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Telomouse—a mouse model with humanlength telomeres generated by a single amino acid change in RTEL1

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Telomeres, the ends of eukaryotic chromosomes, protect genome integrity and enable cell proliferation. Maintaining optimal telomere length in the germline and throughout life limits the risk of cancer and enables healthy aging. Telomeres in the house mouse, *Mus musculus*, are about five times longer than human telomeres, limiting the use of this common laboratory animal for studying the contribution of telomere biology to aging and cancer. We identified a key amino acid variation in the helicase RTEL1, naturally occurring in the short-telomere mouse species *M. spretus*. Introducing this variation into *M. musculus* is sufficient to reduce the telomere length set point in the germline and generate mice with human-length telomeres. While these mice are fertile and appear healthy, the regenerative capacity of their colonic epithelium is compromised. The engineered Telomouse reported here demonstrates a dominant role of RTEL1 in telomere length regulation and provides a unique model for aging and cancer.

Telomeres maintain genome stability and enable cell proliferation by repressing the DNA damage response at the ends of linear chromosomes¹. Human telomeres in most somatic cells shorten with each cell division due to incomplete replication and degradation². In the germline, however, telomeres are elongated by the enzyme telomerase to maintain them within a defined, species-specific range referred to as the "telomere length set point"³. The telomere length set point allows for sufficient, though not unlimited, cell divisions in the soma throughout life. Optimal age-appropriate telomere length is critical for maintaining genome stability and regulating cell proliferation and senescence, the major determinants of carcinogenesis and aging^{4–8}. However, despite decades of research, it is not yet understood how telomere length homeostasis is achieved. The large

difference (-five-fold) in telomere length between two closely related and inter-breedable species of mice, *Mus musculus* and *M. spretus*, was exploited previously to search for genes that determine speciesspecific telomere length^{9–11}. Interestingly, none of the known telomere maintenance genes were identified in these screens. Rather, the difference in telomere length between the two species was attributed to a single locus encoding a DNA helicase, which was therefore termed *'Regulator of Telomere Length'* (*Rtel1*)^{9,11}. However, the specific difference between the *M. musculus* and *M. spretus* RTEL1 proteins that is responsible for the different telomere length set points has not been identified thus far¹².

RTEL1 belongs to a small family of iron-sulfur helicases together with XPD, FANCJ, and ChlR1, which were implicated in the human

¹Department of Genetics, The Silberman Institute of Life Sciences, Safra Campus, The Hebrew University of Jerusalem, Jerusalem 91904, Israel. ²Department of Genetics and Institute for Diabetes, Obesity and Metabolism, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA 19104, USA. ³Division of Gastroenterology and Hepatology, Department of Medicine, Hospital of the University of Pennsylvania, Philadelphia, PA 19104, USA. ⁴Department of Pharmacology and Toxicology, Massey Cancer Center, Virginia Commonwealth University, 401 College St, Richmond, VA 23298, USA. ⁵Present address: Faculty of Biotechnology and Food Engineering, Technion, Haifa 3200003, Israel. e-mail: kaestner@pennmedicine.upenn.edu; tzfati@mail.huji.ac.il inherited diseases Xeroderma pigmentosum, Fanconi anemia, and Warsaw breakage syndrome, respectively^{13,14}. Biallelic germline mutations in human *RTEL1* cause a fatal telomere biology disease named Hoyeraal-Hreidarsson syndrome (HHS)^{15–19}. HHS is characterized by accelerated telomere shortening and diverse clinical symptoms including bone marrow failure, immunodeficiency, and developmental defects²⁰. Other HHS-causing mutations were found in telomerase subunits, or in factors essential for telomerase biogenesis or recruitment to telomeres²¹. While RTEL1 was reported to have multiple roles in maintaining the stability of the genome, and particularly telomeres^{22–27}, it was also suggested to regulate nuclear trafficking of ncRNA²⁸, the localization of telomeric repeat-containing RNA (TERRA)²⁹, and telomere elongation by telomerase^{26,30}.

One of the *RTEL1* missense mutations identified in HHS patients results in a single amino acid change from the highly conserved methionine 492 to isoleucine (Fig. 1a, b)¹⁷. Methionine 492 is conserved nearly uniformly across vertebrates, with only a few species (mostly cats) having a leucine at this position. *M. spretus* is the only vertebrate species having a more radical change at this site, namely to the positively charged amino acid lysine¹⁷. We hypothesized that lysine 492 in *M. spretus* might be responsible for the difference in telomere length between the two mouse species. Here we describe an engineered mouse that confirms this hypothesis and presents an invaluable model for the study of cancer and aging.

Results

Methionine 492 is located at a critical position within the helicase core domain of RTEL1

Methionine 492 of RTEL1 is conserved in vertebrates but found to be altered in human HHS patients and in the short-telomere mouse species *M. spretus*¹⁷. To understand the role of this residue in RTEL1 function, we generated a structural model of the murine RTEL1 helicase domain using the Protein Homology/analogY Recognition Engine V 2.0 (Phyre 2)³¹ and then modeled the complex with single-stranded DNA based on the X-ray crystal structure of the bacterial XPD family helicase DinG in complex with the DNA³². This model revealed that methionine 492 (M492) is buried within the core of the helicase domain in proximity to the single-stranded DNA binding site of the protein, making extensive hydrophobic contacts with the hydrophobic side chains of residues M464, V468, F488, and I494 (Fig. 1c, d)³². Replacing methionine 492 with the larger and positively charged lysine within this hydrophobic patch is predicted to perturb the local structure (Fig. 1e), which in turn may affect the helicase DNA binding, translocation, and unwinding activities of the protein.

A house mouse with short telomeres

To test if the lysine residue at position 492 of *M. spretus* is responsible for its shorter telomeres compared to the house mouse M. musculus, we generated a M. musculus strain in which methionine 492 was changed to a lysine (M492K) using CRISPR-Cas9 assisted genome editing in zygotes. We employed the Cas9 nickase (D10A Cas9 mutant) enzyme and two guide RNAs to generate nicks on both strands of the Rtell gene to facilitate gene conversion using a repair template (Fig. 1f). Successful gene targeting in the resulting offspring was confirmed by DNA sequencing and by allele-specific Taqman probe-based PCR (Fig. 1g, h). This founder mouse was backcrossed to wild-type (WT) C57BL/6 mice to generate F0 mice heterozygous for the Rtel1^{M492K} allele, termed *Rtel1^{M/K}*. FO *Rtel1^{M/K}* mice were intercrossed to generate F1 and F2 *Rtel1^{M/K}*. Homozygosity for the *Rtel1* mutation (*Rtel1^{K/K}*) was established in the third generation (F3). The homozygous *Rtel1*^{K/K} mice were then bred inter se for thirteen additional generations up to F16 to date and were analyzed for telomere length and function. We termed the *Rtel1^{K/K}* mouse model 'Telomouse'.

To determine the effect of the *Rtel1*^{M492K} mutation on telomere length, we employed two experimental paradigms. First, we

established embryonic fibroblast cultures from *Rtel1^{M/K}* and *Rtel1^{K/K}* F3 embryos generated by intercrossing F2 Rtel1^{M/K} mice (Fig. S1). As a control, we employed WT (Rtel1^{M/M}) embryonic fibroblasts generated from intercrossing WT mice which did not have any Rtel1 mutation in their pedigree, to avoid trans-generational inheritance of short telomeres from the heterozygous mice to the WT offspring, as occurred in the fibroblasts derived from the progeny of F2 Rtel1 mutant mice (M/ M** in Fig. S2a). Mouse embryonic fibroblast (MEF) cultures were established by serial passaging and selection for spontaneous immortalization and followed in culture over 250 population doublings (PD; Fig. S1). During growth, the cultures passed one or two transition points in which the MEFs increased their growth rates and changed their cellular morphology, presumably by random inactivation of a tumor suppressor gene or genes such as p16 (Fig. S1 and ref. 33). During the final growth phase, *Rtel1^{M/K}*, *Rtel1^{K/K}*, and *Rtel1^{M/M}* MEFs grew at the same average rate (about one PD/day). We determined the average telomere lengths and the telomeric 3'-overhang length for these three cultures at multiple PD levels by the in-gel hybridization method as follows. We digested high molecular weight genomic DNA samples with the restriction endonuclease Hinfl. separated the restriction fragments by pulse-field gel electrophoresis (PFGE) and dried the gel. First, we hybridized the native DNA in the gel to a telomeric C-rich probe to estimate the average length of the G-rich singlestranded telomeric overhang (see below). Then, the DNA was denatured and hybridized again to the same probe to detect both singleand double-stranded telomeric repeat tracts and estimate the length of the telomeric restriction fragments (TRF; representative gel images are shown in Figs. 2a, S2 and S3). The mean TRF length (MTL) of each sample was repeatedly measured in different gels to increase the measurement precision (Summarized in Supplementary Data 1 and Fig. 2b). The average MTL in WT (*Rtel1*^{M/M}) MEFs was 46.9 kb at PD 10. It shortened to about 40 kb during the initial growth phase (Figs. S1 and 2b), presumably due to the lower telomerase activity present in MEFs prior to immortalization. Indeed, MEF immortalization was reported previously to associate with increased telomerase activity³⁴. Upon immortalization at about PD 70 (indicated by arrows in Fig. 2a, b). telomere length stabilized, reaching an average MTL of 37.8 kb at PD 250. In contrast, the mutant *Rtel1^{M/K}* and *Rtel1^{K/K}* MEFs displayed shorter telomeres at PD 10 (average MTL 34.2 and 30.1 kb, respectively), which continued to shorten gradually even beyond the early growth phase at an average rate of about 50 bp per PD, to 21.6 and 14.7 kb at PD 250, respectively (Fig. 2a, b). Altogether, the average MTL in the Rtel1K/K MEFs shortened by 60% as compared to the WT TRF length after 250 PD and reached a length comparable to that of human fibroblasts immortalized by ectopic expression of telomerase³⁵.

RTEL1 dysfunction was proposed previously to accelerate telomere shortening either by telomere deletion or by reducing telomerase action at telomeres^{25,26,30}. To investigate the cause for telomere shortening in *Rtel1^{K/K}* MEFs, we examined the distinct TRF banding pattern appearing in the *Rtel1^{M/K}* and *Rtel1^{K/K}* MEFs (Figs. 2a, S2 and S3). Such banding pattern was observed previously in telomerase null MEFs³⁶, as well as in mRtel1-deficient mouse embryonic stem cells²⁶. These types of bands were suggested to represent homogenous telomere lengths of individual chromosomes that became clonally fixed within the cell population because they did not undergo significant elongation despite the presence of telomerase²⁶. Furthermore, these TRF bands showed gradual telomere shortening over time in culture, indicating the inability of telomerase to compensate for the telomere shortening that occurs with each round of cell division²⁶. Based on this observation, Uringa and colleagues hypothesized that mRtel1 was required for telomere elongation by telomerase. Similarly, *Rtel1^{M/K}* and Rtel1K/K MEFs maintained several such sharp bands, which gradually shortened with continued passaging (Figs. S2 and S3), indicating that telomerase activity was insufficient to maintain stable telomere length in these cells. Calculating the rates of shortening of individual bands



reveal faster shortening of up to 170 bp per PD during the initial growth phase, and slower shortening of up to 66 bp per PD during the later phase in both heterozygous and homozygous mutant MEFs (Figs. S2b, d and S3d). WT MEFs also displayed sharp bands during the initial growth phase (Figs. 2a, S2e, f and S3c, d). However, these bands did not appear to shorten but rather disappeared upon

immortalization, presumably due to the increased telomerase activity associated with immortalization of MEFs³⁴. These findings indicate that the shortening of telomeres in *Rtel1^{K/K}* and *Rtel1^{M/K}* MEFs was not due to random telomere deletion, which would have caused these bands to disappear, but rather the result of reduced telomerase action at telomere ends.

Fig. 1 | **The RTEL1 helicase domain. a** Map delineating the domains of RTEL1. The N-terminal part of the protein includes helicase domains 1 and 2 (HD1 and HD2), the iron-sulfur coordinating cluster (FeS) and the ARCH domain. The C-terminal part includes the nuclear localization signal (NLS), two harmonin N-like domains (HNL1 and HNL2), the PCNA interacting protein motif (PIP), and a C4C4 ring-finger domain. b Protein sequence alignment showing position 492 with a conserved methionine in *M. musculus* and human (mutated to isoleucine in HHS), and a lysine in *M. spretus.* **c** Predicted structure model of the *M. musculus* RTEL1 helicase domain, formed by residues 3–777 of mRTEL1 in a complex with single-stranded DNA. **d** Enlargement of the helicase domain in the red box in (**c**), showing the contacts of M492 with several hydrophobic residues and distances in Å. **e** Replacing M492 with the larger and charged lysine and overlaying the energy-minimized WT

(green) and mutant (light brown) structures reveals changes in the position of nearby residues, which may affect the interaction with the adjacent single-stranded DNA. **f** Derivation of an *Rtel1*^{M492K} allele using the CRISPR-Cas9 nickase editing strategy. Illustration showing the WT allele, the two positions targeted by gRNAs (blue and red arrowheads), the ssDNA repair template with the mutation, left homology arm (blue), right homology arm (red) and the resulting targeted allele. **g** Sanger sequencing of founder #15 confirming the replacement of the "T" with an "A" at the codon encoding methionine 492, changing it to a lysine (underlined). **h** Genotyping of animals using a custom Taqman SNP Genotyping Assay. Fluorescence intensities of AAG (K) and ATG (M) are plotted for WT (M/M; blue), heterozygotes (M/K; green) and homozygotes (K/K; red) samples, and no-template control (NTC; yellow).



Fig. 2 | **Telomeres of MEFs carrying the** *Rtel1*^{M492K} **mutation shorten progressively with cell division in culture. a** Genomic DNA samples, prepared from WT (M/M) or mutant mouse embryonic fibroblasts (MEF) cultures homozygous (K/K) or heterozygous (M/K) for *Rtel1*^{M492K} at different population doublings (PD), were analyzed repeatedly by PFGE and in-gel hybridization to the denatured DNA. Shown are representative gels for each phenotype out of nine gels performed. The unspliced image of the gels are shown in Fig. S2a, c, e. Mean telomeric restriction fragment length (MTL) values indicated below the lanes were calculated using

*TeloTool*²¹. **b** All MTL values measured in all the gels for each sample (summarized in Supplementary Data 1) were plotted (M/M in blue, M/K in green, and K/K in red). Segmental linear regression was used to draw linear regression lines with a breakpoint set manually to PD 60, because it gave the best fit (R squared values are indicated for each graph). The rate of telomere shortening is indicated above each graph. N.S. indicates no significant difference from a horizontal line (slope 0). The complete change of cell morphology, reflecting cellular immortalization (see Fig. S1c), is indicated by arrows on the gel images (a) and graph (b).

It was proposed previously that the actual length of the telomeric repeat tracts in mouse telomeres is much shorter than the length of the TRF observed by gel electrophoresis due to low sequence complexity and limited representation of restriction sites in the subtelomeric regions^{36,37}. However, the exact length of the subtelomere included in the TRF is still unknown and may vary between chromosome ends and different restriction enzymes used to digest the genomic DNA and generate the TRF. To try to estimate the average length of the

subtelomeric regions within the TRF we examined the correlation between the hybridization signal and MTL. Since equal amounts of genomic DNA were loaded for each sample on the gels and longer telomeres bind more probe, we expected the denatured hybridization signal within each gel to correlate with telomere length. Plotting MTL as a function of the telomeric hybridization signal indeed revealed a linear correlation (Fig. S4). Extrapolation of the regression line to zero signal did not reach zero TRF length as expected if the TRF sequence were entirely telomeric, but rather suggested that the observed TRFs are longer than the telomeric repeat tracks by an average of 10–15 kb (Fig. S4). This analysis confirmed that at least some chromosome ends have long subtelomeric sequences of low complexity that are underrepresented for *Hinfl* restriction sites, contributing to the measured TRF length. Therefore, the actual lengths of the telomeric repeat tracts are significantly shorter than appears by PFGE, as proposed previously^{36,37}. This observation raised the concern that the lengths of TRFs do not reflect accurately the lengths of the telomeric repeat tracts.

To directly compare the actual length of the telomeric repeat tracts between mouse and human telomeres, we used Quantitative Fluorescence In Situ Hybridization (qFISH). We mixed metaphase-arrested *Rtel1^{K/K}* MEFs at PD 250 with human telomerase-positive fibroblasts, spread them onto slides and hybridized them to telomeric (green) and centromeric (red) peptide nucleic acid (PNA) probes (Figs. 3a and S5). *Rtel1^{M/M}* control MEFs prepared and hybridized under the same conditions are shown for comparison in Fig. 3b. The MTL values as calculated by gel electrophoresis for the *Rtel1^{K/K}* MEFs and the human fibroblasts were 14.7 kb and 14 kb, respectively (Figs. 2b and





and *Rtell^{K/K}* MEFs (red). Additional pairs are shown in Fig. S5. For the human samples, between 116 and 324 chromosome ends were quantified per field. For the mouse samples, between 178 and 248 chromosome ends were analyzed per field. The higher number of telomeres per field than the expected 184 (human) and 160 (mouse) may result from more than one metaphase combined in the same field or from aneuploidy. Horizontal lines indicate the median. Unpaired two-tailed t-test showed significant differences in the telomeric signals between the *Rtel1^{K/K}* MEFs PD 250 and human fibroblasts only in pairs 1 and 5. The *P* values are indicated above each pair. A nested one-way ANOVA test for all seven pairs indicates significantly lower signal intensity in the MEFs (P = 0.003). **d** Genomic DNA was extracted from the same cultures shown in (**a**) and (**b**) and analyzed by regular (constant field) gel electrophoresis and in-gel hybridization to the native and denatured DNA.

3d). The centromeric probe enabled us to distinguish the human chromosomes, which are mostly metacentric, from the acrocentric mouse chromosomes. The telomeric signals at each chromosome end were quantified using the Telometer plugin (available at https:// demarzolab.pathology.jhmi.edu/telometer/index.html) of NIH *Imagel*³⁸. The average telomeric signal intensity of the *Rtel1*^{K/K} MEFs at PD 250 was comparable or lower than that of the human telomeres (Figs. 3a, c and S5a, d). In addition, the distribution of individual telomeric signals was significantly wider in the MEFs with a significant portion of telomeres below the detection limit (Fig. 3c). Most of these telomeres displayed very weak hybridization signals, detectable only under prolonged exposure and increased gain, indicating that they were significantly shortened but not completely lost (Fig. S5e). Moreover, regular (constant field) gel electrophoresis followed by in-gel hybridization revealed that a significant fraction of TRF in the Rtel1K/K MEFs PD 250 was shorter than 5 kb (Fig. 3d). In sum, these data are consistent with previous publications suggesting that long subtelomeric regions are included in the TRF due to low sequence complexity and that a significant distance exists between the telomeric repeat tracts and the most distal restriction site^{36,37}. The position of the restriction site likely varies between chromosome ends, thus contributing to the heterogeneity of TRF lengths observed by PFGE. As the telomeres of the *Rtel1^{K/K}* MEFs shorten, the error introduced by the variable subtelomeric portion in the TRF measured by PFGE becomes even more significant in determining the actual length of the telomeres. In addition, some TRF with short subtelomeric regions become so short that they are excluded from the size range detected by PFGE. These conclusions highlight the need for a more accurate method for measuring the actual length of the telomeric repeat tracts.

NanoTelSeq-a novel method for single telomere analysis by long-read Nanopore sequencing

It is well established that the shortest telomeres are the most biologically relevant as they may activate the DNA damage response and determine cellular function and fate³⁹. However, as apparent from the results detailed above, currently used methods based on measuring hybridization signals or telomeric restriction fragments fail to measure the shortest telomeres accurately. A method called TeSLA was designed to PCR-amplify and measure the shortest telomeres⁴⁰. However, it is labor intensive, has significant limitations, and is practically impossible to use for analyzing the ultra-long telomeres of M. musculus. To address this major shortfall, we developed a method for single telomere analysis by long-read Nanopore sequencing and termed it NanoTelSeq. We modified the previously developed single telomere length analysis (STELA) method⁴¹. In STELA, a specific telorette linker with a C-rich telomeric repeat base-pairs with the G-rich telomeric overhang and is ligated to the 5' end of the telomere to enable PCR amplification of the telomere using primers complementary to the telorette oligonucleotide and to a known subtelomeric sequence⁴¹. In our method, a telorette oligonucleotide is ligated in the same way to the 5' end of the telomere but instead of being used for PCR amplification it enables the specific attachment of the Nanopore sequencing adapter to the 5' end of the telomere (Fig. 4a). This telomere endspecific ligation step enriches the sequencing reads for telomeres while avoiding the risk of biased recovery during physical isolation of telomeric DNA fragments. The sequencing adapter enables the sequencing of the telomeric C-rich strand directly from 5' to 3' without any PCR amplification. Since the ligation of the telorette linker requires an intact telomeric 3' overhang, the presence of the telorette immediately at the 5' end of a telomeric sequence and a subtelomeric sequence at the 3' end of the telomere sequence reveals the complete length of the telomeric repeat tracts of individual telomeres. Using NanoTelSeq we sequenced undigested high molecular weight genomic DNA samples prepared from the *Rtel1^{K/K}* and *Rtel1^{M/M}* MEFs at PD 250 as well as human telomerase-positive skin fibroblasts and obtained

several hundreds of telomere reads each (Fig. 4). We first employed a mixture of six telorettes with the six possible permutations of the six nucleotides composing the telomeric repeat at their 3' end. As reported for human telomeres⁴¹, telorette 3 was the most frequently used (60% of the *Rtel1*^{M/M} and 76% of the *Rtel1*^{K/K} and the human telomeric reads), indicating that the 5' strand of most telomeres ends in 3'-CCAATC-5' (Fig. 4b). This specificity can only be expected if the telorette adapter was attached to native telomere ends and not to randomly broken telomeres, thus ensuring that most telomere reads represent intact telomeres. Therefore, for all additional sequencing reactions we employed only telorette 3, which was attached to different barcodes for different samples sequenced together. For long reads that begin with the telorette sequence on the 5' end of the telomeric C-rich strand and terminate with a subtelomeric sequence on the 3' end, we could therefore determine the length of the telomere repeat sequences at base resolution. Telomeric reads were identified in the sequencing output data, processed and presented graphically by a dedicated computer program termed Telomere Analyzer (examples for telomere reads are shown in Fig. 4c). The median read length (RL) of the sequencing reads selected by Telomere Analyzer was 20-27 kb in our various samples (Fig. S6). Since the length of the mouse telomeres is within the same range as the read length, some reads were too short to reach the subtelomere (referred to as 'truncated telomeres'; Supplementary Data 2 and 3), introducing an error in median or average length estimation. Simply discarding truncated telomeres would also introduce a bias due to overrepresentation of shorter complete telomeres. This bias is expected to be greater for the longer and more heterogeneous mouse telomeres than for the shorter human telomeres. To circumvent this obstacle, we examined filtering of the reads based on the read length, excluding all reads below a certain length cutoff regardless of whether the telomere is complete or not. Since the samples differ in telomere length, we could not apply the same length cutoff to all samples and sought to develop a uniform criterion for choosing the optimal cutoff that would vield the largest sample size with minimal bias. To this end, we explored the influence of the read length cutoff on average and median telomere length calculation. We chose the five samples with the largest number of telomeric reads and sorted each sample based on the read length from largest to smallest (Supplementary Data 3). We then calculated 'running' median and 'running' average telomere lengths, meaning that for each telomere read we calculated the median and average telomere length of all reads of equal or greater length (Supplementary Data 3). The 'running' median and 'running' average telomere lengths were plotted as a function of the number of included telomeric reads sorted by decreasing length (Fig. S7). As seen in the graphs, the more telomere reads are included, the smaller the fluctuations in the running median and running average values calculated. However, as expected, including shorter reads eventually reduces the calculated median and average values. The median telomere length values declined significantly only when reads shorter than the corresponding running median were included (Fig. S7, dotted lines). Therefore, we chose reads longer than the running median for calculating median telomere length (Fig. S6, right plot for each sample). The calculated average telomere length may decline earlier, as the inclusion of any truncated telomere introduces an error (Fig. S7).

Using these criteria, we estimated a median telomere length of 5.6 kb for the *Rtel1^{K/K}* MEFs at PD 250, which is 4.9 times shorter than the telomeres of WT MEFs at PD 250 (median telomere length of 27.3 kb), and also shorter than the telomeres of human telomerase-positive fibroblasts (median 9.47 kb; Figs. 4d and S6). Furthermore, 19.5% (40 of 205) of the telomeres of the *Rtel1^{K/K}* MEFs PD 250 were shorter than 3 kb, and 6.3% (13 of 205) were below 2 kb, while only 0.17% (one of 602) of the telomere reads of the *Rtel1^{M/M}* MEFs were shorter than 3 kb and none were shorter than 2 kb (Fig. 4d). Nanopore sequencing thus enables the determination of telomere length at base



short telomeres in the *Rtel1^{K/K}* MEFs. a A scheme showing a telorette oligonucleotide complementary to the G-rich telomeric overhang ligated to the 5' end of the telomere and then to the Nanopore sequencing adapter. b High molecular weight genomic DNA samples prepared from Rtel1^{K/K} and Rtel1^{M/M} MEFs PD 250 and human telomerase-positive fibroblasts (S2 fibroblasts³⁰) were ligated to a mixture of equal amounts of six telorette oligonucleotides, each ending with a different permutation of the C-rich 6 nucleotide telomeric repeat, and then ligated to the Nanopore sequencing adapter and sequenced, each sample separately. The bar graph shows the percentage of the telomeric reads ligated with each telorette in M/ M MEFs PD250 (blue), K/K MEFS PD 250 (red), and human fibroblasts (green). c The

Distribution of telomeric 5' ends 100 90 Human fibroblasts KIK MEFS % of telomeric reads 80 70 <u>=868</u> Telomere 5'-end 60 1 CAATCC-5 50 TCCCAA-5 2 40 CCAATC-5 CCCAAT-5 30 5 ATCCCA-5' 20 AATCCC-5 10 3 2 1 2 4 5 6 2 3 4 5 6 1 - 3 Telorette no.

Length of individual telomeres

b



reads were identified and processed computationally, and presented in graphs of telomeric repeat density (from 0 to 1, within a moving window of 50 nucleotides). Two representative examples are shown and the other reads are summarized in Supplementary Data 2. Indicated are the precise telomere length and read length. d Scatter plots show the length of individual telomeres of the M/M (blue) and K/K (red) MEF cultures at PD 250 and telomerase-positive human fibroblasts (green) for comparison (the same cultures are shown in Fig. 3), with median and quartiles indicated by horizontal lines. Median values in kb are indicated to the right of each scatter plot and the percentage of telomeres below 3 kb and 2 kb to the left of the plots.

short telomeres are functional. The main role of telomeres is to sup-

resolution of all telomeres, long and short. Most importantly, it enables the identification and analysis of very short telomeres, which are not represented in the standard PFGE TRF or qFISH analyses but are nevertheless of utmost importance in determining cell function and fate.

NanoTelSeq also enabled us to determine the telomere lengths at specific mouse chromosome ends using the subtelomeric sequence to map them to the reference genome (Fig. S8). We were only able to map reads to the ends of the long q-arm of the mouse acrocentric chromosomes since the subtelomeres of the p-arms contain repetitive sequences that are not present in the reference genome. Our mapping revealed a large degree of heterogeneity in telomere lengths within and between chromosomes (Fig. S8), the significance of which remains to be investigated in the future.

Functional short telomeres in Rtel1^{K/K} MEFs

The unexpected finding of telomeres below 3 kb (19.5%) and even 2 kb (6.3%) in the *Rtel1^{K/K}* MEFs at PD 250 raised the question whether these press the DNA damage response (DDR) at chromosome ends and to prevent deleterious fusions of chromosomes and cell cycle arrest¹. The presence of DDR foci at telomeres, termed telomere dysfunctioninduced foci (TIF), is a hallmark of a telomere failure to protect the chromosome end. To examine if the telomeres of the Rtel1K/K MEFs retained their protective properties, we studied the presence of TIF using antibodies against the DDR marker yH2AX and the telomere protein TRF1 (Fig. 5a). The overall levels of DDR were high in all MEF cultures, as reported previously for MEFs growing under atmospheric oxygen levels⁴². However, there was no significant difference in the number of genome-wide DDR or TIF foci between WT (M/M) and mutant (K/K) MEFs (Fig. 5a). To identify more accurately all the DDR foci associated with chromosome ends (even those without a detectable telomeric signal) we combined FISH and Immunofluorescence (IF) on metaphase chromosomes (Fig. 5b). This analysis again revealed no significant difference in TIF formation between the genotypes. Furthermore, there was no increase in DDR even at those chromosome



PD

Fig. 5 | The short telomeres of *Rtel1*^{K/K} MEFs largely retain their protective function. a The formation of DDR foci and their localization to telomeres (defined as telomere dysfunction-induced foci, TIF) in interphase nuclei were evaluated by IF with antibodies detecting the DDR marker γ H2AX (red) and the telomere protein TRF1 (green). Scatter plots show the number of DDR foci and TIF per nucleus. A total of 127 and 111 nuclei were counted for M/M and K/K MEFs (PD 102), respectively, in three independent experiments. The mean values are indicated by black lines, and the *P* values for genomic and telomeric damage were calculated by a two-tailed nested t-test. Scale bars equal 40 μ m. **b** Metaphase TIFs were analyzed by

PD

FISH-IF using a γ H2AX (red) antibody and a telomeric C-probe (green). A total of 100 metaphases were counted for each M/M (PD 261) and K/K (PD 250) MEF population in two independent experiments. The mean values are indicated by black lines, and the *P* values for damage at all telomeres and only at signal-free ends were calculated by a two-tailed unpaired t-test. Scale bars equal 5 μ m. **c** The normalized native overhang signal for 77 samples was measured and plotted as described in Fig. S9. The slopes for M/M (blue), M/K (green), and K/K (red) were analyzed by simple linear regression and *P* values were calculated for the deviation from a horizontal line.

PD

ends lacking detectable telomeric hybridization signal ('signal-free ends'; Fig. 5b), indicating that most *Rtel1*^{K/K} telomeres, even the very short ones that are below the limit of detection by FISH, remained functional and able to suppress the DDR.

Shortened telomeric overhangs were found to be associated with telomere dysfunction in HHS patients and heterozygous carriers of the pathogenic M492I mutation in *RTELI*^{17,30,43}. To test whether the *RtelI*^{M492K} mutation affects telomeric overhang maintenance in MEFs, we quantified the native in-gel hybridization signal and normalized it to a WT sample (M/M PD 10) as a measure of the relative average length of the telomeric overhang. The native signal remained constant and comparable to that of WT fibroblasts, indicating no significant impact of the *RtelI*^{M492K} mutation on the overhang maintenance (Figs. 5c and S9).

RTEL1 dysfunction in patient cells and mouse *Rtel1* null cells was reported to induce telomere aberrations observed by telomere FISH on metaphase chromosomes, such as telomere loss, telomere fragility, interstitial telomere insertion (ITS) or telomere fusion^{17,25,30}. To examine the frequency of telomere aberrations, we performed metaphase FISH on MEF cultures at PD 100 and 250 (Fig. 6a–c). The main phenotypes observed were telomere loss and Robertsonian fusions, which increased from PD 100 to 250 in association with the shortening of the telomeres (Fig. 2). Most undetected telomeres by PFGE or qFISH (Figs. 2a and 3a, c) actually had weak telomeric signal observed under increased exposure (see long arrows in Fig. S5e), in line with the short telomeres measured by *NanoTelSeq* (Fig. 4d). The few telomeres without any detectable signal, even under increased FISH exposure,



Fig. 6 | **The chromosomes of** *Rtell*^{K/K} **MEFs display increased telomere loss and Robertsonian fusions. a** Representative images of metaphase chromosomes of M/ M and K/K MEFs at PD 250 hybridized with telomeric (green) and centromeric (red) probes. Scale bars equal 50 μm. Arrows indicate chromosomal aberrations. **b** Aberration types are shown enlarged: telomere loss (TL) (yellow arrows), telomere fragility (blue arrows), interstitial telomere insertions (ITS; red arrows), Robertsonian fusions and other fusions (white arrows). **c** Scatter plots show the average number of each aberration type per metaphase in a field (normalized to the number of chromosomes counted) at PD 100 (top) and PD 250 (bottom). Horizontal lines show mean and standard deviation. Two independent experiments for each M/M PD 100 and K/K PD 100 (*n* = 20 metaphases each experiment), and three

independent experiments for each M/M PD 250 and K/K PD 250 (n = 35 metaphases each experiment) were quantified. The *P*-values indicated on the graphs for each aberration were calculated by two-tailed paired t-test. **d** Chromatid-Orientation (CO)-FISH was performed on K/K and M/M MEFs at PD 105 and PD 104, respectively, labeling the leading and lagging telomeres with green and red using strand-specific PNA probes. Scale bars equal 5 μ m. Two experiments were done, each with M/M and K/K (n = 20 metaphases each). **e** Telomere sister chromatid exchange (T-SCE) events (indicated by arrows in **d**) were counted and presented as a percentage of all telomere ends available for analysis in each metaphase. Horizontal lines show mean

underwent chromosome fusion (short arrows in Fig. S5e). Presumably, Robertsonian fusions between the ends of the short p arms (but not the long q arms) of two chromosomes are tolerated by the cells because of the proximity of the acrocentric centromeres to the p end, and can accumulate unlike other deleterious fusions (Fig. 6b, c). Finally, telomere sister chromatid exchange (T-SCE), indicating elevated telomeric recombination, was reported in RTEL1-deficient patient cells¹⁶. We examined the frequency of T-SCE events in the mutant cells by chromatid-orientation (co)-FISH, differentially labeling leading and lagging telomeres with green and red strand-specific PNA probes. T-SCE events appear as colocalization of the two probes at the same telomere end (Fig. 6d). There was no increase in T-SCE in the K/K mutant compared with the WT MEFs, indicating no increase in homologous recombination at the telomeres of the mutant K/K MEFs. Altogether, although the K/K MEFs displayed dramatically shortened telomeres, these telomeres largely maintained normal telomeric overhangs, which are essential for telomere end structure and

function, and retained the ability to suppress the DDR and homologous recombination. Only Robertsonian fusions significantly accumulated in the K/K MEFs due to increased occasional telomere loss.

and standard deviation. P-value was calculated by two-tailed unpaired t-test.

Short telomeres in Telomice

Telomeres of the *Rtel1*^{K/K} MEFs shortened over time in culture, as described above. However, telomere shortening in cultured fibroblasts does not necessarily imply reduced telomere length in the germline or in somatic tissues of mice. Therefore, we next analyzed telomere length in blood and tail DNA samples taken from *Rtel1*^{K/K} Telomice over 13 generations (F4 to F16). In the blood, the MTL shortened to an average length of 16.7 kb in F16 Telomice, as compared to 39.7 kb in the WT samples (Supplementary Data 4 and Fig. 7a, c). In tail samples, MTL shortened to an average of 19.2 kb in F16 Telomice as compared to 42.6 kb in the WT samples (Supplementary Data 4 and Fig. 7b, d). Importantly, when comparing the average TRF lengths of only the last two generations, F15 and F16, no further shortening was observed,



Length of individual telomeres



Fig. 7 | **Telomouse telomeres progressively shorten over generations.** Genomic DNA samples extracted from blood leukocytes (**a**) or tail (**b**) from mutant (K/K) or WT (M/M) mice at the indicated generations and ages were analyzed by PFGE and in-gel hybridization to the denatured DNA. Representative gels are shown. MTL for each generation was repeatedly measured in **c** blood samples (n = 40 for M/M and n = 99 for K/K in 12 gels) and **d** tail samples (n = 34 for M/M and n = 166 for K/K in 19 gels) and plotted in red for K/K and blue for M/M. The average age for K/K mice analyzed in the gels was 331 days, while the average age for M/M mice was 395 days. All data are summarized in Supplementary Data 4. e Scatter plots show the length of

е

individual telomeres in the indicated mouse blood and tail samples (K/K in red and M/M in blue) and human blood sample (green) as determined by *NanoTelSeq*, with median and quartiles in horizontal lines. Median values in kb are indicated to the right of each scatter plot and the percentage of telomeres below 3 kb and 2 kb to the left of the plots. The details of all individual telomere reads are shown in Supplementary Data 2. The samples were taken from the following mice indicated by their ID numbers and ages (Supplementary Data 4): WT male, 6 months (M-C58), K/K F10 male, 13 months (12820), and K/K F15 male, 6 months (2426).

neither in blood nor in tail samples, suggesting that telomere length had stabilized at a new telomere length set point in F16 mice (Figs. 7c. d and S10). Overall, the TRF lengths shortened in the blood and tail at about 1 kb per generation. In comparison, telomeres of *M. musculus* deleted for telomerase RNA ($mTR^{-/-}$) shortened at a much faster rate of 4.8 ± 2.4 kb per generation³⁶. These results indicate that telomerase activity was not completely eliminated in Telomice but only downregulated to reach a shorter telomere length set point. Consistent with the observation in MEFs, no reduction was observed in the native gel hybridization signal corresponding to the single-stranded G-rich telomeric sequences (Figs. S11 and S12, for tail and blood samples, respectively), indicating that Telomice maintained an intact telomere end structure. Importantly, when analyzing telomere repeat length by NanoTelSeg, we estimated Telomice (F15) to have median telomere lengths of 6.4 kb in the blood and 6.1 kb in tail DNA, which is comparable to that measured in a human blood DNA sample (6.5 kb; Fig. 7e). Only 1.2% of the telomeres (one of 85) in the WT mouse blood sample were shorter than 3 kb and none were shorter than 2 kb (Fig. 7e). In contrast, 4.6% (14 of 302) of the telomeres in the F15 Telomouse blood were shorter than 3 kb and 2.3% (7 of 302) were shorter than 2 kb. Similarly, 5.4% (7 of 129) of the telomeres of the Telomouse F15 tail were shorter than 3 kb and 3.4% (4 of 129) were shorter than 2 kb. These observations suggest that short telomeres may play an important role in Telomice physiology but not in WT mice.

Decreased regeneration potential of Telomouse colonocytes in vivo

Telomerase null mice (mTR^{\prime}) can be intercrossed for only six generations until fertility is lost³⁶. Cells from the fourth *mTR*^{-/-} generation (G₄) onward frequently exhibit chromosome ends devoid of detectable telomere repeats, as well as end-to-end fusions and aneuploidy, correlating with defective spermatogenesis and reduced proliferation capacity of hematopoietic cells in the bone marrow and spleen⁴⁴. In contrast, *Rtel1^{K/K}* Telomice breed readily to at least F16, indicating that telomeres in the germline maintain sufficient length and normal function. To examine if the short telomeres present in Telomice affect highly proliferative somatic tissues, we determined proliferation rates of intestinal stem and progenitor cells, which display the fastest proliferative rate of all cells in the body, by labeling of cells in S-phase using the thymidine analog EdU. We noted a small but significant decrease in the number of EdU-positive epithelial cells in the colon of Telomice (Fig. 8a). To test if this proliferation defect is exacerbated under conditions of rapid epithelial renewal, we treated Telomice and control mice with dextran sodium sulfate (DSS), which causes ulcerative colitis-like pathologies due to its toxicity to colonic epithelial cells and engenders a strong regenerative response once the toxin is removed⁴⁵ (see experimental outline in Fig. 8b). Remarkably, while WT mice showed the expected proliferative response, with ~6 cells in S-phase per crypt cross section, the recovery was significantly blunted in Telomice, with only ~3 replicating cells per crypt cross section (Fig. 8c-e). These results suggested that while in the germline of Telomice telomerase is able to maintain sufficient (even if short) telomere length and normal cell proliferation, somatic cells are compromised in their ability to respond to situations requiring rapid regenerative proliferation, presumably because of their short telomeres and the absence of sufficient telomerase activity in somatic tissues.

Discussion

RTEL1 was reported already in 2004 as a regulator of telomere elongation, based on its genetic association with telomere length in crosses between the long-telomere species *M. musculus* and the closely related, short-telomere species *M. spretus*⁹. However, the specific difference between the RTEL1 proteins from the two mouse species that is responsible for the different telomere length set points has remained unknown to date¹². We derived a novel strain of *M. musculus* with a point mutation in *Rtel1*, replacing methionine 492 with a lysine, which established a new, *human* like telomere length set point, suggesting that this residue of RTEL1 is critical to its function in telomere length control. We termed this new *M. musculus* mouse strain 'Telomouse'. All other mouse species examined had a methionine at position 492. Therefore, the amino acid at this position cannot explain the short telomeres of other wild mouse species, which may rather result from another yet unknown variation in Rtel1 or in another telomere length regulator.

Following the growth of MEFs derived from an *Rtel1^{K/K}* embryo mouse over 250 population doublings and analyzing their telomeres by PFGE revealed that they gradually shortened to an averaged MTL of 14.7 kb at PD 250, as compared to 37.8 kb of the WT MEFs at the same PD (Fig. 2 and Supplementary Data 1). However, detailed analysis of the TRF and qFISH hybridization signals, which correlate with the length of the telomere repeat tracts, indicated that a large subtelomeric region (10-15 kb on average) is included in the TRF and the telomeric repeat tracts are highly heterogeneous in length (Fig. 3), as proposed previously^{36,37}. Precise measurement of individual telomeres by Nanopore sequencing indeed revealed that the actual length of the telomeric repeat tracts is significantly shorter than the MTL measured by PFGE (Fig. 4). The Rtel1K/K MEFs PD 250 telomeres had a median length of 5.6 kb as compared to 27.3 kb of WT MEFs at PD 250, consistent with a subtelomere fraction of about 10 kb that is included in the TRF. Nevertheless, even the short telomeres of the Rtel1K/K MEFs PD 250 largely retained their end-protection function, as indicated by their levels of telomeric and genome-wide DDR foci, FISH aberrations and T-SCE, which were found to be comparable to those in WT MEFs (Figs. 5, 6). Using metaphase FISH, telomere loss, indicated by a telomeric fluorescence hybridization signal below the detection limit, and Robertsonian fusions were elevated in the *Rtel1*^{K/K} MEFs PD 250 (Fig. 6). Since these Robertsonian fusions did not exhibit a telomeric signal at the point of fusion, we assume that this tolerable type of fusion resulted from occasional shortening of telomeres below the critical length⁴⁶. Interestingly, heterozygous Rtell^{M/K} MEFs displayed telomeres nearly as short as the homozygous *Rtel1^{K/K}* MEFs (Fig. 2a, b), suggesting that *Rtel1*^{M492K} is either a dominant mutation with a negative effect on the function of the WT allele or a hypomorphic mutation causing haploinsufficiency. Future research will reveal the mechanism by which the *Rtel1*^{M492K} mutation causes telomere shortening.

Interbreeding Telomice over thirteen generations to F16 revealed that in each generation telomeres shortened until a new telomere length set point was reached. The observed rate of shortening, about 1 kb per generation, was much slower than the rate of shortening in the telomerase null (mTR^{-}) mouse $(4.8 \text{ kb per generation})^{36}$, indicating that telomerase action was not abolished but only reduced in Telomice. Importantly, telomeres did not shorten further from F15 to F16, in both blood or tail (Figs. 7 and S10), suggesting that telomere length had stabilized, or was very close to stabilization, at a new telomere length set point. Together, our results show that the M492K mutation altered the function of RTEL1 in telomere length regulation, in line with other reports suggesting that RTEL1 regulates telomerase action at telomere ends^{9,17,26,27,30}. While we do not understand why the telomeres of M. musculus are ultra-long in the first place, the M492K mutation caused a dramatic reduction in the telomere length to a set point similar to that reached in *M. spretus* during natural evolution, without an apparent effect on the organismal health. This is in contrast to the disease-causing mutation in the same position of human RTEL1 (M492I), which caused genome-wide DNA damage and fatal disease17,25,30,43

To enable precise, base-resolution measurements of telomere length, we developed a novel method for vertebrate telomere length analysis by long read Nanopore sequencing, termed *NanoTelSeq*. Nanopore sequencing has been used successfully for analyzing yeast telomeres⁴⁷, but due to the different and variable yeast telomeric



EdU E-Cad Hoechst

Fig. 8 | Intestinal regeneration following DSS-induced damage to colonocytes is impaired in Telomice. a Intestinal tissues from *Rtel1^{K/K}* mice (F13 males, age 4 months) or C57BL/6 control mice (M/M, male, age 4 months) were analyzed for the homeostatic proliferation rate in the gut epithelium by short-term (120 min) EdU labeling. Crypt stem and progenitor cells in the jejunum of mutant (K/K) mice displayed proliferation rates similar to those observed in control (M/M) mice. In the colon, however, K/K crypt cells divided somewhat less frequently. Blue circles, control (M/M); red squares, mutant (K/K). Each data point represents one animal, with 10 crypts per animal counted and averaged into one data point (n = 4 for each group). Data are presented as mean +/- standard error of the mean, and significance was assessed using an unpaired, two-tailed Student's t-test. Scale bars equal 50 µm. b Experimental timeline for dextran sulfate sodium (DSS) treatment. Rtel1K/K (F13) and control (M/M) mice (males, age 4 months) received 3% DSS in the

drinking water for 144 h, followed by 24 h of regular drinking water to allow for colonic regeneration. Colons were collected and analyzed for crypt cell proliferation rate by short-term (120 min) EdU labeling. c Control (M/M) and d Rtel1^{K/K} (F13)

colons following DSS treatment were stained for EdU labeling (white), E-cadherin (E-Cad; green), and nuclei (Hoechst dye). Note the reduced number of EdU-positive colonocytes in the *Rtell*^{K/K} mice (**d**). Scale bars equal 50 μ m. **e** Quantification of replicating (EdU⁺) cells per crypt in DSS-treated WT (M/M) and Rtel1K/K (F13) colons. Crypt cells in the colon of mutant (K/K) mice (n = 6; red squares) displayed reduced proliferation compared to control (M/M) mice (n = 5; blue circles). Each data point represents one animal, with 10 crypts per animal counted and averaged into one data point. Data are presented as mean and standard error of the mean, and significance was assessed using an unpaired, two-tailed Student's t-test.

repeat, this method could not be employed for studying vertebrate telomeres and a different strategy needed to be developed for ligating the sequencing adapter to the telomere end of vertebrates. The longread sequencing approach developed here provides the distribution, median telomere length and, most importantly, the accurate length and abundance of the shortest telomeres, which cannot be determined by Southern blot, in-gel hybridization analysis, or gFISH. This is

particularly important because the shortest telomeres are those dictating cell function and fate³⁹. In addition, NanoTelSeq can identify and assign telomere sequences to individual chromosome ends (Fig. S8). NanoTelSeq, not only confirmed that the Telomouse telomeres are much shorter than those of WT M. musculus mice, but also enabled the precise analysis of the shortest telomeres not measurable by other methods. NanoTelSeq revealed a significant percentage of telomeres

below 2 kb (but not below 1 kb) in Telomice MEFs, as well as in Telomouse blood and tail (Figs. 4d and 7e). These observations suggest that the critical telomere length in both human and mouse is about 1 kb. Below this length telomeres cease to suppress DDR, arrest cell proliferation, and are depleted from the population of proliferating cells.

Short telomeres are major drivers of the aging-dependent decline in organ function and increase in cancer risk^{5,48}. Unfortunately, to date, these processes have been difficult to study in the common laboratory mouse, M. musculus, due to its exceptionally long telomeres. Attempts to study the implications of short telomeres in mice have thus far been based on telomerase null alleles, which cause ever-shortening telomeres and eventually result in severe pathologies and infertility³⁶, manipulating telomere proteins to induce overt telomere deprotection^{7,49}, or the telomerase haploinsufficient *mTR*^{+/-} strain of *M. musculus* crossed with the *M. castaneus* strain⁵⁰. The resulting hybrid strain is not isogenic and is difficult to maintain because of the required heterozygosity with respect to both mTR and unknown telomere length regulators in the M. castaneus background. The Telomouse model derived here is genetically identical to the common M. musculus laboratory strain C57BL/6 except for the single amino acid difference in RTEL1. Yet, it displays dramatically shorter telomeres with no apparent increase in pathologies under normal physiological conditions, with the exception of a small decrease in the replication rate of the fastest proliferating cells of the colonic epithelium. When Telomouse colonocytes were challenged to mount a regenerative response, Telomouse epithelial cells were compromised in their replicative ability. Since Telomouse telomeres are comparable in length to human telomeres, the accumulation of critically short telomeres in the Telomouse tissues with age provides a unique model for studying the effects of short telomeres on organ function during aging.

The utility of Telomouse depends in part on the ability to introduce and stably maintain additional genetic manipulations. While breeding of Telomice to another strain of mice would introduce a set of chromosomes with long telomeres and may require additional intercrossing to reach short telomeres again, extensive backcrossing may not be needed if the short telomeres inherited from the Telomouse parent are maintained. The telomeres of the heterozygous M/K MEFs shortened almost at the same rate and to the same extent as those of the homozygous K/K MEFs (Fig. 2b) indicating that the M492K mutation is either dominant or causing haploinsufficiency, consistent with the original discovery of Rtel1 as a dominant regulator of telomere length⁹. While this is yet to be examined in the mice, it is plausible that the short telomeres derived from *Rtel1*^K gametes in Telomouse crosses will not elongate in the heterozygous mice. In this case, the effects of manipulating an additional 'gene X' in the Telomouse would occur on the background of the short chromosomes inherited from the Telomouse parent. Alternatively, the genome of the Telomouse can be manipulated by CRISPR/Cas9 assisted genome editing in zygotes to introduce and examine the functional impact of additional loci, instead of crossing it to other strains with long telomeres. Therefore, we believe that the Telomouse is easier to maintain, manipulate and study than the unstable telomerase null or heterozygous strains. Altogether, the Telomouse model will open the way for a broad range of physiological experiments on the organismal level that were not possible previously, and reveal the effects of short telomeres in various tissues on cell proliferation, age-related organ decay, and cancer development.

Methods

The use of animals in this study followed the Guide for the Care and Use of Laboratory Animals, Laboratory Animal Ordinances, and the Animal Welfare Act. This study was approved by the University of Pennsylvania Institutional Animal Care and Use Committee, protocol number 805623.

Generating a three-dimensional structure model of the murine RTEL1 helicase domain

We generated the murine RTEL1 (mRTEL1) 3D structure model consisting of residues 3-770 using the software Protein Homology/analogY Recognition Engine V 2.0 (Phyre 2), which produces a model of the protein of interest based on sequence alignment to known structures³¹. The 3D mRTEL1 structure was modeled using the cryo-EM structure of the yeast TFIIH helicase in the contracted state within the pre-initiation complex (PDB ID: 704K)⁵². TFIIH has 28% sequence identity to the mRTEL1 helicase domain. The presence of a structure with significant sequence identity with the murine gene enabled PHYRE to generate this model with 100% confidence. The model was further refined by applying geometry minimization in UCSF CHIMERA⁵³. We then generated a model of RTEL1 in a complex with single-stranded DNA, using the structure of the highly similar helicase domain of DinG (PDB ID: 6FWS), which belongs to the XPD family of helicases together with RTEL1, FANCJ, and DDX11. The model was further refined by applying geometry minimization in Phenix⁵⁴. The figures were generated in Pymol⁵⁵.

Derivation of *Rtel1*^{M492K} mice

CRISPR guide RNAs were designed as described⁵⁶. For sgRNA preparation, the T7 promoter sequence was added to the sgRNA template by PCR amplification using pX335 (Addgene #42335) with Phusion high-fidelity DNA Polymerase Kit (NEB), Common sgRNA-R and Primer 1 or Primer 2 (see below).

Common sgRNA-R: 5'-AGCACCGACTCGGTGCCACT-3'

Primer 1 with gRNA sequence underlined:

5'-TTAATACGACTCACTATAGGCATCTGCATCTCCAGAGCAA gttttagagctagaaatagc-3'

Primer 2 with gRNA sequence underlined:

5'-TTAATACGACTCACTATAGGCACCTGGAGGTCACAACACT

gttttagagctagaaatagc-3'

PCR products were purified using the QIAquick Gel Extraction kit (Qiagen), followed by in vitro transcription with the T7 High Yield RNA Synthesis kit (NEB). Newly synthesized sgRNAs were purified using the MEGAclear kit (Life Technologies), precipitated and diluted in injection buffer (10 mM Tris/0.1 mM EDTA, pH 7.5) at a concentration of 500 ng/ μ l. The quality of the sgRNAs was confirmed using the total RNA Nano Chip on an Agilent bioanalyzer. The final injection mix was prepared using ssDNA repair template (IDT; 100 ng/ μ l), sgRNA1 (50 ng/ μ l), sgRNA2 (50 ng/ μ l), and Cas9 Nickase mRNA (Trilink; 100 ng/ μ l) in injection buffer. The sequence of the ssDNA repair template is shown below. The mutated A is underlined within codon 492 (in bold).

5'GTTCGTACCCTTATCCTCACCAGCGGTACCCTGGCTC-CACTGTCTTCCTTTGCTCTGGAG**AAG**CAGATGTATGTATGAGTC ACCTGGAGGTCACAACACTAGGAACATGGTGGGTGGGGTTGG-3'

The final mix was spun twice at top speed for 30 min at 4 °C to remove debris to avoid needle clogging. Cytoplasmic injection was done using C57Bl6 zygotes. After injection, SCR7 (Xcessbio) was added to the overnight egg culture at the concentration of 50 μ M. Out of the 17 mice born, two had the targeted allele.

Preparation and culture of MEF

Embryonic day (E)13.5 mouse embryos were dissected with head and inner organs removed, rinsed in HBSS, then digested with 300 μ l papain isolation enzyme (ThermoFisher) for 30 min at 37 °C following the manufacturer's protocol. Embryonic digests were transferred to conical tubes with 1 ml of HBSS, pipetted up and down to achieve single-cell suspension and then spun. The pellets were resuspended in MEF media (DMEM with 20% FCS, Pen/Strep/non-essential amino acids) and plated. MEFs were grown in DMEM media containing 2 mM L-glutamine, penicillin-streptomycin, non-essential amino acids and 20% fetal bovine serum until immortalization (around PD60-70) and

then with 10% fetal bovine serum in the same medium. Media and media supplements were purchased from Biological Industries Inc., Beit Haemek, Israel.

Other cell lines used

Telomerase-positive human primary fibroblasts used in this study were S2 HHS patient cells rescued by ectopic expression of hTERT and WT RTEL1³⁰. These cells were grown for 100 PD in DMEM media supplemented with 10% fetal bovine serum, penicillin-streptomycin, 2 mM L-glutamine, non-essential amino acids, and 30 ng/ml doxycycline to induce ectopic expression of WT RTEL1.

Genomic DNA extraction

Leukocytes were obtained from blood by lysing red blood cells by adding 2 volumes of reticulocyte lysis buffer (155 mM NH₄Cl, 10 mM KHCO₃, and 0.1 mM EDTA pH 8), rotating at room temperature for 10 min, and centrifugation at 300 × *g* for 10 min. The pellet of leukocytes was resuspended in 1 ml of lysis buffer centrifuges as before, and washed with 1 ml of PBS. Leukocytes and MEFs pellets were lysed in 10 mM Tris pH 7.5, 10 mM EDTA, 0.15 M NaCl, 0.5% SDS, and 100 μ g/ml proteinase K overnight at 37 °C. Mouse tail samples were lysed in 100 mM Tris HCl pH 8.5, 5 mM EDTA, 0.1% SDS, 200 mM NaCl and 100 μ g/ml proteinase K overnight at 50 °C. Following cell lysis, high molecular weight genomic DNA was phenol-extracted, ethanol-precipitated, and dissolved in TE. Genomic DNA samples were examined by agarose gel electrophoresis of undigested DNA, and samples suspected to be degraded were excluded from further analysis.

In-gel analysis of telomeric restriction fragments

Genomic DNA was treated with RNase A, digested with the Hinfl restriction endonuclease, and quantified by Qubit fluorometer. Equal amounts $(1-5 \mu g)$ of the digested DNA samples were separated by pulsed-field gel electrophoresis (PFGE) using a Bio-Rad CHEF DR-II apparatus in 1% agarose and 0.5 x TBE, at 14 °C, 200 V and pulse frequency gradient of 1s (initial) to 6s (final) for 20-22h (except for the K/K gel on the right in Fig. 2a, which was electrophoresed in the same parameters but for 18 h), or in regular gel apparatus, 0.7% agarose and 1 x TBE for 1300 V x hour (Fig. 3d). Gels were ethidium-stained, dried, pre-hybridized for 1h at 50 °C in hybridization solution [sodium phosphate buffer (0.5 N⁺, pH 7.2), 1 mM EDTA and 7% (wt/vol) SDS], and hybridized overnight at 50 °C in hybridization solution with a probe mixture containing end-labeled telomeric oligonucleotide (AACCCT)₃, mid-range PFG ladder (NEB Inc.) and 1kb ladder (Gene-Direx Inc.). The ladder probes were digested with Hinfl, dephosphorylated by quick calf intestinal phosphatase (NEB Inc.) and heat inactivated prior to labeling. All probes were 5' end-labeled with $[\gamma^{-32}P]$ -ATP and T4 polynucleotide kinase (NEB Inc.). After hybridization, the gels were washed three times (for 10 min, 30 min, and 50 min) with 4 X SSC at room temperature, and once or twice with pre-warmed 4 X SSC at 40 °C for 30 min. The gels were exposed to a Phosphor Imager. TRF length was quantified using *TeloTool* (corrected mode)⁵¹. Individual band lengths were interpolated from the regression formulas for plotting the log of the ladder band sizes as a function of the migration distance, and native and denatured in-gel hybridization intensity was quantified using ImageQuant-TL (GE Healthcare Inc.).

Immunofluorescence of cultured cells

Cells were seeded onto glass coverslips and grown for 1–4 days. Immunofluorescence was performed as described³⁰ with the following changes: Primary antibodies used were TRF1 (Abcam; AB10579) and γ H2AX (Cell Signaling Technology; 2577S), incubated overnight at 4 °C. Imaging acquisition was by FV-1200 confocal microscope (Olympus, Japan) with a 60X/1.42 oil immersion objective. *NIH ImageJ*³⁸ was used for image analysis and foci quantification.

Metaphase FISH

Metaphase spread and FISH were performed as described³⁰, with the following changes: Cells were arrested in metaphase by 3–4 h incubation with Colcemid. After fixation, both MEFs and human fibroblasts were mixed and dropped on the slides. The PNA probes used were TelC-Alexa488 telomeric probe (F1004, Panagene Inc.) and CENPB-Cy3 centromeric probe (F3002, Panagene, Inc). Imaging acquisition was by FV-1200 confocal microscope (Olympus, Japan), by Olympus IX83 fluorescence microscope, or by a Nikon Eclipse Ti-E microscope equipped with a CoolSNAP Myo CCD camera. The telomeric signal was quantified using *Telometer*⁵⁷.

Meta-TIF assay

We followed the protocol developed by Cesare and colleagues⁵⁸ with a few changes. MEFs were synchronized in metaphase with 0.2 µg/ml Colcemid for 1-2 h, collected by trypsinization, and resuspended in fresh hypotonic solution (0.2% KCl and 0.2% Tri-sodium citrate in DDW) at room temperature (RT). 500 µl of swollen MEFs (200,000 cells/ml) were pipetted into each cytocentrifuge (Thermo Scientific Cytospin 4) funnel and centrifuged for 7-10 min at 1000 rpm with medium acceleration rate onto Superfrost glass slides. The slides were immediately fixed in 1x PBS/4% formaldehyde at RT for 10 min, permeabilized in KCM buffer (120 mM KCl, 20 mM NaCl, 10 mM Tris pH 7.5 and 0.1% Triton X-100) at RT for 10 min, and then blocked with antibody dilution buffer (20 mM Tris pH 7.5, 2% BSA, 0.2% fish gelatin, 150 mM NaCl, 0.1% Triton X-100 and 0.1% sodium azide) containing DNase-free RNase A for 15 min at 37 °C. The slides were incubated overnight at 4 °C, with the primary antibody (yH2AX [Millipore; JBW 301]) diluted 1:1000 in antibody dilution buffer. After washing, the slides were incubated with a secondary antibody (Thermo Scientific, A-11029) diluted 1:500 in antibody dilution buffer for 30 min at RT, washed, and fixed for 10 min in 1x PBS with 4% formaldehyde at RT. The slides were dehydrated using a graded ethanol series (70% for 3 min, 90% for 2 min, and 100% for 2 min) and then air-dried. FISH was performed using the metaphase FISH protocol described above, but with denaturation at 80 °C for 5 min, hybridization overnight at RT using the TelC-Cy3 probe (F1002-5, Panagene), and washing in PNA wash A (70% formamide and 10 mM Tris pH 7.5), then in PNA wash B (50 mM Tris pH 7.5, 150 mM NaCl and 0.08% Tween-20). After rinsing several times with dH₂O, the slides were mounted with Vectashield + DAPI (ThermoFisher Scientific Inc.) and stored at least one overnight before imaging. Images were acquired using the Olympus IX83 fluorescence microscope system (Olympus).

Chromosome-orientation fluorescent in situ hybridization (CO-FISH)

MEFs were cultured in medium supplemented with 7.5 µM bromodeoxyuridine (BrdU) and 2.5 µM bromodeoxycytidine (BrdC) for 10 h, and metaphase spreads were prepared as in the metaphase FISH assay, adding Colcemid (without changing the medium), and incubating for 3 h. Slides were treated with RNase A for 30 min at 37 °C, stained with Hoechst 33258 (0.5 µg/ml in 2x SSC) for 30 min at RT, irradiated with 365-nm UV light for 5.4×10^3 J/m², and then digested with Exonuclease III (9.5 µl of 100 units/µl) at 37 °C for 30 min. Slides were washed in 1x PBS, dehydrated in ice-cold ethanol series of 70%, 85%, and 100% for 2 min each and air-dried in the dark. For PNA hybridization, the same hybridization mix as in the metaphase FISH assay was used, hybridizing 1 h with the TelG probe at RT in the dark, rinsing in wash I (10 mM Tris-HCl pH 7.2, 70% Formamide and 0.1% BSA in 1x PBS) for 2 min, drying briefly by dabbing, then hybridizing for another 1 h with the TelC probe at RT in the dark. After removing the coverslips in 1x PBS, the slides were washed twice in wash I for 15 min each time, then in wash II (0.1 M Tris-HCl pH 7.2, 0.15 M NaCl, 0.08% Tween-20) three times for 5 min each time. Finally, the slides were washed in 1x PBS three times, 5 min

each, dehydrated in ethanol baths (70%, 95%), air dried in the dark, and mounted then imaged as in the meta TIF assay.

Nanopore sequencing of telomeres

Single samples were sequenced separately using six telorette oligonucleotides, each ending with a different permutation of the six base telomeric repeat at its 3' end (Fig. 4a). 10-20 µg of high molecular weight genomic DNA, prepared as described above and treated with RNase A, were ligated overnight at 20 °C with a mix of six 5' phosphorylated telorette oligonucleotides (0.1 µM each) and 2000 units of T4 ligase (NEB Inc.) in a 50 µl reaction. The ligated DNA was then annealed to 1 µM teltail-tether oligonucleotide by heating for 10 min at 65 °C and cooling slowly to room temperature. The DNA was purified using AMPure XP beads (Beckman Coulter Life Sciences), and ligated to the Nanopore sequencing adapter (AMII) following the protocol for genomic DNA ligation sequencing with native barcoding (SQK-LSK109 with EXP-NBD104; Oxford Nanopore Technologies). For multiplexed reactions, 2-6 samples were pooled in a sequencing library using barcoded telorette 3 oligonucleotides. The barcoded telorette oligonucleotides were first annealed to a complementary teltail-tether oligonucleotide and then each double-stranded telorette with a specific barcode $(1 \mu M)$ was ligated to an individual DNA sample $(1-10 \mu g)$ as before. The ligation reactions were stopped by adding 20 µM EDTA and the barcoded samples were combined, purified, ligated to 3 µl sequencing adapter (AMII) and sequenced following the standard protocol for genomic DNA ligation sequencing with native barcoding (SQK-LSK109 with EXP-NBD104; Oxford Nanopore Technologies), except for the library that included the K/K mouse blood sample, which was purified by polyethylene glycol (PEG) precipitation instead of AMPure XP beads following⁵⁹. 0.4–5.5 µg of the Sequencing libraries were loaded onto a Nanopore R9 flow cell. The sequencing library purified by PEG was loaded with SQB buffer (Oxford Nanopore Technologies) without loading beads due to the high viscosity of the library.

Oligonucleotides used for sequencing. Underlined is the telomeric repeat, P. 5' phosphate:

Telorette1: 5'-P-TGCTCCGTGCATCTCCAAGGTTCCTAAC-3' Telorette2: 5'-P-TGCTCCGTGCATCTCCAAGGTTAACCCT-3' Telorette3: 5'-P-TGCTCCGTGCATCTCCAAGGTTCTAACC-3' Telorette4: 5'-P-TGCTCCGTGCATCTCCAAGGTTTAACCC-3' Telorette5: 5'-P-TGCTCCGTGCATCTCCAAGGTTACCCTA-3' Telorette6: 5'-P-TGCTCCGTGCATCTCCAAGGTTCCCTAA-3' Telorette3-NB01n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-CACAAAGACACCGACA ACTTTCTT-CAGCACCT-CTAACC-3' Telorette3-NB02n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-ACAGACGACTACAAA CGGAATCGA-CAGCACCT- CTAACC-3' Telorette3-NB03n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-CCTGGTAACTGGGAC ACAAGACTC-CAGCACCT-CTAACC-3' Telorette3-NB04n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-TAGGGAAACACGATAG AATCCGAA-CAGCACCT-CTAACC-3' Telorette3-NB07n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-AAGGATTCATTCCCAC GGTAACAC-CAGCACCT-CTAACC-3' Telorette3-NB08n: P-5'-TGCTCCGTGCATCTCC-AAGGTTAA-ACGTAACTTGGTTTGTT CCCTGAA-CAGCACCT-CTAACC-3' Teltail-tether: 5'-AACCTTGGAGATGCACGGAGCAAGCAAT-3'

Computational processing of Nanopore sequencing reads Deducing the nucleotide sequence (base calling) from the raw data (Fast5 files) was performed with the Nanopore *Guppy* application at

high or super accuracy mode. The sequences (FastQ files) were filtered and analyzed by a dedicated application termed *Telomere Analyzer*⁶⁰. We noticed a difficulty of the base calling application in distinguishing between purines (A and G) in the context of the telomeric repeats, the repeats on the C-strand were identified by searching for the sequence CCCTRR (R represents a purine) in the six possible permutations. The telomere length and read length were extracted and presented on plots of telomere density along the entire read. Telomere density was calculated in a moving window of 100 nt and describes the portion within each 100 nt sequence that contains telomeric repeats (from 0– no telomeric sequence to 1–fully telomeric). A telomere is identified by a summed telomere density of at least 2 for at least three consecutive windows, each having a telomere density of at least 0.3. Then, the telomere beginning and end are defined by the first and last telomeric repeat within the identified windows or flanking windows.

Mapping of telomere reads to specific chromosome ends

The subtelomeric sequences of telomeric reads were mapped to the *M. musculus* GRCm39 reference genome using the *minimap2* program with the recommended set of parameters optimized for long Nanopore sequencing reads⁶¹. Alignment results were filtered according to the following criteria: (1) At minimum of 70–90% (depending on the sample) of the read's subtelomeric region was aligned. (2) The alignment started within the most distal 10,000 nt of a chromosome end in the reference genome. (3) The read's subtelomere was aligned to the reference sequence in the correct orientation with respect to the chromosome end.

Selection of telomere reads for median calculation

Reads with an overall telomere density of 0.7 were analyzed. A few additional reads with lower density were examined and approved manually. Other reads with lower density or not beginning within 500 nt of the 5' end were excluded. Not all reads were sufficiently long to reach the subtelomere (indicated by asterisks in Supplementary Data 2 and 3). Since excluding these reads alone would create a bias for short telomeres, reads were further selected based on their read length. To do so, telomere reads were initially sorted for read length, from long to short, and a running median for telomere length was calculated from the longest read down to each point. As the ends of the telomere reads contained the telorette sequence (62 nt in length) and the sequencing adapter (exact length is unknown), only reads longer than their corresponding running median telomere length + 100 nt (corresponding to the estimated length of the adapters) were selected for median length calculation. For the reads mapped to chromosome ends, only reads longer than the median + 1000 nt were chosen.

DSS treatment

Dextran sulfate sodium (DSS) (MP Biomedicals, cat. # 0216011001) was administered to *Rtel1^{K/K}* mice (F13) and WT (M/M) mice via drinking water (3% DSS) for 144 h. All mice received regular drinking water for 24 h prior to sacrifice.

Tissue immunohistochemistry and immunofluorescence

For EdU treatment, $150 \,\mu$ l of $10 \,m$ g/ml EdU solution was injected intraperitoneally 2 h before mice were sacrificed. After euthanasia, mouse tissues were rinsed in PBS and fixed with 4% paraformaldehyde overnight. Fixed tissue was rinsed in PBS three times for 30 min per rinse and dehydrated for paraffin embedding and sectioning. EdU was detected according to the manufacturer's protocol (Click-iT Plus EdU Cell Proliferation Kit for Imaging, Alexa Fluor 647 dye, C10640, Invitrogen). Briefly, tissue sections were deparaffinized, and heat antigen retrieval was performed in a pressure cooker with Citrate Buffer, pH 6 (64142-07, Electron Microscopy Sciences). The tissue sections were then incubated in the Click-iT Edu Reaction Cocktail for 30 min at room temperature, washed, and blocked with CAS-Block (008120,

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Invitrogen) for 30 min at room temperature. Tissue sections were stained overnight with a mouse anti-E-Cadherin (610181, 1:500, BD Biosciences) primary antibody. The following day, tissue sections were rinsed in PBS and incubated for 2 h at room temperature with a Cy2 anti-mouse secondary antibody. The tissue sections were washed in PBS and stained with Hoechst (H3570, 1:10,000, Molecular Probes) to visualize nuclei. Images were taken on a Leica laser scanning confocal microscope (TCS SP8, Penn Cell & Developmental Biology Microscopy Core). Images were processed and brightness and contrast were enhanced using FIJI (*ImageJ*, Version 2.1.0, NIH)³⁸.

Mice

DNA from a total of 130 K/K and M/M (100% pure Blk6) mice were analyzed multiple times in separate gels (as detailed in Supplementary Data 4). The average age for K/K mice analyzed in the gels was 331 days, while the average age for M/M mice was 395 days. At each specific generation of breeding, the following mice were analyzed:

F0 (M/M): 24 mice (63-665) days old,

F4 (K/K): one mouse 881 days old,

F5 (K/K): 7 mice (341–895) days old,

F6 (K/K): 13 mice (318-787) days old,

F7(K/K): 2 mice (329-872) days old,

F8 (K/K): 7 mice (233-778) days old,

F9 (K/K): 28 mice (81–402) days old,

F10 (K/K): 7 mice (42–68) days old,

F11 (K/K): 9 mice (68–138) days old, F12 (K/K): 9 mice 118 and 171 days old,

F13 (K/K): 9 mice: 80 days old,

F14 (K/K): 4 mice: 186 and 205 days old,

F15 (K/K): 8 mice: 76, 95, and 182 days old,

F16 (K/K): 2 mice: 144 and 163 days old.

Three mouse embryonic fibroblast cultures were grown over 250 population doublings (PD) and sampled every 10 PD (a total of 121 samples). Samples were analyzed in multiple gels, as detailed in Supplementary Supplementary Data 1.

Statistical analysis

Statistical analysis was performed by Microsoft Excel and GraphPad Prism 8.0, using simple linear or segmental linear regression while setting the break points manually to give the best fit by coefficient of determination (R^2). For statistical tests two-tailed paired and unpaired t-test, nested t-test, and nested one-way ANOVA were used, as indicated for each figure. *P* < 0.05 was considered statistically significant.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Representative gels are shown in Figs. 2, 7, S2–S4, and S9–S12. All measured and calculated values are provided in the Supplementary Data 1 (MEFs) and 4 (mice). The complete un-spliced gel images for Fig. 2a are shown in Fig. S2a, c, e. The complete un-spliced gel images for Fig. 7b, including the native hybridization, are shown in Fig. S11d, h. The complete unprocessed images for all gels used in this study are present in the Source data file provided with this paper. The Nanopore telomeric read characteristics are summarized in Supplementary Data 2 and 3. The nanopore sequencing data and *Telomere Analyzer* output folders are available at the OSF repository⁶². Source data are provided with this paper.

Code availability

The code used for filtering and processing Nanopore sequencing reads, termed *Telomere Analayzer*, is available at GitHub⁶⁰.

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Corresponding authors Y.T. (telomere biology) and K.H.K. (mouse biology): study concept and design; analysis and interpretation of data; writing and revising the manuscript. Y.T. and R.S.: study concept and design; analysis of telomere phenotypes; interpretation of data; writing and revising the manuscript. C.L.M. and H.M.K.: derivation and in vivo analysis of the mouse strains, and writing and revising the manuscript. N.E.: analysis of telomere phenotypes and computational analysis of Nanopore sequencing data. D.L.: writing the Telomere Analyzer code and computational analysis of the mouse RTEL1 structure. Y.R., A.M., V.O., M.R., and M.T.: in vivo analysis of the mouse strains.

Competing interests

The authors declare no competing interests.

Additional information

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