). This kit is based upon an altered tetrazolium salt WST-1, which will be reduced by superoxide anion to form a formazan dye which can then be detected spectrophotometrically at 450 nm wavelength. The presence of SOD will reduce the amount of superoxide anion and thus diminishing the amount of dye produced lowering the absorbance detected at 450 nm. For each aged set of Daphnia, we homogenized 10 individuals in 100 µl of PBS with protease inhibitor cocktail (Sigma-Aldrich, P8340), and then centrifuged at 13,200 rpms for 10 min. The supernatant was kept and used as the samples for the SOD Activity Assay. Note for each experiment 3 biological replicates and 3 technical replicates were performed. To perform the assay, 20 µl of Daphnia homogenate was added to the well of a 96 well plate. Then 200 µl of WST working solution was added to the well and mixed followed by the addition of 20 µl of Enzyme Working Solution. The plate was then incubated for 20 min at 37 °C before being read using a plate reader (Bio-Tek) at 450 nm using Gen5 software that was included as part of the Bio-Tek plate reader. The amount of SOD needed to inhibit 50% of the reduction of the WST-1 dye is defined as one unit of SOD\(^4\). Note that the amount of protein was determined for each sample and the activity of SOD is reported as units of SOD per microgram of protein. To standardize the assay we also performed the experiment using varying amount of commercially available SOD (sigma).

Statistics. To determine statistical significance when analyzing the various assays performed in this study, we executed both a two tailed Student’s T-test, assuming equal variance as well as Nonparametric Wilcoxon test. Each figure legend denotes p values, note that our alpha level was p = 0.05. Importantly, the p-values from both approaches agreed well in every single case.

Data availability
All data generated during this study are included in this article.

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Author contributions
B.U., C.S., and E.K. planned, performed all the experiments and interpreted the results; R.C.P. and J.L.D. supervised the work and B.U., C.S., J.L.D., and R.C.P. wrote the manuscript.

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