DNA mutations in somatic cells have been implicated in the causation of aging, with longer-lived species having a higher capacity to maintain genome sequence integrity than shorter-lived species. In an attempt to directly test this hypothesis, we used single-cell whole-genome sequencing to analyze spontaneous and bleomycin-induced somatic mutations in lung fibroblasts of four rodent species with distinct maximum life spans, including mouse, guinea pig, blind mole-rat, and naked mole-rat, as well as humans. As predicted, the mutagen-induced mutation frequencies inversely correlated with species-specific maximum life span, with the greatest difference observed between the mouse and all other species. These results suggest that long-lived species are capable of processing DNA damage in a more accurate way than short-lived species.

RESULTS
Spontaneous somatic SNV and INDEL frequencies
In view of the high cost of whole-genome sequencing of multiple cells from multiple species, we selected four rodent species with the largest differences in life span: mouse (Mus musculus), guinea pig (Cavia porcellus), blind mole-rat (Nannospalax galli; i.e., Upper Galilee mountains blind mole-rat), and naked mole-rat (Heterocephalus glaber), as well as human (Homo sapiens), with maximum life spans of approximately 4, 12, 21, and 37 years for the rodents, respectively, and...
115 years for human (10–13). These specific rodent species were also selected because of the availability of relatively high-quality reference genome sequences (table S1).

To identify somatic mutations and accurately estimate their frequency, single-cell whole-genome sequencing is necessary, because each single cell acquires its unique set of somatic mutations during the lifetime of the organism. We performed single-cell whole-genome sequencing using the single-cell multiple displacement amplification method that we developed earlier (see Materials and Methods) (6). This method substantially reduces potential artificial mutations occurring during the whole-genome amplification process. We applied the method on early passage, primary lung fibroblasts from each species, isolated from young adults (Fig. 1A) (14), with single-cell amplicons that passed our locus dropout test selected for whole-genome sequencing (table S2) (6). We also sequenced DNA of bulk fibroblast populations of the same individual animals/humans to filter out germline variations. Whole-genome sequencing was performed using the Illumina HiSeq X Ten or NovaSeq platform, reaching an average sequencing depth of 27.6x across the genome (table S3). Somatic SNVs and INDELs of each cell were identified after filtering out germline variants identified from bulk DNA based on an updated version of SCcaller (see Materials and Methods) (6), capable of identification of INDELs in addition to SNVs.

Somatic mutation frequency was defined as the ratio of the number of somatic mutations to the number of base pairs covered in the genome with at least 20x sequencing depth, with adjustments for genome ploidy (i.e., diploid) and mutation calling sensitivity (see Materials and Methods and table S4). To determine de novo spontaneous somatic mutation frequency, we analyzed whole-genome sequences of each of three cells per individual animal/human participant from a total of 12 individuals, 4 animals for the mouse, and 2 for each other species. As shown in Fig. 1B, the median spontaneous somatic SNV frequency of mouse single cells [3.6 × 10⁻⁷ base pair (bp⁻¹)] is about twofold higher than that of any other species, i.e., guinea pig, blind mole-rat, naked mole-rat, and human (a median SNV frequency of 1.7 × 10⁻⁷, 0.9 × 10⁻⁷, 1.4 × 10⁻⁷, and 1.7 × 10⁻⁷ bp⁻¹ for the four species, respectively; P = 0.0061, two-tailed Student’s t test), among which no significant differences were found. Notably, these results for mouse and human cells confirm the previously observed difference between somatic mutation frequencies in dermal fibroblasts from these two species (15). The median frequencies of spontaneous somatic INDELs (Fig. 1C) were found to be about four to five times lower than those of SNVs but showed a similar pattern of a significantly higher frequency in mouse cells (5.7 × 10⁻⁸ bp⁻¹) than in cells of the other species (4.6 × 10⁻⁸, 3.3 × 10⁻⁸, 3.4 × 10⁻⁸, and 2.6 × 10⁻⁸ bp⁻¹ for guinea pig, blind mole-rat, naked mole-rat, and human cells, respectively). Notably, it occurred to us that inbred mice might have different somatic mutation rates than outbred animals, which is why we also analyzed cells from two four-way cross mice (UM-HET3). We did not observe a significant difference between mutation frequencies in lung fibroblasts from the two mouse strains (P = 0.535 and P = 0.155 for SNVs and INDELs, respectively). These results indicate a significantly higher spontaneous mutation frequency in lung fibroblasts from the mouse than in the cells of same type from any of the other species analyzed; among the latter, no significant differences were found.

### Spontaneous somatic SNV and INDEL spectra

We then studied the somatic mutation spectra for possible differences between cells of these species. The most obvious species-to-species difference was found between mouse and the other species (Fig. 2A). Mouse fibroblasts had significantly more somatic SNVs at A/T bases than other species: 12, 26, and 27% of total somatic SNVs in mouse cells are T > A, T > C, and T > G, respectively, while only 9, 17, and 8% of somatic SNVs in cells of the other species are of these types (P = 0.004, P = 7.7 × 10⁻⁵, and P = 1.23 × 10⁻⁹, respectively, two-tailed Student’s t test). Notably, the most significant difference was in T > G transversions. We did not observe significant differences in mutation spectra between the inbred and outbred mouse strain. Notably, the above differences reflect the fractions of mutations of the total number of mutations in the cells. In absolute numbers, the mutations at C bases are also different between species. For example,
mouse cells have the highest absolute numbers of spontaneous C > T mutations (100 per cell on average), which are likely due to deamination of 5-methylcytosine, followed by guinea pig (69 per cell; \( P = 0.115 \), two-tailed Student’s \( t \) test compared to mouse), blind mole-rat (65 per cell; \( P = 0.035 \)), naked mole-rat (52 per cell; \( P = 0.008 \)), and human (87 per cell; \( P = 0.379 \)). These numbers inversely correlate with the maximum life span within the rodent group.

By taking the two flanking bases of each mutation into consideration (Fig. 2B), we found somatic SNVs in mice, but not in the other species, to occur more frequently at TT bases. A similar mutational pattern of TT bases has been observed in human lymphoid cells [i.e., signature 9 in the Catalogue of Somatic Mutations in Cancer (COSMIC) database], which was considered the result of the error-prone DNA polymerase \( \varepsilon \) during somatic hypermutation (16). This same mutational signature was previously observed in our study of somatic mutations in normal human memory B lymphocytes, which suggests a possible shared mechanism (17). Because a mutation spectrum is a linear combination of one or more mutational signatures, we can compare our mutation spectra with known mutational signatures using correlation coefficients. On the basis of cosine similarity, the mutation spectrum found in mouse cells has the highest correlation with signature 9 (cosine correlation = 0.76), while its correlations with other COSMIC signatures are all less than 0.7. In addition, the correlations between the spectra of cells of the other four species and signature 9 are only in the range of 0.49 to 0.58.

To further test whether difference in DNA polymerase \( \varepsilon \) gene, \( Polh \), is a cause of above unique mutation spectrum in mouse, we compared the protein sequences of the \( Polh \) orthologs from the five species, using multiple sequence alignment by Clustal Omega (18) and neighbor joining tree by Jalview (19). The mouse ortholog was

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Fig. 2. Mutational spectra of spontaneous somatic mutations. (A) Mutational spectra of spontaneous somatic SNVs. (B) Mutational spectra of spontaneous somatic SNVs in the context of their two flanking base pairs. This classified mutations into 96 categories. On the x axis, the 96 categories were sorted according to their alphabetic order. For example, the first bar from the left indicates ACA to AAA mutation, and the first from the right indicates TTT to TGT mutation. The y axis indicates the fraction of each category out of the total number of mutations. (C) A phylogeny tree of protein sequences of the \( Polh \) orthologs of the five species. Multiple sequence alignment was performed using Clustal Omega, and the phylogeny tree was calculated using the neighbor joining algorithm using Jalview based on the PID (i.e., the percentage identity between the two sequences at each aligned position) score, which indicates the number of identical residues per 100 residues. (D) Mutational spectra of spontaneous somatic INDELs. Error bars in (A) and (D) indicate SD.
found to be the outlier (Fig. 2C), which is in obvious contrast with the species phylogeny. Although we cannot exclude other underlying causes, these results suggest that DNA polymerase eta may be an important factor in the elevated mutation frequency in mouse somatic cells.

For spontaneous somatic INDELs, the mutational spectra between different species were much more similar than those of somatic SNVs. As shown in Fig. 2D, 1-bp deletion accounts for about 50% of total INDELs across cells of all species, with mouse fibroblasts having the most. The second most common INDEL is 1-bp insertion, accounting for about 20% of these events across species. This indicates that, for somatic INDELs, unlike somatic SNVs, species specificity in mutation accumulation is only observed for frequency but not spectrum.

**Bleomycin-induced somatic SNVs and INDELs**

To further investigate the capability of maintaining genome sequence integrity in the different rodent and human species, we analyzed mutation frequencies and spectra in the primary lung fibroblasts after treatment with three doses of bleomycin: 10, 40, and 80 μM (see Materials and Methods). Bleomycin has a cytotoxic effect and can induce DNA single-strand break and DNA DSB with a ratio of approximately 6:1, which may vary depending on the dose (20, 21). Small INDELs and SNVs are induced as a consequence of errors during DNA DSB repair (22). After bleomycin treatment, the cells were allowed to grow for two population doublings (PDs) to allow repair of the DNA damage and fixation of the mutations (Fig. 1A). A substantial difference between the species was already apparent in their proliferation abilities after the bleomycin treatment. Although bleomycin delayed cell growth in cells from all species (fig. S1, A to E), mouse cells were not even able to reach two PDs after 40 or 80 μM bleomycin treatment. Mouse cells were found to carry a substantially higher proportion (approximately 40%) of apoptotic cells compared to cells from the longer-lived species or humans under the same treatment conditions (fig. S2, A to D).

To determine somatic mutation frequency, we then isolated single fibroblasts for the three doses of bleomycin treatment for all individual animals/humans of all five species studied (for mouse only after 10 μM bleomycin) and performed whole-genome amplification, sequencing, and mutation calling as described above and in Materials and Methods. We first analyzed the resulting dataset using a linear mixed-effects model: Mutation frequency was modeled as a function of (i) bleomycin dose (fixed effect), (ii) the ranking of species-specific life span (mouse, 1; guinea pig, 2; blind mole-rat, 3; naked mole-rat, 4; human, 5) (fixed effect), and (iii) animal/human individual (random effect; see Materials and Methods). We found both bleomycin dose and the life-span rank to be significant factors affecting somatic SNV frequencies (P = 0.011 and P = 0.010, respectively, ANOVA; Fig. 3A) and somatic INDEL frequencies (P < 0.0001 and P = 0.0052, respectively, ANOVA; Fig. 3B). In addition, as with spontaneous mutations, we did not observe a significant difference between the two mouse strains in response to bleomycin treatment (P = 0.832 and P = 0.137 for SNVs and INDELs, respectively, sANOVA).

The cell-to-cell variation in mutation frequency appeared to be generally higher in mouse, guinea pig, and blind mole-rat than in the naked mole-rat or human. Examples of outliers found among cells of the first three species are indicated in Fig. 3 (A and B). Although the number of cells analyzed is not sufficient to draw firm conclusions, these results may suggest that mutation frequency among cells from longer-lived species is more stable than in those of shorter-lived species and therefore also point toward greater genome maintenance capacity.

We also analyzed mutational spectra but did not observe significant differences in spectra before and after bleomycin treatment (fig. S3, A and B). This strongly suggests that mechanisms involved in repair of bleomycin-induced DNA damage are similar to those acting upon spontaneous DNA damage.

We then tested whether the extent of the bleomycin-induced mutation frequency increase also inversely correlated with species-specific life span. Specifically, we estimated the elevated mutation frequency per 1 μM bleomycin for each species separately, using linear regression models, and compared them with the life-span rankings of the species. The results show that, for somatic SNVs, mouse cells acquired the highest number of mutations per 1 μM bleomycin, with guinea pig cells the second highest, and cells from the other species jointly at the lowest level (Fig. 3C and fig. S4A). We tested whether such a pattern of negative correlation with species-specific life span was statistically significant by a permutation test, randomizing the species that each animal/human individual belongs to. After 2000 repeats, we found that not a single repeat reflected such a pattern, corresponding to a P < 0.0005 (permutation test; 2000 repeats; one-tailed). For somatic INDELs, a similar pattern was observed with the exception that blind mole-rat cells were about the same as guinea pig cells (P = 0.0035, permutation test; 2000 repeats; one-tailed; Fig. 3D and fig. S4B). The results above indicate that genome maintenance accuracy is the highest in somatic cells of humans and naked mole-rat, followed by those of blind mole-rat and guinea pig, with that in mouse cells the lowest (Fig. 3E). Notably, the largest interspecies difference remained between mice and all other species, as was found for spontaneous mutations. This could be interpreted either in terms of the mouse as a unique species, different from all others, or as a possible plateau in maintaining genome sequence integrity readily reached in longer-lived species.

**DISCUSSION**

In a study conducted well before the advent of high-throughput sequencing, germline mutation rates during evolution were estimated from interspecies DNA sequence differences analyzed by thermal stability of DNA hybrids or neutral base pair substitutions in the relatively few coding regions sequenced. The results indicated different rates of DNA change among different phylogenetic groups, with the slowest rates observed for higher primates and faster rates for short-lived rodents, including mouse, rat, and hamster (23). These results are unlikely to be explained by differences in generation time and may instead reflect less-accurate DNA replication systems in shorter-lived rodents than in longer-lived primates (23, 24). Within primates, the rate of DNA sequence change in the human lineage compared to other primate lineages is the lowest. This so-called hominoid slowdown (25) coincides with the longevity increase during evolution of primates (26). Hence, the evidence suggests that genome instability as controlled by accuracy of replication and repair is a major factor in determining species-specific life span.

Evidence that germline mutation rate is a factor in the aging process has also come from studies within the human species. Recently, age-adjusted mutation rates were determined in 61 women and 61 men from the Utah CEPH (Centre d’Etude du Polymorphisme
Humain) families, with higher mutation rates notably associated with higher all-cause mortality in both sexes and a shorter reproductive life span in women (27).

Germline mutation frequencies are not necessarily the same as somatic mutation frequencies. While it is reasonable to assume that the selection processes responsible for varying germline mutation rates between species have the same effect on somatic mutation rates, the latter have been shown to differ between tissues within one species (28). Moreover, somatic mutation frequencies, corrected for the difference in the number of cell divisions per generation, has been shown to be at least an order of magnitude higher than the germline mutation frequency in both mouse and human (15). Unfortunately, somatic mutation burdens are different from cell to cell and cannot be assessed by bulk sequencing. For this reason, we previously developed a single-cell mutation analysis method (6). Single-cell or clone whole-genome sequencing provides not only accurate estimation of mutational frequency per cell (17, 29–31) but also the future possibility of studying interactions among mutations within a cell, which may also have substantial functional effects relevant to aging and/or age-related diseases (32, 33).

In this present work, we directly studied spontaneous and mutagen-induced mutation frequency in primary lung fibroblasts from short- and long-lived rodent species and in human cells of the same type. The results show that cells from short-lived mice have a significantly higher frequency of somatic mutations than the same cell type from longer-lived rodents or humans. On the basis of our observation of excess mutations at TT bases in the mouse cells alone and its association with DNA polymerase eta, we found that the latter is a phylogenetic outlier among the species studied. Hence, the observed increase in SNV frequency in mouse cells can reasonably be explained, at least in part, by the more error-prone DNA polymerase eta, of which the protein sequence in mouse deviates
from that in all other species. While this could still be longevity-related and DNA polymerase eta may also play a role in SNV induction in other similarly short-lived rodent species, the main conclusion remains that the life-span differences between the other species do not correlate with differences in spontaneous mutation rates.

Stress-related aging and longevity differences are often only uncovered after challenging animals or cells. Here, we treated the primary lung fibroblasts from each species with the mutagen bleomycin to provide such a challenge. The results confirm that, in mouse cells, bleomycin-induced mutation frequency is also significantly higher than in cells of other longer-lived rodent species or in human cells. This also could be explained in part by the more error-prone polymerase eta. However, for bleomycin-induced mutations, we find induced mutation loads to correlate with species-specific life span across all species tested. This result cannot be explained by polymerase eta alone and suggests that other mechanisms contribute to lower mutation frequency in long-lived animals. Multiple mechanisms could be involved from enhanced enzymatic function of DNA repair enzymes to elevated expression levels. Expression of genome maintenance genes has been found to correlate with life span in different rodent species (34, 35). Genome maintenance is complex, and it will be very difficult to tease out the species-specific pathways responsible for variation in mutagenic response. Meanwhile, this first evidence that accuracy of maintaining genome sequence integrity is correlated with species-specific life span could serve as a model for more extensive studies. These studies will become feasible once the currently still high-sequencing costs have come down and robust reference genomes are available for most rodent species.

**MATERIALS AND METHODS**

**Animal subjects and human individuals**

All rodent experiments were performed according to procedures approved by the University of Rochester Committee on Animal Resources. In this study, we used cells from four mice (M. musculus), including two inbred mice (C57BL/6; 6 months old) from the Gorbunova laboratory and two four-way cross mice (UM-HET3; 4 months old) from the Gladyshev laboratory. Two guinea pigs (C. porcellus), two blind mole-rats (N. galili; i.e., Upper Galilee mountains blind mole-rat), and three naked mole-rats (H. glaber) were used in this study, which were young adults available in the Gorbunova laboratory. On the basis of body size and weight, they are also young adults. Two frozen human (H. sapiens) primary fibroblast samples were obtained from the American Type Culture Collection (ATCC) (HS-A; 14 years old) and Lonza (HS-B; 12 years old). All species are approximately matched by specific fractions of their maximum life span (age at about 10% of the maximum life span).

**Lung fibroblast isolation and cell culture**

Primary lung fibroblasts were isolated following a cell isolation protocol adapted from Seluanov *et al.* (14). Briefly, the lung from each rodent species (young adult) was minced and incubated in Dulbecco’s modified Eagle’s medium (DMEM) F-12 medium with Liberase Blendzyme 3 (0.13 U/ml) and 1× penicillin/streptomycin (P/S) at 37°C for 40 min. Dissociated cells were washed, plated in cell culture dishes with complete DMEM F-12 medium and 15% fetal bovine serum (FBS), and cultured at 37°C, 5% CO₂, and 3% O₂ (32°C for naked mole rat). When reaching confluence, cells were split and replated in Eagle’s minimum essential medium (EMEM; ATCC, 30-2003) supplemented with 15% FBS and 1% P/S (100 U/ml). Lung fibroblasts were purified by further passaging in the same medium. Human lung fibroblasts were obtained from ATCC and Lonza. The human cells were cultured in the complete fibroblast growth medium (FGM; Lonza, CC-3132) as recommended by the instructions.

**Bleomycin treatment**

Early-passage primary lung fibroblasts were cultured in EMEM (ATCC, 30-2003) with 15% FBS and P/S (100 U/ml). When reaching confluence, cells were starved in low-serum EMEM (0.1% FBS) for cell cycle synchronization at 37°C (32°C for naked mole rat cells) for 24 hours. Then, the cells were washed twice with phosphate-buffered saline (PBS) and treated, in plain EMEM (with 1% P/S), with different doses of bleomycin (0, 10, 40, and 80 μM) at 37°C (32°C for naked mole rat cells) for 1 hour. The bleomycin (EMD Millipore Corporation, 203401-10MG) was dissolved in nuclease-free water. The cells were trypsinized, harvested, and counted after treatment. Then, the cells were replated at 0.2 million as the starting number in the complete EMEM, allowing DNA damage repair and mutation fixation over two rounds of cell replications. Notably, human cells were starved in fibroblast growth basal medium (Lonza, CC-3131) with 1% P/S without supplemented FBS. The complete FGM was used for human cell culture.

**Cell replication and apoptosis**

Cell growth was monitored by counting cell numbers with the Cellometer Auto T4 (Nexelom Bioscience) twice or thrice until cells reaching two PDs after replated. The cell PD time was estimated on the basis of the cell growth curve for each species under different doses of bleomycin treatment. The results of PDs are shown in fig. S1.

The cell apoptosis was detected using the Guava Annexin Red Kit (Luminex, FCCH100108) and Guava easyCyte flow cytometers (Millipore) according to the manufacturer’s instructions. The percent of apoptotic cells was analyzed by GuavaSoft software with flow cytometer. Briefly, the cells were harvested at 0, 24, 48, 72 and 96 hours after cell treatment and replating. The cells were counted and prepared at a cell concentration of approximately between 2 × 10⁵ and 1 × 10⁶ cells/ml. Then, 100 μl of each cell solution was transferred into one well of 96-well plate and mixed with 100 μl of annexin reagent from the kit. The mixture was incubated at room temperature for 20 min in the dark. Then, the plate with samples was loaded in a flow cytometer, and the program “Nexin assay plus” was run. All the annexin V-positive cells were calculated as apoptotic cells. The percentage of apoptotic cells from different species at each time point was shown in fig. S2. The setting of the Nexin program and analysis software for samples from different species, conditions, or time points was consistent.

**Single-cell isolation**

The cells were harvested when the number reached two PDs and prepared for single-cell isolation. Single-lung fibroblasts were isolated using the CellRaft array (Cell Microsystems), as described previously (6). Briefly, we first wetted an array by adding 2 ml of cell culture medium and removing after 3 min, which was repeated three times. We then added approximately 5000 fibroblasts in 3 ml of medium to the array and incubated at 37°C, 3% O₂, and 10% CO₂. After 3 hours, the fibroblasts elongated and attached to the array. We then removed the medium and washed the array with the
attached fibroblasts twice with 1 ml of PBS. We then added 3 ml of fresh complete medium. We identified the rafts in the array that contained one cell by microscopy. Using a magnetic wand supplied with the CellRaft system, we then transferred each of the rafts with one cell into a 0.2-ml polymerase chain reaction (PCR) tube containing 2.5 µl of PBS. We validated with a magnifier that each tube contained a single raft, froze them on dry ice, and kept them at −80°C until usage.

**Single-cell whole-genome amplification**

We amplified the isolated single cells using the Single-Cell Multiple Displacement Amplification protocol reported previously (6). Briefly, for each single cell, we added 1 µl of exo-resistant random primer (Thermo Fisher Scientific) and 3 µl of lysis buffer (400 mM KOH, 100 mM dithiothreitol, and 10 mM EDTA) and incubated on ice for 10 min. We neutralized the lysis buffer by adding 3 µl of stop buffer [400 mM HCl and 600 mM tris-HCl (pH 7.5)]. We then added 32 µl of master mix containing 30 µl of multiple displacement amplification reaction buffer and 2 µl of Phi29 polymerase (REPLI-g UltraFast Mini Kit, QIAGEN), incubated for 1.5 hours at 30°C and 3 min at 65°C, and held at 4°C until purification. We purified the ampiclings using AMPure XP beads (Beckman Coulter) and quantified DNA concentration with the Qubit High-Sensitivity dsDNA Kit (Thermo Fisher Scientific). Simultaneously, we amplified 1 ng of genomic DNA in 2.5 µl of PBS as positive control and 2.5 µl of PBS without any template as negative control. We performed the locus dropout test as described previously (6) with primers designed for each species separately (table S2). Three single-cell ampiclings per individual per experimental condition that passed the locus drop-out test were prepared for whole-genome sequencing.

**DNA library preparation and whole-genome sequencing**

Sequencing libraries of the bulk DNA and single-cell ampiclings were constructed using the TruSeq Nano DNA HT Sample Prep Kit (Illumina) by our laboratory or by Novogene Inc. The libraries were purified using AMPure XP beads (Beckman Coulter) and subjected to quality control using Bioanalyzer 2100 (Agilent) and real-time PCR. The libraries were sequenced on the Illumina HiSeq X Ten or NovaSeq S4 sequencing platforms for $2 \times 150$-bp paired-end reads by Novogene.

**Bulk DNA extraction**

Bulk DNA, i.e., DNA of bulk fibroblast populations, of each individual was extracted from primary lung fibroblasts before the bleomycin treatment experiments, using the DNeasy Blood and Tissue Kit (QIAGEN) following the manufacturer’s instructions. We determined their concentration with the Qubit High-Sensitivity dsDNA Kit (Thermo Fisher Scientific) and their quality with 1% agarose gel electrophoresis.

**Sequence alignment**

For raw sequencing reads of each single-cell or bulk DNA sample, we performed adapter and quality trimming using Trim Galore (version 0.6.4). Reads before and after trimming were subjected to quality control using FastQC (version 0.11.8). We aligned the trimmed sequences to their species-specific reference genomes (see table S1 for the version of reference genomes used) using Burrows-Wheeler Aligner-Maximal Exact Match (BWA-MEM, version 0.7.17) (36) and removed PCR duplicated sequences using SAMtools (version 1.9) (37).

To perform INDEL realignment and base quality score recalibration, known INDELs and single-nucleotide polymorphisms (SNPs) are required but are not available for species except for human and mouse. To handle this, we followed the Genome Analysis Toolkit (GATK) instructions to use INDELs and SNPs called from our data: We called INDELs and SNPs using GATK (version 3.5.0) (38) HaplotypeCaller from the sequence alignment (obtained above) of each bulk DNA sample and used them as a database for INDEL realignment and base quality score recalibration for all bulk DNA and single-cell samples, which were done also using GATK (version 3.5.0). We kept chromosome contigs with a minimum length of 1 Mbp for mutation analysis to avoid potential bias in mutation calling, in short, incompletely assembled contigs in some species. The resulted sequence alignments were used for finding somatic mutations as described below.

**Calling somatic SNVs and INDELS**

We identified somatic SNVs and INDELS, i.e., SNVs and INDELS observed only in a single cell and not in its corresponding bulk DNA by SCcaller (version 2.0.0) that we developed for variant calling of single cells previously (6). The version 2.0.0 of SCcaller was based on the same principle of its 1.0.0 version (for SNV calling only), with the additional function to call INDELS (here, we focused on 1 to 10 bp of small INDELS) in single-cell sequencing (freely available at https://github.com/biosimodx/Scaller). Briefly, we first called heterozygous germline SNPs using the GATK HaplotypeCaller and used them in SCcaller to correct for potential amplification bias occurred during the single-cell whole-genome amplification. We then used SCcaller to call somatic autosomal SNVs and INDELS only observed in a single cell but not in its corresponding bulk, requiring 20× depth in both the samples. For INDELS, we additionally required a genotype calling quality of ≥30 and a maximum size of 10 bp to ensure the identification of true mutations instead of false positives. Germline heterozygous SNPs and INDELS of the same single cells, i.e., those mutations also present in bulk DNA sample, were also identified using SCcaller under the same criteria. This enabled us to estimate the sensitivity of variant calling, defined as the ratio of number the germline heterozygous SNPs or INDELS observed in the single cells to the number of germline heterozygous SNPs or INDELS observed in their corresponding bulk DNA sequences. Because the number of germline heterozygous SNPs and INDELS is limited in the inbred mouse strain, C57BL/6, sensitivity in these cells is much harder to estimate than that for noninbred subjects or strains. So, we approximated the sensitivities of C57BL/6 mouse cells as the average sensitivity of all the other single cells. Somatic SNV and INDEL frequencies were determined after correcting the sensitivity, the genome coverage ($≥20×$ and autosomal requirements), and the ploidy of a single-fibroblast genome (table S4).

**Analyzing mutation frequency with linear mixed-effects regression**

When analyzing mutation frequency as a function of bleomycin dose and rank of species-specific life span (mouse, 1; guinea pig, 2; blind mole-rat, 3; naked mole-rat, 4; human, 5), we used a linear mixed-effect regression model: bleomycin dose and rank of life span with fixed effects and different individuals within each species with random effects. This was performed using the “lme” function in the R package “nlme”: lme(mutation freq ~ life-span rank + bleomycin dose, random = ~1|individual) (39).
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