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A pro longevity role for cellular senescence

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A pro longevity role for cellular senescence

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Abstract

Cellular senescence is a fundamental process that may play positive or detrimental roles for the organism. It is involved in tissue development and in tumor prevention although during aging is becoming a detrimental process contributing to the decline of tissue functions. In previous investigations, we have uncovered a better capacity to detect DNA damage in cells from long-lived mammals. Here we report that cultured cells derived from long-lived species have a higher propensity to undergo senescence when challenged with DNA damage than cells derived from short-lived species. Using a panel of cells derived from six mammals which range in lifespan from 2-3 years up to 100 years we examined cell cycle response, apoptosis, and induction of senescence. All species exhibited a cell cycle arrest while induction of apoptosis was variable. However, a significant positive correlation was found between the relative percent of cells within a population which entered senescence following damage and the lifespan of the species. We suggest that cellular senescence may have a positive role during development allowing it to contribute to the evolution of longevity.

Key words:

Cellular senescence, apoptosis, longevity, body mass, SA β -gal.

Abbreviation list:

ETO: etoposide; NCS: neocarzinostatin; LBB: little brown bat; Flow cytometry: FCM; Senescence-associated β -galactosidase: SA β -gal; quantitative real time PCR (qRT-PCR)

1. Introduction

Evolutionary pressure on reproductive fitness is thought to indirectly dictate the lifespan of the species through the selection of traits that may be beneficial to survival but detrimental to longevity (Kirkwood & Rose, 1991). Many cellular mechanisms have been proposed to contribute to the evolution of longevity; some of these mechanisms can be species or taxa specific, while other mechanisms are common among species (Crawford et al., 2012; Croco et al., 2017; Tian et al., 2017). One fundamental cellular mechanism that has been implicated in the evolution of lifespan is DNA damage recognition and repair. We have previously shown that an enhanced capacity to recognize DNA double strand breaks is positively correlated with longevity, which appears to be related to the recognition of DNA-ends by the Ku80-Ku70 heterodimer (Lorenzini et al., 2009), as well as a greater capacity to form DNA damage foci of the phosphorylated histone H2AX (γ H2AX) and of 53BP1 (Fink et al., 2011; Croco et al., 2017).

An improved capacity to detect DNA damage would be expected to facilitate repair. Indeed we have observed lower levels of genomic damage in cultured cells derived from longer-lived species compared with short lived-species when the cultures were treated with similar amounts of genotoxic substances (Croco et al., 2017). Additional mechanisms related to DNA damage may also contribute to enhanced longevity. For example, resistance to DNA damage could also be coupled with an enhanced capacity for the induction of apoptosis or cellular senescence. In fact, these cell responses bear functional similarities with DNA repair, facilitating the removal of damaged/dysfunctional cells to preserve the integrity of the cell population within a tissue. Apoptosis acts through the elimination of irreparably damaged cells, while cellular senescence blocks proliferation of damaged cells, permitting clearance by the immune system.

The aim of the present work is to examine the correlation between apoptosis, cellular senescence and species longevity using primary fibroblast cultures of six mammalian species in which body mass is not correlated with lifespan, when challenged with DNA damage.

Adult body mass is a known confounding variable in any correlation with lifespan, as clearly explained by Speakman (Speakman, 2005), and correlation between damage responses and body mass were also considered.

2. Materials and methods

2.1. Cell cultures

Fibroblast strains from rat, mouse, little brown bat (LBB), dog and cow were established as previously described (Lorenzini et al., 2005). Cells were maintained in Minimum Essential Medium (MEM) with Earle's Salts and L-glutamine containing 10% fetal bovine serum (FBS), minimum essential medium vitamins, amino acids and penicillin-streptomycin (all from Sigma Aldrich, St. Louis, Mo, USA). Rat cells were from Sprague Dawley rats. Mouse cells were from a first filial generation hybrid mice offspring of a cross between C57BL/6J females (B6) and 129S1/SvImJ males (129S). All strains displayed replicative senescence with the exception of LBB and were used for all experiments before two third of their

expected replicative life span, see (Lorenzini et al., 2005). LBB strains reached more than 160 population doubling (PD), indicating probable immortalization. Consequently, for this species, cultures were used for experiments between 6 and 11 PD, to exclude the possibility of using immortalized cultures. Fibroblasts were passaged weekly. For the experiments, cells from confluent monolayers were seeded at 10000 cells / cm².

2.2. Genotoxic treatments

To challenge the cultures, cells were treated with a combination of concentrations and exposure times of genotoxic drugs so to cause damage only to a fraction of the cell population. After the challenge, the cells were allowed to recover for different amount of time in fresh media. This choice was dictated by the desire of studying cellular responses to mild and chronic insults as the one caused by the wear and tear of the aging process instead of studying the response to acute stressors. We used etoposide (ETO, an antineoplastic agent and well-known apoptosis inducer) at 100 µM for 30min and neocarzinostatin (NCS, a chemotherapeutic antibiotic) at 0.13 µM for 2h, both in serum free media to avoid possible interaction with serum albumin. ETO and NCS were both from Sigma Aldrich. After ETO treatment, the highest level of cell death (measured as percentage of necrotic plus apoptotic cells, see next paragraph) reached 38% at 48h from treatment (**Supplemental figure 1a**). With NCS treatment, the highest percentage of cell death (measured by FCA Guava ViaCount assay) was around 30% at 120h from the beginning of treatment (**Supplemental method and figure 1b**). See also the vitality results in our previously published work where we used also higher concentrations of ETO and NCS (Croco, Marchionni, Bocchini, et al., 2017).

2.3. Cell cycle analysis by Flow cytometry (FCM)

Cell-cycle analysis was performed by flow cytometry (FCM, Guava easyCyte 5HT equipped with a class IIIb laser operating at 488 nm, Merck Millipore, Darmstadt, Germany) and analyzed with Guava Cell-Cycle software, 24 and 48h after treatment with 100 µM ETO for 30min. Cells were fixed and permeabilized with ice-cold 70% ethanol and washed with PBS (Sigma Aldrich). Fixed cells were then resuspended in Guava Cell-Cycle Reagent (Merck Millipore) containing the dye propidium iodide (PI) and each sample was incubated in dark at room temperature for 30 min. The PI is able to penetrate the membrane of permeabilized cells, bind covalently to DNA and emit red fluorescence. Cells in G₂/M phase have double fluorescence compared to those in G₀/G₁ phase, while cells in S phase have intermediate fluorescence. The obtained results were calculated as the percentage of cells in the different stages of cell cycle in treated cultures relative to that present in the control cultures and expressed as fold induction.

2.4. Apoptosis quantitation by FCM

The percentage of viable, apoptotic and necrotic cells was assessed by FCM and analyzed with Guava Nexin software (Merck Millipore). After 24 and 48h from treatment with 100 µM ETO for 30min, Guava Nexin Reagent (Merck Millipore) was added to the cells, the reagent contain two dyes, 7-aminoactinomycin D (7-AAD) and Annexin-V-PE. 7-AAD allows the discrimination between live and dead cells, while Annexin-V-PE allows identification of apoptotic cells by binding to phosphatidylserine and emitting yellow fluorescence. In particular, live cells are 7-AAD and Annexin-V-PE negative, apoptotic cells are 7-AAD negative and Annexin-V-PE positive and necrotic cells are 7-AAD and Annexin-V-PE positive. Briefly, cells were incubated for 20 min at room temperature in the dark and samples analyzed. The obtained results were calculated as the percentage of cells in

apoptosis in treated cultures relative to that present in the untreated cultures and expressed as fold induction.

2.5. Histochemical Staining of Senescence Cells

After 24h and 120h from treatment with NCS 0.13 μ M for 2h, cells were fixed and stained for senescence-associated β -galactosidase (SA β -gal). 24h experiments were performed in AL laboratory at the University of Bologna, 120h experiments were performed in CS laboratory at Drexel University College of Medicine with different cell strains. Cells staining was performed using the Senescence Cells Histochemical Staining Kit (Sigma Aldrich). The Kit contains X-gal (5-bromo-4-chloro-3-indolyl-b-D-galactopyranoside) which in presence of β -galactosidase produces galactile and an insoluble blue compound. Briefly, cells were fixed with fixation buffer for 6-7 minutes at room temperature, washed with PBS and incubated with staining mixture at 37°C without CO₂ for 24h. The percentage of senescent cells was evaluated by microscopy at 200X magnification using Olympus IX50 fluorescence microscope (Tokyo, Japan). A total number of 400 cells were counted for each sample. The obtained results were calculated as the percentage of stained cells in treated cultures relative to that present in the control cultures and expressed as fold induction.

2.6. p21 quantitation by FCM

The mean fluorescence intensity value of p21 protein was analyzed in mouse and human strains, after 24h and 48h from treatment with NCS 0.13 μ M for 2h, by FCM with Guava Incyte software (Merck Millipore). Cells were fixed in PBS plus formaldehyde 4% and permeabilized in 90% cold methanol. Cells were then incubated with Anti-p21 antibody (#AHZ0422, Thermo Fisher, Waltham, MA, USA) and washed, incubated with AlexaFluor488-conjugated goat anti-mouse secondary antibody (#4408S; Cell Signaling, Danvers, MA, USA) and then analyzed. The obtained results were expressed as fold induction: mean fluorescence intensity values of cells in treated cultures relative to control culture values.

2.7. p21 mRNA quantitation by qRT-PCR

Total RNA extraction was performed 72h after NCS 0.13 μ M/2h treatment using TRIzol reagent (Thermo Fischer) and according to the manufacturer's instructions. RNA samples were purified by RNeasy Mini kit (Qiagen, Hilden, Germany). 200 ng of total RNA was used for qRT-PCR amplification of p21. p21 specific primers for all the species were designed by using NCBI Primer-BLAST software. The primer sequences are provided as supplemental information (**S2**). The expression of housekeeping gene GAPDH was used as an internal control to normalize the variability in the expression levels. qRT-PCR was performed by using Verso SYBR Green 1-Step qRT-PCR Low ROX Mix kit (Thermo Fisher). The run was set according to the manufacturer's instructions: cDNA synthesis (50°C for 15 minutes, 1 cycle), Thermo-Start activation (95°C for 15 minutes, 1 cycle), denaturation (95°C for 15 seconds), annealing (60°C for 30 seconds) and extension (72°C for 30 seconds) (40 cycles). Reactions were run in triplicate. Expression data were normalized to the geometric mean of housekeeping gene GAPDH to control the variability in expression levels.

2.8. p21 visualization by immunofluorescence

Mouse and human fibroblasts were seeded onto glass coverslips and treated with NCS 0.13 μ M for 2h. Cells were fixed at 24 and 48h after damage in 4% paraformaldehyde for 10min and permeabilized in 0.2% Triton X-100 in phosphate-buffered saline (PBS) for 5min. Cells were washed once in PBS and blocked for 30min in 4% bovine serum albumin (BSA)

in PBS containing Tween-20 (PBST), after which they were incubated with the p21 primary antibody (#AHZ0422, Thermo Fisher) in 1% BSA–PBST buffer for 2h at room temperature in a humidified chamber. Slides were washed three times in PBST and incubated with secondary antibody (#4408S; Cell Signaling) in 0.1 % BSA–PBST for 1h. Cells were washed three times, stained with 4',6-diamidino-2'-phenylindole dihydrochloride (DAPI, Sigma-Aldrich), and mounted with Vectashield mounting medium (Vector Laboratories, Burlingame, CA, USA) before analysis. Images were captured using Olympus IX50 fluorescence microscope equipped with a Canon camera G16 (Tokyo, Japan).

2.9. Statistical Analysis

All results are expressed as mean \pm standard error of the mean (SEM).

For the analysis of the relation between cell cycle or apoptosis with longevity and body mass we used linear regression. For the analysis of the relations between senescence with longevity, body mass, sexual maturity and post-natal growth rate, we used linear regression, correlation and partial correlation. To compare apoptosis induction among species, we used the 1-way Analysis of Variance (ANOVA) for unpaired data, followed by Tukey test for multiple post comparison. For p21 analysis between mouse and human, we used unpaired t-test.

All the statistical analyses were performed using Prism Software 7, with the exception of partial correlations that were performed using the VassarStats website.

2.9. Species information

Species data are all from AnAge database (Tacutu et al., 2018). Maximum longevity in the text is referred simply as longevity. Body mass is taken from weight value in AnAge. For sexual maturity, we used the average between male and female data. Growth rate in AnAge is calculated by fitting empirical data taken from published growth curves to sigmoidal growth functions (Table 1).

Table 1: Traits of analyzed species

SPECIES	Common name	Sample Size*	Maximum Longevity (years)	Body Mass (g)	Sexual Maturity (days)	Postnatal growth rate (days⁻¹)
<i>Rattus norvegicus</i>	Rat	Large	3.8	300	80	0.0207**
<i>Mus musculus</i>	House Mouse	Large	4.0	20.5	42	0.0298
<i>Bos taurus</i>	Domestic cattle; cow	Large	20	750000	548	0.0031
<i>Canis familiaris</i>	Dog (Beagle)	Large	24	40000	510	0.0244
<i>Myotis lucifugus</i>	Little brown bat (LBB)	Medium	34	10	210	0.116

<i>Homo sapiens</i>	Human	Huge	90***	62000	4927	0.0005
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Table Legend:, Values are means (except for maximum longevity) obtained from the AnAge database, available at: <http://genomics.senescence.info/species/> (Tacutu et al., 2018). * Medium, 100-1000; Large, over 1000; Huge, to indicate the special status of the human data. ** For the growth rate of rat (*rattus norvegicus*), value absent in AnAge, was used instead the value from *rattus rattus*. *** Maximum human longevity is 122.5 years. In our analysis, we consider a maximum longevity of 90 years to account for the fact that, for the others species, only small cohorts were used to determine maximum life span, and 90 years seem a more realistic estimate for a random sample of humans.

3. Results

3.1. Cell cycle analysis

We initially sought to determine if cells, from species which vary in their lifespan have a similar capacity to control and arrest the cell cycle when faced with DNA damage. In order to ascertain this, cell cycle profiling was performed following damage with etoposide (ETO), a topoisomerase II inhibitor known to induce an S phase arrest. A short pulse of ETO was used to induce damage specific to S phase, allowing visualization of G2/M arrest as a readout of cell cycle arrest.

The percent of cells in G2/M was compared at time points 24h and 48h following damage in untreated cultures and treated cultures, **Figures 1a** and **1b**. Cultured cells derived from mouse, dog, cow, little brown bat and human were examined. In these comparisons, there was an increase in G2/M phase in all species, demonstrating that all are capable of cell cycle arrest in response to DNA damage.

An analysis of the relationship of the magnitude of the cell cycle arrest as determined by the percent of the population in G2/M following this response with longevity gives a negative result (**Figure 1c** $r^2 = 0.03$, $p = n.s.$; **Figure 1d** $r^2 = 0.04$, $p = n.s.$). Although not significant, a relationship was observed between cell cycle arrest and adult body mass (**Figure 1e** $r^2 = 0.63$, $p = n.s.$; **Figure 1f** $r^2 = 0.20$, $p = n.s.$).

3.2. Induction of Apoptosis

Next, we sought to determine if longevity correlates with the intrinsic capacity to induce apoptosis. Using ETO as a damaging agent, the fold induction of apoptosis was determined for each species. A subset of species showed little response to the concentrations of ETO used (mouse and dog) while in the others (rat, cow, LBB and human) the response was evident. The fold induction levels for apoptosis in cultures after 24h and 48h following DNA damage are reported in **Figures 2a** and **2b**. Regression analysis of the results, revealed no meaningful relationship between fold induction of apoptosis and either longevity or body mass (**Figures 2c, 2d** and **2e, 2f**). We also investigated the possibility that a relationship may exist between longevity or body mass and the total number of dying cells (apoptotic + necrotic cells). Even when these parameters are combined, no relationship was detected between these data and either longevity or body mass (data not shown).

Apoptosis is a fundamental cellular process and the results may suggest that apoptosis represents a species-specific or taxa-specific longevity-assurance strategy. For example, Zhang et al. have reported positive selection for p53, a key mediator of apoptosis, in a species of vesper bat, *Myotis davidii* (Zhang et al., 2013). Our data is consistent with the concept that p53 mediated apoptosis is a longevity-assurance mechanism for the LBB (an extremely long-lived species for its size) since only this species shows a strong response. This interpretation however, is weakened by the high variation of this species response, and an ANOVA analysis failed to show a significant difference among the species in terms of apoptotic response. We also analyzed LBB cells at high PD (between 80 and 90), and although these data were not used in our comparative analyses, the response was intense and similar to the response of LBB at low PD.

3.3 Induction of stress induced senescence

In order to assess the relative capacity to induce senescence following genotoxic damage in each species, we used an antibiotic that has effects on DNA which are biologically similar to the effects of ionizing radiation: neocarzinostatin (NCS) (Goldberg, 1987) followed by an evaluation of the percentage of cells positive for SA- β -gal.

All species responded to the genotoxic stressor with an increase in the percentage of SA β -gal stained cells. The increase varied from a minimum of 1.31 fold for mouse cultures at 120h to a maximum of 3.52 fold for human cultures also at 120h. Regression analysis demonstrated a positive, strong and highly significant relationship with longevity both at 24h and 120h (**Figure 3c**, $r^2=0.97$, $p<0.001$; **Figure 3d**, $r^2=0.87$, $p<0.01$). No significant relationship was found with body mass at either time point (**Figure 3e**, $r^2=0.18$, $p=n.s.$; **Figure 3f**, $r^2=0.04$, $p=n.s.$). The lack of relationship with body mass is important as it is well known that body mass and longevity are related. For example, De Magalhães and colleagues, analyzed data from 856 mammals and reported that body mass explained 66% of the longevity variation (Magalhães et al., 2007). However, body mass is not significantly related to lifespan among the six species we used in this analysis (**Figure 3d**). To further explore the possible role of body mass in the relationship between senescence and longevity, we performed correlation and partial correlation analysis (**Table 2**). This exploration confirms that longevity is the parameter primarily related to the ability to induce cellular senescence.

Table 2. Correlations and Partial Correlations

Correlation 24h	r of correlation	Controlling for	r of partial correlation
Senescence vs Longevity	0,983	Body Mass	0,98
Senescence vs Body Mass	0,425	Longevity	-0,20
Longevity vs Body Mass	0,466	Senescence	0,29
Correlation 120h	r of correlation	Controlling for	r of partial correlation
Senescence vs Longevity	0,938	Body Mass	0,97
Senescence vs Body Mass	0,212	Longevity	-0,73

Longevity vs Body Mass	0,466	Senescence	0,79
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Table Legend: Senescence is presented as fold induction of DNA damage stimulated senescence (data used are plotted in **Figure 3**).

As an additional proof for our finding, we used p21 as a further marker for cellular senescence. In the presence of unresolved DNA damage, the ATM kinase activates p53 and its transcriptional target p21, which arrests proliferation by inhibiting cell-cycle-dependent kinases [reviewed in (Bitto et al., 2014)]. Stimuli that elicit a DNA damage response induce senescence primarily through the p53-p21 pathway (Campisi & d'Adda di Fagagna, 2007). p21, additionally, seems essential for developmentally programmed senescence (Storer et al., 2013). We assayed, p21 protein expression in human and mouse cells using an antibody that recognized an epitope conserved between these species. Direct comparisons with other species was not possible since few antibodies recognize an epitope conserved across multiple species. The results of this analysis, is presented in **Figure 4c** and quantified in **Figure 4a**. Human cells increased p21 expression to a significantly greater extent than mouse cells at both 24h and 48h following DNA damage. Additionally, to compensate for the lack of a single antibody able to recognize p21 in all species, we performed quantitative real time PCR (qRT-PCR) to measure the relative abundance of p21 mRNA. Data presented in **Figure 4b**, shows the relative abundance of p21 mRNA for the species in which we were able to design primers that gave reliable results. We observed the same trend for the steady state p21 mRNA levels as we did with SA β -gal, i.e. greater induction of senescence in long-lived species.

A positive relationship between longevity and slow development, is well supported by large comparative analyses (Magalhães et al., 2007; Ricklefs, 2010). Consequently we decided to use available data on two developmental characteristic of species to see if a regression analysis will show a relationship between development and the capacity to induce senescence. **Supplemental Figure 2a** and **2b** shows that we found a significant positive relationship between the capacity to induce cellular senescence and time to sexual maturity. For growth rate, we observed a negative trend although the relationship is not significant (**Supplemental Figure 2c** and **2d**).

4. Discussion

The evolution of large body mass requires the adaptation of cellular mechanisms to allow the high number of cell divisions required to maintain the larger body mass (reviewed in Croco et al., 2017). Accordingly, species with higher body mass tend to have a more accurate erythropoiesis (Croco et al., 2016). Similarly, the evolution of greater longevity should also be accompanied by an adaptation of cellular functions that support the longer period of functionality associated with enhanced longevity. For example, it was observed that cells from long lived species retain higher viability when challenged with different stressors (Kapahi et al., 1999). In this study, we sought to determine if the evolution of longevity was accompanied by an enhanced regulation of cell cycle progression, apoptosis, or senescence using equivalent levels of damage as inducers.

4.1. Longevity vs inhibition of growth and induction of apoptosis

Our analysis reveals no evident relationship between the capacity to arrest the cell cycle or to induce apoptosis with either species longevity or body mass.

There are several potential explanations for these negative results. Cell cycle regulation and apoptosis are a highly conserved cellular mechanisms present in all metazoan, important for maintaining a homeostatic balance within tissues during development and in adulthood (Crawford et al., 2012; Meier et al., 2000). Consequently, all mammalian species are equipped with effective cell cycle regulation and apoptotic machineries and it may be that these processes are too fundamental to allow significant variation beyond that required for evolutionary fitness.

An alternative hypothesis, is that enhanced cell cycle control and/or apoptosis may be associated with longevity in a subset of long-lived species having been associated with selective pressure for reproduction in a few instances. This hypothesis is supported by reports of a remarkably efficient apoptotic response in elephants (Abegglen et al., 2015; Vazquez, et al., 2018).

There are several caveats to this study. The small number of species involved, and the fact that for cow, dog, and the laboratory rodent strains used, we cannot ignore the potential confounding impact of human based artificial selection to which they were exposed. Consequently, the possibility exists that our study is not sufficiently powered to detect a weak correlation with longevity. In this regard, it is noteworthy that there is a tendency for higher capacity to induce apoptosis in species with higher longevity (**Figure 2c and 2d**).

4.2. Longevity is positively associated with stress induced senescence

Using the SA β -gal staining method, we detected a highly significant relationship between longevity and the capacity to induce senescence following DNA damage. The expression of the senescence marker p21 is also higher in longer lived species, reinforcing this observation. Although correlation does not imply causation, the relevance of this finding is underscored by the fact that SA β -gal staining is an established marker for cellular senescence (Dimri et al., 1995) and represents a reliable, early sign of replicative senescence (Maier et al., 2007), while multiple studies have demonstrated that this marker is capable of detecting stress induced senescence (Debacq-Chainiaux et al., 2005; Dumont et al., 2000; Fripiat et al., 2001; Zhao et al., 2018). In addition, stress induced senescence appears to be a more universal response than replicative senescence, which can be absent in both short lived and long lived species (Zhao et al., 2018). Furthermore, cells of the long lived rodent, the naked mole rat, favor senescence over apoptosis when compared to mouse cells (Zhao et al., 2018).

4.3. A possible role for stress induced senescence during development

Because cellular senescence is considered one of the hallmarks of the aging process (López-Otín et al., 2013) and data are available demonstrating an increase in senescent markers in individuals of increasing age (Lawrence et al., 2018; Waaijer et al., 2016), the positive relationship between the capacity to induce senescence and longevity that we have observed may appear counter intuitive.

Our hypothesis is that spontaneous DNA damage may be induced at any time including early development, the moment in life when tissues require exponential growth and when, consequently, DNA replication errors would be expected to be more frequent. The appearance of a damaged but surviving cell during development poses a great challenge if the cell is not repaired or eliminated by apoptosis. However, the rate of apoptosis may be limited by other evolutionary constraints. In this case, a compromised cell could retain the capacity to proliferate with the consequence that all descendants will carry the same inherent genomic damage. For this reason, efficiently inducing cellular senescence, to avoid the accumulation of a sub-population of damaged cells, may represent a pro longevity assurance mechanism. Consistent with this notion, a physiological role for senescence during early development has been demonstrated in both humans and in mice (Chuprin et al., 2013; Storer et al., 2013).

Our previous studies have led us to consider developmental time as an important factor in the theory of aging (Lorenzini et al., 2011). It is possible that cellular mechanisms supporting functional integrity during development may be more significant than those of adult life in explaining the functional declines of aging. Ameer and colleagues, for example, have observed that most somatic mtDNA mutations do not result from DNA damage buildup in adult life, but occur during development (Ameer et al., 2011). With a reduced rate of development, cells damaged by errors such as replication fork collapse or transcription/replication fork collisions may be able to undergo a more prolonged arrest, to either repair the damage or undergo apoptosis or senescence.

Argüelles and colleagues propose that efficient apoptosis could be considered a pleiotropic aging factor, positive during development but negative during adulthood (Argüelles et al., 2019). We suggest that the same logic applies to senescence: enhanced senescence may be a positive feature during development although accumulation of senescence cells in adulthood remains pro aging.

Finally, Peto's paradox suggests that species with higher body mass and great longevities should succumb to cancer more often than small short-lived species due to the requirement for increased cell division (representing more potential targets for oncogenic transformation) and greater time available to accumulate mutations. The paradox is that the opposite is observed (Peto, 2015). Our findings represent an additional explanation to this puzzle.

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Figure legends

Figure 1. Cell cycle arrest following DNA damage: relationship with longevity and body mass. Fibroblast cultures were treated with 100 μ M ETO for 30 min; data are expressed as fold induction in each cell cycle phase 24h (**a**) and 48h (**b**) following treatment relative to control cultures; species are ordered by increasing longevity. Fold induction specifically in the G2/M phase are plotted against maximum longevity (**c** for 24h data; **d** for

48h data) and against body mass in (**e** for 24h data; **f** for 48h data). Linear regression trend lines, coefficient of determination and significance of the regression are shown. The experiment was performed twice in duplicates.

Figure 2. Induction of apoptosis relative to longevity and body mass. Fibroblast cultures were treated with 100 μ M ETO for 30 min; the percentage of cells in apoptosis after 24h and 48h following treatment is reported relative to the percentage in untreated cultures (**a** and **b** respectively); species are ordered by increasing longevity. The fold induction of apoptosis are plotted against longevity (**c** for 24h and **d** for 48h) and against body mass (**e** for 24h and **f** for 48h). Linear regression trend lines, coefficient of determination and significance of the regression are shown. Experiments were performed three times in duplicates.

Figure 3. Induction of senescence and its relationship with longevity and body mass. The percentage of SA β -gal stained cells relative to the percentage in untreated cultures, 24h and 120h following treatment with NCS (0,13 μ M for 2h) is reported in **a** and **b** respectively; species are ordered for increasing longevity. The fold induction of SA β -gal stained cells were plotted against longevity (**c** for 24h and **d** for 120h) and against body mass (**e** for 24h and **f** for 120h). Linear regression trend lines, coefficient of determination and significance of the regression are shown. 24h experiments were performed at least three times in duplicates, 120h experiments were performed twice in duplicates.

Figure 4. Expression of p21 following DNA damage. FCM determination of p21 expression in mouse and human cells 24h and 48h from treatment with NCS (0.13 μ M for 2h) or untreated is shown as fold induction relative to untreated cells. In **c** representative histogram plots are presented. In **b** qRT-PCR values for p21 expression among different species normalized to GAPDH are presented. In **d**, pictures of treated human and mouse cells after immunofluorescence staining with the p21 antibody are presented. FCM and immunofluorescence experiments were performed twice in duplicates. qRT-PCR experiment was performed once in triplicates.

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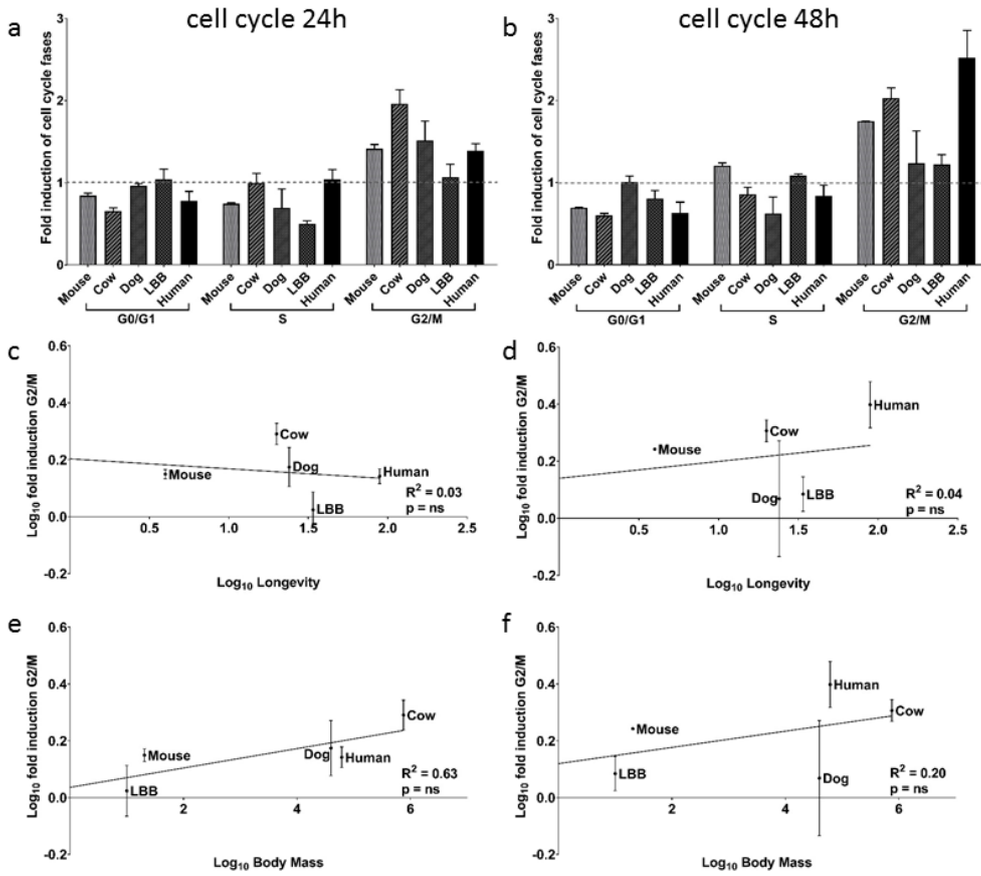
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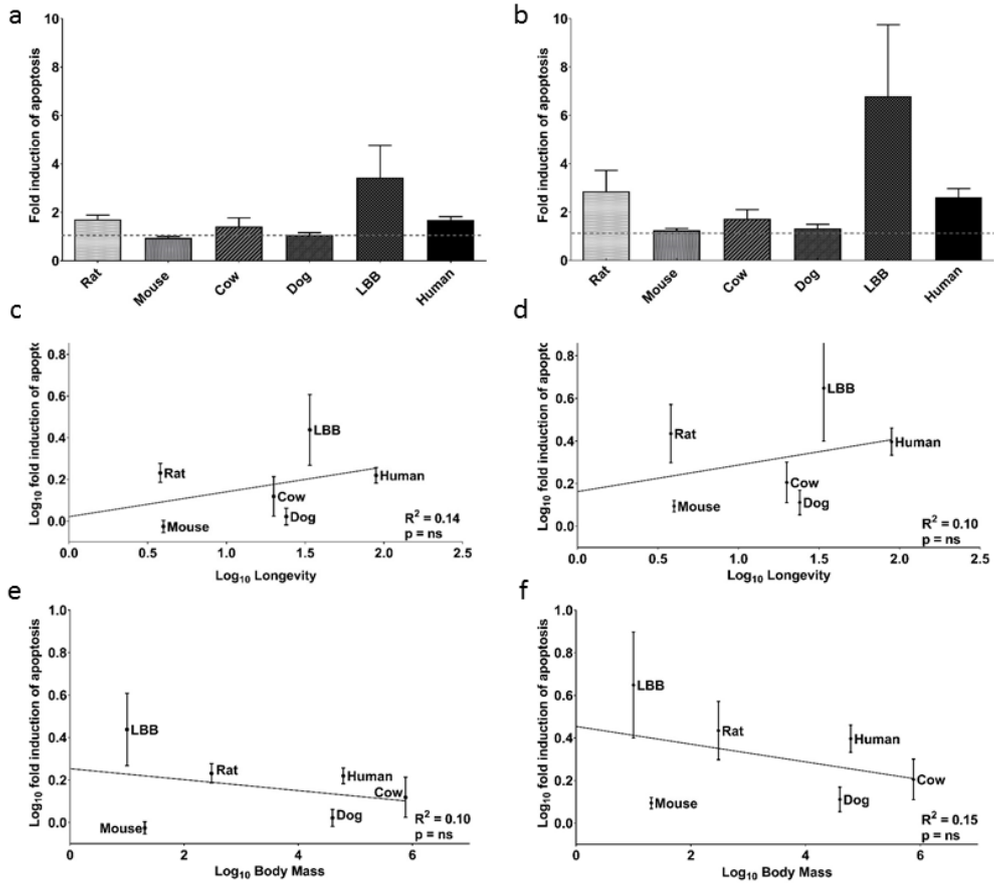
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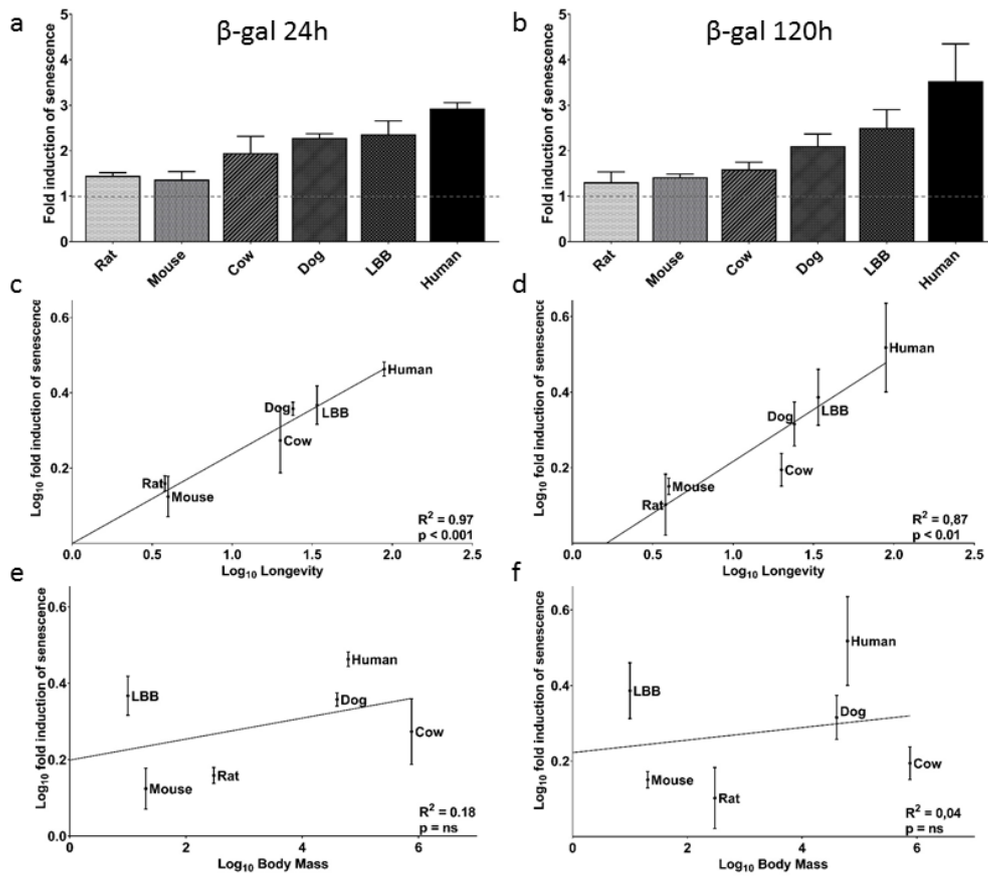
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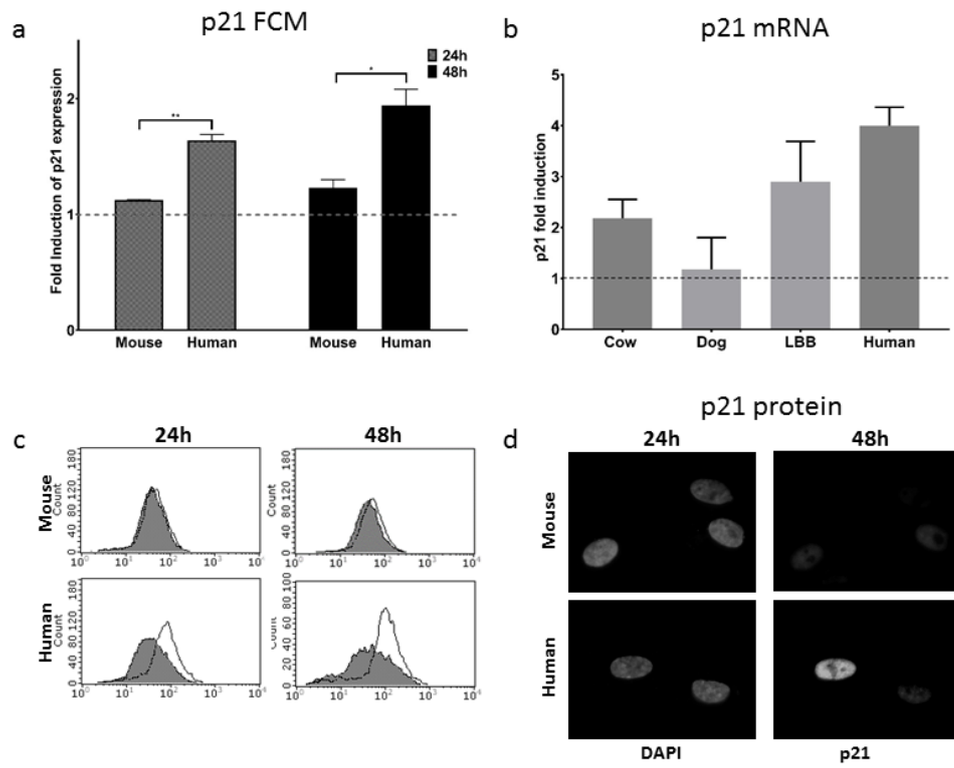
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Supplemental Information

A pro longevity role for cellular senescence

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Supplemental methods

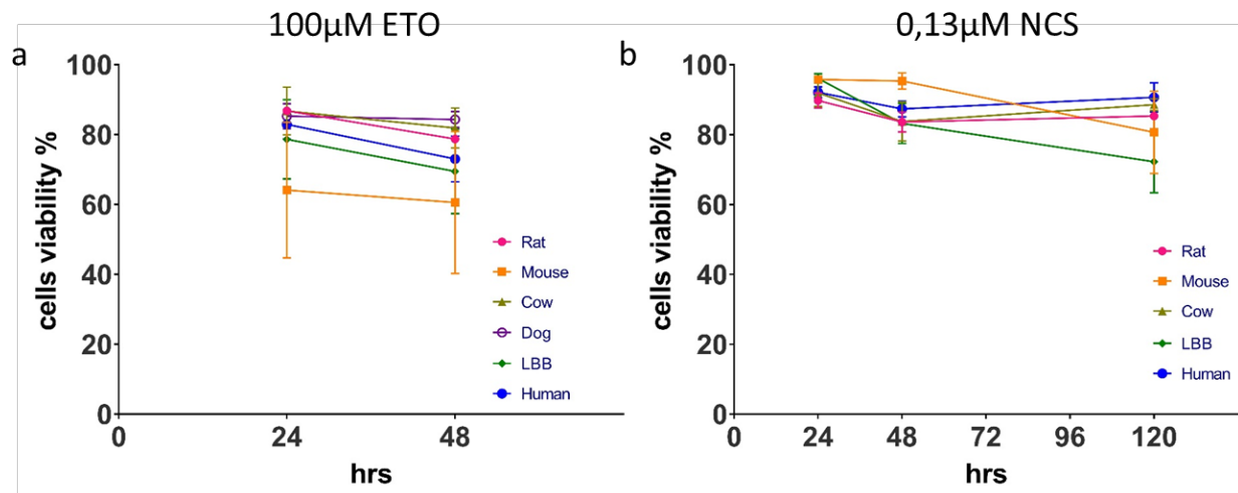
S1. Cell viability after NCS treatment

Viability was measured using the ViaCount Kit for Guava FCM (Merck Millipore). Cells growing in 12-wells plates, were treated with NCS 0.13 μ M for 2h, trypsinized and resuspended in 50 μ l of media and 450 μ l of ViaCount solution. The FCM was set to count a thousand cells for each sample. Each species was analyzed in triplicates.

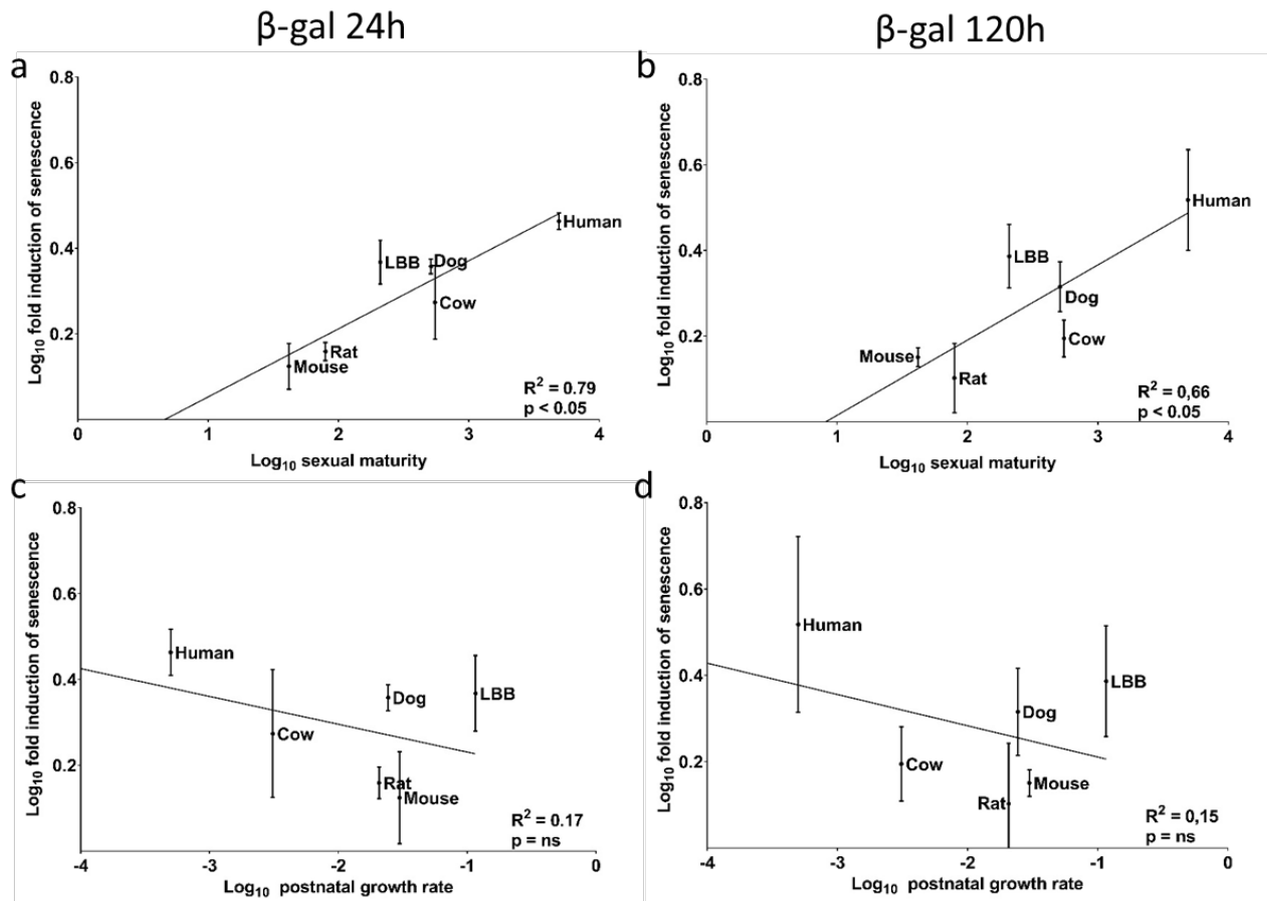
S2. Primers used for qRT-PCR

SPECIES	SEQUENCE (5' > 3')		PRODUCT LENGTH
RAT	Forward	TTGTGATATGTACCAGCCACAGG	115
	Reverse	TCAACTGCTCACTGTCCACG	
MOUSE	Forward	GACATTCAGAGCCACAGGCAC	80
	Reverse	GACAACGGCACACTTTGCTC	
DOG	Forward	GGCAGACCAGCATGACAGATTT	910
	Reverse	CACTAAGCTGGGGGAACAGG	
BOVINE	Forward	CCATGCCCTTTCCCCTTAGT	107
	Reverse	CTACGCCTAGCATGTCCCTC	
LBB	Forward	AGACTGCGATGCTCTGATGG	370
	Reverse	GCGTTTGGAGTGATAAAAATCTGTC	
HUMAN	Forward	TGCCGAAGTCAGTTCCTTGT	86
	Reverse	GTTCTGACATGGCGCCTCC	

Supplemental Figures



Supplemental Figure 1. Cell viability assay after genotoxic treatments. Fibroblast cultures were treated with ETO at 100 µM for 30min (a), or NCS 0.13 µM for 2h (b) and viability was measured after different times by FCM.



Supplemental Figure 2. Relationship between senescence induction values and developmental schedules. The relationship between longevity and body mass for the 6 species analyzed is plotted in **a**. Fold induction of senescence for all the species assayed were plot against time to sexual maturity (**b** for 24h and **c** for 120h) and against growth rate (**d** for 24h and **c** for 120h). Linear regression trend line, coefficient of determination and significance of these regressions are shown.