Concurrent Evolution of Antiaging Gene Duplications and Cellular Phenotypes in Long-Lived Turtles

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Abstract

There are many costs associated with increased body size and longevity in animals, including the accumulation of genotoxic and cytotoxic damage that comes with having more cells and living longer. Yet, some species have overcome these barriers and have evolved remarkably large body sizes and long lifespans, sometimes within a narrow window of evolutionary time. Here, we demonstrate through phylogenetic comparative analysis that multiple turtle lineages, including Galapagos giant tortoises, concurrently evolved large bodies, long lifespans, and reduced cancer risk. We also show through comparative genomic analysis that Galapagos giant tortoises have gene duplications related to longevity and tumor suppression. To examine the molecular basis underlying increased body size and lifespan in turtles, we treated cell lines from multiple species, including Galapagos giant tortoises, with drugs that induce different types of cytotoxic stress. Our results indicate that turtle cells, in general, are resistant to oxidative stress related to aging, whereas Galapagos giant tortoise cells, specifically, are sensitive to endoplasmic reticulum stress, which may give this species an ability to mitigate the effects of cellular stress associated with increased body size and longevity.

Key words: comparative genomics, longevity, cancer biology, turtle, DevoEvo, evolutionary medicine.

Significance

Although all animals are sensitive to developing cancer, some are remarkably resistant to developing cancer during their lifetimes. Identifying how these species evolved to be cancer resistant can provide new insights into basic cancer biology and evolution. We found that long-lived turtles and tortoises evolved enhanced cancer resistance, at least in part by gaining extra copies of cancer-fighting genes. These observations may help the development of new cancer treatment and prevention strategies.

Introduction

Body size and longevity are fundamental life history traits that vary tremendously across vertebrates. Maximum body mass in vertebrates ranges from 0.5 g in the red-backed salamander (Plethodon cinereus) (Moore et al. 2001) to 136,000 kg in the blue whale (Balaenoptera musculus) (Magalhães and Costa 2009), whereas maximum lifespan ranges from 8 weeks in the pygmy goby (Eviota sigillata) (Depczynski and Bellwood 2005) to over 400 years in the Greenland shark (Somniosus...
shorter-lived ones (Peto 2015). Although this correlation exists organisms should be at greater risk of cancer than smaller, performing cells is central to the aging process (López-Otín et al. 2013). Thus, if cancer-suppressing mechanisms play a critical role in cancer resistance by removing potentially cancerous and toxic and cytotoxic stress, and apoptosis also plays a primary role in cancer resistance by removing potentially cancerous cells (Verfaillie et al. 2013). At the same time, apoptosis, which is the programmed destruction of unfit or damaged cells, is reduced in older individuals (Salminen et al. 2011). This decline in cell performance in combination with a decreased ability to remove poor-performing cells is central to the aging process (López-Otín et al. 2013). Similarly, cancer can arise from cumulative genotoxic and cytotoxic stress, and apoptosis also plays a primary role in cancer resistance by removing potentially cancerous cells (Verfaillie et al. 2013). Thus, if cancer-suppressing mechanisms are similar across species, then larger, longer-lived organisms should be at greater risk of cancer than smaller, shorter-lived ones (Peto 2015). Although this correlation exists within species, for example, cancer incidence increases with increasing adult height for most cancer types in humans (Green et al. 2011) and overall body mass in dogs (Dobson 2013), there is no such correlation between species—an observation often referred to as “Peto’s paradox” (Peto 2015).

The molecular and cellular mechanisms underlying the evolution of large bodies and long lifespans have been explored in mammals such as elephants (Sulak et al. 2016; Vazquez et al. 2018), whales (Seim et al. 2014), bats (Foley et al. 2018; Gorbunova et al. 2020), and naked mole rats (Salón et al. 2008; Gorbunova et al. 2014), but are less well studied in other vertebrates. Reptiles are an excellent system in which to study the evolution of body size and longevity because diverse lineages have repeatedly evolved large body sizes and long lifespans (Chiari et al. 2018). Turtles, in particular, have lower rates of neoplasia than snakes and lizards (Garner et al. 2004; Sykes and Trupkiewicz 2006), are especially long-lived, and are “slower aging” than other reptiles (Hoekstra et al. 2020). Most notably, Galapagos giant tortoises (Chelonoidis niger species complex; hereafter referred to as C. niger) and Aldabra giant tortoises (Aldabrachelys gigantea) can live over 150 years (3–5 times longer than their closest relatives) and weigh over 200 kg (50–100 times heavier than their closest relatives) (Caccone et al. 1999; Palkovacs et al. 2002; Poulakakis et al. 2012; Chiari 2020). Galapagos giant tortoises also appear to have evolved a suite of cellular traits that may contribute to their longevity, such as a slower rate of telomere shortening and extended cellular lifespans compared with mammals (Goldstein 1974).

Here, we explore the evolution of body size and lifespan in turtles by integrating several approaches (fig. 1): 1) phylogenetic comparative analysis of body size, lifespan, and intrinsic cancer risk in turtles; 2) gene duplication analysis of aging and cancer-related genes across available turtle genomes; and 3) cell-based assays of apoptosis and necrosis in multiple turtle species varying in body size and lifespan. We show that species with remarkably long lifespans, such as Galapagos giant tortoises, also evolved reduced cancer risk. We also confirm that the Galapagos giant and desert tortoise genomes encode numerous duplicated genes with tumor suppressor and anti-aging functions (Quesada et al. 2019). Our comparative genomic analysis further suggests that cells from large, long-lived species may respond differently to cytotoxic stress, including endoplasmic reticulum (ER) and oxidative stress. The combined genomic and cellular results suggest that at least some turtle lineages evolved large bodies and long lifespans, in part, by increasing the copy number of tumor suppressors and other antiaging genes and undergoing changes in cellular phenotypes associated with cellular stress.

Results

Repeated Evolution of Large Body Size in Turtles

We found substantial independent accelerations in the rate of body size evolution in several turtle lineages, including a 29× rate increase (386% increase in carapace length) in the stem-lineage of sea turtles (Chelonioidae) and a further 103× rate increase (757% increase in carapace length) in leatherback sea turtles (Dermochelys coriacea), a 37× rate increase (35% increase in carapace length) in the stem-lineage of soft-shell turtles (Trionychidae), and a 200× rate increase (364% increase in carapace length) in the stem-lineage of narrow-headed soft-shell turtles (Chitra chitra and Chitra indica), and a 463× rate increase (364% increase in carapace length) in Cantor’s giant softshell turtle (Pelochelys cantonii) (fig. 2A).

Among the more notable groups with increased rates of body size evolution were the “giant” tortoises, including a 81× rate increase (10% increase in carapace length) in the stem-lineage of recently extinct Mascarene giant tortoises (Cylindraspis spp.), a 137× rate increase (55% increase in carapace length) in the stem-lineage of Aldabra giant tortoises (Aldabrachelys spp.) and a 87× rate increase (383% increase in carapace length) in A. gigantea, and a series of rate accelerations in the ancestral lineages of Galapagos giant tortoises, including a 20× rate increase (19% increase in carapace length) in the stem-lineage of Geochelone and Chelonoidis,
a 20x rate increase (27% increase in carapace length) in the stem-lineage of Chelonoidis, and a 49x rate increase (81% increase in carapace length) in the stem-lineage of C. niger. These data indicate that gigantism evolved independently in multiple lineages of turtles, and step-wise in the evolution of giant tortoises with several rate accelerations in lineages ancestral to C. niger.

Reduction of Intrinsic Cancer Risk in Turtles

In order to account for a relatively constant prevalence of cancer across species (Dorn et al. 1968; Abegglen et al. 2015; Boddy et al. 2020), intrinsic cancer risk must coevolve with changes in body size and lifespan across species. For example, a 100-year retrospective study of neoplasia in zoo reptiles identified only six neoplasms in 490 turtle necropsies, which ranged in size from the West African mud turtle (Pelusios castaneus, carapace length ~25–28 cm) to the spiny softshell turtle (Apalone spinifer spinifer, carapace length ~54 cm) (Sykes and Trupkiewicz 2006). As expected, relative intrinsic cancer risk (RICR) in turtles also varies with changes in body size and lifespan (fig. 2A). We estimated a 73-log2 decreased RICR in the stem-lineage of sea turtles (Chelonioidae), a 129-log2 decreased RICR in leatherback sea turtles (Dermochelys coriacea), a 14-log2 decrease in RICR in the stem-lineage of soft-shell turtles (Trionychidae), a 34-log2 decreased RICR in the stem-lineage of narrow-headed softshell turtles (Chitra chitra and Chitra indica), and a 140-log2 decreased RICR in the stem-lineage of Cantor’s giant softshell turtle (Pelochelys cantorii).

Among the “giant” tortoises, we estimated a 97-log2 decreased RICR in the stem-lineage of Mascarene giant tortoises (Cylindraspis spp.), a 154-log2 decreased RICR in the stem-lineage of Aldabrachelys gigantea (Aldabrachelys spp.) and a 50-log2 decreased RICR in A. gigantea. In the lineages ancestral to Galapagos giant tortoises, we estimated a 67-log2 decreased RICR in the stem-lineage of C. niger a 27-log2 decrease RICR in the stem-lineage of the Chelonoidis, and a 19-log2 decreased RICR in the stem-lineage of Geochelone and Chelonoidis. Thus, turtles coevolved large bodies and reduced intrinsic cancer risk, including step-wise reductions in the lineages ancestral to C. niger.

Identification of Tumor Suppressor and Antiaging Gene Duplications in Turtle Genomes

Previous studies have shown that large-bodied cancer-resistant species such as elephants (Sulak et al. 2016; Vazquez and Lynch 2021) and whales (Keane et al. 2015) evolved an increased number of tumor suppressors, suggesting that the same may be possible in giant, long-lived turtles. A previous study of Galapagos giant tortoises, for example, identified several gene duplications in pathways that might be related to body size evolution and reduced cancer risk (Quesada et al. 2019). Therefore, we reanalyzed the Galapagos giant tortoise genome and other turtle genomes to identify gene duplications and used maximum likelihood-based ancestral state reconstruction (ASR) to determine lineages in which genes were duplicated. We identified ~86 duplications in the stem-lineage of Testudines, 245 in the stem-lineage of Pleurodira, 33 in the stem-lineage of tortoises (AncTortoise), 259 in Chelonoidis abingdonii, 201 in Gopherus agassizii, 273 in Terrapene carolina, 315 in C. picta, and 270 in Pelodiscus sinensis (fig. 3A).

Consistent with previous studies which observed duplication of tumor suppressor and other antiaging genes in large, long-lived species, we found that 12% of the pathways enriched among Galapagos giant tortoises were related to cancer and aging biology, whereas only 0–6% of the pathways that were enriched among gene duplications in other lineages were related to cancer and aging biology (fig. 3B).

Next, we identified GO cellular component terms that were
enriched among gene duplications in each lineage. Although gene duplications in some lineages were enriched in GO terms related to cancer biology and aging, significantly more ontology terms in Galapagos giant tortoises were related to cancer and aging biology (fig. 3C). Enriched pathway and ontology terms in Galapagos (fig. 3D) tortoises included “apoptosis,” “programmed cell death,” “cell death signaling via NRAGE, NRIF and NADE,” “dual incision in GG-NER” and “formation of incision complex in GG-NER,” and “reduction of cytosolic Ca++ levels.” We also observed that “regulation of insulin-like growth factor (IGF) transport and uptake by Insulin-like Growth Factor Binding Proteins (IGFBPs)” was an enriched pathway term among Galapagos giant tortoise gene duplications, which may be related to the regulation of body size. Among the GO cellular component terms exclusively enriched among Galapagos giant tortoise gene duplications were “ER membrane protein complex,” “endoplasmic reticulum membrane” and “nuclear outer membrane-endoplasmic reticulum membrane network,” and “anaphase-promoting complex.”
Desert tortoise-specific gene duplications were enriched in pathways related to cancer biology and aging, particularly compared with other turtles. For example, 9.4% of the pathways enriched among desert tortoise duplicates were related to cancer and aging biology, significantly more than other turtle lineages but less than Galapagos giant tortoises (fig. 3).

Similarly, significantly more GO terms in desert tortoises were related to cancer and aging biology (fig. 3C). Enriched in GO terms included “integral component of mitochondrial outer membrane,” “nucleotide-excision repair complex,” and “autophagosome.” These data suggest that desert tortoises have evolved gene duplications that may also contribute to cancer resistance and the evolution of longevity.

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Similarly, significantly more GO terms in desert tortoises were related to cancer and aging biology (fig. 3C). Enriched pathway and ontology terms in desert tortoises (fig. 3E) were related to DNA damage and repair including “Formation of TC-NER Pre-Incision Complex,” “Gap-filling DNA repair synthesis and ligation in TC-NER,” “Dual incision in TC-NER,” “Apoptotic cleavage of cell adhesion proteins,” and “Transcription-Coupled Nucleotide Excision Repair (TC-NER).” Enriched in GO terms included “integral component of mitochondrial outer membrane,” “nucleotide-excision repair complex,” and “autophagosome.” These data suggest that desert tortoises have evolved gene duplications that may also contribute to cancer resistance and the evolution of longevity.

Turtle Cells Have Unique Responses to Genotoxic and Cytotoxic Stress

Our observation that gene duplications in the Galapagos and desert tortoise genomes are enriched in pathways and GO terms related to the biology of aging, apoptosis, cell cycle regulation, DNA damage repair, and mitochondrial oxidative DNA damage protection (fig. 3D), suggests that cells from these species may have different cellular responses to genotoxic and cytotoxic stress than cells from other turtles. To test this hypothesis, we treated primary fibroblasts from C. niger, Geochelone platynota, Gop. agassizii, Homopus aerolatus, and T. carolina (fig. 2B) with drugs to induce different types of stress including: 1) tunicamycin, which induces ER stress and the unfolded protein response (UPR) through an accumulation of unfolded and misfolded proteins (Banerjee et al. 2011; Guha et al. 2017); 2) etoposide, which forms a ternary complex with DNA and topoisomerase II and prevents religation of replicating DNA strands leading to single- and double-stranded DNA breaks.
We found that tunicamycin induced a dose-dependent increase in apoptosis in cells from most species (fig. 4A). However, at 24 h and 25 μM, C. niger cells had an apoptotic response that was at least double that of other species, whereas G. platynota cells were insensitive to tunicamycin (fig. 4A).

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(Wozniak and Ross 1983); and 3) paraquat, which causes oxidative stress through the production of reactive oxygen species and S-phase cell cycle arrest (Salmon et al. 2008). We quantified the kinetics of cell death using the RealTime-Glo™ Annexin V Apoptosis and Necrosis assay (RTG) every 30 min for 48 h.
necrosis in cells from most species but did induce a strong apoptotic response in Gop. agassizii cells (fig. 4B). Turtle cells were also variably sensitive to paraquat, but cells from all species induced apoptosis in response to paraquat treatment (fig. 4C). Thus, C. niger cells are more sensitive to ER and UPR stress than other species, Gop. agassizii cells are more sensitive to DNA damage induced by etoposide than other species, and cells from all species were sensitive to oxidative stress induced by paraquat.

**Discussion**

Reptiles in general, and turtles specifically, are an excellent system for studying the mechanisms underlying variation in body size, lifespan, and cancer resistance (Chiari et al. 2018; Hoekstra et al. 2020). Adult body size among turtle species differs by three orders of magnitude, ranging from the speckled dwarf tortoise (Chersobius signatus; 100 g) to Aldabra giant tortoises (A. gigantea; >300 kg), whereas longevity across species varies from tens of years to >150 years in giant tortoises. Yet, even the smallest turtles live relatively long (up to 30 years) compared with most other vertebrates. Turtles also have lower estimated cancer rates (~1.2%) compared with mammals (~12.5%), suggesting that they have evolved the means to delay aging and reduce cancer susceptibility (Garner et al. 2004; Sykes and Trupkiewicz 2006; Abegglen et al. 2015; Boddy et al. 2020). Although increased lifespan and cancer prevalence in turtles could, in part, be due to reduced damage resulting from their lower metabolic rates, previous genomic and cellular data suggest that there may also be molecular differences that enable extremes in body size, longevity, and cancer resistance (Goldstein 1974; Quesada et al. 2019).

Here, we found that body size rapidly increased in multiple turtle lineages independently, including a dramatic increase in both body size and the rate of body size evolution in Galapagos giant tortoises. When body size, longevity, and intrinsic cancer rates are analyzed together within an explicit phylogenetic context, large-bodied lineages evolved reduced intrinsic cancer risk. Among the most dramatic decreases in cancer risk among turtles was in the stem-lineage of Galapagos giant tortoises, suggesting this lineage evolved genetic and cellular mechanisms that reduce cancer risk. Consistent with the evolution of reduced intrinsic cancer risk in this lineage, we found that gene duplications in the Galapagos lineage are enriched in ER stress associated pathways is similar to a previous analysis by Quesada et al. (2019), who found that genes under positive selection in Galapagos giant tortoises are also enriched for ER function and stress pathways. ER stress dysregulates protein homeostasis, which is one of the hallmarks of aging and reduced animal lifespan (López-Otín et al. 2013; Morimoto and Cuervo 2014; Kaushik and Cuervo 2015). The impairment of protein production, folding, and degradation associated with ER stress can impact many cellular processes and lead to the buildup of toxic protein aggregates that are linked to many age-related diseases and cancers (Vilchez et al. 2014). Our observations that duplicate genes are enriched in anticancer and antiaging pathways is similar to several previous studies (Caulin et al. 2015; Tollis et al. 2020; Tejada-Martinez et al. 2021; Vazquez and Lynch 2021), suggesting that increased copy number of these genes may be a general trend in large, long-lived species.

Our data suggest a mechanistic connection between genes related to the ER, enhanced responses to ER stress, and the UPR in Galapagos giant tortoise cells. Indeed, we found that tunicamycin, which induces ER stress by activating the UPR and ultimately inducing apoptosis (Nami et al. 2016), causes an immediate and pronounced apoptotic response in Galapagos giant tortoise cells, whereas cells from other species were much slower to respond. These results indicate that Galapagos giant tortoise cells have an extremely sensitive ER stress response. In contrast, Galapagos giant tortoise cells did not exhibit a heightened apoptotic response to paraquat and etoposide, which induce oxidative stress, DNA damage, and apoptosis independently of the ER stress/UPR-signaling pathway (Mizumoto et al. 1994; Suntres 2002). These data suggest that changes in the ER stress and UPR-signaling pathways may contribute to the evolution of long lifespans, large bodies, and augmented cancer resistance in Galapagos giant tortoises.

Unexpectedly, we also found that cells from most species were unresponsive to etoposide treatment, even at the highest dose and longest exposure time. The sole exception was Gop. agassizii cells, which were similar to other species in their response to tunicamycin and paraquat, but markedly more sensitive to etoposide treatment. These data suggest that turtle cells can either export intracellular etoposide before it can induce DNA damage, are generally insensitive to DNA damage induced by etoposide, or can rapidly repair etoposide-induced DNA damage. Regardless of these, and potentially other mechanisms that alter sensitivity to etoposide, Gop. agassizii cells respond differently than cells from the other species tested. The functional and organismal consequences of this altered sensitivity are unclear, but is similar to elephant cells which have evolved to induce apoptosis at relatively low levels of DNA damage (Sulak et al. 2016; Vazquez et al. 2018; Vazquez and Lynch 2021). This reduced threshold for inducing apoptosis in response to cellular stress may clear cells that have been exposed to the kinds of stresses that eventually lead to cancer before transformation into cancer cells.

Although medical literature has often portrayed high levels of apoptosis as a maladaptive response to cellular stress contributing to aging (Szegedi et al. 2006; Chadwick and Lajoie 2019), previous work in elephants showed that heightened apoptotic responses to genetic and cellular damage can actually be adaptive and beneficial response linked to increased body size and longevity (Abegglen et al. 2015; Sulak et al. 2016; Vazquez et al. 2018). This is because rapid
and effective clearance of damaged or injured cells can help maintain tissue integrity (Baar et al. 2017; de Keizer 2017), especially in organisms with many cells and large amounts of cell turnover. Thus, our results are compatible with ER stress as a potential factor in the evolution of large, long-lived turtles.

Caveats and Limitations

There are several limitations to our findings that could be the subject of future research. Although all cellular assays were performed on a common cell type, fibroblasts, the tissue origin of these cells differed among species. We note, however, that for Galapagos giant tortoises, tunicamycin response was not confounded with the site of fibroblast origin. We were also unable to obtain the age of source animals, which could affect results since apoptosis can decline during senescence (Salminen et al. 2011). We also have not classified duplicate genes into functional categories, such as those that have preserved ancestral functions, evolved new ones, or evolved new expression domains because these classifications are extremely difficult to predict based on sequence characteristics alone. Finally, the taxon sampling can be expanded to include cells from closely related species with large differences in body size or lifespan. This would enable better resolution in isolating the evolutionary origins of enhanced responses to genetic size or lifespan. This would enable better resolution in isolating the evolutionary origins of enhanced responses to genetic and cellular stress, for example, by including the closest living relative of Galapagos giant tortoises (Chaco tortoises; Chelonioides chilensis). We relied on primary cell lines that were currently available in frozen zoos and commercial biobanks; but future cellular work could attempt to sample fresh cells from the same tissue type and life stage across species, which is logistically challenging.

Conclusions

Although numerous studies have found comparative genomic signatures associated with the evolution of body size, longevity, and cancer resistance (Keane et al. 2015; Herrera-Álvarez et al. 2021; Babarinde and Saitou 2020), there have been few attempts to validate these findings experimentally at the cellular level (Jimenez et al. 2018). Furthermore, most previous cellular studies on these subjects focus almost exclusively on placental mammals. The work presented here utilizes turtle cell lines, which is much more feasible than studying body size and aging phenotypes in these long-lived animals. Our most salient finding is that Galapagos giant tortoises are much more sensitive at inducing apoptosis in response to ER stress compared with other turtle species, and also have genomic and phylogenetic signatures of rapid evolutionary increases in body size, lifespan, and cancer resistance. We also found more generally that all turtle cell lines were resistant to oxidative stress induced by paraquat. This supports previous oxidative stress studies in turtles, and indicates that turtles, in general, may be a promising model system in which to study resistance to stress from long lifespans (Lutz et al. 2003). However, although the resistance of turtle cells to oxidative stress may contribute to the generally long lifespans of turtles, the sensitivity of Galapagos giant tortoise cells to ER stress may increase their resistance to oncogenic transformation thus promoting healthy aging and larger body sizes.

Materials and Methods

Intrinsic Cancer Risk Estimation

The dramatic increase in body size and lifespan in some turtle lineages, and the relatively constant rate of cancer across species of diverse body sizes and lifespan (Leroi et al. 2003), would predict an increase in cancer risk concurrent with an increase in body size or lifespan. In order to identify lineages with exceptional changes in body size (with total carapace length used here as a proxy for body size), longevity, or intrinsic cancer risk, we jointly estimated these parameters across turtles and reconstructed ancestral states within a phylogenetic framework. Using phylogenetic and body size data from Colston et al. (2020), which includes 357 extant turtle species five species of extinct Cylindraspis, and longevity data from AnAge (Magalhães and Costa 2009), and following Peto’s (2015) model of cancer risk (Vazquez and Lynch 2021), we estimated the intrinsic cancer risk (K) as the product of risk associated with body size and lifespan (lifespan × body size). In order to determine (K) across species and at ancestral nodes, we first estimated body size at each node. We used a generalization of the Brownian motion model that relaxes assumptions of neutrality and gradualism by considering increments to evolving characters to be drawn from a heavy-tailed stable distribution (the stable model) implemented in StableTraits (Elliot and Mooers 2014). The stable model allows for large jumps in traits and has previously been shown to out-perform other models of body size evolution, including standard Brownian motion models, Ornstein–Uhlenbeck models, early burst maximum likelihood models, and heterogeneous multirate models (Prang 2019). We used phylogenetic generalized least-square regression (Grafen and Hamilton 1989; Martins and Hansen 1997; Pagel 1997) using a Brownian covariance matrix as implemented in the R package ape (Paradis and Schliep 2019) to infer ancestral lifespans across turtles using our estimates for body size (Colston et al. 2020) and reported maximum lifespans for each species (Magalhães and Costa 2009). Fold-change in cancer susceptibility was estimated between all ancestral (K1) and descendant (K2) nodes. The fold-change in cancer risk between a node and its ancestor was then defined as K2/K1 (supplementary fig. 2—source data 1, Supplementary Material online).

We note that two variables that are correlated with each other, such as body mass and lifespan, will be correlated with the product of these variables, such as intrinsic cancer risk. Thus, we may have reached similar conclusions about
changes in inferred cancer risk across turtle evolution using either body size or lifespan data alone. But, we would have not inferred the magnitudes of those changes correctly for most species and may have inferred the wrong direction for some species that decrease dramatically in body size but increase dramatically in lifespan (although there are few such species).

Identification of Duplicated Genes and Reconstruction of Ancestral Copy Numbers

Following Caulin (2015), we identified duplicated genes in the genomes of the Pinta Island Galapagos giant tortoise (C. abingdonii; ASM359739v1), Goode’s thornscrub tortoise (Gopherus evgoodei; rGopEvg1_v1.p) and Agassiz’s desert tortoise (Gop. agassizii; ASM289641v1), Chinese softshell turtle (Pelodiscus sinensis; PelSin_1.0), painted turtle (Chrysemys picta bellii; Chrysemys_picta_bellii-3.0.3), three-toed box turtle (T. carolina triunguis; T_m_triunguis-2.0), chicken (Gallus gallus; GRCg6a), and eastern brown snake (Pseudonaja textilis; EBS10Xv2-PRI) using the Ensembl (Genes 103) BioMart web-based tool to extract same-species paralogies (within_species_paralog) from each genome; same-species paralogies are identified by Ensembl Compara (Howe et al. 2021) using gene tree species tree reconciliation. Specifics of the Ensembl Compara (Howe et al. 2021) method can be found in the Ensembl Compara documentation: https://useast.ensembl.org/info/genome/compara/index.html. Genes that were classified as pseudogenes on Ensembl were not included. Duplicate genes identified in Goode’s thornscrub tortoise (Gop. evgoodei; rGopEvg1_v1.p) were manually verified in Agassiz’s desert tortoise (Gop. agassizii; ASM289641v1) using reciprocal best BLAT using default parameters: https://useast.ensembl.org/Help/View?id=451#toc6. Interested readers are referred to these documents for specifics about the Compara and BLAST/BLAT methods as a complete description is beyond the scope of this manuscript.

We used maximum likelihood-based ASR to determine when in the evolution of turtles each gene was duplicated. We encoded the copy number of each putatively functional gene for each species as a discrete trait, with state 0 for one gene copy and state 1 for two or more copies. We used IQ-TREE to select the best-fitting model of character evolution (GTR2) with character state frequency optimized (FO) by maximum-likelihood from the data. We next inferred gene duplication events with the empirical Bayesian ASR method implemented in IQ-TREE (Kalyaanamoorthy et al. 2017; Minh et al. 2020), the best-fitting model of character evolution (GTR2+FO), and the unrooted species tree for turtles (fig. 3A). We considered ASRs to be reliable if they had Bayesian posterior probability (BPP) greater than or equal to observed state frequency from the alignment; less reliable reconstructions were excluded from further analyses.

Gene Duplication Pathway Enrichment Analysis

To determine if gene duplications were enriched in particular biological pathways, we used Enrichr (Chen et al. 2013, p. 5; Kuleshov et al. 2016) to perform overrepresentation analysis (ORA) of the Reactome database. Furthermore, we used the Panther GO enrichment analysis tool (Mi et al. 2021) to perform ORA on GO cellular component terms (Ashburner et al. 2000; Gene Ontology Consortium 2021). Gene duplicates in each lineage were used as the foreground gene set, and the initial query set was used as the background gene set. Enrichr uses a hypergeometric test for statistical significance of pathway overrepresentation, whereas the Panther GO enrichment analysis tool uses a binomial test for statistical significance of GO cellular component term overrepresentation.

Cell Culture

Experimental cellular phenotypes were generated to compare results from the body size, longevity, and intrinsic cancer risk analysis and the gene duplication and enrichment analysis. Apoptosis was chosen as the major cellular endpoint of interest because of its central role in aging and cancer through the removal of damaged or cancerous cells (Salminen et al. 2011; Verfaillie et al. 2013).

We cultured cell lines from four tortoise taxa (C. niger, G. platynota, Gop. agassizii, H. aerolatus) obtained from the San Diego Frozen Zoo and one turtle species (T. carolina) obtained from the American Type Culture Collection (table 1). All cells were primary fibroblasts derived from either the heart (T. carolina), trachea (Gop. agassizii, G. platynota, C. niger), or eye (H. aerolatus). Turtle cells have previously been shown to grow within a range of 23–30 °C (Clark and Karzon 1967; Clark et al. 1970; Goldstein 1974). Therefore, cells were incubated at 25 °C, with 5% CO2. Cells were cultured in minimum essential medium (MEM; Gibco) with 10% fetal bovine serum (Gibco) and 1% penicillin–streptomycin antibiotic (Gibco) in standard T75 flasks (Thermo Fisher Scientific). Media was changed every 3 days. Cells were passaged before reaching 90% confluency, approximately every 7–9 days. For passaging, cell plates were rinsed with one volume of 37 °C DPBS (Gibco) and cells detached with 0.25% trypsin-EDTA (Gibco). We note that cells detached quickly with the assistance of gentle tapping of plates and without incubation. The cell suspension was transferred to a 15-ml conical tube (Thermo Fisher Scientific) with an equal volume of complete media to stop trypsinization. Cells were then centrifuged at 500×g for 5 min, and then the pellet was resuspended in 1 ml of complete media. Cell viability was determined using a TC10 Automated Cell Counter (Bio-Rad Laboratories) and was >75% for all cell lines throughout the experiment.
We selected various cytotoxic drugs to induce different types of cellular stress, including etoposide (Cayman Chemical Company), which induces single- and double-stranded DNA breaks (Wozniak and Ross 1983), paraquat (Sigma–Aldrich), which induces oxidative stress through production of reactive oxygen species and s-phase cell cycle arrest (Salmon et al. 2008), and tunicamycin (Cayman Chemical Company), which induces ER stress and the UPR by causing an accumulation of unfolded and misfolded proteins (Banerjee et al. 2011; Guha et al. 2017).

Kinetic Measurements of Cell Death

The RTG visualizes the kinetics of apoptosis over a given period of time and differentiates secondary necrosis occurring during late apoptosis from necrosis caused by other cytotoxic events (Landreman et al. 2019). RTGs were performed for each species by seeding 5,000 cells per well into an opaque bottomed 96-well plate with three replicates per treatment per species and one empty column with no cells (background control). Cells were left to adhere for 24 h, after which seeding media was aspirated off and serial dilutions of Tunicamycin (0, 10, 50, 100 µM), Etoposide (0, 100, 500, 1,000 µM), or Paraquat (0, 100, 1,000, 5,000 µM) were applied. All drug treatments were made using Fluorobrite DMEM media. The “0 µM” control treatments consisted of the vehicle used (DMSO or PBS) at the concentration matching the highest drug concentration, whereas the background control (NoCell) consisted of the drug treatment and assay reagents with no cells. As per the RTG product protocol, the 500-fold dilution of reagents in Fluorobrite DMEM media was added to wells immediately after drug treatments were applied (effectively reducing the initial drug dilutions in half). Readings were then taken every 30 min for 48 h using a GloMax Luminometer (Promega Corporation).

Statistical Analysis

All statistical analyses reported in this paper are estimation statistics, including effect sizes, 95% confidence intervals (CIs) of the effect size, and P values. Effect sizes and 95% CIs are reported as: effect size (CI width lower bound; upper bound) with 5,000 bootstrap samples; the CI is bias-corrected and accelerated. P values reported are the likelihood of observing the effect sizes if the null hypothesis of zero difference is true. For each permutation P value, 5,000 reshuffles of the control and test labels were performed (Ho et al. 2019). All statistical analyses were performed in RStudio (RStudio Team 2020).

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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Author Contributions

S.G.: Conceptualization, Funding Acquisition, Investigation, Methodology, Project Administration, Supervision, Visualization, Writing—Original Draft Preparation. S.E.B.: Data curation, Formal Analysis, Investigation, Writing—Original Draft preparation. J.M.V.: Data curation, Formal Analysis, Investigation, Visualization, Software, Writing—Review and Editing. Y.C.: Conceptualization, Funding Acquisition, Investigation, Project Administration, Supervision, Writing—Review and Editing. V.J.L.: Conceptualization, Formal Analysis, Funding Acquisition, Investigation, Methodology, Project Administration, Supervision, Visualization, Software, Writing—Original Draft Preparation.
Data Availability

All data are either publicly available or provided as Supplementary Materials and Data Sets.

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