UCLA UCLA Previously Published Works

Title

Universal DNA methylation age across mammalian tissues.

Permalink

https://escholarship.org/uc/item/5jw70238

Journal

Nature Aging, 3(9)

Authors

Lu, A Fei, Z Haghani, A <u>et al.</u>

Publication Date

2023-09-01

DOI

10.1038/s43587-023-00462-6

Peer reviewed

nature aging

Resource

Universal DNA methylation age across mammalian tissues

Received: 29 September 2022

Accepted: 21 June 2023

Published online: 10 August 2023

Check for updates

A list of authors and their affiliations appears at the end of the paper

Aging, often considered a result of random cellular damage, can be accurately estimated using DNA methylation profiles, the foundation of pan-tissue epigenetic clocks. Here, we demonstrate the development of universal pan-mammalian clocks, using 11,754 methylation arrays from our Mammalian Methylation Consortium, which encompass 59 tissue types across 185 mammalian species. These predictive models estimate mammalian tissue age with high accuracy (r > 0.96). Age deviations correlate with human mortality risk, mouse somatotropic axis mutations and caloric restriction. We identified specific cytosines with methylation levels that change with age across numerous species. These sites, highly enriched in polycomb repressive complex 2-binding locations, are near genes implicated in mammalian development, cancer, obesity and longevity. Our findings offer new evidence suggesting that aging is evolutionarily conserved and intertwined with developmental processes across all mammals.

Aging is associated with multiple cellular changes that are often tissue specific¹. Cytosine methylation, however, stands out, as it allows for the development of pan-tissue aging clocks (multivariate age estimators) that are applicable to all human tissues²⁻⁴. The subsequent development of similar pan-tissue clocks for mice and other species suggests a conserved aspect to the aging process⁵⁻⁷, thereby challenging the belief that aging is solely driven by random cellular damage accumulated over time. To investigate this, we sought to (1) develop universal age estimators applicable to all mammalian species and tissues (pan-mammalian clocks) and (2) identify and characterize cytosines with methylation levels that change with age across all mammals. For this purpose, we employed the mammalian methylation array, which we recently developed to profile methylation levels of up to 36,000 CpG sites with flanking DNA sequences highly conserved across the mammalian class⁸. We employed such profiles from 11,754 samples from 59 tissue types, originating from 185 mammalian species across 19 taxonomic orders (Supplementary Data 1.1-1.4 and Supplementary Notes 1 and 2) with ages ranging from prenatal to 139 years old (bowhead whale, Balaena mysticetus)9. These data are a subset from our Mammalian Methylation Consortium, which characterized maximum lifespan⁹. As we were interested in developing pan-mammalian clocks, we restricted the analysis to animals with known ages.

Results

Universal pan-mammalian epigenetic clocks

In separate articles, we described the application of the mammalian methylation array to individual mammalian species¹⁰⁻¹⁹. These studies already demonstrate that one can build dual-species epigenetic age estimators (for example, human–naked mole rat clocks)¹⁰⁻¹⁷, in contrast to first- and second-generation clocks that measure human age^{4,20,21} and mortality risk^{22,23}, respectively. However, it is not yet known whether one can develop a mathematical formula to estimate age in all mammalian species. Here we present three such pan-mammalian age estimators.

The first, basic clock (clock 1), regresses log-transformed chronological age on DNA methylation levels of all available mammals. Although such a clock can directly estimate the age of any mammal, its usefulness could be further increased if its output were adjusted for differences in the maximum lifespan of each species as well, as this would allow biologically meaningful comparisons to be made between species with very different lifespans. To this end, we developed a second universal clock that defines individual age relative to the maximum lifespan of its species; generating relative age estimates between 0 and 1. Because the accuracy of this universal relative age clock (clock 2) could be compromised in species for which knowledge of maximum lifespan is inaccurate, we developed a third universal clock, using age at sexual maturity (ASM) and gestation time instead of maximum lifespan,

e-mail: shorvath@mednet.ucla.edu

as these traits are better established and explain over 69% of maximum lifespan variation on the log scale (Supplementary Data 2). This third clock is referred to as the universal log–linear age clock (clock 3). The non-linear mathematical function underlying the age transformation of clock 3 reflects the fact that epigenetic clocks tick faster during development, an observation that led to the establishment of the first pan-tissue clock for humans⁴ (Extended Data Fig. 1a,b,d,e).

Performance of universal epigenetic clocks across species

To evaluate the clocks' accuracy, we employed leave-one-fraction-out (LOFO) and leave-one-species-out (LOSO) cross-validation analyses. Each analysis divides the dataset differently for validation: LOFO into ten fractions with similar proportions of species and tissue types; LOSO excludes one species per iteration. The final models of the clocks use less than 1,000 CpG sites each (Supplementary Data 3.1-3.3), with 401 common genes proximal to CpG sites in both clock 2 and clock 3 (Supplementary Data 3.5). LOFO cross-validation reveals the universal clocks as highly accurate estimators of chronological age ($r \approx 0.96 - 0.98$) with a median absolute error (MAE) of <1 year between chronological age and DNA methylation (DNAm)-based age estimate (DNAmAge) and a relative error of <3.3% (Figs. 1a, c and 2, Extended Data Fig. 2a, Supplementary Table 1 and Supplementary Data 4.1-4.3). Despite the mammalian array mapping fewer CpG sites to marsupials⁸, clocks 2 and 3 maintain their accuracy when analysis is confined to marsupials (for example, r = 0.91, median MAE < 0.80 year for clock 2; Fig. 1b). Moreover, our monotreme study (n = 15) produced encouraging results (for example, r = 0.85 for clock 2; Supplementary Data 4.1).

Using LOSO cross-validation, the clocks displayed age correlations as high as r = 0.941 (Supplementary Table 1), suggesting their applicability to species not included in the training set. However, for certain species, such as bowhead whales, the basic clock's predicted epigenetic age poorly aligns with chronological age (Extended Data Fig. 2a).

For the basic clock 1, the mean discrepancy between LOSO DNAmAge and chronological age (Delta.Age) is negatively correlated with species maximum lifespan (r = -0.84, $P = 1.0 \times 10^{-19}$) and ASM (r = -0.75, $P = 7.9 \times 10^{-14}$; Extended Data Fig. 2c, d). Here, the strengths of clocks 2 and 3 come to fore as they adjust for these species characteristics during their construction (Extended Data Fig. 1).

Universal clocks 2 and 3, arguably more biologically meaningful than clock 1, achieve a correlation of $r \ge 0.95$ between DNAm transformed age and observed transformed age, respectively (Fig. 1d, f). We will focus on them in the following text. They are pan-tissue clocks offering comparable accuracy in LOFO estimates across numerous tissue types (Fig. 1 and Supplementary Data 4.2). For instance, clock 2 yielded high age correlations in humans (LOFO estimate of r = 0.959 across 20 tissue types), mice (r = 0.948, 26 tissues) and bottlenose dolphins (r = 0.945, two tissues). Fig. 2 displays circle plots for the age correlation estimates in different species sorted by maximum lifespan.

Visual inspection indicates no relationship between age correlation from clocks 2 and 3 and maximum lifespan (dashed line, Fig. 2, circle). While accurately predicting age for the humpback whale and other mammals, the clocks sometimes underestimated bowhead whale reported age (mammalian species index 4.11.1 in Fig. 1a,c), possibly due to overestimation of older whales' ages by aspartic acid racemization.

Clocks 2 and 3 provide similarly accurate LOSO age estimates between evolutionarily distant species (Supplementary Data 5.2), including dogs (n = 742, 93 breeds, r = 0.94, MAE < 2.28 years), African elephants (r = 0.96, MAE < 4.0 years) and flying foxes (r = 0.97, MAE < 2.3 years) (Fig. 1j–1). Such accuracy demonstrates these clocks' broad relevance, tapping into conserved age-related mechanisms across mammals, including species not in the training data (Supplementary Data 5.1–5.2).

The three universal clocks performed well for 114 species with fewer than 15 samples each ($r \approx 0.90$, MAE ≈ 1.2 years for clocks 1–3; Extended Data Fig. 3a–c), exhibiting strong correlation for relative age (r = 0.91 for clock 2; Extended Data Fig. 3d).

Pan-mammalian universal clocks across tissues

The significantly distinct epigenomic landscape across tissue types^{24,25} prompted an assessment of these clocks' performance in different tissues. We assessed the tissue-specific accuracy of clock 2 for estimating relative age (r = 0.95, Fig. 1d) across 33 distinct tissue types, observing a median correlation of 0.91 and a median MAE for relative age of 0.027 (Supplementary Data 4.3). High age correlation was consistently observed in brain regions: whole brain (r = 0.991), cerebellum (r = 0.963), cortex (r = 0.957), hippocampus (r = 0.954) and striatum (r = 0.935; Extended Data Fig. 5a,d,f,g,i and Supplementary Data 4.3) as well as in organs: spleen (r = 0.982), liver (r = 0.963) and kidney (r = 0.963; Extended Data Fig. 5b,c,e). Blood and skin also showed high estimates of relative age correlations across different species: blood (r = 0.952, MAE = 0.022, 124 species) and skin (r = 0.942, MAE = 0.027, 92 species; Extended Data Fig. 5h,k).

Tissue-specific pan-mammalian clocks

The universal pan-mammalian clocks, derived from multiple tissue types, are essentially pan-tissue clocks. We also constructed analogous clocks solely based on blood (Universal BloodClock 2 and Universal BloodClock 3) and skin (Universal SkinClock 2 and Universal SkinClock 3), the tissues most readily accessible across all species. These tissuespecific clocks tend to demonstrate slightly higher accuracy than the pan-tissue clocks when analyzing their respective tissues. Both the blood and skin clocks exhibit robust age correlations ($r \approx 0.983-0.987$ for blood and $r \approx 0.951-0.968$ for skin; Extended Data Fig. 4c,g).

Human mortality risk, clinical biomarkers and lifestyle factors

Retrospective studies indicate that human epigenetic clocks can predict mortality risk and time to death, even when adjusted for chronological age and other risk factors^{23,26,27}. We tested whether this applies to pan-mammalian methylation clocks, using data from the Framingham Heart Study Offspring cohort (FHS, n = 2,544) and the Women's Health Initiative (WHI, n = 2,107). We devised a method to impute mammalian methylation array data from human Infinium array data (Supplementary Note 5). Our meta-analysis demonstrates that both clocks 2 and 3 can predict human mortality risk after adjusting for age and other confounders. The hazard ratio (HR) for 1 year of epigenetic age acceleration was significantly associated with all-cause mortality (HR = 1.03 and $P = 6.0 \times 10^{-19}$ for clock 2 and HR = 1.03, $P = 5.3 \times 10^{-11}$ for clock 3; Fig. 3a,b), although less pronounced than specialized human clocks designed to estimate human mortality risk^{22,23,28}.

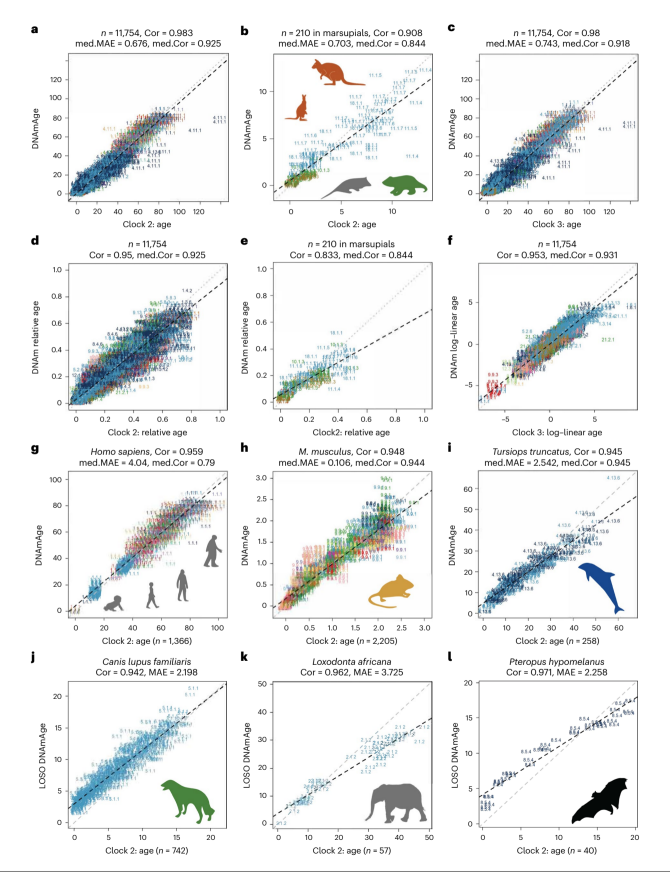
We evaluated the cross-sectional associations of lifestyle factors and clinical biomarkers with clocks 2 and 3 in the same cohorts. Robust correlation analysis (biweight midcorrelation (bicor)²⁹) revealed associations of both clocks with inflammation (C-reactive protein, bicor = 0.12, $P = 9.9 \times 10^{-16}$) and dyslipidemia (triglyceride

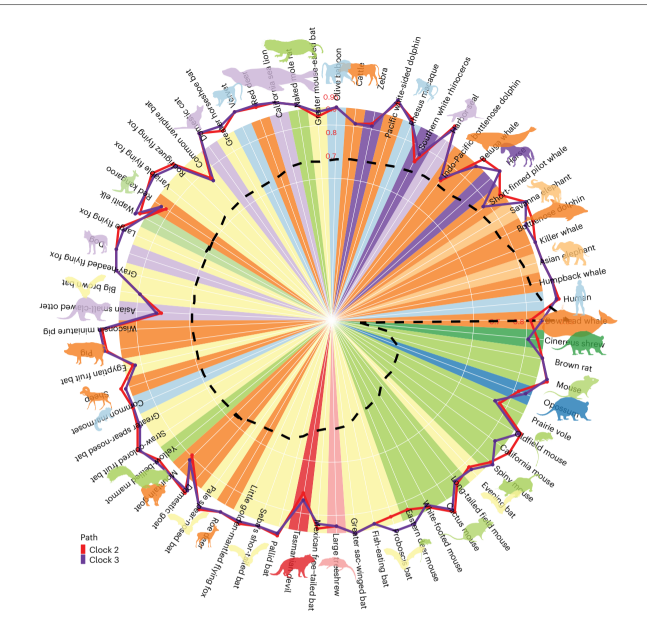
Fig. 1 | **Universal clocks for transformed age across mammals.** The figure displays relative age estimates of universal clock 2 (clock 2) and log–linear-transformed age of universal clock 3 (clock 3). Relative age estimates incorporate maximum lifespan and assume values between 0 and 1. Log–linear age is formulated with ASM and gestational time. **a**–**i**, Age estimated by LOFO cross-validation for clock 2 and clock 3. **j**–**l**, Age estimated via LOSO cross-validation for clock 2. The DNAm estimates of age (*y* axes) of **a**–**c** are transformations of relative age (clock 2) or log–linear age (clock 3) into units of years. **b**, **e**, Only marsupials

(nine species). Each panel reports a Pearson correlation (Cor) coefficient. The gray and black dashed lines correspond to the diagonal line (y=x) and the regression line, respectively. Median correlation (med.Cor) and median of MAE (med.MAE) are calculated across species ($\mathbf{a}-\mathbf{f}$) or across species–tissue ($\mathbf{g}-\mathbf{l}$). All correlation *P* values are highly significant ($P < 1.0 \times 10^{-22}$). Each sample is labeled by mammalian species index and indicated by tissue color (Supplementary Data 1.3–1.4). All *P* values reported are unadjusted and two sided. levels, $P = 3.2 \times 10^{-7}$; Supplementary Table 2). Less significant associations were for fasting glucose levels (P = 0.0093), body mass index (P = 0.011), smoking status (P = 0.027) or physical exercise (P = 0.0064). While these are nominally significant, they are far weaker than those observed with custom clocks for human mortality risk^{23,28}.

Heritability analysis in humans

To investigate whether genetic control within a species influences the epigenetic aging rates measured by pan-mammalian clocks, we used human pedigree data from the FHS. Pedigree-based polygenic models of epigenetic age, adjusted for age and sex, yielded significant





I Primates
1.1.1 Human
1.1.2 Gorilla
1.1.3 Chimpanzee
1.1.4 Orangutan
1.2.1 Diademed sifaka
1.2.2 Golden-crowned sifaka
1.2.3 Potto
1.3.1 White-headed lemur
1.3.2 Crowned lemur
1.3.3 Brown lemur
1.3.4 Black lemur
1.3.5 Mongoose lemur
1.3.6 Red-bellied lemur
1.3.7 Collared brown lemur
1.3.8 Blue-eyed black lemur
1.3.9 Red lemur
1.3.10 Sanford's brown lemur
1.3.11 Bamboo lemur
1.3.12 Ring-tailed lemur
1.3.13 Red ruffed lemur
1.3.14 Red ruffed lemur
1.4.1 Vervet
1.4.2 Rhesus macaque
1.4.3 Olive baboon
1.5.1 Slender loris
1.5.2 Slow loris
1.5.3 Pygmy slow loris
1.5.4 Potto
1.6.1 Fat-tailed dwarf lemur

1 Primates

1.7.1 South African galago
1.7.2 Greater galago
1.8.1 W.F. marmoset
1.8.2 Common marmoset
1.9.1 Aye-aye
2 Proboscidea
2.1.1 Asian elephant
2.1.2 Savanna elephant
3 Lagomorpha
3.1.1 European rabbit
3.1.2 Cotton rabbit
4 Artiodactyla
4.1.1 Addax
4.1.2 Impala
4.1.3 Springbok
4.1.4 Cattle
4.1.5 Domestic goat
4.1.6 White-bearded gnu
4.1.7 Thomson's gazelle
4.1.8 Slender H. gazelle
4.1.9 Roan antelope
4.1.10 Sable antelope
4.1.11 Nile lechwe
4.1.12 Gerenuk
4.1.13 Dama gazelle
4.1.14 Grant's gazelle
4.1.15 Soemm.'s gazelle
4.1.16 Mountain goat

1.6.2 Gray mouse lemur

1.6.3 N. giant mouse lemu

4.1.17 Sheep 4.1.18 Eland 4.1.19 Nyala 4.1.20 Bongo 4.1.21 Lesser kudu 4.1.22 Sitatunga 4.1.22 Greater kudu 4.2.1 Alpaca 4.3.1 Roe deer 4.31 fkoe deer 4.32 Red deer 4.33 Indian muntjac 4.35 Wapiti elk 4.41 Giraffe hybrid 4.42 Okapi 4.71 Pig 4.72 Wi mini. pig 4.72 Wi mini. pig 4.72 Wi mini. pig 4.713 Devheda whale 4.133 Chorme-s dolphin 4.132 Hector's dolphin 4.133 Erketor's dolphin 4.134 PAC ws. dolphin 4.13.4 PAC w.s. dolphin 4.13.5 Killer whate 4.13.6 Bottlenose dolphin 4.13.9 IP. bottlenose dolphin 4.13.10 R-toothed dolphin 4.13.11 Maui dolphin 4.13.12 Common dolphin 4.17.1 Harbor porpoise 4.19.1 Beluga whale

5.1.1 Dog 5.1.3 Maned wolf 5.2.1 Cheetah 5.2.2 Domestic cat 5.2.4 Lion 5.2.5 Tiger 5.2.6 Florida panther 5.2.6 florida panther 5.5.1 Spotted hyena 5.6.1 Spotted hyena 5.6.2 Southern sea otter 5.6.3 Førret 5.7.1 Pacific walrus 5.8.1 Steller sea lion 5.8.2 AU sea lion 5.8.2 AU sea lion 5.8.3 California sea lion 5.9.1 Harbor seal 5.9.2 Harp seal 5.11.1 Florida black bear **G Perissodarylia** 6 Perissodactyla 6.1.1 Horse 6.1.2 Grevy's zebra 6.1.3 Zebra 6.1.4 Somali wild ass 6.2.1 White rhino 6.2.2 E. black thinoceros 6.2.3 Greater o b. rhino. 6.2.3 Greater o.h. rhino 8 Chiroptera 8.4.1 Jamaican fruit bat 9 Rodentia 9.1.2 Cape mole rat 9.1.3 Naked mole rat

5 Carnivora

5.1.5 Red fox

8.4.2 Seba's s.t. bat 8.4.3 Common vampire bat 8.4.4 Lesser l.n. bat 8.4.5 Pale spear-nosed bat 8.4.6 Greater s.n. bat 8.5.1 Lesser s.n. fruit bat 8.5.2 Straw-colored fruit bat 8.5.1 Lesser S.n. Truit Dat 8.5.2 Straw-colored fruit bat 8.5.3 kina-colored fruit bat 8.5.4 Variable flying fox 8.5.5 Cerey-headed flying fox 8.5.6 Little gun. flying fox 8.5.7 Rodriguez flying fox 8.5.8 Large flying fox 8.5.8 Large flying fox 8.5.8 Large flying fox 8.5.8 Large flying fox 8.5.7 Brandts bat 8.7.7 Brandts bat 8.7.8 Noctule 8.7.4 Evening bat 8.7.8 Noctule 8.7.8 Frandts bat 8.7.9 Greater me, bat 8.7.19 Greater me, bat 8.7.19 Greater me, bat 8.7.19 Crish-eating bat 8.2.7 Plassing bat 8.2.7 Plassing bat 8.8.2 Pallas's mastiff bat 8.8.3 Mexican f.t. bat 8.17.1 Greater s.w. bat 8.17.2 Proboscis bat mole rat 11.1.7 Red kangaroo 11.4.1 Koala





narrow-sense heritability estimates for clock $2(h^2 = 0.44, P = 3.4 \times 10^{-8})$ and clock 3 ($h^2 = 0.41$, $P = 4.0 \times 10^{-7}$). These heritability estimates for pan-mammalian clocks are on par with that of Horvath's human pantissue clock $(h^2 = 0.39, P = 4.0 \times 10^{-7})^4$.

Epigenetic reprogramming reverses epigenetic age

Epigenetic clocks, such as the human pan-tissue clock, suggest that cellular reprogramming based on the Yamanaka factors (collectively termed as OSKM: OCT4, SOX2, KLF4, and c-MYC) induces age reversal^{4,30}. Fig. 2|Accuracy of universal clocks are independent of species lifespan. The circle plot displays Pearson correlation between age and DNAmAge estimated by universal clocks 2 (clock 2) and 3 (clock 3) for various species. Of the 185 species, correlation analysis was performed on 69 species (with $n \ge 15$ in a single tissue) across 12 taxonomic orders. We took log transformation of maximum lifespans of species and divided them by log (211), which is the maximum lifespan of bowhead whales. Values of the resulting ratios ranged from 0.12 (cinereus shrew) to 1 (bowhead whales). These ratios are displayed in descending order in the circle plot marked by the black dashed line, starting with the bowhead whale (1) and human (0.90) and ending with the cinereus shrew (0.12), in counterclockwise direction. In the background, circumferences with increasing radii represent increasing correlation levels up to 0.9. These correlations between age and DNAmAge were estimated by clock 2 (red path line) and clock 3 (purple path line) for each species. Colors within the circle represent the taxonomic order of the corresponding species, as listed below the circle. The median of correlation

across species is 0.926 for clock 2 and 0.918 for clock 3. Straw-colored fruit bats exhibit the highest correlation (*r* = 0.985) based on clock 2, and Wisconsin miniature pigs have the second highest correlation (*r* = 0.984) based on clock 3. A majority of species with their circle lines located outside the background indicates that their correlation estimates are greater than 0.9. The text at the bottom lists the 185 species under their corresponding taxonomic order. Each taxonomic order is marked by the same color matching with the circle plot. The numbers after the first and second decimal points enumerate the taxonomic family and species, respectively. AU, Australian; Comme., Commerson's; E., eastern; f.t., free-tailed; g.m., golden-mantled; H. (gazelle), Horn gazelle; Hoff., Hoffman's; IP, Indo-Pacific; L.'s, Linne's; l.n., long-nosed; m.e., mouse-eared; mini., miniature; N., northern; o.h., one horned; s.c., small-clawed; PAC w.s., Pacific white-sided; R.-toothed, Rough-toothed; Soemm., Soemmerring's; S.finn., Short-finned; s.n., short nosed; s.t., short-tailed; s.w., sac-winged; W. western; W.F., White-fronted; WI mini., Wisconsin miniature.

To examine whether the universal clocks show a similar age-reversal pattern during reprogramming, we applied clock 2 and clock 3 to a previously published reprogramming dataset in human dermal fibroblasts³¹. We imputed the mammalian methylation array data on the basis of the existing human Infinium array data. Both clocks suggest age reversal after OSKM transduction (Fig. 3c, d). Notably, universal clock 2 showed a decrease in epigenetic age in partially reprogrammed cells after 11 d (Fig. 3c), mirroring observations with human epigenetic clocks^{4,30,32}.

Transgenic mice for studying the somatotropic axis

Growth hormone, generated by somatotropic cells, stimulates body tissue growth, including bone. The somatotropic axis (growth hormone and insulin-like growth factor 1 (IGF-1) levels and their cognate receptors) is central to aging and longevity studies³³. Decreased growth hormone–IGF-1 signaling extends longevity in various species, including mice³⁴. A full-body growth hormone receptor-knockout (KO) (GHRKO) mouse holds the official record for being the longest-lived representative of *Mus musculus*, living 1 week shy of 5 years³³.

We examined whether reduced growth hormone–IGF-1 pathway activity slows universal pan-mammalian clocks, using three mouse models: (1) Snell dwarf mice, lacking growth hormone production and hence living longer^{35,36}, (2) full-body GHRKO mice with increased lifespan³⁷ and (3) liver-specific GHRKO mice, showing lowered serum IGF-1 levels but not lifespan increase.

Clock 2 and 3 analyses revealed that Snell dwarf mice exhibit a significantly lower epigenetic age across all considered tissues than wild-type mice (cerebral cortex, Student's *t*-test, $P = 2.0 \times 10^{-8}$; kidney, $P = 6.0 \times 10^{-10}$; liver, $P = 1.0 \times 10^{-7}$; tail, $P = 1.0 \times 10^{-6}$; blood, $P = 2.0 \times 10^{-3}$; spleen, P = 0.03; Fig. 3e, f). Similarly, full-body GHRKO mice showed lower epigenetic age in several tissues (liver, $P = 3.0 \times 10^{-5}$; kidney, $P = 2.0 \times 10^{-5}$; cerebral cortex, P = 0.02; Fig. 3e, f).

Growth hormone receptor signaling stimulates IGF-1 liver synthesis, suggesting that dwarf mice's epigenetic age reversal may be

Fig. 3 | **Applications of universal pan-mammalian clocks in human cohorts, reprogramming experiment and murine anti-aging studies. a,b**, Forest plots representing the fixed effect (FE) model meta-analysis, combining HRs from Cox regression models for time to death, based on epigenetic age acceleration measures of clock 2 (AgeAccelClock2) and clock 3 (AgeAccelClock3) across different ethnic groups within two epidemiological cohorts. Each row indicates an HR for a 1-year increase in the age acceleration (AgeAccel) measure, along with a 95% confidence interval (Cl). **c,d**, DNAmAge estimates of human dermal fibroblasts during OSKM-induced reprogramming. The *y* axes are DNAmAge estimates of clock 2 and clock 3 at day 0, 3, ..., 42 and 49, respectively, during reprogramming³¹. **e**, Evaluations of mouse anti-age interventions: (1) age-matched Snell dwarf mutation study: 48 normal and 47 dwarf mice with ages of approximately 0.52 (mean \pm s.d. = 0.52 \pm 0.01) years, (2) age-matched whole-body GHRKO experiment 1 (Exp.1) with 36 normal and 35 GHRKO mice (mean \pm s.d. = 0.65 \pm 0.06 years), (3) age-matched GHRKO due to lower circulating IGF-1 levels. This hypothesis, however, is not supported by our epigenetic age measurements of liver-specific GHRKO mice, which exhibit a non-significant difference from the wild-type controls (Fig. 3e). Both clocks 2 and 3 show that the liver-specific GHRKO mice are not epigenetically younger than wild-type mice (Fig. 3e). Unlike full-body GHRKO mice, liver-specific GHRKO mice do not possess a longevity advantage^{38,39}.

Caloric restriction in mice

Caloric restriction (CR), which also slows the somatotrophic axis (growth hormone–IGF-1), is associated with prolonged lifespan in several mouse strains^{40,41}. Previous studies using mouse clocks have shown that CR reduces the rate of epigenetic aging in liver samples⁵⁻⁷. Using existing methylation data from a murine study of CR⁴², we find that clocks 2 and 3 yield a reduced epigenetic age for mouse liver samples ($P = 6.0 \times 10^{-12}$ for clock 2, $P = 7.0 \times 10^{-15}$ for clock 3; Fig. 3e,f). These results for pan-mammalian clocks align with those obtained with mouse-specific clocks^{5,43,44}.

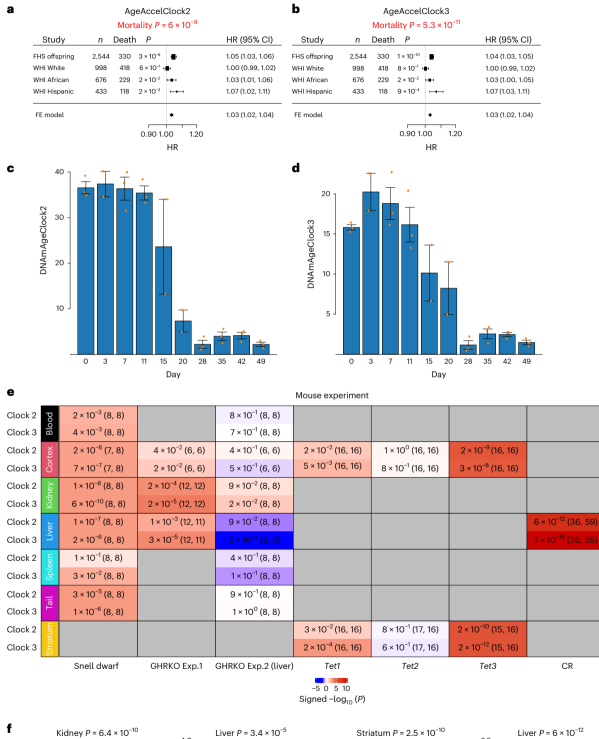
TET enzyme-KO studies in mice

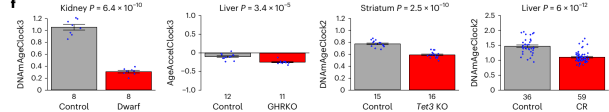
TET enzymes are instrumental in active DNA demethylation. Because hydroxymethylation mediated by TET enzymes is prevalent in brain tissue, we applied the universal clocks to brain tissue samples from *Tet1-, Tet2-* and *Tet3-*KO mice. Analysis with our universal clocks revealed that *Tet3-*KO mice exhibit a reduced rate of epigenetic aging (cerebral cortex, $P = 3.0 \times 10^{-9}$ and striatum, $P = 2.0 \times 10^{-12}$; Fig. 3e, f). By contrast, significant epigenetic age-reversal effects in brain tissue were relatively weak for *Tet1* (cerebral cortex, $P = 6.0 \times 10^{-3}$ and striatum, $P = 2.0 \times 10^{-4}$; Fig. 3e) and could not be observed for *Tet2-*KO mice (P > 0.6; Fig. 3e).

The differential effect of *Tet3* KO versus *Tet1* or *Tet2* KO in neurons echoes the results of an epigenetic reprogramming study in mouse retinal ganglion cells (*Oct4, Sox2* and *Klf4* (ref. 45)).

experiment 2 with GHRKO in livers only with 48 normal and 48 GHRKO genotypes (mean ± s.d. = 0.51 ± 0.03 years old), (4) Tet gene-KO study with all samples at age 0.5 years (*Tet1*, 32 controls and 32 *Tet1* KO; *Tet2*, 33 controls and 32 *Tet2* KO; *Tet3*, 31 controls and 32 *Tet3* KO) and (5) CR study in livers (59 in CR versus 36 control mice with all ages at 1.57 years old). Comparisons in experiments 2 and 3 were based on AgeAccel measures. The color gradient is based on the sign of *t*-test for controls versus experimental mice, with a positive sign indicating that the mice in the control group exhibit higher age acceleration than the mice in the experimental group. **f**, Bar plots for selective tissue types and clocks across Snell dwarf mice (eight normal and eight dwarf mice) GHRKO experiment 1 (12 normal and 11 GHRKO mice), *Tet3*-KO mice (15 normal and 16 *Tet3*-KO mice) and the entire CR experiment, respectively. The orange dots in **c** and **d** and the blue dots in **e** correspond to individual observations. The *y* axes of the bar plots depict the mean of one standard error. All *P* values reported are two sided and are unadjusted for multiple testing. Meta epigenome-wide association study of age across species Universal clocks, founded on penalized regression models, consist solely of CpG sites that are most predictive of age. Consequently, most other age-related CpG sites are not included in the final regression models.

To identify all age-related CpG sites, we carried out two-stage metaanalysis across species and tissues in eutherians (98% of the samples). Our epigenome-wide association study (EWAS) of age indicated that CpG sites becoming increasingly methylated with age (positively correlated with age) are conserved across tissues and species (Fig. 4a).





CR

Imposing a stringent unadjusted significance threshold of $\alpha = 10^{-200}$ limited our analysis to fewer than 1,000 CpG sites across all eutherian species and tissues (Fig. 4a and Supplementary Data 6.1). Of the 832 resulting age-related CpG sites, those most significantly associate with age were cg12841266 ($P = 1.4 \times 10^{-1.001}$) and cg11084334 ($P = 2.6 \times 10^{-891}$), both located in exon 2 of *LHFPL4* (hg38). Notably, cg12841266 exhibited a correlation ≥ 0.8 in 28 species (Supplementary Data 7; three examples are shown in Fig. 4b–d). Another CpG, cg09710440, resides in exon 1 of *LHFPL3* ($P = 5.0 \times 10^{-787}$), a paralog of *LHFPL4* (Fig. 4a, Extended Data Fig. 6 and Supplementary Data 6.1–6.7). As *LHFPL4* and *LHFPL3* are in human chromosomes 3 and 7, respectively, their consistent age-related gain of methylation is not due to physical proximity.

Beyond *LHFPL4* and *LHFPL3*, other significant gene pairs among the top 30 age-related CpG sites include *ZIC1* (chromosome 3) and *ZIC2* (chromosome 13), *PAX2* (chromosome 10) and *PAX5* (chromosome 9) and *CELF6* (chromosome 15) and *CELF4* (chromosome 18; Supplementary Data 6.1). Located on separate chromosomes, their shared age-related methylation changes cannot be due to physical proximity, indicating a likely functional role in aging. Intriguingly, each gene pair encodes proteins with activities in development.

We observed that numerous cytosines change during the initial 6 weeks of murine postnatal development. In particular, *LHFPL4* cg12841266 displayed a positive correlation (r > 0.6) with age across murine tissues, especially in the brain and muscle (Fig. 5a–g). High age correlations were also evident in older mice (ranging from 0.2 years to 2.5 years; Fig. 5h–o).

We obtained a broad overview of age association across different temporal domains by repeating our two-stage meta-EWAS for young, middle and old-age groups (Fig. 6a–c). Importantly, methylation changes related to age in young animals strongly align with those seen in middle-aged or old animals, refuting the idea that these changes are purely tied to organismal development (Fig. 6a–c). This observation is further reinforced by visualizing the mean methylation levels (β values) of age-related CpG sites relative to their distances from transcriptional start sites (TSS; Fig. 6d).

EWAS of age in marsupials and monotremes

We extended the age-related EWAS analysis to marsupials and monotremes. The top age-related CpG sites for marsupials were found near genes involved in development, including *GRIK2* ($P = 8.8 \times 10^{-21}$; Supplementary Data 6.8), encoding a neurotransmitter-associated glutamate receptor, and *ZIC4* ($P = 2.7 \times 10^{-19}$), encoding a zinc finger protein. The age-related EWAS in monotremes implicated cg22777952 in *FOXB1* ($P = 8.1 \times 10^{-10}$; Supplementary Data 6.9), encoding a forkhead box protein. Moderate positive correlation with eutherian age-related methylation changes was observed (r = 0.295 in marsupials, Fig. 4e; r = 0.227in monotremes, Fig. 4f), in part due to the lower sample numbers in these groups. However, the age effect on methylation of cg11084334

Fig. 4 | Meta-analysis of methylation change in function of chronological age across species and tissues. a-d,g,h, Eutherian EWAS of age. a, Meta analysis -log10 (Pvalues) for age-related CpG sites (annotated by proximal genes) on chromosomes (x axis in hg38). Top and bottom, CpG sites that gain or lose methylation with age, respectively. CpG sites in red and blue denote highly significant positive and negative age correlation ($P < 10^{-200}$), respectively. The most significant CpG (cg12841266, $P = 1.41 \times 10^{-1.001}$) resides in exon 2 on the LHFPL4 gene in humans and most mammals, followed by cg11084334 $(P = 2.59 \times 10^{-891})$. These two CpG sites and cg097720 $(P = 4.97 \times 10^{-787})$ located in the paralog gene LHFPL3 are marked with purple diamonds. b-d, Scatterplots of cg12841266 versus chronological age (years) in mini pigs (Sus scrofa minusculus) (b), Oldfield mice (Peromyscus polionotus) (c) and horses (Equus caballus) (d). Tissue samples are labeled by the mammalian species index and colored by tissue type as detailed in Supplementary Data 1.1-1.4. e,f, Correlation analysis between Z scores of EWAS of age in eutherians versus marsupials (e) and eutherians versus monotremes (f). g,h, Annotations of the top 1,000 CpG sites

(not cg12841266) in *LHFPL4* is preserved in marsupials ($P = 4.8 \times 10^{-7}$; Fig. 4e) and monotremes ($P = 2.4 \times 10^{-5}$; Fig. 4f), despite these limitations.

Meta-analysis of age-related CpG sites across specific tissues

To understand age-related CpG sites across species and tissues, we focused on six tissues with many available species: brain (whole and cortex), blood, liver, muscle and skin. We performed an EWAS metaanalysis on 935 whole brains (18 species-brain tissue categories, eight species), 391 cortices (six species), 4,513 blood samples (56 species), 1,063 livers (ten species), 354 muscle samples (five species) and 2,363 skin samples (65 species; Supplementary Data 1.6–1.11).

Consistently across all tissues, CpG sites with positive age correlations outnumbered those with negative correlations (Extended Data Fig. 6). While many age-related cytosines were either specific to individual organs (Supplementary Data 6.2–6.7) or shared between several organs, 51 CpG sites (48 positively and three negatively age related) were common to all five organs (Fig. 4g and Supplementary Table 3). In total, 35 genes were proximal to the 48 positive CpG sites, and three genes were proximal to the three negative CpG sites. Interestingly, 20 of these 35 genes encode transcription factors (TFs), including 11 homeobox proteins, seven zinc finger TFs and two paired box proteins, involved in developmental processes including embryonic development (Supplementary Table 3). The relevance of this becomes evident below, where the chromatin state, function and tissue-specific accessibility associated with the location of age-related CpG sites are described.

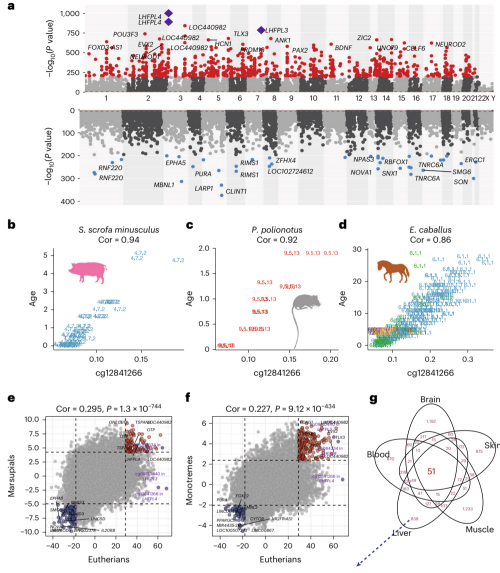
Analyses of chromatin states of DNA bearing age-related cytosines

We observed that 57% of the top 1,000 positively age-related CpG sites were situated in a CpG island (human genome), while only 2% of the top 1,000 negatively age-related CpG sites resided there (EWAS of age across all tissues; Supplementary Data 6.1).

To understand the epigenetic context of age-related CpG sites, we accessed a detailed universal chromatin state annotation of the human genome. This resource, derived from 1,032 experiments mapping 32 chromatin marks across 100+ human cell and tissue types⁴⁶ (Fig. 4h, Extended Data Fig. 7 and Supplementary Data 8.2-8.9), allowed us to overlay the positions of the top 1,000 age-related CpG sites. We found that positively age-related CpG sites were significantly enriched in states associated with polycomb repressive complex 2 (PRC2)-binding sites (states BivProm1, BivProm2, ReprPC1), These CpG sites localized to PRC2-binding sites, as defined by embryonic ectoderm development (EED), enhancer of zeste 2 PRC2 subunit (EZH2) and PRC2 subunit (SUZ12) binding (the first row of Fig. 4h). This PRC2 enrichment could be observed for all tissue types collectively (odds ratio (OR) = 22.8, hypergeometric $P = 1.9 \times 10^{-449}$) and when analyzed individually: blood (OR = 29.8, $P = 2.9 \times 10^{-510}$), liver (OR = 14.3, $P = 7.3 \times 10^{-338}$), skin $(OR = 14.3, P = 9.9 \times 10^{-337})$, cortex $(OR = 6.5, P = 3.7 \times 10^{-163})$ and brain

with increased or decreased methylation with age that were identified in EWAS meta-analysis across all species and tissues (results in **a**) (brain, cortex, blood, liver, muscle and skin tissues). **g**, The overlap of age-associated CpG sites across various organs, based on the top 1,000 CpG sites showing positive or negative age correlation in EWAS. The Venn diagram includes 51 age-associated CpG sites shared across all organs, adjacent to 38 genes (35 with positive and three with negative age correlation) categorized by protein family. The 35 positive genes are color coded based on their protein family: two in LHFPL, 12 in homeobox, three in paired box or T-box, three in bHLH, seven in zinc finger and eight in others. **h**, Selected universal chromatin state and polycomb group protein enrichment results. ORs (*P* values) are presented in each cell. The color gradient is based on $-\log_{10}$ (hypergeometric *P* value) times sign of OR > 1. The complete results are listed in Extended Data Fig. 7. State annotation can be found in Supplementary Data 8.2. HET denotes heterochromatin. Except for the hypergeometric analysis in **h**, all figure *P* values are unadjusted and two sided. (OR = 3.2, $P = 9.7 \times 10^{-57}$). Indeed, the majority of the top 1,000 positively age-related CpG sites were significantly enriched in PRC2-binding sites: 80.8% (808 CpG sites) in blood, 67.5% in liver and 67.2% in skin (Supplementary Data 8.1).

PRC2, a transcriptional repressor complex, is a key contributor to H3K27 methylation, a chromatin modification linked to transcriptional repression⁴⁷. Importantly, PRC2-mediated histone 3 lysine 27 (H3K27) methylation is crucial for establishing bivalent promoters, which house





h																		
••					Increa	ised wit	th age			Decreased with age								
	PRC2		22.8 (1.9 × 10 ⁻⁴⁴⁹)	29.8 (2.9 × 10 ⁻⁵¹⁰)	14.2 (9.9 × 10 ⁻³³⁷)	14.3 (7 × 10 ⁻³³⁸)	2 (5 × 10 ⁻¹⁷)	3.2 (9.7 × 10 ⁻⁵⁷)	6.5 (3.7 × 10 ⁻¹⁶³)	0.02 (5.8 × 10 ⁻⁶²)	0.04 (2.4 × 10 ⁻⁵⁷)	0.03 (8.4 × 10 ⁻⁵⁹)	0.1 (3.2 × 10 ⁻³⁸)	0.2 (2.1 × 10 ⁻³¹)	0.04 (5.8 × 10 ⁻⁵⁶)	0.03 (8.4 × 10 ⁻⁵⁹)		
	PRC1		1.4 (9.5 × 10 ⁻²)	1.2 (0.27)	1.2 (0.27)	1.5 (4 × 10 ⁻²)	0.06 (4.8 × 10 ⁻⁷)	0.3 (5.1 × 10 ⁻⁴)	0.2 (1.3 × 10 ⁻⁴)	0.6 (7.6 × 10 ⁻²)	0.6 (7.6 × 10 ⁻²)	0.6 (4.3 × 10 ⁻²)	0.5 (1.1 × 10 ⁻²)					
	BivProm1		9 (8.7 × 10 ⁻¹²¹)	7.6 (6.1 × 10 ⁻⁹⁷)	7.9 (2.8 × 10 ⁻¹⁰²)	10.1 (2.2 × 10 ⁻¹³⁹)	1.8 (5.7 × 10 ⁻⁶)	2.6 (2.5 × 10 ⁻¹⁴)	4 (4.7 × 10 ⁻³⁶)	0 (9 × 10 ⁻¹⁹)	0 (9 × 10 ⁻¹⁹)	0 (9 × 10 ⁻¹⁹)	0.05 (8.7 × 10 ⁻¹⁶)	0.1 (9 × 10 ⁻¹²)	0.05 (8.7 × 10 ⁻¹⁶)	0.1 (1.2 × 10 ⁻¹²)		
	BivProm2		15.7 (1.4 × 10 ⁻²⁴⁹)	16.5 (1.2 × 10 ⁻²⁶¹)	13.7 (3 × 10 ⁻²¹⁹)	10.3 (5 × 10 ⁻¹⁶⁰)	2.3 (9.4 × 10 ⁻¹²)	3.1 (3.7 × 10 ⁻²⁴)	5 (5.8 × 10 ⁻⁵⁹)			0 (4.5 × 10 ⁻²²)	0.02 (2.4 × 10 ⁻²⁰)	0.06 (1.1 × 10 ⁻¹⁷)	0.06 (1.1 × 10 ⁻¹⁷)	0 (4.5 × 10 ⁻²²)		
	ReprPC1		8.2 (1.6 × 10 ⁻⁹⁹)	9.5 (2.5 × 10 ⁻¹²¹)	6.6 (2.5 × 10 ⁻⁷⁴)	4 (1.2 × 10 ⁻³²)	2 (1.3 × 10 ⁻⁶)	2.9 (4.7 × 10 ⁻¹⁷)	5.2 (3.3 × 10 ⁻⁵¹)	0.2 (1.7 × 10 ⁻⁹)	0.2 (1.7 × 10 ⁻⁹)	0.2 (8.2 × 10 ⁻⁹)	0.03 (2.2 × 10 ⁻¹⁵)	0 (5.5 × 10 ⁻¹⁷)	0.3 (1.4 × 10 ⁻⁷)	0.3 (1.7 × 10 ⁻⁶)		
	Quies1		0 (5.5 × 10 ⁻⁵)	0 (5.5 × 10 ⁻⁵)	0.1 (6.1 × 10 ⁻⁴)	0 (5.5 × 10 ⁻⁵)	1.3 (0.26)	0.4 (3.5 × 10 ⁻²)	0.3 (1.2 × 10 ⁻²)	5.8 (5.8 × 10 ⁻²⁰)	3.4 (5.1 × 10 ⁻⁸)	6.4 (3.9 × 10 ⁻²³)	1.3 (0.26)	0.2 (3.4 × 10 ⁻³)	0.2 (3.4 × 10 ⁻³)	0.1 (6.1 × 10 ⁻⁴)		
	Quies2		0 (7.4 × 10 ⁻⁴)	0.6 (0.16)	0.3 (2.6 × 10 ⁻²)	0.4 (7.4 × 10 ⁻²)	5.3 (2.7 × 10 ⁻¹³)	2.8 (1.1 × 10 ⁻⁴)	4.9 (7.4 × 10 ⁻¹²)	1.6 (0.1)	0.1 (6.2 × 10 ⁻³)	0.3 (2.6 × 10 ⁻²)	0.6 (0.16)					
	HET1		0 (0.15)	0 (0.15)	0 (0.15)	0 (0.15)	0.5 (0.43)	0.5 (0.43)	0 (0.15)	5.3 (1.1 × 10 ⁻⁴)	3.4 (1.2 × 10 ⁻²)	7.5 (3.7 × 10 ⁻⁷)	0.5 (0.43)	0 (0.15)	0 (0.15)	0.5 (0.43)		
	HET7		0 (0.37)	0 (0.37)	0 (0.37)	0 (0.37)	0 (0.37)	0 (0.37)	0 (0.37)	7.1 (4.1 × 10 ⁻⁴)	1 (0.63)	7.1 (4.1 × 10 ⁻⁴)	1 (0.63)	0 (0.37)	1 (0.63)	3.2 (7.6 × 10 ⁻²)		
	TxEx4		0 (1.7 × 10 ⁻⁴)	0 (1.7 × 10 ⁻⁴)	0.5 (6.9 × 10 ⁻²)	0.1 (1.6 × 10 ⁻³)	0 (1.7 × 10 ⁻⁴)	0 (1.7 × 10 ⁻⁴)	0 (1.7 × 10 ⁻⁴)	3 (6.7 × 10 ⁻⁶)	2.6 (1.7 × 10 ⁻⁴)	0.7 (0.25)	4.1 (2.1 × 10 ⁻¹⁰)	6.7 (3.7 × 10 ⁻²²)	14.5 (1.7 × 10 ⁻⁶⁰)	12.9 (1.6 × 10 ⁻⁵²)		
	c	0.0	All	Blood	Skin	Liver	Muscle	Brain	Cortex	All	Blood	Skin	Liver	Muscle	Brain	Cortex		
PRC2-binding proportion																		
		Exo HET				Polycomb repressed Quiescent PRC1 and PRC2 TF-binding regions weak promot												

Resource

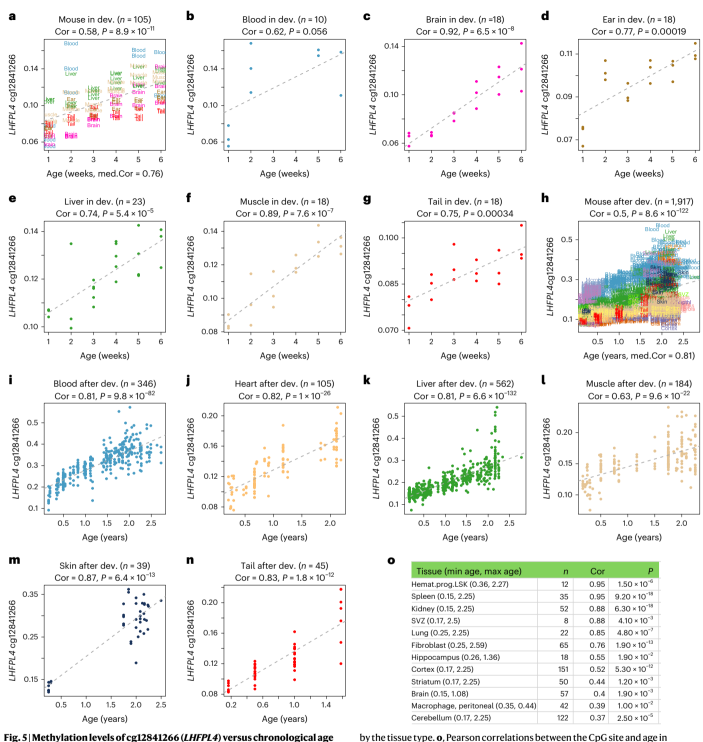
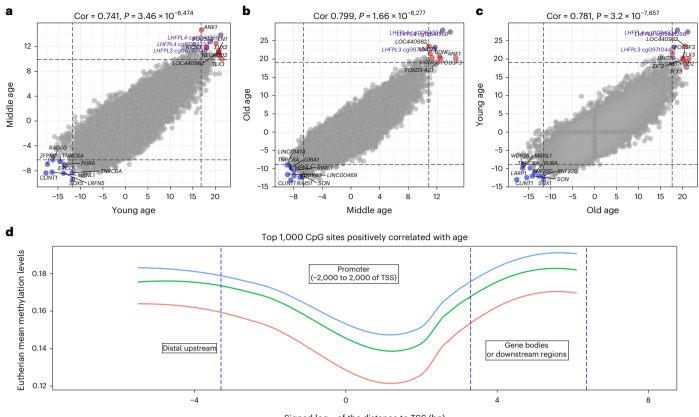


Fig. 5 | Methylation levels of cg12841266 (*LHFPL4*) versus chronological age in mouse tissues. Results are reported for different tissues and age groups. a-g, Postnatal development (dev.) (from 1 week to 6 weeks). h-o, Age effects in adult mice. Mean \pm s.d.⁹⁶ of chronological age is 3.5 \pm 1.7 (1.0–6.0) weeks in the developmental age group and 1.12 \pm 0.72 (0.15–2.78) years in the postdevelopmental group. a, h, All tissues combined. Each dot (sample) is colored

histones with both H3K27 trimethylation (H3K27me3) and histone 3 lysine 4 trimethylation (H3K4me3). As such, it is consistent that positively age-related CpG sites are also found to be enriched in bivalent promoter states (rows 3 and 4 of Fig. 4h). They show even greater presence in a bivalent state associated with more H3K27me3 than H3K4me3 (BivProm2) than in BivProm1, associated with more balanced levels of these histone additional mouse tissues and cell types from the Mammalian Methylation Consortium. Hemato.prog.LSK, hematopoietic progenitor cells with lineage⁻Sca⁻¹⁺c-Kit⁺ phenotype; max, maximum; min, minimum; *n*, sample size; SVZ, subventricular zone. Pearson correlation coefficients and nominal (unadjusted) two-sided correlation test *P* values are shown.

modifications⁴⁶. The top EWAS hit, *LHFPL4* cg12841266, in a bivalent state (BivProm2) and PRC2-binding region (EED-, EZH2-, SUZ12-binding sites), exemplifies this (Supplementary Data 8.1). These mammalian results echo those from human studies^{48,49}, in which tissue-independent age-related gain of methylation is characterized by cytosines that are located in PRC2-binding sites and bivalent chromatin domains.



Signed \log_{10} of the distance to TSS (bp)

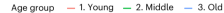


Fig. 6 | **EWAS of age in three different age groups.** For each species, the age groups were defined with respect to the average ASM obtained from the Animal Aging and Longevity Database (AnAge) (de Magalhaes et al.⁸⁶). We defined the three age groups using intervals defined by multiples of ASM: young age is defined as age <1.5 × ASM, middle age is defined as age between 1.5ASM and 3.5ASM, and old age is defined by age \geq 3.5ASM. Each axis reports *a Z* score from the meta-analysis EWAS of age across all mammalian species and tissues. Each dot corresponds to a CpG site. Labels are provided for the top ten hypermethylated or hypomethylated CpG sites according to the product of *Z* scores in *x* and *y* axes. CpG sites that are located in *LHFPL4* and *LHFPL3* are colored in purple. The Pearson correlation coefficient and corresponding nominal (unadjusted) two-sided correlation test *P* value can be found in the title. **a**, EWAS of age in young animals versus EWAS in niddle-aged animals. **b**, EWAS of age in middle-aged animals versus EWAS in old animals. **c**, EWAS of age in young animals versus EWAS

of age in old animals. The high pairwise correlations indicate that conserved aging effects in mammals are largely preserved in different age groups. Many of the top CpG sites for conserved aging effects in young mammals remain the top CpG sites for conserved aging effects in old mammals. Specifically, we analyzed the mean methylation levels in eutherians across the three age groups. **d**, Mean methylation (y axis) across the top 1,000 CpG sites positively correlated with age according to the EWAS across all mammalian tissue types (Fig. 4a). The x axis denotes the distance to the closest TSS in a log₁₀ scale of bp. The positive TSS indicates the direction from 5' to 3', and the negative TSS indicates from the direction from 3' to 5'. The horizontal phase is categorized into three regions: distal upstream \rightarrow promoter \rightarrow gene bodies. The mean methylation levels have higher propensity to increase with age.

We found that ORs for the overlap between positively age-related CpG sites and PRC2-binding sites were markedly higher in proliferative tissues (blood, skin, liver) than in non-proliferative tissues (skeletal muscle, brain, cerebral cortex; Fig. 4h). The distinction between proliferative and non-proliferative tissues also manifested when considering negatively age-related CpG sites (those that lose methylation levels with age). In highly proliferative tissues (blood, skin), age-related loss of methylation was seen in CpG sites located in select heterochromatin (HET1, HET7), which are marked by histone 3 lysine 9 trimethylation, or inactive chromatin states (Quies1, Quies2), as listed in Supplementary Data 8.2 and Vu & Ernst⁴⁶. Conversely, in non-proliferative tissues, agerelated methylation loss could be seen in the exon- and high-expression-associated transcription state TxEx4 (OR = $12.9, P = 1.6 \times 10^{-52}$ in the cerebral cortex and OR = 6.7, $P = 3.7 \times 10^{-22}$ in skeletal muscle). TxEx4 is far less enriched with age-related cytosines that lose methylation in proliferative tissues such as blood (OR = 2.6, $P = 1.7 \times 10^{-4}$) or skin (OR = 0.7, P = 0.25).

Overlap with late-replicating domains

Our chromatin state analysis of age-related loss of methylation demonstrated that it is important to distinguish proliferating tissues (blood, skin) from non-proliferative tissues (brain, muscle). Consequently, we examined the correlation between DNA replication and methylation. Late-replicating genome domains, prone to partial methylation, show pronounced methylation loss in solo WCGW cytosines (CpG sites flanked by A or T on either side⁵⁰). We overlaid the top 1,000 agerelated CpG sites (positive or negative) on the reported late-replicating domains, which are enriched with partially methylated domains (PMDs)⁵⁰. As previously reported for human tissues⁵⁰, we observed age-related loss of methylation in PMDs and solo WCGW sites in mammalian tissues that proliferate, such as blood and skin (Extended Data Fig. 8 and Supplementary Data 9). Notably, the top 1,000 negatively agerelated CpG sites overlap significantly with CpG sites that are both common PMDs and solo WCGW sites (hg19): skin (OR = 7.9, $P = 1.6 \times 10^{-90}$), blood (OR = 5.3, $P = 1.5 \times 10^{-50}$) and all tissues (OR = 7.3, $P = 4.4 \times 10^{-81}$;

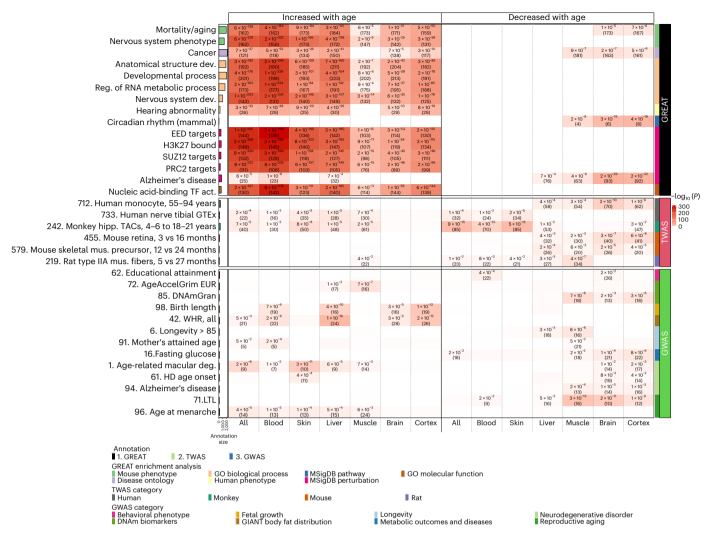


Fig. 7 | **Biological pathways and functional gene sets enriched in age-related CpG sites.** Selected results from (1) genomic region-based GREAT functional enrichment (top), (2) gene-based EWAS–TWAS enrichment analysis (middle) and (3) genomic region-based EWAS–GWAS enrichment analysis (bottom). All enrichment analyses were based on hypergeometric tests with background based on the mammalian array. The bar plots in the first column report the total number of genes at each studied gene set adjusted based on the background. The left and right parts of the *x* axis list the top 1,000 CpG sites that increased or decreased with age from meta-EWAS of age across all blood, skin, liver, muscle, brain and cerebral cortex tissues, respectively. On the right side, the first column color band depicts the three types of enrichment analyses. The second column color band depicts (1) six ontologies in the GREAT analysis, (2) four species in our TWAS collections and (3) seven categories of human complex traits in the GWAS as described in the legend. The heatmap color codes $-\log_{10}$ (hypergeometric P values). Unadjusted hypergeometric P values (number of overlapped genes) are reported in the heatmap provided (1) false discovery rate < 0.05, P < 0.001 and the number of overlapped genes \geq 3 for GREAT analysis, (2) P < 0.05 for EWAS-TWAS and (3) P < 0.05 for EWAS-GWAS. Comprehensive results can be found in Supplementary Data 10, 12 and 13. Abbreviations: act., activity; deg., degeneration; AgeAccelGrim, epigenetic age acceleration derived from the mortality clock: GrimAge²³; DNAmGran, DNAm granulocyte (Supplementary Note 5); GIANT, Genetic Investigation of ANthropometric Traits; GTEx, Genotype–Tissue Expression; HD, Huntington's disease; hipp., hippocampal; LTL, leukocyte telomere length; MSigDB, Molecular Signatures Database; mus., muscle; OPCs, oligodendrocyte precursor cells; reg., regulation; TACs, transiently amplifying progenitor cells; WHR, waist-to-hip ratio.

Extended Data Fig. 8). Contrastingly, non-proliferative tissues, such as the brain, show a different pattern: CpG sites losing methylation with age are enriched in highly methylated domains (HMDs, OR = 3.3, $P = 1.9 \times 10^{-74}$) over PMDs (OR = 0.2, $P = 4.9 \times 10^{-64}$). CpG sites gaining methylation with age show weaker overlap with both PMDs and highly methylated domains. Similar findings were observed in late-replicating mouse genome domains (mm10; Extended Data Fig. 8). In summary, pan-mammalian CpG sites losing methylation with age are enriched in late-replicating regions of highly proliferative tissues.

Functional enrichment analysis of age-related CpG sites

We used the Genomic Regions Enrichment of Annotations Tool (GREAT) to annotate the potential function of *cis* regulatory regions

of age-related CpG sites⁵¹. We sought to identify biological processes and pathways potentially associated with the top 1,000 positively and negatively age-related CpG sites (Fig. 7 and Supplementary Data 10.1–10.17). To avoid array-design bias, we used mammalian array CpG sites as a background set in our hypergeometric enrichment test.

Analysis of CpG sites positively correlated across all tissues revealed 'nervous system development' as a highly significant gene ontology (GO) term ($P = 1.3 \times 10^{-203}$). This term was consistent across blood ($P = 1.9 \times 10^{-224}$), liver ($P = 2.6 \times 10^{-137}$), muscle ($P = 3.4 \times 10^{-14}$), skin ($P = 1.7 \times 10^{-145}$), brain ($P = 6.4 \times 10^{-35}$) and cortex ($P = 1.0 \times 10^{-78}$). Other significant GO terms included 'developmental process', 'regulation of RNA metabolic process', 'nucleic acid-binding TF activity', 'pattern specification' and 'anatomical structure development' (Fig. 7). The GREAT analysis also indicated that a significant proportion of the top 1,000 positively age-related CpG sites are located in PRC2 target sites ($P = 8.3 \times 10^{-212}$), which was also true for individual core PRC2 subunits (SUZ12, EED or EZH2; Fig. 7). It follows that these CpG sites were also enriched in promoters with the H3K27me3 modification in embryonic stem cells for all tissues ($P = 1.9 \times 10^{-269}$), blood ($P = 2.8 \times 10^{-202}$), liver ($P = 3.4 \times 10^{-132}$), muscle ($P = 9.0 \times 10^{-17}$), skin ($P = 7.8 \times 10^{-202}$), brain ($P = 1.4 \times 10^{-54}$) and cortex ($P = 2.7 \times 10^{-115}$; Fig. 7). As PRC2 plays a critical role in development, these results reinforce the epigenetic link between development and aging. This connection is supported by observations that developmentally compromised mice, due to growth hormone receptor (GHRKO) ablation or anterior pituitary gland removal (Snell mice), show reduced rates of epigenetic aging in multiple tissues, as measured by universal epigenetic age clocks (Fig. 3e).

While positively age-related CpG sites (across all tissues) were enriched in 2,961 GO or Molecular Signatures Database terms at a false discovery rate of 0.05 (Supplementary Data 10.1), negatively agerelated CpG sites were enriched in only three. Negatively age-related CpG sites in brain and muscle were enriched in genes associated with circadian rhythm (brain, $P = 3.3 \times 10^{-15}$; cerebral cortex, $P = 4.0 \times 10^{-19}$; muscle, $P = 2.3 \times 10^{-8}$; Fig. 7) and Alzheimer's disease-related gene sets (for example, $P = 1.8 \times 10^{-29}$ in brain and $P = 2.4 \times 10^{-22}$ in the cerebral cortex in Fig. 7). These CpG sites also overlapped with gene sets related to mitochondrial function in brain, cerebral cortex and muscle (for example, $P = 3.6 \times 10^{-7}$; Supplementary Data 10.2).

The GREAT analysis showed enrichment of both positively and negatively age-related CpG sites in mortality or aging gene sets, cancer (Fig. 7) and targets of three Yamanaka factors: SOX2, MYC and OCT4 (Supplementary Data 10.3). Of the 341 genes proximal to positively age-related CpG sites, 162 were implicated in mortality or aging $(P = 6.3 \times 10^{-138}; \text{Fig. 7})$. Similar enrichments were seen in specific tissues: blood ($P = 3.8 \times 10^{-184}$), liver ($P = 2.7 \times 10^{-112}$), muscle ($P = 6.2 \times 10^{-5}$), skin ($P = 9.1 \times 10^{-84}$), combined brain tissues ($P = 1.2 \times 10^{-21}$) and the cerebral cortex ($P = 5.0 \times 10^{-50}$).

As inflammation increases with aging, we assessed the overlap with inflammation-related gene sets (Supplementary Data 10.4). Positively age-related CpG sites are enriched in the gene set associated with inflammation in the murine pancreas (all tissues, $P = 8.4 \times 10^{-21}$ and skin, $P = 9.4 \times 10^{-20}$). Negatively age-related CpG sites are enriched in Toll-like signaling (G0:0034121) genes (muscle, $P = 9.2 \times 10^{-8}$).

Both positively and negatively age-related CpG sites are enriched in immunologic signature gene sets associated with interleukin (IL; for example, IL-6, IL-23) and transforming growth factor (TGF)- β 1 exposure in type 17 helper T cells (Supplementary Data 10.4) for notably brain ($P_{\text{negative}} = 6.1 \times 10^{-11}$) and cerebral cortex ($P_{\text{negative}} = 9.1 \times 10^{-8}$) and, to a lesser extent, skin ($P_{\text{positive}} = 4.0 \times 10^{-4}$) tissues.

Concerns that these highly significant enrichments may be a result of potential biases in the mammalian methylation array platform could be discounted after sensitivity analysis, as reported in Supplementary Note 3.

Fig. 8 | scATAC-seq analysis in human bone marrow and mouse HSCs.

a–**i**, Results using human BMNCs. **j**, Murine HSCs. **a**, scATAC-seq results for 17 of the 35 genes (listed in Supplementary Table 3) that show a called ATAC peak in the region overlapping with our top CpG sites with positive age correlation. The *y* axis lists the gene symbol. The *x* axis reports the Pearson correlation between chronological age and the percentage of cells with an scATAC-seq signal overlapping the respective CpG site (labeled by the adjacent gene). The genes are ordered by correlation estimate (from the most negative). A negative correlation estimate indicates that the accessibility of the CpG site decreases with chronological age. Each dot presents a gene. Seven genes with *P* < 0.05 are marked with a solid shape. **b**, scATAC-seq analysis results for *LHFPL4*. The *y* axis depicts chronological age, and the *x* axis shows the percentage of cells with an scATAC-seq signal. **c**, Percentage of cells identified containing scATAC-seq signal in one of the seven significantly associated genes averaged across all samples. Cells are split into the called identities using the scRNA-seq measurement

TF binding

We used the CellBase⁵² and ENCODE databases⁵³ to annotate CpG sites with binding sites for 68 TFs identified through chromatin immunoprecipitation followed by sequencing (ChIP-seq) in 17 cell types. If a CpG site overlapped with the binding site of a TF (hg19) in at least one cell type, it was assigned to that TF. Analysis of the most significant age-related CpG sites across mammals showed that the REST TF was the most significant TF for the top 1,000 positively age-related CpG sites across all tissues (OR = 8.4, $P = 3.1 \times 10^{-54}$), especially in proliferative tissues such as blood (OR = 5.8, $P = 2.7 \times 10^{-32}$), skin (OR = 8.7, $P = 6.8 \times 10^{-59}$) and liver (OR = 5.4, $P = 1.5 \times 10^{-28}$). REST TF enrichment was less significant in non-proliferative tissues such as muscle (OR = 1.8, $P = 2.2 \times 10^{-3}$), cerebral cortex (OR = 1.6, P = 0.01) and brain (OR = 1.4, P = 0.09; Extended Data Fig. 9 and Supplementary Data 11).

REST TF ChIP-seq analysis was performed on five cell lines, including a human embryonic stem cell line (Supplementary Data 11.1). REST is known for repressing neuronal genes in non-neuronal tissues, which could explain the weak enrichments in brain regions. Notably, CpG cg12841266 near *LHFPL4* is within the REST-binding region.

Substantial binding enrichments were observed for transcription factor 12 (TCF12) and histone deacetylase 2 (HDAC2). TCF12 is part of the basic helix–loop–helix (bHLH) E-protein family, associated with neuronal differentiation, and top positively age-related CpG sites are proximal to another bHLH gene, *NEUROD1* (Supplementary Table 3 and Supplementary Data 11). Lower enrichments were noted for CCCTCbinding factor (CTCF) and Nanog homeobox (NANOG). For the top 1,000 negatively age-related CpG sites, fewer significant TF binding enrichments emerged, withJUN (c-Jun) in blood (OR = $2.8, P = 2.6 \times 10^{-9}$) and brain (OR = 1.5, P = 0.024; Extended Data Fig. 9) being exceptions.

Age-related CpG sites and age-related transcriptomic changes We studied whether the top 1,000 positively and negatively agerelated CpG sites neighbor genes with age-correlated mRNA levels. Using GenAge⁵⁴ and Enrichr^{55,56} databases, we scrutinized age-specific transcriptome-wide association studies (TWAS) in four mammalian species. The EWAS–TWAS overlap analysis (Fig. 7, Extended Data Fig. 10a and Supplementary Data 12) indicates significant overlaps between age-related CpG sites and transcriptomic age changes in several species, including Genotype–Tissue Expression (GTEx) human tibial nerve samples, normal monkey hippocampal samples ($P = 9 \times 10^{-15}$) and various rat and mouse tissues. However, the age-related EWAS and TWAS overlap is generally weak and tissue specific.

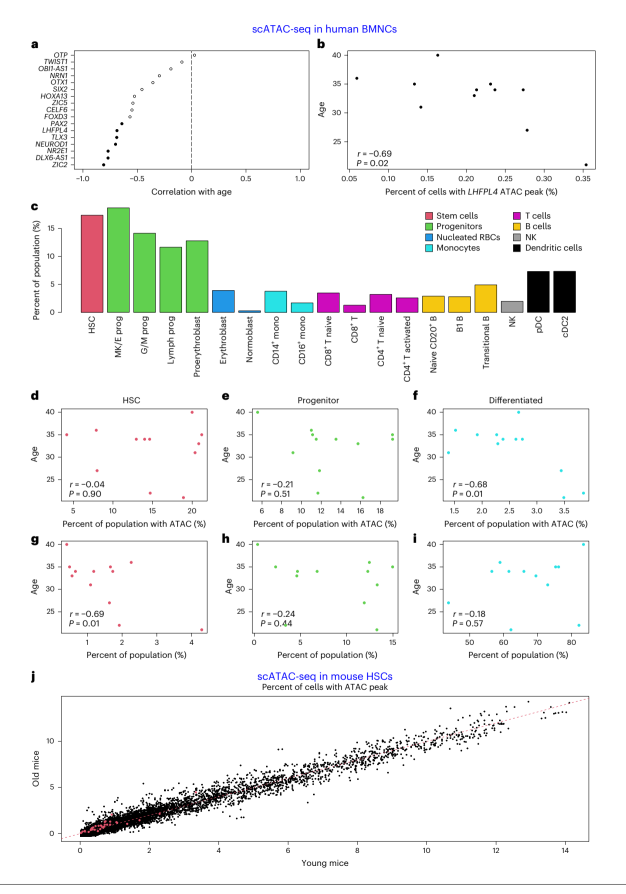
$\label{eq:second} \mbox{Age-related CpG} sites and genome-wide association studies of human traits$

We compared proximal genes of the top 1,000 positively and negatively age-related CpG sites with the top 2.5% of genes implicated in various human genome-wide association studies (GWAS). Notable enrichments were seen in genes associated with waist-to-hip ratio for positively age-related CpG sites in livers ($P_{\text{positive}} = 1.0 \times 10^{-16}$), and with human

including HSCs, the various progenitors and differentiated cells. DC, dendritic cell; mono, monocyte; MK/E prog, megakaryocyte-erythroid progenitor; G/M prog, granulocyte-monocyte progenitor; NK, natural killer; prog, progenitor; RBC, red blood cell. **d**–**f**, The percentage of these three cell populations (HSC (**d**), progenitor (**e**) and differentiated cell type (**f**)) that contain at least one ATAC-seq signal in any of the seven significant genes, plotted against the age of each individual (*y* axis). **g**–**i**, The percentage of these three cell populations per individual (HSC (**g**), progenitor (**h**) and differentiated cell type (**i**)), plotted against the age of each individual. **j**, The percentage of cells with called ATAC peaks overlapping with our mammalian CpG sites in young mouse (10-week) versus old mouse (20-month) HSCs. The red dots denote 33 of the top 35 positively age-related CpG sites (listed in Supplementary Table 3) that map to the mouse genome. The red dashed line corresponds to the diagonal line (*y*=*x*). All *P* values reported are unadjusted and two sided.

length at birth for positively age-related CpG sites in the cortex ($P_{\text{positive}} = 1.0 \times 10^{-12}$) and liver ($P_{\text{positive}} = 2.0 \times 10^{-10}$; Fig. 7). Significant enrichments (defined here as nominal $P < 5.0 \times 10^{-4}$) were also seen with genes linked to mother's longevity (mother attained age;

 $P_{\text{positive}} = 2.0 \times 10^{-4}$; Fig. 7, Extended Data Fig. 10b and Supplementary Data 13.1–13.7), human longevity for negatively age-related CpG sites in muscle ($P_{\text{negative}} = 8.0 \times 10^{-6}$), epigenetic age acceleration on the mortality clock (GrimAge $P_{\text{positive}} = 7.0 \times 10^{-7}$ in muscle), age-related



macular degeneration ($P_{\text{positive}} = 2.0 \times 10^{-8}$ in all tissues), Alzheimer's disease ($P_{\text{negative}} = 1.0 \times 10^{-4}$ in brain), leukocyte telomere length ($P_{\text{negative}} = 3.0 \times 10^{-13}$ in muscle and $P_{\text{negative}} = 2 \times 10^{-11}$ in brain) and age at menarche ($P_{\text{positive}} = 4.0 \times 10^{-5}$ in all tissues). Overall, our GWAS overlap analysis indicates that pan-mammalian age-related CpG sites are proximal to genes influencing human development (birth length, menarche), obesity and longevity.

Single-cell ATAC-seq analysis in human bone marrow

Low-methylated regions distant from TSS correlate with open chromatin, TF binding and enhancers⁵⁷. Hence, our top positively age-related pan-mammalian CpG sites (initially low in methylation, gaining methylation with age) could imply a gradual loss of these open chromatin regions. To validate this, we examined the association between the top 35 positively age-related CpG sites (Supplementary Table 3) and chromatin accessibility in single cells from human bone marrow mononuclear cells (BMNCs). Single-cell assay for transposase-accessible chromatin with sequencing (scATAC-seq) data from a recent study⁵⁸ employed 10x Multiome technology to profile both ATAC and gene expression within the same cell across ten donors of varying age. Overlaying the genomic regions of the top 35 CpG sites (Supplementary Table 3) with the called ATAC peaks within the BMNC dataset identified 17 genes, including *LHFPL4* (Supplementary Data 14.1 and Fig. 8a).

We calculated the percentage of cells per individual with the respective peak. A strong, statistically significant negative correlation (Fig. 8b) was found between age and the number of cells with the ATAC peak overlapping cg12841266 in *LHFPL4*. This shows that, with age (as methylation increases), open chromatin cell number decreases. Of 17 gene regions, 16 correlated negatively with age, with seven being statistically significant (P < 0.05; Fig. 8a). The hypermethylated sites were highly enriched for this age-associated accessibility loss (P < 0.001; Fig. 8b). The significant genes (*LHFPL4*, *TLX3*, *ZIC2*, *PAX2*, *NR2E1*, *NEUROD1*, *DLX6-AS1*) are related to developmental processes (Supplementary Table 3). *ZIC5*, another Zic family gene, also showed a nearly significant negative age correlation (r = -0.54, P = 0.07; Supplementary Data 14.1). No scATAC-seq signal was detected in the cg09710440 region of *LHFPL3*, possibly due to proximity to a bivalent gene's TSS (232 bp).

We examined whether the seven significant ATAC peaks identified a particular cell type subset. Due to the sparsity of scATAC-seq data, we determined the fraction of each cell group containing at least one of these regions. We found that stem cell–progenitor populations had a higher proportion of open chromatin at these sites than differentiated cells (mean of 14.9% versus mean of 2.9%; Fig. 8c). This suggests that the observed age-related reduction of open chromatin states could be due to the loss (for example, death or differentiation) of progenitor cells in the tissue.

We studied three cell groups: hematopoietic stem cells (HSCs), progenitor cells and differentiated cells. Age showed a negative correlation with the percentage of HSCs (r = -0.69, P = 0.01) but no significant correlation with progenitor or differentiated cells (Fig. 8g–i). Next, we analyzed the correlation between age and the proportion of cells containing an ATAC peak in at least one of the seven significant CpG regions (Fig. 8d–f). Differentiated cells demonstrated a significant loss of ATAC signal in these regions with age (r = -0.68, P = 0.01; Fig. 8f), whereas no change was seen in HSCs or progenitor cells (Fig. 8d,e). This suggests that these regions, gaining methylation and losing accessibility with age, belong to a differentiated cell population. Lastly, analyzing increasing lists of positively age-related CpG sites, we noted that the percentage of cells with an ATAC peak at these locations decreasing with age in human BMNCs (median correlation < -0.2 across the top 500 or 1,000 positively age-related CpG sites).

scATAC-seq analysis in murine HSCs

We tested whether our human HSC findings extended to murine HSCs by analyzing another public scATAC-seq dataset from murine HSCs with

four replicates each in young (10-week) and old (20-month) mice⁵⁹. This dataset provided access to our age-related CpG sites in 4,492 young and 3,300 old HSCs. Of the top 35 positively age-related CpG sites, 33 overlapped with ATAC peaks (Supplementary Data 14.2). We then calculated the proportion of HSCs in each age group with the respective peak. The proportion of old HSCs with a peak near *Lhfpl4* was not significantly different from that of young HSCs (OR = 0.94, P = 0.7), implying no observable age-related chromatin compactification in murine HSCs. This was also true for the other 32 CpG sites and their associated peaks. Contrarily, the proportion of old HSCs with an ATAC peak was significantly higher than that of young HSCs for five CpG sites (near *Bdnf, Isl1, Twist1, Nr2e1, Sall1*; Fisher exact *P* value < 0.05; Supplementary Data 14.2), indicating age-related chromatin opening (Fig. 8j), aligning with Itokawa et al.'s report⁵⁹.

Discussion

The consistent age-related alterations in DNA methylation profiles across mammalian species challenges the view that aging is simply due to the random accumulation of cellular damage. Our Mammalian Methylation Consortium investigated this question with an extensive set of DNA methylation profiles from 348 species⁹, using 174 eutherian, nine marsupial and two monotreme species in this study.

We found a set of CpG sites in DNA sequences conserved across mammals consistently changing with age, predominantly gaining methylation. These CpG sites are often in PRC2-binding sites and the bivalent chromatin states BivProm1 and BivProm2, regulating the expression of genes involved in the process of development^{47,60,61}, which is one of the most conserved biological processes that threads through all mammalian species. Examples of age-related CpG sites include those near *LHFPL4* and *LHFPL3*. The known function of *LHFPL4* in synaptic clustering of γ -aminobutyric acid (GABA) receptors does not provide a clear connection to aging across tissues. Nevertheless, the specificity of their methylation change with age is clear, considering their distinct chromosomal locations, as observed with gene pairs such as *LHFPL3*– *LHFPL4*, *ZIC2–ZIC5*, *PAX2–PAX5* and *CELF4–CELF6*.

The scATAC-seq analysis of BMNCs revealed that age-correlated CpG sites are located in regions that lose chromatin accessibility with age in differentiated cells but not in progenitor cells. This suggests that methylation likely instigates such chromatin compaction⁶², hindering PRC2 access to its target sites. We observed this phenomenon in human bone marrow, where (1) top age-related PRC2 targets are open in substantially more progenitor cells than differentiated cells and (2) the percentage of progenitor cells with open age-related PRC2 targets did not diminish with age. Similarly, the percentage of murine HSCs with open age-related PRC2 targets did not diminish with age. By contrast, the percentage of differentiated human bone marrow cells with open PRC2 targets diminished with age, underscoring the need for further research into other differentiated cell types.

When it comes to age-related gain of methylation, it is important to distinguish proliferative tissues from non-proliferative tissues such as the brain and muscle. The overlap between PRC2-binding sites and positively age-related changes is far more pronounced in proliferative tissues than in non-proliferative tissues (Fig. 4h). The dichotomy between proliferative and non-proliferative tissues is even more pronounced when it comes to characterizing age-related loss of methylation.

In proliferative tissues, negatively age-related CpG sites are often located in quiescent chromatin states, heterochromatin and PMDs. Interestingly, PMDs are in late DNA-replication regions. As methylation of replicated DNA is slow and only completed very late in S and G2 phases, late-replicated regions of the DNA are particularly disadvantaged in this regard. Indeed, progressive methylation loss in PMDs is exploited as a mitotic clock, which also correlates very well with chronological age⁵⁰. As such, their identification as pan-mammalian negatively age-related CpG sites is entirely consistent with studies observed in human cells. Interestingly, this late-replication effect on DNA methylation can be prevented by the binding of histone 3 lysine 36 trimethylation (H3K36me3) to these regions⁵⁰. This appears to be mediated by H3K36me3 recruitment of DNA methyltransferase 3 (DNMT3) to unmethylated and newly replicated DNA. Conversely, functional loss of NSD1, the enzyme that generates H3K36me3, leads to hypomethylation of DNA and accelerated epigenetic aging^{63,64}. Age-related loss of methylation in non-proliferative tissues (brain and muscle), on the other hand, is observed at CpG sites located in an exon-associated transcription state (TxEx4), which is the most highly enriched state for transcription termination sites and is associated with the highest gene expression levels across many cell and tissue types⁴⁶.

Unlike CpG sites that gain methylation with age, CpG sites that lose methylation are typically not related to developmental genes. Instead, they are related to genes of circadian rhythm and mitochondria, the functions of which are progressively eroded with age. Finally, the *LARP1* gene, which is proximal to the highest-ranked hypomethylated cytosine in the liver and second across all tissues, encodes an RNA-binding protein that is involved in several processes, including post-transcriptional regulation of gene expression and translation of downstream targets of mammalian target of rapamycin (mTOR)⁶⁵. mTOR has very well-documented links with aging and longevity⁶⁶ and is also linked to epigenetic aging^{67,68}. Overall, we provide collective evidence that the methylated mammalian age-related CpG sites that we identified are not merely stochastic marks accrued with age. They are instead methylation changes that capture multiple facets of mammalian aging.

The deterministic features of these age-related changes on the mammalian epigenome make a compelling case that aging is not solely a consequence of random cellular damage accrued in time. It is instead a pseudo-programmed process that is also intimately associated with mammalian development that begins to unfold from conception. This is supported by and is consistent with the finding that genes proximal to age-related CpG sites were also identified by GWAS of human development features such as length at birth and age at menarche. A large body of literature including those by Williams in 1957 (refs. 69,70) has suggested a connection between growth and development and aging. More recently, several authors have suggested epigenetics to be the link between these two processes^{69,71-80}. This notion is further supported by the recent demonstration of age reversal through the expression of Yamanaka factors^{45,81-84}, which can also be observed for our universal pan-mammalian clocks (Fig. 3c,d).

According to the pseudo-programmatic theory of aging, the process of aging is very much a consequence of the process of development, and the ticking of the epigenetic clock reflects the continuation of developmental processes^{69,80}. As predicted by the epigenetic clock theory of aging, universal epigenetic clocks provide a continuous readout of age from early development to old age in all mammals, as this feature underlies the continuous and largely deterministic process of aging from conception to tissue homeostasis⁷⁴. Consistent with this theory, pan-mammalian methylation clocks are slowed by conditions that delay growth and/or development including Snell mice and full-body GHRKO mice. The successful construction of universal clocks is a compelling mathematical demonstration of the deterministic element in the process of aging that transcends species barriers within the mammalian class. Indeed, the centrality of PRC2, which is also present in non-mammalian classes, implies that the process of aging that is uncovered here is likely to be shared by vertebrates in general. Our human epidemiological studies and mouse interventional studies show that pan-mammalian clocks relate to human and mouse mortality risk, respectively. Cross-sectional epidemiological studies in humans reveal that the pan-mammalian clocks correlate with markers of inflammation (C-reactive protein) and dyslipidemia (triglyceride levels).

Our study has certain limitations. The study primarily focuses on highly conserved DNA sequences, thus limiting our examination to approximately 36,000 CpG sites of the tens of millions that exist in most mammalian genomes. Additionally, our array platform exhibits a slight bias, featuring more probes that align with eutherian genomes than with marsupial genomes⁸.

Overall, our results demonstrate that select epigenetic aging effects are universal across all mammalian species and capture multiple processes and manifestations of age that have thus far been thought to be unrelated to each other. We expect that the availability of panmammalian epigenetic clocks will open the path to uncovering interventions that modulate conserved aging processes in mammals.

Methods

Ethics

All local ethical guidelines were followed, and necessary approvals from respective human ethical review boards and animal ethical committees were duly obtained. Details can be found in Supplementary Notes 1, 2 and 4.

Statistics and reproducibility

Data collection and analysis were not performed blind to the conditions of the experiments. In the ensuing sections, we delineate the quality-control measures for our samples and the statistical methods employed in each analysis, with additional details provided in Supplementary Notes 1 and 5.

Tissue samples

We used a subset of the data from the Mammalian Methylation Consortium for which age information was available⁹. The tissue samples are described in Supplementary Data 1.1–1.4, and related citations are listed in Supplementary Notes 1 and 2. We used the SeSAMe normalization method⁸⁵.

Quality controls for establishing universal clocks

Our epigenetic clocks were trained and evaluated on samples with highly confident age assessments (less than 10% error). We focused on typical aging patterns, hence excluding tissues from preclinical anti-aging or pro-aging intervention studies.

Species characteristics

Species characteristics such as maximum lifespan (maximum observed age) and ASM were obtained from an updated version of AnAge⁸⁶ (https://genomics.senescence.info/species/index.html). To facilitate reproducibility, we have posted this modified and updated version of AnAge in Supplementary Data 1.13.

Three universal pan-mammalian clocks

We applied elastic net regression models to establish three universal mammalian clocks for estimating chronological age across all tissues (n = 11,754 from 185 species) in eutherians (n = 11,439 from 174 species), marsupials (n = 210 from nine species) and monotremes (n = 15 from two species). The three elastic net regression models, implemented using the glmnet 4.1-7 package in R, corresponded to different outcome measures described in the following:

- 1. log-transformed chronological age: log(Age + 2), where an offset of 2 years was added to avoid negative numbers in case of prenatal samples,
- 2. -log(-log(RelativeAge)) and
- 3. log-linear transformed age.

DNAmAge estimates of each clock were computed via the respective inverse transformation. Age transformations used for building universal clocks 2 and 3 incorporated a selection of three species characteristics: gestational time (GestationT), age at sexual maturity (ASM) and maximum lifespan (MaxLifespan). All of these species variables surrounding time are measured in units of years. **loglog transformation of relative age for clock 2.** Our measure of relative age leverages gestation time and maximum lifespan. We define relative age (RelativeAge) and apply the double logarithmic loglog transformation:

$$RelativeAge = \frac{Age + GestationT}{MaxLifespan + GestationT}$$
(1)

$$loglogAge = -log(-log(RelativeAge)).$$
(2)

By definition, RelativeAgeis between 0 and 1, and loglogAge is positively correlated with age. The incorporation of gestation time is not essential. We simply include it to ensure that RelativeAge takes on positive values. We used the double logarithmic transformation to link relative age to the covariates (cytosines) for the following reasons. First, the transformation maps the unit interval to the real line. Second, this transformation ascribes more influence on exceptionally high and low age values (Extended Data Fig. 1a–c). Third, this transformation is widely used in the context of survival analysis. Fourth, this non-linear transformation worked better than the identity transformation in terms of age correlation and calibration.

The regression model underlying universal clock 2 predicts loglogAge. To arrive at the DNAmAge, one needs to apply the inverse transformation to loglogAge based on the double exponential transformation:

$$DNAmAge = exp(-exp(-loglogAge))$$

$$\times (MaxLifespan + GestationT) - GestationT.$$
(3)

All species characteristics (for example, maximum lifespan, gestational time) come from our updated version of AnAge. We were concerned that the uneven evidence surrounding the maximum age of different species could bias our analysis. While billions of people and many mice have been evaluated for estimating the maximum age of humans (122.5 years) or mice (4 years), the same cannot be said for any other species. To address this concern, we made the following assumption: the true maximum age is 30% higher than that reported in AnAge for all species except for humans and mice. Therefore, we multiplied the reported maximum lifespan of non-human or non-mouse species by 1.3. Our predictive models turn out to be highly robust with respect to this assumption.

Transformation based on log-linear age for clock 3. Our measure of log-linear age leverages ASM. The transformation has the following properties: it takes the logarithmic form when the chronological age is young, and it takes the linear form otherwise. It is continuously differentiable at the change.

First, we define a ratio of the age relative to ASM, termed RelativeAdultAge, as the following:

$$RelativeAdultAge = \frac{Age + GestationT}{ASM + GestationT},$$
 (4)

where the addition of GestationT ensures that the RelativeAdultAge is always positive. To model a faster rate of change during development, we used a log–linear transformation on RelativeAdultAge based on a function that generalizes the original transformation proposed for the human pan-tissue clock⁴:

$$y = f(x;m) = \begin{cases} \frac{x}{m} - 1, \ \frac{x}{m} \ge 1\\ \log \frac{x}{m}, \ \frac{x}{m} < 1 \end{cases}$$
(5)

$$f^{-1}(y;m) = \begin{cases} m(y+1), \ y \ge 0\\ me^y, \ y < 0 \end{cases}$$
(6)

In the function f(x;m), x denotes RelativeAdultAge, m represents a parameter and f represents the log-linear transformation. The output, y, is the results of applying the function f to x and m. This transformation is designed to reflect a higher rate of change for younger RelativeAdultAges when $x \le m$. This transformation ensures continuity and smoothness at the change point at x = m.

In the following, we describe the estimation of the parameter *m*. To ensure that the maximum value of *y* is the same across all species, the parameter *m* should be proportional to the maximum of *x* for each species, that is, the best value for *m* would be the oracle value

$$m^* = c_1 \left(\frac{\text{MaxLifespan+GestationT}}{\text{ASM+GestationT}}\right)$$
 (Extended Data Fig. 1d).

The proportionality constant c_1 controls the distribution of y. We chose the value of c_1 so that y follows approximately a normal distribution with mean zero. Because we wanted to define clock 3 without using MaxLifespan, we opted to use the ratio $\frac{\text{GestationT}}{\text{ASM}}$ as a surrogate for the oracle value m^* . We achieved this approximation by fitting the following regression model with all mammalian species available in our AnAge database,

$$\log \frac{\text{MaxLifespan} + \text{GestationT}}{\text{ASM} + \text{GestationT}} \approx 2.92 + 0.38 \times \log \frac{\text{GestationT}}{\text{ASM}}.$$
 (7)

The two log variables in equation (7) have moderate correlation (r = 0.5). Subsequently, we defined \hat{m} as follows:

$$\hat{m} = c_2 \left(\frac{\text{GestationT}}{\text{ASM}}\right)^{0.38},\tag{8}$$

where $c_2 = c_1 e^{2.92}$. We chose $c_2 = 5.0$ so that log-linear age termed y in equation (5) follows approximately a normal distribution with mean zero (median = 9.0×10^{-4} , skewness = -0.02; Extended Data Fig. 1f). Setting x=RelativeAdultAge in equation (5) results in

$$f(\text{RelativeAdultAge}; \hat{m}) = \begin{cases} \frac{\text{RelativeAdultAge}}{\hat{m}} - 1, \text{ RelativeAdultAge} \ge \hat{m} \\ \log \frac{\text{RelativeAdultAge}}{\hat{m}}, \text{ RelativeAdultAge} < \hat{m} \end{cases}$$
(9)

Universal clock 3 predicts loglinearAge (denoted as y). To arrive at an age estimate, we employ both equations (4) and (6):

$$DNAmAge = \begin{cases} \hat{m} \times (ASM + GestationT) \times (y+1) - GestationT, \ y \ge 0\\ \hat{m} \times (ASM + GestationT) \times e^{y} - GestationT, \ y < 0 \end{cases}$$
(10)

Statistics for performance of model prediction. To validate our model, we used DNAmAge estimates from LOFO and LOSO analyses, respectively. At each type of estimate, we computed Pearson correlation coefficients and MAE between DNAm-based and observed variables across all samples. Correlation and MAE were also computed at the species level, limited to the subgroup with $n \ge 15$ samples (within a species). We reported the medians for the correlation estimates (median correlation) and the medians for the MAE estimates (med.MAE) across species. Analogously, we repeated the same analysis at the species–tissue level, limited to the subgroup with at least 15 samples (within a species–tissue category).

For Extended Data Fig. 2, we evaluated the difference Delta.Age (Δ Age) between the LOSO estimate of DNAmAge and chronological age at half the maximum lifespan (0.5 × MaxLifespan). As expected, Δ Age = LOSO DNAmAge – (0.5 × MaxLifespan) is negatively correlated with species maximum lifespan.

Epigenetic age acceleration

To adjust for age, we defined epigenetic age acceleration (AgeAccel) as the raw residual resulting from regressing DNAmAge (from universal clocks 2 and 3) on chronological age. By definition, the resulting AgeAccel measure is not correlated with chronological age.

Human epidemiological cohort studies

We applied our universal clocks 2 and 3 to 4,651 individuals from (1) the FHS Offspring cohort (n = 2,544 Caucasians, 54% women)⁸⁷ and (2) the WHI cohort^{88,89} (n = 2,107,100% women; Supplementary Note 4). Methylation levels were profiled in blood samples using the Illumina 450k arrays. The FHS cohort had a mean (s.d.) age of 66.3 (8.9) years at blood draw, with 330 deaths during an average follow-up of 7.8 years. The WHI cohort, which enrolled postmenopausal women 50–79 years in age, consisted of three ethnic groups: 47% of European ancestry, 32% African Americans and 20% of Hispanic ancestry. These groups exhibited similar age distributions, with a mean (s.d.) age of 65.4 (7.1) years, and a mean (s.d.) follow-up time of 16.9 (4.6) years. During the follow-up, 765 women died.

Mortality analysis for time to death. Our mortality analysis was performed as follows. First, we applied our Array Converter algorithm (Supplementary Note 5) to yield the imputed mammalian arrays and to estimate DNAmAge values based on our universal clocks. Second, we computed AgeAccel for each cohort. Third, we applied Cox regression analysis for time to death (as a dependent variable) to assess the predictive ability of our universal clocks for all-cause mortality. The analysis was adjusted for age at blood draw and for sex in the FHS. We stratified the WHI cohort by ethnic or racial groups and combined a total of four results across FHS and WHI cohorts by fixed-effect models weighted by inverse variance. The meta-analysis was performed with the R 'metafor' function.

Human epidemiological cohort studies for lifestyle factors. We performed a robust correlation analysis (bicor²⁹) between (1) our AgeAccel measures from clocks 2 and 3 and (2) 59 variables spanning diet, clinically relevant measurements and lifestyle factors. Comprehensive details of these variables and our analytical approaches, inclusive of meta-analysis, are elucidated in Supplementary Note 5.

Polygenic models for heritability analysis. We calculated the narrowsense heritability of our clocks by employing polygenic models as defined in SOLAR⁹⁰ and its R interface solarius⁹¹ as detailed in Supplementary Note 5.

OSKM reprogramming cells in human dermal fibroblasts. We applied our universal clock 2 and clock 3 to a previously published dataset (GSE54848)³¹ in which the authors had transfected human dermal fibroblasts with the Yamanaka factors (OSKM) over a 49-d period. The successfully transformed cells were collected and profiled on the human Illumina 450k arrays. Similar to the applications for the FHS and WHI cohorts, we applied our Array Converter algorithm (Supplementary Note 5) to yield the imputed mammalian arrays and to estimate DNAmAge based on our universal clocks. The clocks were applied to a total of n = 27 samples across experiment days 0, 3, 7, 11, 15, 20, 28, 35, 42 and 49, respectively.

Murine anti-aging studies

None of the samples from the murine anti-aging studies were used in the training set of the universal clocks, that is, these are truly independent test data. Clocks 2 and 3 were evaluated in five mouse experiments (independent test data): (1) Snell dwarf mice (n = 95), (2) GHRKO experiment 1(GHRKO, n = 71 samples), (3) GHRKO experiment 2 (n = 96samples), (4) three Tet experiments: *Tet1* KO (n = 64), *Tet2* KO (n = 65)and *Tet3* KO (n = 63) and (5) CR (n = 95). Details can be found in Supplementary Note 6.

Meta-analysis for EWAS of age

In our primary EWAS of age, we focused on samples from eutherians (n = 65 species) for which each species has at least 15 samples from the

Our meta-analysis for EWAS of age in eutherian species combined Pearson correlation test statistics across species-tissue strata that contained at least 15 samples each. The minimum sample size requirement resulted in 143 species-tissue strata from 65 eutherian species (Supplementary Data 1.5). To counter the dependency patterns resulting from multiple tissues from the same species, the meta-analysis was carried out in two steps. First, we meta-analyzed the EWAS of different tissues for each species separately. These tissue-specific summary statistics were combined within the same species to represent the EWAS results at species level. Second, we meta-analyzed the resulting 65-species EWAS results across species to arrive at the final meta-EWAS of age. In each meta-analysis step, we used the unweighted Stouffer's method as implemented in R. In more detail, we gathered 68 blood samples from 27 distinct lemur species and 23 skin samples from 23 distinct lemur species, each species-tissue stratum with at most three samples. We therefore combined those 68 blood samples to perform blood EWAS in lemurs. Similarly, we combined the 23 skin samples for skin EWAS in lemurs. As listed in Supplementary Data 1.5, the combined species in lemurs was denoted by Strepsirrhine in the column 'Species Latin Name'.

EWAS of age in marsupials was based on a two-step meta-analysis in which we relaxed the threshold of sample size in the species–tissue category to $n \ge 10$ (Supplementary Data 1.12). Due to a small sample size in monotremes (n = 15), we combined all monotreme samples into a single dataset.

Brain EWAS. We applied the two-step meta-analysis approach to the brain EWAS results based on more than 900 brain tissues (cerebellum, cortex, hippocampus, hypothalamus, striatum, subventricular zone and whole brain) from eight species including human, vervet monkey, mice, olive baboon, brown rat and pig species (Supplementary Data 1.6).

EWAS of a single tissue. For the cerebral cortex brain region, we simply combined tissue-specific EWAS results across different species using the unweighted Stouffer's method (Supplementary Data 1.7). Similarly, we carried out the one-step meta-analysis EWAS of blood, liver, muscle and skin (Supplementary Data 1.8–1.11). Details can be found in Supplementary Note 5.

All the Manhattan plots were generated based on a modified version of the gmirror function in R.

Stratification by age groups. To assess whether the age-related CpG sites in young animals relate to those in old animals, we split the data into three age groups: young-age (age < 1.5ASM), middle-age (age between 1.5ASM and 3.5ASM) and old-age (age \geq 3.5ASM) groups. The threshold of sample size in species–tissue was relaxed to $n \geq$ 10. The age correlations in each age group were meta-analyzed using the above-mentioned two-step meta-analysis approach.

Polycomb repressive complex

Polycomb repressive complex annotations were defined based on the binding of at least two transcriptional factor members of polycomb repressor complex 1 (PRC1 with subgroups RING1, RNF2, BMI1) or PRC2 (with subgroups EED, SUZ12 and EZH2) in 49 available ChIP-seq datasets from ENCODE⁵³.

We identified 640 and 5,287 CpG sites in the array that were located in regions bound by PRC1 and PRC2, respectively. We performed a onesided hypergeometric analysis to study both enrichment (OR > 1) and depletion (OR < 1) patterns for our age-related markers based on the top 1,000 CpG sites increased with age and the top 1,000 CpG sites decreased with age from the EWAS of age.

Universal chromatin state analysis

To annotate our age-related CpG sites based on chromatin states, we assigned a state for all our mammalian CpG sites based on a recently published universal ChromHMM chromatin state annotation of the human genome⁴⁶. The underlying hidden Markov model was trained with over 1,000 datasets of 32 chromatin marks in more than 100 cell and tissue types. This model then produced a single chromatin state annotation per genomic position that is applicable across cell and tissue types, as opposed to producing an annotation that is specific to one cell or tissue type. A total of 100 distinct states were generated and categorized into 16 major groups according to the parameters of the model and external genome annotations⁴⁶ (described in Supplementary Data 8.2).

We performed a one-sided hypergeometric analysis to study both enrichment (OR > 1) and depletion (OR < 1) patterns for our age-related markers based on the top 1,000 CpG sites with a positive correlation with age and the top 1,000 CpG sites with a negative correlation with age across different eutherian species.

Analysis of late-replicating domains

The annotation of late-replicating domains (hg19 and mm10) was obtained from Zhou et al. 50 , as described in Supplementary Note 5.

GREAT enrichment analysis

We applied the GREAT analysis software tool⁵¹ to the top 1,000 positively age-related and the top 1,000 negatively age-related CpG sites from the EWAS of age. GREAT implemented foreground–background hypergeometric tests over genomic regions where we input all CpG sites of the mammalian array as background and the genomic regions of the 1,000 CpG sites as foreground. This approach yielded hypergeometric *P* values that were not confounded by the number of CpG sites within a gene (Supplementary Note 5).

EWAS-TWAS overlap analysis

Our EWAS–TWAS-based overlap analysis related the gene sets found by our EWAS of age with the gene sets from our in-house TWAS database. The TWAS database, along with our analytical approaches, is described in Supplementary Note 5.

EWAS-GWAS overlap analysis

Our EWAS–GWAS overlap analysis linked the gene sets discovered in our EWAS of age with those identified in published large-scale GWAS studies of various phenotypes (Supplementary Note 5).

Transcription factor binding analysis

We used the CellBase database⁵², incorporating ENCODE⁵³ TF binding sites for our analysis (Supplementary Note 5).

Single-cell ATAC-seq of human bone marrow

Recent advances have enabled the sequencing of ATAC profiles within single cells, enabling assessment of the proportion of cells containing an open chromatin region⁵⁸. We cross-referenced the top 35 CpG sites with positive age correlation across mammalian tissues with publicly available scATAC-seq data (Supplementary Table 3). We downloaded 10x Multiome count data in AnnData format as H5AD from the Gene Expression Omnibus (accession number GSE194122). The ATAC array data were managed using the Python package anndata⁹². hg38 ATAC peak locations were extracted from the metadata '.var' section using anndata. Peak locations were overlapped with probe locations using GenomicRanges⁹³ for the top 35 CpG sites. The overlapping peaks were then used to extract the processed counts for each cell. The proportion of cells containing an ATAC peak for each individual sample was calculated. A correlation was calculated by comparing this value against the age of each individual sample. The cell type for each barcode was extracted from the observable object. We subsequently computed the proportion of each cell type containing an ATAC peak in one of the seven significantly correlated regions (*LHFPL4*, *TLX3*, *ZIC2*, *PAX2*, *NR2E1*, *NEUROD1* and *DLX6-AS1*). Progenitor cells were grouped as MK/E progenitors, G/M progenitors, lymph progenitors and proerythroblasts, and differentiated cells were grouped as CD14⁺ monocytes, CD16⁺ monocytes, CD8⁺T naive, CD8⁺T, CD4⁺T naive, CD4⁺T activated, naive CD20⁺ B, B1 B, transitional B and NK. The percentage of each of the three populations (HSC, progenitor and differentiated cells) was calculated, and the proportion of cells containing an ATAC peak in one of the seven significantly correlated regions was calculated. To confirm enrichment for the hypermethylated sites showing decrease in chromatin accessibility with age, we randomly selected 1,000 sets of 17 ATAC peaks and compared the mean correlation with age of the selected regions to the 1,000 sampled sets of regions.

Mouse single-cell ATAC-seq in hematopoietic stem cells. We downloaded the publicly available data (H5, meta and fragment files of Illumina HiSeq 1500 array data) from Itokawa et al.⁵⁹ (GSE162662).

scATAC-seq data were profiled in four biological replicates in young (10-week) and old (20-month) mice. The ATAC-seq data were managed and analyzed with R Signac⁹⁴. We applied Fisher's exact test to ascertain whether locations with differential accessibility between young and old animals were enriched with the 33 top positively agerelated CpG sites (OR > 1 indicates a higher proportion in the old group). Further analytical details, including ATAC-seq data quality controls, are presented in Supplementary Note 5.

URLs

The following URLs are available: AnAge (https://genomics.senescence. info/species/index.html), GREAT (http://great.stanford.edu/public/ html/), late-replicating domains (https://zwdzwd.github.io/pmd), UCSC Genome Browser (http://genome.ucsc.edu/index.html).

Reporting summary

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

Data availability

The individual-level data from the Mammalian Methylation Consortium can be accessed from several online locations. All data from the Mammalian Methylation Consortium are posted on Gene Expression Omnibus (complete dataset, GSE223748), Subsets of the datasets can also be downloaded from accession numbers GSE174758, GSE184211, GSE184213, GSE184215, GSE184216, GSE184218, GSE184220, GSE184221, GSE184224, GSE190660, GSE190661, GSE190662, GSE190663, GSE190664, GSE174544, GSE190665, GSE174767, GSE184222, GSE184223, GSE174777, GSE174778, GSE173330, GSE164127, GSE147002, GSE147003, GSE147004, GSE223943 and GSE223944. Additional details can be found in Supplementary Note 2. The mammalian data can also be downloaded from the Clock Foundation webpage: https://clockfoundation.org/MammalianMethylationConsortium. The mammalian methylation array is available through the non-profit Epigenetic Clock Development Foundation (https://clockfoundation.org/). The manifest file of the mammalian array and genome annotations of CpG sites can be found on Zenodo (https://doi.org/10.5281/zenodo.7574747). All other data supporting the findings of this study are available from the corresponding author upon reasonable request.

Code availability

The chip manifest files, genome annotations of CpG sites and the software code for universal pan-mammalian clocks can be found on GitHub⁹⁵ at https://github.com/shorvath/MammalianMethylation-Consortium/tree/v2.0.0. The individual R code for the universal pan-mammalian clocks, EWAS analysis and functional enrichment studies can be also found in the Supplementary Code.

References

- 1. Ferrucci, L. et al. Measuring biological aging in humans: a quest. Aging Cell **19**, e13080 (2020).
- 2. Bell, C. G. et al. DNA methylation aging clocks: challenges and recommendations. *Genome Biol.* **20**, 249 (2019).
- Field, A. E. et al. DNA methylation clocks in aging: categories, causes, and consequences. *Mol. Cell* 71, 882–895 (2018).
- 4. Horvath, S. DNA methylation age of human tissues and cell types. *Genome Biol.* **14**, R115 (2013).
- Petkovich, D. A. et al. Using DNA methylation profiling to evaluate biological age and longevity interventions. *Cell Metab.* 25, 954–960 (2017).
- 6. Stubbs, T. M. et al. Multi-tissue DNA methylation age predictor in mouse. *Genome Biol.* **18**, 68 (2017).
- 7. Cole, J. J. et al. Diverse interventions that extend mouse lifespan suppress shared age-associated epigenetic changes at critical gene regulatory regions. *Genome Biol.* **18**, 58 (2017).
- Arneson, A. et al. A mammalian methylation array for profiling methylation levels at conserved sequences. *Nat. Commun.* 13, 783 (2022).
- 9. Haghani, A. et al. DNA methylation networks underlying mammalian traits. *Science* **381**, eabq5693 (2023).
- 10. Prado, N. A. et al. Epigenetic clock and methylation studies in elephants. *Aging Cell* **20**, e13414 (2021).
- Sugrue, V. J. et al. Castration delays epigenetic aging and feminizes DNA methylation at androgen-regulated loci. *eLife* 10, e64932 (2021).
- Schachtschneider, K. M. et al. Epigenetic clock and DNA methylation analysis of porcine models of aging and obesity. *GeroScience* 43, 2467–2483 (2021).
- Robeck, T. R. et al. Multi-species and multi-tissue methylation clocks for age estimation in toothed whales and dolphins. *Commun. Biol.* 4, 642 (2021).
- 14. Wilkinson, G. S. et al. DNA methylation predicts age and provides insight into exceptional longevity of bats. *Nat. Commun.* **12**, 1615 (2021).
- Horvath, S. et al. DNA methylation clocks tick in naked mole rats but queens age more slowly than nonbreeders. *Nat. Aging* 2, 46–59 (2022).
- 16. Horvath, S. et al. DNA methylation aging and transcriptomic studies in horses. *Nat. Commun.* **13**, 40 (2022).
- Larison, B. et al. Epigenetic models developed for plains zebras predict age in domestic horses and endangered equids. *Commun. Biol.* 4, 1412 (2021).
- 18. Horvath, S. et al. DNA methylation clocks for dogs and humans. Proc. Natl Acad. Sci. USA **119**, e2120887119 (2022).
- 19. Kerepesi, C. et al. Epigenetic aging of the demographically nonaging naked mole-rat. *Nat. Commun.* **13**, 355 (2022).
- 20. Hannum, G. et al. Genome-wide methylation profiles reveal quantitative views of human aging rates. *Mol. Cell* **49**, 359–367 (2013).
- Bocklandt, S. et al. Epigenetic predictor of age. PLoS ONE 6, e14821 (2011).
- 22. Levine, M. E. et al. An epigenetic biomarker of aging for lifespan and healthspan. *Aging* **10**, 573–591 (2018).
- 23. Lu, A. T. et al. DNA methylation GrimAge strongly predicts lifespan and healthspan. *Aging* **11**, 303–327 (2019).
- 24. Kundaje, A. et al. Integrative analysis of 111 reference human epigenomes. *Nature* **518**, 317–330 (2015).
- Illingworth, R. et al. A novel CpG island set identifies tissuespecific methylation at developmental gene loci. *PLoS Biol.* 6, e22 (2008).
- 26. Levine, M. E. et al. Menopause accelerates biological aging. *Proc. Natl Acad. Sci. USA* **113**, 9327–9332 (2016).
- 27. Marioni, R. E. et al. DNA methylation age of blood predicts allcause mortality in later life. *Genome Biol.* **16**, 25 (2015).

- 28. Lu, A. T. et al. DNA methylation GrimAge version 2. Aging **14**, 9484–9549 (2022).
- 29. Langfelder, P. & Horvath, S. WGCNA: an R package for weighted correlation network analysis. *BMC Bioinformatics* **9**, 559 (2008).
- 30. Gill, D. et al. Multi-omic rejuvenation of human cells by maturation phase transient reprogramming. *eLife* **11**, e71624 (2022).
- 31. Ohnuki, M. et al. Dynamic regulation of human endogenous retroviruses mediates factor-induced reprogramming and differentiation potential. *Proc. Natl Acad. Sci. USA* **111**, 12426–12431 (2014).
- 32. Olova, N., Simpson, D. J., Marioni, R. E. & Chandra, T. Partial reprogramming induces a steady decline in epigenetic age before loss of somatic identity. *Aging Cell* **18**, e12877 (2019).
- Basu, R., Qian, Y. & Kopchick, J. J. Mechanisms in endocrinology: lessons from growth hormone receptor gene-disrupted mice: are there benefits of endocrine defects? *Eur. J. Endocrinol.* **178**, R155– R181 (2018).
- 34. Fontana, L., Partridge, L. & Longo, V. D. Extending healthy life span—from yeast to humans. *Science* **328**, 321–326 (2010).
- Flurkey, K., Papaconstantinou, J., Miller, R. A. & Harrison, D. E. Lifespan extension and delayed immune and collagen aging in mutant mice with defects in growth hormone production. *Proc. Natl Acad. Sci. USA* **98**, 6736–6741 (2001).
- Dominick, G. et al. Regulation of mTOR activity in Snell dwarf and GH receptor gene-disrupted mice. *Endocrinology* 156, 565–575 (2015).
- Coschigano, K. T., Clemmons, D., Bellush, L. L. & Kopchick, J. J. Assessment of growth parameters and life span of GHR/BP genedisrupted mice. *Endocrinology* 141, 2608–2613 (2000).
- List, E. O. et al. Liver-specific GH receptor gene-disrupted (LiGHRKO) mice have decreased endocrine IGF-I, increased local IGF-I, and altered body size, body composition, and adipokine profiles. *Endocrinology* **155**, 1793–1805 (2014).
- Nagarajan, A., Srivastava, H., Jablonsky, J. & Sun, L. Y. Tissuespecific GHR knockout mice: an updated review. *Front. Endocrinol.* 11, 579909 (2020).
- 40. Everitt, A. V., Rattan, S. I., Couteur, D. G. & de Cabo, R. *Calorie Restriction, Aging and Longevity* (Springer Science & Business Media, 2010).
- Kennedy, B. K., Steffen, K. K. & Kaeberlein, M. Ruminations on dietary restriction and aging. *Cell. Mol. Life Sci.* 64, 1323–1328 (2007).
- 42. Acosta-Rodríguez, V. et al. Circadian alignment of early onset caloric restriction promotes longevity in male C57BL/6J mice. *Science* **376**, 1192–1202 (2022).
- 43. Wang, T. et al. Epigenetic aging signatures in mice livers are slowed by dwarfism, calorie restriction and rapamycin treatment. *Genome Biol.* **18**, 57 (2017).
- 44. Thompson, M. J. et al. A multi-tissue full lifespan epigenetic clock for mice. *Aging* **10**, 2832–2854 (2018).
- 45. Lu, Y. et al. Reprogramming to recover youthful epigenetic information and restore vision. *Nature* **588**, 124–129 (2020).
- 46. Vu, H. & Ernst, J. Universal annotation of the human genome through integration of over a thousand epigenomic datasets. *Genome Biol.* **23**, 9 (2022).
- 47. Margueron, R. & Reinberg, D. The polycomb complex PRC2 and its mark in life. *Nature* **469**, 343–349 (2011).
- Rakyan, V. K. et al. Human aging-associated DNA hypermethylation occurs preferentially at bivalent chromatin domains. *Genome Res.* 20, 434–439 (2010).
- 49. Teschendorff, A. E. et al. Age-dependent DNA methylation of genes that are suppressed in stem cells is a hallmark of cancer. *Genome Res.* **20**, 440–446 (2010).
- 50. Zhou, W. et al. DNA methylation loss in late-replicating domains is linked to mitotic cell division. *Nat. Genet.* **50**, 591–602 (2018).

Resource

- 51. McLean, C. Y. et al. GREAT improves functional interpretation of cis-regulatory regions. *Nat. Biotechnol.* **28**, 495–501 (2010).
- Bleda, M. et al. CellBase, a comprehensive collection of RESTful web services for retrieving relevant biological information from heterogeneous sources. *Nucleic Acids Res.* 40, W609–W614 (2012).
- 53. Davis, C. A. et al. The Encyclopedia of DNA Elements (ENCODE): data portal update. *Nucleic Acids Res.* **46**, D794–D801 (2018).
- Tacutu, R. et al. Human Ageing Genomic Resources: new and updated databases. *Nucleic Acids Res.* 46, D1083–D1090 (2018).
- 55. Chen, E. Y. et al. Enrichr: interactive and collaborative HTML5 gene list enrichment analysis tool. *BMC Bioinformatics* **14**, 128 (2013).
- Kuleshov, M. V. et al. Enrichr: a comprehensive gene set enrichment analysis web server 2016 update. *Nucleic Acids Res.* 44, W90–W97 (2016).
- 57. Ziller, M. J. et al. Charting a dynamic DNA methylation landscape of the human genome. *Nature* **500**, 477–481 (2013).
- Luecken, M. D. et al. A sandbox for prediction and integration of DNA, RNA, and proteins in single cells. In 35th Conference on Neural Information Processing Systems Datasets and Benchmarks Track (Round 2) (eds Vanschoren, J. & Yeung, S.) (NeurIPS, 2021); https://openreview.net/forum?id=gN35BGa1Rt
- Itokawa, N. et al. Epigenetic traits inscribed in chromatin accessibility in aged hematopoietic stem cells. *Nat. Commun.* 13, 2691 (2022).
- Boyer, L. A. et al. Polycomb complexes repress developmental regulators in murine embryonic stem cells. *Nature* 441, 349–353 (2006).
- 61. Lynch, M. D. et al. An interspecies analysis reveals a key role for unmethylated CpG dinucleotides in vertebrate polycomb complex recruitment. *EMBO J.* **31**, 317–329 (2012).
- Choy, J. S. et al. DNA methylation increases nucleosome compaction and rigidity. J. Am. Chem. Soc. 132, 1782–1783 (2010).
- 63. Martin-Herranz, D. E. et al. Screening for genes that accelerate the epigenetic aging clock in humans reveals a role for the H3K36 methyltransferase NSD1. *Genome Biol.* **20**, 146 (2019).
- Jeffries, A. R. et al. Growth disrupting mutations in epigenetic regulatory molecules are associated with abnormalities of epigenetic aging. *Genome Res.* 29, 1057–1066 (2019).
- Fonseca, B. D. et al. La-related protein 1 (LARP1) represses terminal oligopyrimidine (TOP) mRNA translation downstream of mTOR complex 1 (mTORC1). J. Biol. Chem. 290, 15996–16020 (2015).
- Kennedy, B. K. & Lamming, D. W. The mechanistic target of rapamycin: the grand conducTOR of metabolism and aging. *Cell Metab.* 23, 990–1003 (2016).
- 67. Horvath, S., Lu, A. T., Cohen, H. & Raj, K. Rapamycin retards epigenetic ageing of keratinocytes independently of its effects on replicative senescence, proliferation and differentiation. *Aging* **11**, 3238–3249 (2019).
- 68. Lu, A. T. et al. Genetic variants near *MLST8* and *DHX57* affect the epigenetic age of the cerebellum. *Nat. Commun.* **7**, 10561 (2016).
- 69. de Magalhaes, J. P. Programmatic features of aging originating in development: aging mechanisms beyond molecular damage? *FASEB J.* **26**, 4821–4826 (2012).
- 70. Williams, G. C. Pleiotropy, natural selection, and the evolution of senescence. *Evolution* **11**, 398–411 (1957).
- 71. Bowles, J. T. The evolution of aging: a new approach to an old problem of biology. *Med. Hypotheses* **51**, 179–221 (1998).
- Blagosklonny, M. V. Aging and immortality: quasi-programmed senescence and its pharmacologic inhibition. *Cell Cycle* 5, 2087–2102 (2006).
- 73. Booth, L. N. & Brunet, A. The aging epigenome. *Mol. Cell* **62**, 728–744 (2016).

- 74. Horvath, S. & Raj, K. DNA methylation-based biomarkers and the epigenetic clock theory of ageing. *Nat. Rev. Genet.* **19**, 371–384 (2018).
- 75. Mayne, B., Berry, O., Davies, C., Farley, J. & Jarman, S. A genomic predictor of lifespan in vertebrates. *Sci. Rep.* **9**, 17866 (2019).
- Mitteldorf, J. An epigenetic clock controls aging. *Biogerontology* 17, 257–265 (2016).
- 77. Rando, T. A. & Chang, H. Y. Aging, rejuvenation, and epigenetic reprogramming: resetting the aging clock. *Cell* **148**, 46–57 (2012).
- Sen, P., Shah, P. P., Nativio, R. & Berger, S. L. Epigenetic mechanisms of longevity and aging. Cell 166, 822–839 (2016).
- 79. Yang, J.-H. et al. Erosion of the epigenetic landscape and loss of cellular identity as a cause of aging in mammals. Preprint at SSRN https://doi.org/10.2139/ssrn.3461780 (2019).
- 80. Gems, D. The hyperfunction theory: an emerging paradigm for the biology of aging. *Ageing Res. Rev.* **74**, 101557 (2022).
- 81. Ocampo, A. et al. In vivo amelioration of age-associated hallmarks by partial reprogramming. *Cell* **167**, 1719–1733 (2016).
- Rodríguez-Matellán, A., Alcazar, N., Hernández, F., Serrano, M. & Ávila, J. In vivo reprogramming ameliorates aging features in dentate gyrus cells and improves memory in mice. *Stem Cell Reports* 15, 1056–1066 (2020).
- Sarkar, T. J. et al. Transient non-integrative expression of nuclear reprogramming factors promotes multifaceted amelioration of aging in human cells. *Nat. Commun.* 11, 1545 (2020).
- 84. Takahashi, K. & Yamanaka, S. Induction of pluripotent stem cells from mouse embryonic and adult fibroblast cultures by defined factors. *Cell* **126**, 663–676 (2006).
- 85. Zhou, W., Triche, T. J. Jr, Laird, P. W. & Shen, H. SeSAMe: reducing artifactual detection of DNA methylation by Infinium BeadChips in genomic deletions. *Nucleic Acids Res.* **46**, e123 (2018).
- de Magalhaes, J. P., Costa, J. & Church, G. M. An analysis of the relationship between metabolism, developmental schedules, and longevity using phylogenetic independent contrasts. *J. Gerontol. A Biol. Sci. Med. Sci.* 62, 149–160 (2007).
- Dawber, T. R., Meadors, G. F. & Moore, F. E. Jr. Epidemiological approaches to heart disease: the Framingham Study. *Am. J. Public Health Nations Health* 41, 279–281 (1951).
- 88. Anderson, G. L. et al. Implementation of the Women's Health Initiative study design. *Ann. Epidemiol.* **13**, S5–S17 (2003).
- The Women's Health Initiative Study Group. Design of the Women's Health Initiative clinical trial and observational study. Control. Clin. Trials 19, 61–109 (1998).
- Almasy, L. & Blangero, J. Multipoint quantitative-trait linkage analysis in general pedigrees. Am. J. Hum. Genet. 62, 1198–1211 (1998).
- Ziyatdinov, A. et al. solarius: an R interface to SOLAR for variance component analysis in pedigrees. *Bioinformatics* **32**, 1901–1902 (2016).
- 92. Virshup, I., Rybakov, S., Theis, F. J., Angerer, P. & Wolf, F. A. anndata: annotated data. Preprint at *bioRxiv* https://doi.org/ 10.1101/2021.12.16.473007 (2021).
- 93. Lawrence, M. et al. Software for computing and annotating genomic ranges. *PLoS Comput. Biol.* **9**, e1003118 (2013).
- Stuart, T., Srivastava, A., Madad, S., Lareau, C. A. & Satija, R. Single-cell chromatin state analysis with Signac. *Nat. Methods* 18, 1333–1341 (2021).
- 95. Horvath, S. et al. DNA methylation studies of mammalian species. *Github* https://github.com/shorvath/Mammalian MethylationConsortium/ (2022).
- 96. Peters, M. J. et al. The transcriptional landscape of age in human peripheral blood. *Nat. Commun.* **6**, 8570 (2015).

Acknowledgements

This work was mainly supported by the Paul G. Allen Frontiers Group (S.H.). Additional support was also provided by the Open Philanthropy–Silicon Valley Fund (S.H. and K.R.). J.E. was supported by UCLA Jonsson Comprehensive Cancer Center and Eli and Edythe Broad Center of Regenerative Medicine and Stem Cell Research Ablon Scholars Program. J.A.M. and the NHP Core were supported by the Intramural Research Program, National Institute on Aging, NIH. Plains zebra sample collection was supported by National Geographic Society grant 8941-11. We acknowledge the Museum of Vertebrate Zoology and C.J. Conroy from the University of California, Berkeley. We acknowledge R. Miller and his laboratory (http://www. richmillerlab.com/long-lived-mutants) for providing dwarf mice and controls (Snell dwarf mouse, GHRKO experiments). Lemur sample collections were supported by Duke Lemur Center. N.C.B. was funded by a DST-NRF SARChI chair of Mammalian Behavioural Ecology and Physiology (GUN 64756). V.N.G., A.S. and V.G. were supported by NIA grants. V.G. was supported by the Milky Way Research Foundation. D.T.O. was supported by European Research Council (788937) and Cancer Research UK (20412). The FHS is funded by National Institutes of Health contracts NO1-HC-25195 and HHSN268201500001I. The laboratory work for this investigation was funded by the Division of Intramural Research, National Heart, Lung, and Blood Institute, National Institutes of Health. The analytical component of this project was funded by the Division of Intramural Research, National Heart, Lung, and Blood Institute and the Center for Information Technology, National Institutes of Health. The WHI program is funded by the National Heart, Lung, and Blood Institute, National Institutes of Health, U.S. Department of Health and Human Services through contracts HHSN268201600018C, HHSN268201600001C, HHSN268201600002C, HHSN268201600003C and HHSN268201600004C. We thank the WHI investigators and staff for their dedication and the study participants for making the program possible. A full listing of WHI investigators can be found at https://www.whi.org/doc/WHI-Investigator-Long-List.pdf. The views expressed in this study are those of the authors and do not necessarily represent the views of funding bodies such as the National Heart, Lung, and Blood Institute, the National Institutes of Health or the U.S. Department of Health and Human Services.

Author contributions

A.T.L., Z.F., C.L., J.A.Z. and S.H. developed the universal clocks. A.T.L., A.H., R.L., Q.Y., S.B.K., C.E.B., M.J.T., M.P., H.V., W.Z. and S.H. carried out additional bioinformatic analyses. A. Arneson, J.E. and S.H. designed the mammalian methylation array. A.T.L., Z.F., A.H., R.L., Q.Y., K.R. and S.H. drafted the first version of the article. The remaining authors contributed tissues or DNA samples or helped with the data-generation process. All authors helped with editing the article and data interpretation. S.H. conceived the study and design and supervised this work.

Competing interests

The Regents of the University of California filed a patent application (publication number WO2020150705) related to this work on which S.H., A. Arneson and J.E. are named inventors. S.H. and R.T.B. are founders of the non-profit Epigenetic Clock Development Foundation, which has licensed several patents from UC Regents, and distributes the mammalian methylation array. The remaining authors declare no competing interests.

Additional information

Extended data is available for this paper at https://doi.org/10.1038/s43587-023-00462-6.

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s43587-023-00462-6.

Correspondence and requests for materials should be addressed to S. Horvath.

Peer review information *Nature Aging* thanks Payel Sen for their contribution to the peer review of this work.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons. org/licenses/by/4.0/.

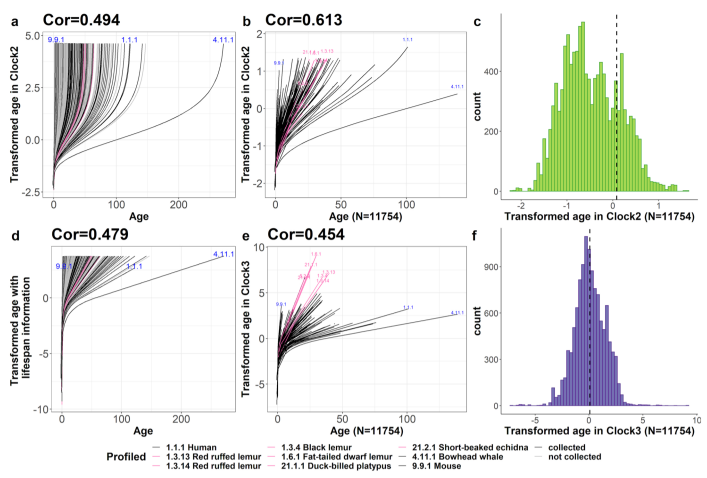
© The Author(s) 2023, corrected publication 2023

A. T. Lu^{® 1,2,137}, Z. Fei^{3,4,137}, A. Haghani^{® 1,2}, T. R. Robeck⁵, J. A. Zoller^{® 3}, C. Z. Li^{® 3}, R. Lowe⁶, Q. Yan², J. Zhang^{® 1}, H. Vu^{7,8}, J. Ablaeva⁹, V. A. Acosta-Rodriguez^{® 10}, D. M. Adams^{® 11}, J. Almunia^{® 12}, A. Aloysius^{® 13}, R. Ardehali¹⁴, A. Arneson^{7,8}, C. S. Baker^{® 15}, G. Banks¹⁶, K. Belov¹⁷, N. C. Bennett¹⁸, P. Black¹⁹, D. T. Blumstein^{© 20,21}, E. K. Bors¹⁵, C. E. Breeze^{® 22}, R. T. Brooke²³, J. L. Brown²⁴, G. G. Carter²⁵, A. Caulton^{26,27}, J. M. Cavin²⁸, L. Chakrabarti²⁹, I. Chatzistamou³⁰, H. Chen³¹, K. Cheng³², P. Chiavellini³³, O. W. Choi³⁴, S. M. Clarke^{® 26}, L. N. Cooper^{® 35}, M. L. Cossette^{® 36}, J. Day^{® 37}, J. DeYoung³⁴, S. DiRocco³⁸, C. Dold³⁹, E. E. Ehmke⁴⁰, C. K. Emmons^{® 41}, S. Emmrich^{® 42}, E. Erbay⁴³, C. Erlacher-Reid^{38,44}, C. G. Faulkes^{® 45}, S. H. Ferguson^{® 46,47}, C. J. Finno⁴⁸, J. E. Flower⁴⁹, J. M. Gaillard⁵⁰, E. Garde⁵¹, L. Gerber⁵², V. N. Gladyshev^{® 53}, V. Gorbunova^{® 42}, R. G. Goya^{® 33}, M. J. Grant⁵⁴, C. B. Green¹⁰, E. N. Hales⁴⁸, M. B. Hanson⁴¹, D. W. Hart^{® 18}, M. Haulena⁵⁵, K. Herrick⁵⁶, A. N. Hogan⁵⁷, C. J. Hogg^{® 17}, T. A. Hore^{® 58}, T. Huang^{® 59,60}, J. C. Izpisua Belmonte², A. J. Jasinska^{® 34}, G. Jones^{® 61}, E. Jourdain⁶², O. Kashpur⁶³, H. Katcher⁶⁴, E. Katsumata⁶⁵, V. Kaza⁶⁶, H. Kiaris^{® 66,67}, M. S. Kobor⁶⁸, P. Kordowitzki^{® 69,70}, W. R. Koski^{® 71}, M. Krützen^{® 72}, S. B. Kwon⁷⁸, B. Larison^{73,74}, S. G. Lee⁵³, M. Lehmann³³, J. F. Lemaitre^{® 50}, A. J. Levine⁷⁵, C. Li^{76,77}, X. Li⁷⁸, A. R. Lim¹, D. T. S. Lin⁷⁹, D. M. Lindemann³⁸, T. J. Little⁸⁰, N. Macoretta⁴², D. Maddox⁸¹, C. O. Matkin⁸², J. A. Mattison⁸³, M. McClure⁸⁴, J. Mergl⁸⁵, J. J. Meudt⁸⁶, G. A. Montano³⁹, K. Mozhui^{® 7,88}, J. Munshi-South⁸⁹, A. Naderi^{® 67}, M. Nagy^{® 90}, P. Narayan^{® 54}, P. W. Nathanielsz^{76,77}, N. B. Nguyen¹⁴,

C. Niehrs^{91,92}, J. K. O'Brien³⁷, P. O'Tierney Ginn @ ^{63,93}, D. T. Odom^{94,95}, A. G. Ophir @ ⁹⁶, S. Osborn⁹⁷, E. A. Ostrander @ ⁵⁷, K. M. Parsons⁴¹, K. C. Paul⁷⁵, M. Pellegrini⁹⁸, K. J. Peters @ ^{72,99}, A. B. Pedersen¹⁰⁰, J. L. Petersen @ ¹⁰¹, D. W. Pietersen @ ¹⁰², G. M. Pinho @ ⁷³, J. Plassais @ ⁵⁷, J. R. Poganik @ ⁵³, N. A. Prado¹⁰³, P. Reddy^{2,104}, B. Rey @ ⁵⁰, B. R. Ritz @ ^{105,106,107}, J. Robbins @ ¹⁰⁸, M. Rodriguez¹⁰⁹, J. Russell @ ⁵⁶, E. Rydkina⁴², L. L. Sailer @ ⁹⁶, A. B. Salmon @ ¹¹⁰, A. Sanghavi @ ⁶⁴, K. M. Schachtschneider^{111,112,113}, D. Schmitt @ ¹¹⁴, T. Schmitt⁵⁶, L. Schomacher @ ⁹¹, L. B. Schook @ ^{111,115}, K. E. Sears @ ^{73,98}, A. W. Seifert¹³, A. Seluanov @ ⁴², A. B. A. Shafer¹¹⁶, D. Shanmuganayagam @ ^{86,117}, A. V. Shindyapina⁵³, M. Simmons⁴⁰, K. Singh¹¹⁸, I. Sinha⁷³, J. Slone⁵⁹, R. G. Snell @ ⁵⁴, E. Soltanmaohammadi⁶⁷, M. L. Spangler @ ¹⁰¹, M. C. Spriggs¹⁹, L. Staggs³⁸, N. Stedman¹⁹, K. J. Steinman¹¹⁹, D. T. Stewart¹²⁰, V. J. Sugrue⁵⁸, B. Szladovits @ ¹²¹, J. S. Takahashi @ ^{10,122}, M. Takasugi⁴², E. C. Teeling @ ¹²³, M. J. Thompson @ ⁹⁸, B. Van Bonn¹²⁴, S. C. Vernes^{125,126}, D. Villar @ ¹²⁷, H. V. Vinters¹²⁸, M. C. Wallingford ^{63,129}, N. Wang^{130,131}, R. K. Wayne⁷³, G. S. Wilkinson @ ¹¹, C. K. Williams⁷⁵, R. W. Williams⁸⁸, X. W. Yang @ ^{130,131}, M. Yao³, B. G. Young @ ¹³², B. Zhang⁵³, Z. Zhang⁴², P. Zhao @ ^{14,133}, Y. Zhao⁴², W. Zhou^{134,135}, J. Zimmermann @ ¹³⁶, J. Ernst @ ⁷⁸, K. Raj @ ^{6,137} & S. Horvath @ ^{1,2,3,137}

¹Department of Human Genetics, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA, USA. ²Altos Labs, San Diego Institute of Science, San Diego, CA, USA. ³Department of Biostatistics, Fielding School of Public Health, University of California, Los Angeles, Los Angeles, CA, USA. ⁴Department of Statistics, University of California, Riverside, Riverside, CA, USA. ⁵Zoological SeaWorld Parks and Entertainment, Orlando, FL, USA. ⁶Altos Labs, Cambridge Institute of Science, Cambridge, UK. ⁷Bioinformatics Interdepartmental Program, University of California, Los Angeles, CA, USA. 8Department of Biological Chemistry, University of California, Los Angeles, Los Angeles, CA, USA. 9Department of Biology, University of Rochester, Rochester, NY, USA. ¹⁰Department of Neuroscience, Peter O'Donnell Jr. Brain Institute, University of Texas Southwestern Medical Center, Dallas, TX, USA. ¹¹Department of Biology, University of Maryland, College Park, MD, USA. ¹²Loro Parque Fundacion, Puerto de la Cruz, Spain. ¹³Department of Biology, University of Kentucky, Lexington, KY, USA. ¹⁴Division of Cardiology, Department of Internal Medicine, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA, USA.¹⁵Marine Mammal Institute, Oregon State University, Newport, OR, USA.¹⁶School of Science and Technology, Clifton Campus, Nottingham Trent University, Nottingham, UK. ¹⁷School of Life and Environmental Sciences, the University of Sydney, Sydney, New South Wales, Australia.¹⁸Department of Zoology and Entomology, University of Pretoria, Hatfield, South Africa.¹⁹Busch Gardens Tampa, Tampa, FL, USA. ²⁰Department of Ecology and Evolutionary Biology, University of California, Los Angeles, Los Angeles, CA, USA. ²¹Rocky Mountain Biological Laboratory, Crested Butte, CO, USA. 22 Altius Institute for Biomedical Sciences, Seattle, WA, USA. 23 Epigenetic Clock Development Foundation, Los Angeles, CA, USA.²⁴Center for Species Survival, Smithsonian Conservation Biology Institute, Front Royal, VA, USA.²⁵Department of Evolution, Ecology and Organismal Biology, The Ohio State University, Columbus, OH, USA. 26 AgResearch, Invermay Agricultural Centre, Mosgiel, New Zealand. ²⁷Department of Biochemistry, University of Otago, Dunedin, New Zealand. ²⁸Gulf World, Dolphin Company, Panama City Beach, FL, USA. ²⁹School of Veterinary Medicine and Science, University of Nottingham, Nottingham, UK. ³⁰Department of Pathology, Microbiology and Immunology, School of Medicine, University of South Carolina, Columbia, SC, USA. ³¹Department of Pharmacology, Addiction Science and Toxicology, the University of Tennessee Health Science Center, Memphis, TN, USA. ³²Medical Informatics, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA, USA. ³³Biochemistry Research Institute of La Plata, Histology and Pathology, School of Medicine, University of La Plata, La Plata, Argentina. ³⁴Center for Neurobehavioral Genetics, Semel Institute for Neuroscience and Human Behavior, Department of Psychiatry and Biobehavioral Sciences, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA, USA.³⁵Department of Anatomy and Neurobiology, Northeast Ohio Medical University, Rootstown, OH, USA. ³⁶Department of Environmental and Life Sciences, Trent University, Peterborough, Ontario, Canada. ³⁷Taronga Institute of Science and Learning, Taronga Conservation Society Australia, Mosman, New South Wales, Australia, ³⁸SeaWorld of Florida, Orlando, FL, USA, ³⁹Zoological Operations, SeaWorld Parks and Entertainment, Orlando, FL, USA, ⁴⁰Duke Lemur Center, Durham, NC, USA, ⁴¹Conservation Biology Division, Northwest Fisheries Science Center, National Marine Fisheries Service, National Oceanic and Atmospheric Administration, Seattle, WA, USA. ⁴²Departments of Biology and Medicine, University of Rochester, Rochester, NY, USA. ⁴³Altos Labs, San Francisco, CA, USA. ⁴⁴SeaWorld Orlando, Orlando, FL, USA.⁴⁵School of Biological and Behavioural Sciences, Queen Mary University of London, London, UK.⁴⁶Fisheries and Oceans Canada, Freshwater Institute, Winnipeg, Manitoba, Canada. ⁴⁷Department of Biological Sciences, University of Manitoba, Winnipeg, Manitoba, Canada. ⁴⁸Department of Population Health and Reproduction, University of California, Davis School of Veterinary Medicine, Davis, CA, USA.⁴⁹Mystic Aquarium, Mystic, CT, USA. 50 Universite de Lyon, Universite Lyon 1, CNRS, Laboratoire de Biometrie et Biologie Evolutive, Villeurbanne, France. 51 Greenland Institute of Natural Resources, Nuuk, Greenland. ⁵²Evolution and Ecology Research Centre, School of Biological, Earth and Environmental Sciences, UNSW Sydney, Sydney, New South Wales, Australia. 53 Division of Genetics, Department of Medicine, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA. ⁵⁴Applied Translational Genetics Group, School of Biological Sciences, Centre for Brain Research, the University of Auckland, Auckland, New Zealand. 55 Vancouver Aquarium, Vancouver, British Columbia, Canada. 56 SeaWorld of California, San Diego, CA, USA. 57 Cancer Genetics and Comparative Genomics Branch, National Human Genome Research Institute, National Institutes of Health, Bethesda, MD, USA. 58 Department of Anatomy, University of Otago, Dunedin, New Zealand. 59 Division of Human Genetics, Department of Pediatrics, University at Buffalo, Buffalo, NY, USA. 60 Division of Genetics and Metabolism, Oishei Children's Hospital, Buffalo, NY, USA.⁶¹School of Biological Sciences, University of Bristol, Bristol, UK.⁶²Norwegian Orca Survey, Andenes, Norway. 63 Mother Infant Research Institute, Tufts Medical Center, Boston, MA, USA. 64 Yuvan Research, Mountain View, CA, USA. ⁶⁵Kamogawa Sea World, Kamogawa, Japan.⁶⁶Peromyscus Genetic Stock Center, University of South Carolina, Columbia, SC, USA.⁶⁷Department of Drug Discovery and Biomedical Sciences, College of Pharmacy, University of South Carolina, Columbia, SC, USA. 68 Edwin S.H. Leong Healthy Aging Program, Centre for Molecular Medicine and Therapeutics, University of British Columbia, Vancouver, British Columbia, Canada. 69 Institute of Animal Reproduction and Food Research of the Polish Academy of Sciences, Olsztyn, Poland. 70 Institute for Veterinary Medicine, Nicolaus Copernicus University, Torun, Poland. ⁷¹LGL Limited, King City, Ontario, Canada. ⁷²Evolutionary Genetics Group, Department of Evolutionary Anthropology, University of Zurich, Zurich, Switzerland. 73Department of Ecology and Evolutionary Biology, UCLA, Los Angeles, CA, USA. 74Center for Tropical Research, Institute for the Environment and Sustainability, UCLA, Los Angeles, CA, USA.⁷⁵Department of Neurology, David Geffen School of Medicine, University of California, Los Angeles, Los

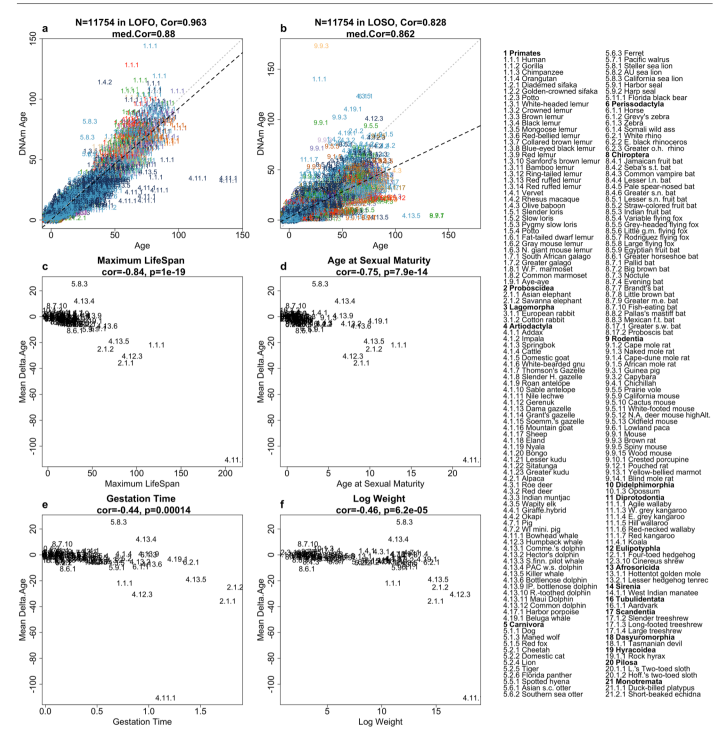
Angeles, CA, USA.⁷⁶Texas Pregnancy and Life-course Health Center, Southwest National Primate Research Center, San Antonio, TX, USA.⁷⁷Department of Animal Science, College of Agriculture and Natural Resources, Laramie, WY, USA. 78 Technology Center for Genomics and Bioinformatics, Department of Pathology and Laboratory Medicine, University of California, Los Angeles, Los Angeles, CA, USA. 79 Centre for Molecular Medicine and Therapeutics, BC Children's Hospital Research Institute, University of British Columbia, Vancouver, British Columbia, Canada. 80 Institute of Ecology and Evolution, School of Biological Sciences, University of Edinburgh, Edinburgh, UK.⁸¹White Oak Conservation, Yulee, FL, USA.⁸²North Gulf Oceanic Society, Homer, AK, USA. ⁸³Translational Gerontology Branch, National Institute on Aging Intramural Research Program, National Institutes of Health, Baltimore, MD, USA.⁸⁴ABS Global, DeForest, WI, USA. 85 Marineland of Canada, Niagara Falls, Ontario, Canada. 86 Biomedical and Genomic Research Group, Department of Animal and Dairy Sciences, University of Wisconsin–Madison, Madison, WI, USA. ⁸⁷Department of Preventive Medicine, University of Tennessee Health Science Center, College of Medicine, Memphis, TN, USA. 88 Department of Genetics, Genomics and Informatics, University of Tennessee Health Science Center, College of Medicine, Memphis, TN, USA.⁸⁹Louis Calder Center—Biological Field Station, Department of Biological Sciences, Fordham University, Armonk, NY, USA. 90 Museum fur Naturkunde, Leibniz Institute for Evolution and Biodiversity Science, Berlin, Germany. 91 Institute of Molecular Biology, Mainz, Germany. 92 Division of Molecular Embryology, DKFZ-ZMBH Alliance, Heidelberg, Germany. 93 Department of Obstetrics and Gynecology, Tufts University School of Medicine, Boston, MA, USA. ⁹⁴Cancer Research UK Cambridge Institute, University of Cambridge, Cambridge, UK. ⁹⁵Division of Regulatory Genomics and Cancer Evolution, Deutsches Krebsforschungszentrum, Heidelberg, Germany, ⁹⁶Department of Psychology, Cornell University, Ithaca, NY, USA. 97 SeaWorld of Texas, San Antonio, TX, USA. 98 Department of Molecular Cell and Developmental Biology, University of California, Los Angeles, Los Angeles, CA, USA. 99 School of Earth, Atmospheric and Life Sciences, University of Wollongong, Wollongong, Australia. 100 Institute of Evolutionary Biology, School of Biological Sciences, University of Edinburgh, Edinburgh, UK.¹⁰¹Department of Animal Science, University of Nebraska, Lincoln, NE, USA. ¹⁰²Mammal Research Institute, Department of Zoology and Entomology, University of Pretoria, Hatfield, South Africa. ¹⁰³Department of Biology, College of Arts and Science, Adelphi University, Garden City, NY, USA. ¹⁰⁴Salk Institute for Biological Studies, La Jolla, CA, USA. ¹⁰⁵Department of Epidemiology, UCLA Fielding School of Public Health, Los Angeles, CA, USA. ¹⁰⁶Department of Environmental Health Sciences, UCLA Fielding School of Public Health, Los Angeles, CA, USA. 107 Department of Neurology, UCLA David Geffen School of Medicine, Los Angeles, CA, USA. 108 Center for Coastal Studies, Provincetown, MA, USA.¹⁰⁹Miami Seaquarium, Miami, FL, USA.¹¹⁰The Sam and Ann Barshop Institute for Longevity and Aging Studies and Department of Molecular Medicine, UT Health San Antonio and the Geriatric Research Education and Clinical Center, South Texas Veterans Healthcare System, San Antonio, TX, USA. 111 Department of Radiology, University of Illinois at Chicago, Chicago, IL, USA. 112 Department of Biochemistry and Molecular Genetics, University of Illinois at Chicago, Chicago, IL, USA. 113 National Center for Supercomputing Applications, University of Illinois at Urbana-Champaign, Urbana, IL, USA.¹¹⁴College of Agriculture, Missouri State University, Springfield, MO, USA.¹¹⁵Department of Animal Sciences, University of Illinois at Urbana-Champaign, Champaign, IL, USA.¹¹⁶Department of Forensic Science, Environmental and Life Sciences, Trent University, Peterborough, Ontario, Canada. 117 Department of Surgery, University of Wisconsin School of Medicine and Public Health, Madison, WI, USA. 118 Shobhaben Pratapbhai Patel School of Pharmacy and Technology Management, SVKM'S NMIMS University, Mumbai, India.¹¹⁹Species Preservation Laboratory, SeaWorld San Diego, San Diego, CA, USA. ¹²⁰Biology Department, Acadia University, Wolfville, Nova Scotia, Canada. ¹²¹Department of Pathobiology and Population Sciences, Royal Veterinary College, Hatfield, UK. ¹²²Howard Hughes Medical Institute, Department of Neuroscience, University of Texas Southwestern Medical Center, Dallas, TX, USA. ¹²³School of Biology and Environmental Science, University College Dublin, Dublin, Ireland. ¹²⁴John G. Shedd Aquarium, Chicago, IL, USA. ¹²⁵School of Biology, the University of St Andrews, Fife, UK. ¹²⁶Neurogenetics of Vocal Communication Group, Max Planck Institute for Psycholinguistics, Nijmegen, the Netherlands. 127 Blizard Institute, Faculty of Medicine and Dentistry, Queen Mary University of London, London, UK.¹²⁸Department of Pathology and Laboratory Medicine, David Geffen School of Medicine at UCLA, Los Angeles, CA, USA.¹²⁹Division of Obstetrics and Gynecology, Tufts University School of Medicine, Boston, MA, USA. 130 Center for Neurobehavioral Genetics, Jane and Terry Semel Institute for Neuroscience and Human Behavior, University of California, Los Angeles, Los Angeles, CA, USA. ¹³¹Department of Psychiatry and Biobehavioral Sciences, David Geffen School of Medicine at UCLA, Los Angeles, CA, USA.¹³²Fisheries and Oceans Canada, Winnipeg, Manitoba, Canada.¹³³Eli and Edythe Broad Center of Regenerative Medicine and Stem Cell Research, University of California, Los Angeles, CA, USA. ¹³⁴Center for Computational and Genomic Medicine, Children's Hospital of Philadelphia, Philadelphia, PA, USA. ¹³⁵Department of Pathology and Laboratory Medicine, University of Pennsylvania, Philadelohia, PA, USA, ¹³⁶Department of Mathematics and Technology, University of Applied Sciences Koblenz, Koblenz, Germany, ¹³⁷These authors contributed equally: A. T. Lu, Z. Fei, K. Raj, S. Horvath. 🖂 e-mail: shorvath@mednet.ucla.edu

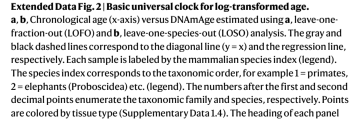


Extended Data Fig. 1 | **Transformed age in universal clocks.** The plot displays transformed age in universal Clock 2 (**a**-**c**) and universal Clock 3 (**d**-**f**). (**a**, **b**) Loglog transformation of Relative Age (y-axis) versus age in universal Clock 2 and (**d**, **e**) log-linear age (y-axis) versus age in our universal Clock 3. Of the 969 mammalian species with available gestation time, age at sexual maturity and maximum lifespan in AnAge database, 339 species are available in our collection. We multiplied the reported maximum lifespan of non-human or non-mouse species by 1.3. Transformed ages were calculated for all the 969 species with simulated age ranging from gestation time through the modified maximum lifespan. The columns (**a**, **d**) display all the 969 species with the simulated ages. In panel d, we proposed the log-linear age with the parameter **m** formulated with maximum lifespan as the information is available for all species

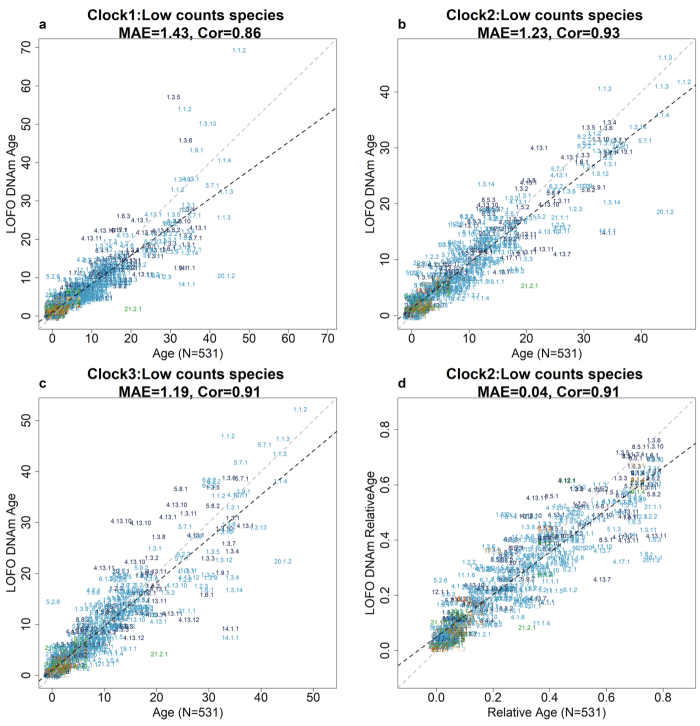
 $(m^*=c_1*\frac{MaxLifespan+GestationT}{ASM+GestationT}$ in Methods). Of the 339 species, 185 species with age

information of high confidence and known tissue types were used in training universal clocks. The columns (**b**, **e**) empirically display these 185 species with the age variable (x-axis) based on the observed ages from all the samples in our collection (N = 11,754). In panel e, we applied the log-linear age formulated without knowing maximum lifespan to train Clock 3 (formula (5) in Methods). Each line represents a species marked by gray for non-profiled and marked by black or pink for profiled species in our collection, as listed in the legend. Some species such as lemurs with relatively short gestation time in regressing m^* (formula (7) in Methods) exhibiting high log-linear ages in (e) are marked in pink. Each panel reports the Pearson correlation coefficient. (c, f) display the histograms of transformed ages based on all samples from the 185 species with vertical lines presenting at means.



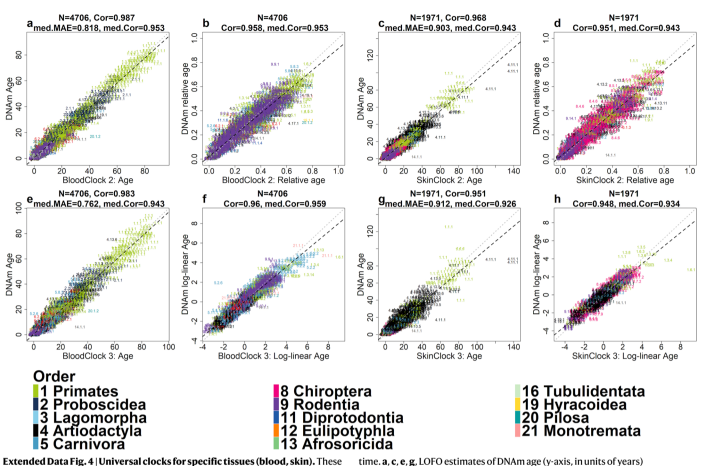


reports the Pearson correlation (cor) across all samples. Here med. Cor denotes the median value across species that contain at least 15 samples. **c**-**f**, The y-axis reports the mean difference between the LOSO estimate of DNAm age and chronological age evaluated at a fixed age defined as half the maximum lifespan (denoted as Mean Delta.Age). The scatter plots depict mean delta half lifespan per species (y-axis) versus c, maximum lifespan observed in the species, d, average age at sexual maturity e, gestational time (in units of years), and f, (log-transformed) average adult body mass in units of grams. All P-values reported are unadjusted and are based on two-sided tests.

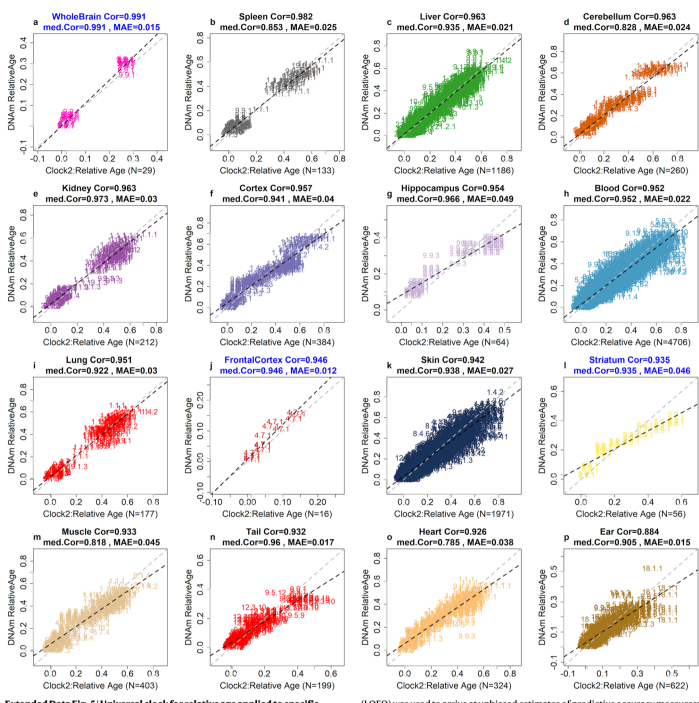


Extended Data Fig. 3 | **Universal clocks applied to species with fewer than 15 samples.** The title of each panel lists the type of universal clock: **a**, Clock 1 = basic universal clock based on log(Age + 2), b, d, Clock 2 = universal clock for relative age, c, Clock 3 = universal clock for log-linear age. Leave-one-fraction-out

(LOFO) methylation estimates versus a-c, chronological age or d, relative age for clock 2. The respective inverse transformations were applied to arrive at DNA methylation-based estimates of chronological age in years or relative age (y-axis).

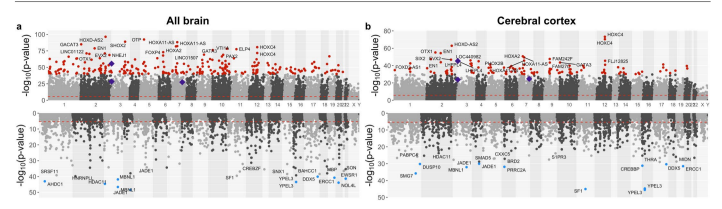


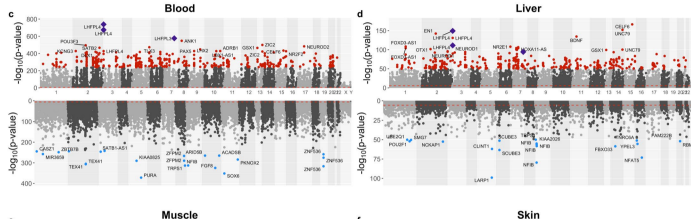
Extended Data Fig. 4 | **Universal clocks for specific tissues (blood, skin).** These tissue-specific universal clocks were constructed in an analogous fashion to the pan-tissue clocks described in the main text. The panels show leave-one-fractionout (LOFO) estimates (y-axis) of four clocks: universal blood clock 2 (Universal BloodClock 2) which estimates relative age, universal blood clock 3 (Universal BloodClock 3) which estimates log-linear transformation of age. Analogously, we defined Universal SkinClock2 and Universal SkinClock3. Relative age estimation incorporates maximum lifespan and gestational age and assumes values between 0 and 1. Log-linear age is formulated with age at sexual maturity and gestational time. **a**, **c**, **e**, **g**, LOFO estimates of DNAm age (y-axis, in units of years) based on transforming relative age (Clock 2) or log-linear age (Clock 3). **b**, **f**, **d**, **h**, transformed age (x-axis) versus corresponding DNAm estimates (y-axis). The title of each panel reports the Pearson correlation coefficient across all data points and the median correlation (med.Cor) and median of median absolute error (med.MAE) across all species. Each sample is labeled by mammalian species index (explained in Fig. 2) and colored by taxonomic order. The legend reports the taxonomic order and the mammalian order index as a prefix.

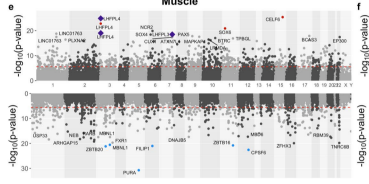


Extended Data Fig. 5 | **Universal clock for relative age applied to specific tissues. a**–**p**, DNA methylation-based estimates of relative age (y-axis) versus actual relative age (x-axis). The specific tissue or cell type is reported in the title of each panel. Each sample is labeled by mammalian species index and colored by tissue type (Supplementary Data 1.3–1.4). The analysis is restricted to tissues that have at least 15 samples available. Leave-one-fraction-out cross-validation

(LOFO) was used to arrive at unbiased estimates of predictive accuracy measures: median absolute error (MAE) and age correlation based on relative age. 'Cor' denotes the Pearson correlation coefficient based on all available samples. 'med. Cor' denotes the median values across all species for which at least 15 samples were available. Title is marked in blue if a tissue type was collected from a single species.







Extended Data Fig. 6 | Meta-analysis of chronological age in mammalian samples across specific tissue types. Meta-analysis p-value (-log base 10 transformed) versus chromosomal location (x-axis) according to human genome assembly 38 (hg38) in (a), brain tissues (across multiple brain regions), (b) cerebral cortex, (c) blood, (d) liver, (e) muscle and (f) skin tissues. The upper and lower panels of the Manhattan plot depict the CpG sites that gain/ lose methylation with age. In panel a, P values were calculated via two-stage meta-analysis that combined EWAS results across strata formed by species/ brain-tissue (with n ≥ 15 samples, Methods). CpGs are colored in red and blue if

400-

they exhibit highly significant positive and negative age correlations according to a meta analysis P < 1.0×10^{-40} , 1.0×10^{-30} , 1.0×10^{-250} , 1.0×10^{-50} , 1.0×10^{-20} and 1.0×10^{-150} for a – f, respectively. Red dashed horizontal lines denote Bonferroni correction. Gene names are annotated for the top 20 CpGs with positive and negative associations, respectively. CpGs are labeled by adjacent genes. Purple color and diamond shapes mark CpGs of particular interest: cg12841266 and cg11084334 in *LHFPL4* and cg09710440 in *LHFPL3*. All P-values presented in this figure are unadjusted and computed using two-sided tests.

		Inc	creas	sed w	/ith a	ge			De	crea	sed v	vith a	age		
GapArtf1 (20)		1	1	12.2	1	15.7	19.7		1	!	!	1	1	!	1
Quies1 (360)	0	0	0.1	0				5.8	3.4	6.4				0.1	
Quies2 (265)	0	0	0	0				5.3	2.8	4.9					
Quies3 (625)	0	0	0	0			0.1	2.1	2.1	2.4			0.1	0.2	
HET1 (71)							1	5.3		7.5		1			
HET2 (206)										3.1					
HET7 (37)		0.5	0.0	4	0	2.0	5.0	7.1	0.0	7.1	0.00	0	0.0	0.0	-
ReprPC1 (1356)	8.2	9.5	6.6	4	2	2.9 2.5	5.2 2.5	0.2	0.2	0.2	0.03	0	0.3 0.2	0.3	
ReprPC2 (444)	0	0	0	0		2.5	2.5	0.2				1	0.2	0.2	
ReprPC4 (605)	0	0	0	0	1	1	1	2.5	1	2.8	1	0.2	0.00	0.2	
ReprPC5 (366)	0	0.1	Ŭ	0				2.0		2.0		0.2	0	0	
ReprPC6 (433)	Ō	0	0	0.2								1	0.08	0.08	
ReprPC7 (422)	0	0	0	0.09								1			
ReprPC8 (110)										3.7					
Acet1 (76)		1	1	1	1	1	1			6.2	1	1	1	1	
Acet2 (263)	0	0	1	0	1	1	1	4.3	2.7	4.3		1			
Acet3 (221)		l	l		1		1	2.7	4.1	3.3	1	1	1		
Acet5 (108)							1	6.4	5.9	4.2					-
EnhWk2 (266)	0	0	0	0.00								4.4	0.0		
EnhWk3 (603)	0	0	0.06	0.06	i	1	1	2.3	2.3	2.6		0.5	0.2	0.2	signed -log10P
EnhWk4 (1563) EnhWk8 (642)	0	0	0.04 0	0 0.06				2.3	2.3	2.6		0.5 0.06	0.2 0.06	0.2	6 00
EnnVVK8 (642) EnhA1 (281)	0	0	0	0.06			_	2.5	3.1	۷.۱		3.3	3.1	0.2	400
EnhA2 (712)	0	0	0.1	0.1				2.3	3.4		2.9	5.5	5.1		200
EnhA3 (243)		U	0.1	0.1				2.0	4.6		2.0		4.1		0
EnhA4 (132)									5.5						_ -200
EnhA5 (255)	0	0	0	0				3.5	3.5	3	3.5				
EnhA6 (683)	0	0	0.05	0.1				2.1							ChromHMM/PRC
EnhA9 (122)												4.8			group
EnhA12 (85)									4.3						others
EnhA16 (95)									5.3						quescient HET
EnhA17 (1088)	0.07	0	0.1	0.3	1.9	1.9			1.8			0.4	0.2	0.1	polycomb repressed
EnhA18 (353)	0	0	0		4							0		0.1	acetylations
EnhA19 (343) TxEnh3 (181)	0.1	0			4									3.8	weak enhancers
TxEnh4 (804)	0	0		0.04		0.3	0.2		0.3	0.4	2.1	5	7.2	8.2	enhancers
TxEnh5 (756)	0	0	0.1	0.04		0.0	0.2		0.0	0.4	2.1	0	1.2	2.1	transcribed and enhancer
TxEnh6 (256)	0		0												weak transcription
TxEnh7 (193)											2.9				transcription
TxEnh8 (288)	0	0					0				3.7		3.7	3.5	exon weak promoters
TxWk1 (617)	0	0	0.06	0									0.2	0.3	promoters
TxWk2 (772)	0	0.05	0.05	0			0.2	2.1		2.2					TSS
Tx4 (272)	0	0	0	0		0	0								PRC1&2 TFs binding regions
Tx5 (449)	0	0	0	0.2	0.2	0.07	0								
Tx6 (525)	0	0 0.04	0 0.04	0 0	0.2 0.3	0.07 0.1	0 0.3	0.2	0.3	0.3			2		
Tx7 (837) Tx8 (494)	0	0.04	0.04	0	0.5	0.1	0.5	0.2	0.5	0.5			2		
TxEx1 (1029)	0	0	0.2	0	0.4	0.2	0.07			0.4	1.9	2.7	5	5.3	
TxEx2 (1511)	0	0	0.02	0.05	0.4	0.2	0.07		0.2	0.4	1.0	2.1	3.2	3.3	
TxEx3 (862)	0	0	0.1	0.2	0.2	0.3	0.2		0.3	011			0.1	2	
TxEx4 (320)	0	0			0	0	0	3	2.6		4.1	6.7	14.5	12.9	
BivProm1 (1502)	9	7.6	7.9	10.1	1.8	2.6	4	0	0	0	0.05	0.1	0.05	0.1	
BivProm2 (1770)	15.7	16.5	13.7	10.3	2.3	3.1	5	0.04	0.04	0	0.02	0.06	0.06	0	
BivProm3 (930)							1.8	0.3			0.4	0.4	0.3	0.3	
BivProm4 (905)		0	6			1.9	2.3				0.5	1.9	0.1	0.5	-
PromF2 (382)	0	0	0	0.00		0.09	0.09	0.0		0.0	2.5	3.1	3.4	2.5	
PromF3 (474)	0	0.08	0	0.08	0	0.08	0.2	0.2	0	0.2			0	0	
PromF4 (1083)	0 2.3	0 2.2	0 1.9	0.03	0	0	0	0	0 0.03	0 0.06	0.2	0.4	0.09	0.09	
PromF5 (1184)	2.3	2.2	1.9	3.1				0	0.03	0.00	4.9	0.4	4.9	4.9	
TSS1 (829)	0.1	0.09	0.2	0.09	0.2	0.09	0.04	0	0.04	0.04	т.э		0.09	0.04	
TSS2 (655)	0.1	0.00	0.2	2.8	0.2	0.00	0.04	0.1	0.04	0.04		0.05	0.03	0.04	
PRC1 (640)	1.4	1.4	1.4	1.4	1.2	1.2	1.5	0.06	0.3	0.00	0.6	0.6	0.6	0.5	
PRC2 (5287)	22.8	29.8	14.2	14.3	2	3.2	6.5	0.02	0.04	0.03	0.1	0.2	0.04	0.03	
0.5	All	Blood	Skin	Liver	Muscle		Cortex	All	Blood	Skin	Liver	Muscle	Brain	Cortex	
PRC2 binding	propotio	ו													
			-	-											

Polycomb repressive complex & ChromHMM state annotation

 $\label{eq:constraint} Extended \, Data \, Fig. \, 7 \, | \, See \, next \, page \, for \, caption.$

Extended Data Fig. 7 | **Chromatin state analysis of age-related CpGs.** The heatmap color-codes the hypergeometric overlap analysis between age-related CpGs (columns) and two groupings of CpGs (1) universal chromatin states analysis¹ and (2) binding by polycomb repressive complex 1 and 2 (PRC1, PRC2) defined based on ChIP-Seq datasets in ENCODE⁵³, see the last two rows. The first column shows a bar plot that reports the proportion of CpGs that are known to be bounded by PRC2 that ranges from zero to one (PRC2). Note that chromatin states that contain a high proportion of PRC2 bound CpGs overlap significantly with the top 1,000 CpGs that increased with age across tissues and mammal species. For each row (chromatin state or PRC annotation), the table reports odds ratios (OR) from hypergeometric test results for the top 1,000 CpGs that increased with age across all, blood,

skin, liver, muscle, brain and cerebral cortex tissues, respectively. Unadjusted hypergeometric P values based on one-sided are listed in Supplementary Data 8.3–8.9. The heatmap color gradient is based on –log10 (unadjusted hypergeometric P value) multiplied by the sign of OR greater than one. Red colors denote OR greater than one in contrast with blue colors for OR less than one. Legend lists states based on their group category and PRC group. The y-axis lists state or PRC name and number of mammalian array CpGs inside parentheses. The left/right panel lists the results based on the top 1,000 CpGs with positive/ negative age correlation. We displayed 63 universal chromatin states that show significant enrichment/depletion at *P* < 0.001 in any of the tissues. HET, heterochromatin; exon, transcription and exons; weak promoters, bivalent promoters; promoters, promoter flanking.

				Increa	ased wit	h age					Decre	ased wi	th age		
Hg19 (13589)	commonHMD	0.2 (8.3e-95)	0.2 (8.5e-85)	0.2 (5.5e-66)	0.3 (1.3e-42)	1.1 (0.11)	0.6 (4.4e-14)	0.4 (8.7e-29)	0.3 (1.6e-43)	0.4 (5.5e-31)	0.3 (3.6e-64)	1.6 (2.6e-13)	2.3 (1.8e-39)	3.3 (6.6e-76)	2.9 (1.4e-61)
mm10 (10593)	comme	0.2 (2e-68)	0.2 (5.7e-73)	0.3 (1.2e-46)	0.4 (4.5e-31)	0.9 (9e-2)	0.6 (8.5e-14)	0.4 (1.1e-30)	0.3 (4e-49)	0.3 (9.3e-44)	0.2 (6.4e-66)	1.4 (8.8e-9)	2.2 (6.6e-36)	3.3 (4e-76)	3.1 (5.5e-68)
Hg19 (12527)	commonPMD	1.8 (2e-21)	1.8 (2.6e-18)	2 (4.5e-25)	1.4 (3.2e-7)	0.9 (0.2)	1.2 (9e-4)	1.3 (3.3e-5)	3.8 (1.3e-92)	3 (3.1e-63)	4.5 (2.4e-115)	0.9 (2.5e-2)	0.4 (4.8e-31)	0.2 (2.1e-62)	0.3 (7.7e-56)
mm10 (2615)	commo	1.3 (9.4e-3)	1.4 (2e-3)	1.6 (7.6e-6)	1.5 (1.9e-5)	1 (0.38)	1.4 (1.4e-3)	1.5 (4.5e-5)	3.2 (1.2e-40)	2.5 (1.2e-24)	3.2 (3.9e-41)	1.2 (5.5e-2)	0.3 (1.5e-14)	0.3 (4.7e-13)	0.4 (4e-12)
Hg19 (10087)	Neither	2.2 (8.4e-32)	2.2 (1.8e-31)	1.7 (4e-16)	2 (2.7e-26)	1.1 (0.12)	1.6 (6.7e-12)	1.9 (9.3e-21)	0.5 (2.5e-14)	0.7 (1e-5)	0.5 (2.6e-21)	0.7 (1e-5)	0.9 (0.12)	0.8 (9.7e-5)	0.8 (7.1e-3)
mm10 (15306)	Nei	3.3 (2.1e-64)	3.3 (1.9e-65)	2.4 (4.7e-38)	2 (1.1e-26)	1.2 (6.2e-4)	1.6 (1.6e-14)	2 (5.4e-27)	1.4 (1.3e-8)	1.7 (4.8e-17)	1.5 (6.3e-9)	0.7 (2.5e-6)	0.6 (6.7e-12)	0.4 (9.1e-45)	0.4 (5.9e-42)
Hg19 (1286)	solo-WCGW ommonPMDS	0.03 (1.6e-14)	0.03 (1.6e-14)	0.08 (3.8e-12)	0 (4.3e-16)	0.7 (8.1e-2)	1.2 (0.18)	0.9 (0.26)	7.3 (1e-81)	5.3 (1.3e-50)	7.9 (1.6e-90)	2.8 (2.7e-15)	0.7 (2.4e-2)	0.3 (6.6e-6)	0.2 (4.1e-8)
mm10 (281)	solo-V commo	0 (6.2e-5)	0 (6.2e-5)	0.1 (6.7e-4)	0.1 (6.7e-4)	0.1 (6.7e-4)	1.5 (9.6e-2)	1.2 (0.35)	8.6 (1.2e-32)	5.4 (1.1e-16)	9.1 (1.7e-35)	3.1 (1.2e-6)	0.3 (1.4e-2)	0.2 (3.7e-3)	0.4 (3.8e-2)
Hg19 (3031)	solo-WCGW	0.07 (6e-29)	0.07 (6e-29)	0.08 (7.8e-28)	0.1 (8.9e-26)	0.8 (1.8e-2)	1 (0.4)	0.7 (9.4e-3)	4.4 (1.6e-69)	3.7 (4.3e-53)	4.2 (4.3e-65)	2.6 (2.7e-24)	1.3 (2.1e-2)	1.1 (0.13)	0.9 (0.19)
mm10 (2089)	solo-V	0.1 (1.9e-22)	0.1 (1.9e-22)	0.1 (9.3e-20)	0.1 (9.3e-20)	0.8 (4.8e-2)	1.2 (8.5e-2)	0.9 (0.27)	4.1 (2.6e-55)	3.3 (3.5e-38)	3.9 (4.4e-51)	2.6 (2.9e-21)	1.1 (0.3)	1 (0.46)	0.9 (0.13)
_		All	Blood	Skin	Liver	Muscle	Brain	Cortex gned -log		Blood	Skin	Liver	Muscle	Brain	Cortex

Late-replicating domain annotation

signed -log10P

-100 -50 0 50 100 150

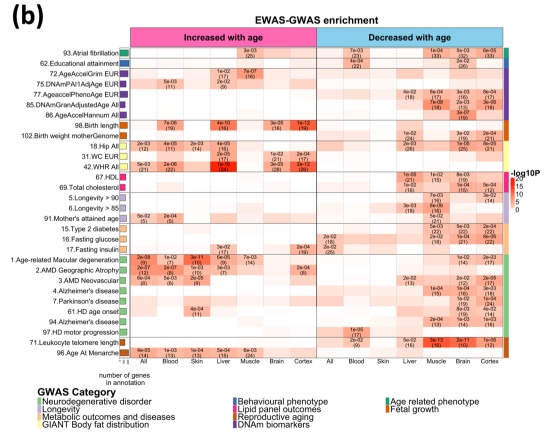
Extended Data Fig. 8 | **Overlap with late-replicating domains.** The heatmap color-codes the hypergeometric overlap analysis between age-related CpGs (columns) and CpGs related to late-replicating domains in hg19 and mm10 assembly⁵⁰, respectively. Two groups of late-replicating domains were analyzed (1) common PMD/HMD structures: highly methylated domains (commonHMD), partially methylated domains (commonPMD), and neither (Neither), and (2) solo-WCGW structures: genome-wide (solo-WCGW) and those in the common PMD regions (solo-WCGW commonPMDs). The y-axis lists categories of late-replicating domains and number of mammalian array CpGs inside parentheses

for Hg19 and mm10 genome, respectively. For each row, the table reports odds ratios (OR) from hypergeometric test results for the top 1,000 CpGs that increased/decreased with age from meta-EWAS of age across all, blood, skin, liver, muscle, brain, and cerebral cortex tissues, respectively. The heatmap color gradient is based on -log10 (unadjusted hypergeometric P value) multiplied by the sign of OR greater than one. Red colors denote OR greater than one in contrast with blue colors for OR less than one. The left/right panel lists the results based on the top 1,000 CpGs with positive/negative age correlation. Unadjusted P values are reported and derived from one-sided hypergeometric tests.

		Ir	ncreas	sed w	ith ag	e	Decreased with age									
CJUN (671)		 							2.6e-9 (2.8)				2.4e-2 (1.5)			
CTCF (3125)	8.2e-5 (1.5)	8.9e-6 (1.6)	1.1e-2 (1.3)	1.4e-5 (1.5)										3.3e-2 (1.2)		
EGR1 (1423)	6e-6 (1.9)		4.2e-3 (1.5)	2e-9 (2.2)		2.7e-3 (1.5)	1.6e-6 (1.9)									
GATA2 (74)		1.4e-2 (3.2)										1.4e-2 (3.2)	2.7e-5 (5.7)	1.4e-2 (3.2)		
HDAC2 (427)	1.6e-20 (5.4)	2.8e-14 (4.3)	6.2e-15 (4.5)	1.2e-17 (4.9)	5.7e-4 (2.2)	3.9e-2 (1.6)	2.8e-3 (2)									
INI1 (753)	1.4e-6 (2.3)	3.2e-6 (2.3)		7.9e-12 (3)			6.4e-4 (1.9)									
MAX (2856)		1		6.7e-5 (1.5)		1					1					
NANOG (116)	2.8e-4 (3.9)		1.3e-2 (2.7)	8.4e-8 (5.9)	1.1e-3 (3.5)	1.3e-2 (2.7)										
REST (637)	3.1e-54 (8.2)	2.7e-32 (5.8)	6.8e-59 (8.7)	1.5e-28 (5.4)	2.2e-3 (1.8)		1.3e-2 (1.6)									
RAD21 (2613)	7.8e-9 (1.8)	2.8e-7 (1.7)	2.5e-6 (1.7)	3.9e-11 (2)												
SIN3AK20 (944)	3e-8 (2.4)	5.4e-4 (1.8)	9.2e-6 (2)	6.3e-14 (3)												
SP1 (1164)	3.6e-13 (2.7)	2.4e-6 (2)	1.9e-11 (2.6)	7e-17 (3)			2.4e-4 (1.7)									
TAF1 (3254)	5.1e-7 (1.6)	3.7e-6 (1.6)	2.3e-3 (1.4)	8.5e-7 (1.6)												
TCF12 (374)	2.2e-16 (5)	2.7e-14 (4.6)	1.5e-18 (5.5)	4.3e-17 (5.2)	3.7e-7 (3.1)	4.5e-2 (1.6)	8.1e-5 (2.5)									
USF1 (979)	3.5e-13 (2.9)	2.9e-12 (2.8)	2.6e-6 (2.1)	1.3e-16 (3.2)		3.4e-4 (1.8)	6.3e-11 (2.7)									
YY1 (2533)	8.9e-16 (2.3)	1.3e-9 (1.9)	1e-14 (2.2)	5.5e-20 (2.5)			1.8e-2 (1.3)									
number of genes	All in annotation	Blood	Skin	Liver	Muscle	Brain	Cortex	All	Blood	Skin	Liver	Muscle	Brain	Cortex		

Extended Data Fig. 9 | **Enrichment with Transcription factor binding regions.** We studied the overlapping genomic regions between (1) the CpG sites located in the binding regions of 68 transcription factors (TF) in hg19 and (2) the top 1000 CpGs that increased/decreased with age from EWAS of age across mammalian tissues. TF results (y-axis, rows) versus mammalian EWAS of age are stratified by tissue type (x-axis, columns). The left/right panels of the x-axis list the top 1000 CpGs that increased/decreased with age from meta-EWAS of age across all tissues, blood only, skin only, liver, muscle, brain and cerebral cortex, respectively. The y-axis lists the names of transcription factors and number of mammalian array CpGs located in the binding sites. Background in hypergeometric tests was based on the genes present in our mammalian array. The bar plots in the first column report the total number of genes at each TF according to the background. The heatmap color codes -log10 (unadjusted hypergeometric P value). Unadjusted, one-sided hypergeometric P values (odds ratio) are listed on the heatmap provided *P* < 0.05.

(a)					E٧	VAS-	TWAS	6 enri	chme	nt				
(a)			Increa	ised w	ith age				0	Decrea	ased w	vith ag	e	
603.Human liposarcoma 41vs70 yr	1e-04	3e-04 (17)	3e-04	4e-04			1e-02 (17)			4e-03				
362.Human liposarcoma 45vs75 yr	(19) 2e-04 (20)	5e-04 (18)	(19) 4e-03 (18)	(20) 9e-04 (21)			(17)	3e-04 (28)	3e-02	(22) 1e-05				
350.Human brain (frontal cortex) 55vs82 yr	7e-04 (22)	1e-03 (20)	9e-04 (23)	1e-02 (21)				9e-03 (28)	(20) 5e-02 (23)	(31) 7e-03 (28)			3e-02 (25)	1e-03 (31)
712.Human monocyte 55 to 94 yr	(22)	(20)	(23)	(21)				(20)	(23)	(20)	4e-04 (58)	3e-04 (54)	2e-10 (70)	1e-06 (62)
733.Human nerve tibial GTEx	2e-04 (22)	1e-02 (16)	4e-05 (25)	1e-05 (28)	7e-04 (30)			1e-04 (32)	1e-02 (24)	2e-05 (34)	(50)	(54)	(70)	(02)
242.Monkey hipp. TACs 4-6 to 18-21 yr	7e-05	1e-02	8e-08	1e-05	2e-05			9e-15	4e-10	5e-15	1e-02			3e-02
249.Monkey hipp. microglia 4-6 to 18-21 yr	(40) 4e-04 (21)	(30)	(50) 6e-05 (24)	(48) 4e-03 (21)	(61) 2e-04 (31)			(85) 5e-14 (50)	(70) 7e-08 (37)	(85) 1e-11 (46)	(53) 2e-02 (26)		3e-02 (23)	(47) 4e-02 (23)
430.Mouse striatum 6vs21 mon	4e-05	3e-04	9e-06	1e-05	1e-03			(00)	(57)	(40)	(20)		(23)	(23)
752.Mouse diaphragm skeletal mus. satellite cell 3 to 24 mon	(21)	(18)	(23)	(25)	(26)						3e-02	6e-05	7e-13	6e-08
788.Mouse marrow late pro-b cell 3 to 24 mon											(97) 4e-02 (90)	(102) 2e-05 (99)	(128) 3e-12 (121)	(111) 2e-10 (114)
771.Mouse limb mus. skeletal mus. satellite cell 3 to 24 mon											1e-03 (126)	3e-07 (130)	6e-11 (142)	8e-06 (121)
568.Mouse hipp. 3vs16 mon								4e-02 (25)			(120)	2e-04 (33)	2e-10 (46)	3e-08 (41)
749.Mouse brain non myeloid oligodendrocyte 3 to 24 mon								(20)				2e-02 (85)	3e-10 (118)	1e-06 (104)
790.Mouse marrow naive b cell 3 to 24 mon											4e-02 (71)	1e-05 (81)	1e-09 (93)	1e-08 (89)
811.Mouse trachea endothelial cell 3 to 24 mon											(71) 2e-03 (86)	3e-04 (82)	2e-09 (100)	1e-06 (89)
794.Mouse pancreas pancreatic b cell 3 to 24 mon											(86) 4e-03 (66)	1e-03 (62)	2e-09	1e-06 (72)
791.Mouse marrow precursor b cell 3 to 24 mon											(00)	5e-06 (77)	(81) 2e-09 (87)	2e-08 (83)
455.Mouse retina 3vs16 mon											4e-03 (32)	2e-03 (30)	3e-07 (40)	6e-08 (41)
789.Mouse marrow macrophage 3 to 24 mon											1e-02 (31)	2e-03 (31)	4e-08 (43)	2e-07 (41)
559.Mouse brain 4vs22 mon											(0.)	(0.)	5e-09 (36)	5e-07 (32)
769.Mouse limb mus. endothelial cell 3 to 24 mon											9e-06 (74)	1e-05 (67)	2e-08 (75)	4e-06 (67)
773.Mouse liver hepatocyte 3 to 24 mon											1e-04		4e-03 (36)	1e-02 (34)
579.Mouse skeletal mus. precursor 12vs24 mon											(46) 2e-04 (26)	6e-03 (20)	2e-05 (26)	4e-03 (20)
382.Mouse liver 5vs24 mon											2e-04 (15)	(20)	(20)	(20)
586.Mouse neocortex 5vs30 mon											(10)	1e-09 (37)	1e-03 (24)	1e-03 (24)
510.Mouse gastrocnemius 5vs30 mon											4e-02 (20)	3e-08 (33)	1e-02 (20)	3e-02 (18)
774.Mouse lung adventitial cell 3 to 24 mon											(20) 9e-03 (64)	3e-07 (75)	6e-03 (58)	4e-02 (52)
202.Rat brain excitatory neurons 5vs27 mon								2e-07 (12)	4e-04 (8)	1e-05 (10)			1e-02 (6)	1e-02 (6)
219.Rat typell mus. fibers 5vs27 mon					4e-02 (22)			1e-02 (23)	8e-03 (22)	4e-02 (21)	3e-03 (27)	4e-07 (34)	(0)	(0)
428.Rat extensordigitorumlongusmus.s 6vs30 mon					(=2)			(20)	(=2)	(= 1)	(21)	(34) 5e-06 (28)		
0.001	All	Blood	Skin	Liver	Muscle			All	Blood	Skin	Liver	Muscle	Brain	Cortex
Annotatio	on size					AS Ca uman	Monk	ey 🛛 N	louse	Rat				



Extended Data Fig. 10 | See next page for caption.

Extended Data Fig. 10 | EWAS-TWAS and EWAS-GWAS enrichment. Panel (a) illustrates the overlap between genes identified in transcriptome-wide association studies (TWAS) across various cell types or species, and the top 1,000 CpGs that have increased/decreased with age in EWAS across mammalian tissues. TWAS results are stratified by tissue type, including all tissues, blood, skin, liver, muscle, brain, and cerebral cortex. Overlapping genes with P < 0.05 are reported. Similarly, Panel (b) demonstrates the overlaps between the top 2.5% genes implicated in genome-wide association studies (GWAS) of human complex traits, and the top 1,000 CpGs that have increased/decreased with age in EWAS across mammalian tissues. GWAS results are also stratified by tissue type, with significant overlaps reported where P < 0.05. Both panels utilize unadjusted, one-sided hypergeometric P values, with a background for hypergeometric tests derived from genes (panel a) or genomic regions (panel b) in our mammalian array. The heatmap color encodes -log10 P values. The right-side annotation indicates (a) the species categories for TWAS collections and (b) phenotype categories for GWAS collections. Further details for TWAS and GWAS indices are available in Supplementary Data 12 & 13. Abbreviations: (a) hipp.=hippocampus, MPNST = malignant peripheral nerve sheath tumor, mus.=muscle, TACs = transiently amplifying progenitor cells. (b) All = All ancestries, EUR = European ancestry, AFR = African American ancestry, FTD = frontotemporal dementia, WHR = waist to hip ratio.

nature portfolio

Corresponding author(s): Steve Horvath Last updated by author(s): $_{6/01/2023}$

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our <u>Editorial Policies</u> and the <u>Editorial Policy Checklist</u>.

Statistics

For	all st	atistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Cor	firmed
	\boxtimes	The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
		A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	\boxtimes	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
		A description of all covariates tested
		A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
		A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
		For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable</i> .
\boxtimes		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
		Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
		Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information	about availability of computer code
Data collection	Collections of Mammalian samples are described in Supplementary Information Note 1. Mammalian samples are maintained as Excel spread sheets or Rdata generated in R.
Data analysis	Matlab_2017b: MAGENTA analysis for yielding gene-level p values based on GWAS SNP associations in EWAS-GWAS analysis Python 3 10.3: package appdata 0.8.0 for managing human single cell ATAC (sc-ATAC) array data

- R_4.0.2: Programming language for statistical computing
- R_sesame_1.3.0: Normalize Illumina Infinium DNA methylation array data
- R_glmnet_4.1-7 Fit penalized generalized linear models
- R_WGCNA_1.69: Weighted correlation network analysis for analysis and graphics.
- R_rGREAT_1.22.0: Enrichment analysis using Genomic Regions Enrichment of Annotations Tool
- $\ensuremath{\mathsf{R}_g}\xspace$ was vcf_0.1.1: Manage GWAS summary datasets in VCF format
- R_Signac: Perform quality controls (R_Signac/CreateChromatinAssay function) and data management for mouse sc-ATAC analysis
- R_GenomicRanges: Overlap locations between peak calling locations from sc-ATAC and probe locations in our Mammalian array.
- R_ggplot2, R_ComplexHeatmap_2.6.2 and R_gmirror: for figures including box plots, heat maps, and Manhattan plots.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Policy information about availability of data

- All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
 - Accession codes, unique identifiers, or web links for publicly available datasets
 - A description of any restrictions on data availability
 - For clinical datasets or third party data, please ensure that the statement adheres to our policy

The data for all species from the Mammalian Methylation Consortium can be downloaded from Gene Expression Omnibus (GEO) using the accession number GSE223748. To facilitate comparative analyses across species, the consortium applied a single measurement platform (the mammalian methylation array, GPL28271) to n=15,216 DNA samples derived from 70 tissue types of 348 different mammalian species (331 eutherian-, 15 marsupial-, and 2 monotreme species). The 11,754 samples used for training our universal clocks are part of the samples, which are available for age information.

Subset of the data can be accessed from multiple online locations. First, the data are available from https://clockfoundation.org/

MammalianMethylationConsortium. Second the data can be downloaded from GEO using the following accession numbers: GSE174758, GSE184211, GSE184213, GSE184215, GSE184216, GSE184218, GSE184220, GSE184221, GSE184224, GSE190660, GSE190661, GSE190662, GSE190663, GSE190664, GSE174544, GSE190665, GSE174767, GSE184222, GSE184223, GSE174777, GSE174778, GSE173330, GSE164127, GSE147002, GSE147003, GSE147004. Additional details can be found in Supplementary information note 2. The mammalian methylation array is available through the non-profit Epigenetic Clock Development Foundation (https:// clockfoundation.org/).

Human research participants

Policy information about studies involving human research participants and Sex and Gender in Research.

Reporting on sex and gender	The Sex/Gender variable in the Framingham Heart Study (FHS) Cohort, https://www.framinghamheartstudy.org/index.php, is based on self-report. The FHS data are available in dbGaP (accession number: phs000363.v16.p10 and phs000724.v2.p9). The Women's Health Initiative (WHI) is a national study that enrolled postmenopausal women aged 50-79 years into the clinical trials (CT) or observational study (OS) cohorts between 1993 and 1998.
Population characteristics	We applied our universal Clocks 2 and 3 on 4,651 individuals from (a) the Framingham Heart Study (FHS) offspring cohort (n=2,544 Caucasians, 54% females)and (b) Women's Health Initiative cohort (WHI, n=2107, 100% woman, Supplementary Information, note 4). Methylation levels were profiled in blood samples using Illumina 450k arrays. The FHS cohort had a mean (SD) age of 66.3 (8.9) years at blood draw, with 330 deaths during an average follow-up of 7.8 years. The WHI cohort, which enrolled postmenopausal women aged 50-79 years, consisted of three ethnic groups: 47% of European ancestry (Caucasians), 32% African Americans, and 20% of Hispanic ancestry. These groups exhibited similar age distributions, with a mean (SD) age of 65.4 (7.1) years, and a mean (SD) follow-up time of 16.9 (4.6) years. During the follow-up, 765 women died.
Recruitment	We did not recruit any human beings in this study. Rather, we used existing data. The FHS cohort is a large-scale longitudinal study initiated in 1948, originally designed to explore the common factors and characteristics contributing to cardiovascular disease (CVD) (https://www.framinghamheartstudy.org/index.php). This study initially enrolled participants from the town of Framingham, Massachusetts, who were asymptomatic for overt CVD, heart attack, or stroke at the time of enrollment. In 1971, the FHS Offspring Cohort was established to encompass a second generation of participants, specifically the adult children and their spouses of the original cohort (n=5124), for similar examinations. Participants from the FHS Offspring Cohort were included in our study if they had attended the 8th examination cycle and consented to the use of their molecular data for research purposes. We utilized data from 2,544 participants with available DNA methylation profiles (measured at exam 8), obtained from the group providing Health/Medical/Biomedical consent (IRB, MDS). The FHS data can be accessed through the db GaP (accession numbers: phs000363.v16.p10 and phs000724.v2.p9). The Women's Health Initiative, a national landmark study, recruited postmenopausal women aged between 50-79 years into the clinical trials (CT) or observational study (OS) cohorts between 1993 and 199866,67. We incorporated data from 2,017 WHI participants from "Broad Agency Award 23" (WHI BA23) which had available phenotype and DNA methylation array data. WHI BA23 is dedicated to the identification of miRNA and genomic biomarkers of coronary heart disease (CHD), with an aim to integrate these biomarkers into diagnostic and prognostic predictors of CHD and associated phenotypes. This study encompasses three WHI sub-cohorts, namely GARNET, WHIMS, and SHARE.
Ethics oversight	The Framingham Heart Study is funded by National Institutes of Health contract N01-HC-25195 and HHSN268201500001I. The laboratory work for this investigation was funded by the Division of Intramural Research, National Heart, Lung, and Blood Institute, National Institutes of Health. The analytical component of this project was funded by the Division of Intramural Research, National Heart, Lung, and Blood Institute, and the Center for Information Technology, National Institutes of Health, Bethesda, MD.

The Women's Health Initiative program is funded by the National Heart, Lung, and Blood Institute, National Institutes of Health, U.S. Department of Health and Human Services through contracts HHSN268201600018C, HHSN268201600001C, HHSN268201600002C, HHSN268201600003C, and HHSN268201600004C. The authors thank the WHI investigators and staff for their dedication, and the study participants for making the program possible. A full listing of WHI investigators can be found at: http://www.whi.org/researchers/Documents%20%20Write%20a%20Paper/WHI%20Investigator%20Long% 20List.pdf. The views expressed in this manuscript are those of the authors and do not necessarily represent the views of funding bodies such as the National Heart, Lung, and Blood Institute; the National Institutes of Health; or the U.S. Department of Health and Human Services.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences

Behavioural & social sciences 🛛 🛛 Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>

Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description	Observational data based on existing samples stored in freezers. We generated 11,754 methylation arrays from over 57 tissue-types derived from 185 mammalian species. We aimed to profile animals from the entire age range: from very young to very old. Roughly uniform distribution. We only analyzed tissues from animals whose ages were known with 90% confidence.
Research sample	I. We employed a custom methylation array (HorvathMammalMethylChip40) that profiles methylation levels of 36k CpGs with flanking DNA sequences that are highly-conserved across the mammalian class. II. We obtained such profiles from 11,754 samples from 59 tissue types, derived from 185 mammalian species, representing 19 taxonomic orders and ranging in age from prenatal to 139 years old (bowhead whale).
	III. The tissue samples are described in the Supplement and related citations as listed in Supplementary Information, Note 1. IV. To enhance the reproducibility of our findings we include our updated version of the animal age (anAge) database, which is reported in the supplementary data.
	V. Below are the list of ipapers that describe specific species:
	1. Horvath, S. et al. Pan-primate DNA methylation clocks. bioRxiv, 2020.11.29.402891 (2021).
	2. Horvath, S. et al. Epigenetic clock and methylation studies in the rhesus macaque. GeroScience (2021).
	3. Jasinska, A.J. et al. Epigenetic clock and methylation studies in vervet monkeys. GeroScience (2021).
	4. Horvath, S. et al. DNA methylation age analysis of rapamycin in common marmosets. GeroScience (2021).
	5. Schlabritz-Loutsevitch, N.E. et al. Metabolic adjustments to moderate maternal nutrient restriction. British journal of nutrition 98, 276-284 (2007).
	6. Kavitha, J.V. et al. Down-regulation of placental mTOR, insulin/IGF-I signaling, and nutrient transporters in response to maternal nutrient restriction in the baboon. FASEB journal : official publication of the Federation of American Societies for Experimental Biology 28, 1294-1305 (2014).
	7. Schlabritz-Loutsevitch, N.E. et al. Development of a system for individual feeding of baboons maintained in an outdoor group social environment. Journal of Medical Primatology 33, 117-126 (2004).
	8. Zehr, S.M. et al. Life history profiles for 27 strepsirrhine primate taxa generated using captive data from the Duke Lemur Center. Scientific Data 1, 140019 (2014).
	9. Morgello, S. et al. The National NeuroAIDS Tissue Consortium: a new paradigm in brain banking with an emphasis on infectious disease. Neuropathol Appl Neurobiol 27, 326-35. (2001).
	10. Horvath, S. et al. HIV, pathology and epigenetic age acceleration in different human tissues. Geroscience (2022). 11. Horvath, S. et al. Perinatally acquired HIV infection accelerates epigenetic aging in South African adolescents. AIDS (London, England) 32, 1465-1474 (2018).
	12. Horvath, S. & Ritz, B.R. Increased epigenetic age and granulocyte counts in the blood of Parkinson's disease patients. Aging (Albany NY) 7, 1130-42 (2015).
	13. Kabacik, S., Horvath, S., Cohen, H. & Raj, K. Epigenetic ageing is distinct from senescence-mediated ageing and is not prevented by telomerase expression. Aging (Albany NY) 10, 2800-2815 (2018).
	14. Ross, C.N. et al. The development of a specific pathogen free (SPF) barrier colony of marmosets (Callithrix jacchus) for aging research. Aging (Albany NY) 9, 2544 (2017).
	15. Sailer, L.L. et al. Pair bonding slows epigenetic aging and alters methylation in brains of prairie voles. bioRxiv, 2020.09.25.313775 (2020).
	16. Ophir, A.G. Navigating Monogamy: Nonapeptide Sensitivity in a Memory Neural Circuit May Shape Social Behavior and Mating Decisions. Frontiers in Neuroscience 11(2017).
	 Horvath, S. et al. Methylation studies in Peromyscus: aging, altitude adaptation, and monogamy. GeroScience 44, 447-461 (2022). Horvath, S. et al. DNA methylation aging and transcriptomic studies in horses. Nat Commun 13, 40 (2022).
	19. Burns, E.N. et al. Generation of an equine biobank to be used for Functional Annotation of Animal Genomes project. Animal genetics 49, 564-570 (2018).
	20. Horvath, S. et al. DNA methylation clocks tick in naked mole rats but queens age more slowly than nonbreeders. Nature Aging 2, 46-59 (2022).
	21. Ke, Z., Vaidya, A., Ascher, J., Seluanov, A. & Gorbunova, V. Novel husbandry techniques support survival of naked mole rat

(Heterocephalus glaber) pups. J Am Assoc Lab Anim Sci 53, 89-91 (2014).

22. Tan, L. et al. Naked Mole Rat Cells Have a Stable Epigenome that Resists iPSC Reprogramming. Stem cell reports 9, 1721-1734 (2017).

23. Sugrue, V.J. et al. Castration delays epigenetic aging and feminizes DNA methylation at androgen-regulated loci. eLife 10, e64932 (2021).

24. Schachtschneider, K.M. et al. Epigenetic clock and DNA methylation analysis of porcine models of aging and obesity. GeroScience (2021).

25. Robeck, T.R. et al. Multi-Tissue Methylation Clocks for Age and Sex Estimation in the Common Bottlenose Dolphin. Frontiers in Marine Science 8(2021).

26. Robeck, T.R. et al. Multi-species and multi-tissue methylation clocks for age estimation in toothed whales and dolphins. Commun Biol 4, 642 (2021).

27. Bors, E.K. et al. An epigenetic clock to estimate the age of living beluga whales. Evolutionary Applications (2020).

28. Raj, K. et al. Epigenetic clock and methylation studies in cats. GeroScience (2021).

29. Prado, N.A. et al. Epigenetic clock and methylation studies in elephants. Aging Cell 20, e13414 (2021).

30. Pinho, G.M. et al. Hibernation slows epigenetic ageing in yellow-bellied marmots. Nature Ecology & Evolution 6, 418-426 (2022). 31. Lemaître, J.-F. et al. DNA methylation as a tool to explore ageing in wild roe deer populations. Molecular Ecology Resources n/ a(2021).

32. Larison, B. et al. Epigenetic models developed for plains zebras predict age in domestic horses and endangered equids. Communications Biology 4, 1412 (2021).

33. Harley, E.H., Knight, M.H., Lardner, C., Wooding, B. & Gregor, M. The Quagga project: progress over 20 years of selective breeding. African Journal of Wildlife Research 39, 155-163 (2009).

34. Horvath, S. et al. Reversing age: dual species measurement of epigenetic age with a single clock. bioRxiv, 2020.05.07.082917 (2020).

35. Horvath, S. et al. Epigenetic clock and methylation studies in dogs. PNAS In Press(2022).

36. Plassais, J. et al. Whole genome sequencing of canids reveals genomic regions under selection and variants influencing morphology. Nature Communications 10, 1489 (2019).

37. Plassais, J. et al. Analysis of large versus small dogs reveals three genes on the canine X chromosome associated with body weight, muscling and back fat thickness. PLOS Genetics 13, e1006661 (2017).

38. The American Kennel Club. The Complete Dog Book: 20th Edition, (Howell Book House, New York, NY, 2006).

39. Wilcox, B. & Walkowicz, C. The Atlas of Dog Breeds of the World, (T.F.H. Publications, 1995).

40. Wilkinson, G.S. et al. DNA methylation predicts age and provides insight into exceptional longevity of bats. Nature Communications 12, 1615 (2021).

41. Kordowitzki, P. et al. Epigenetic clock and methylation study of oocytes from a bovine model of reproductive aging. Aging Cell 20, e13349 (2021).

42. Mozhui, K. et al. Genetic loci and metabolic states associated with murine epigenetic aging. eLife 11, e75244 (2022). 43. Lu, A.T. et al. DNA methylation study of Huntington's disease and motor progression in patients and in animal models. Nature Communications 11, 4529 (2020).

44. Coschigano, K. et al. Deletion, but not antagonism, of the mouse growth hormone receptor results in severely decreased body weights, insulin, and insulin-like growth factor I levels and increased life span. Endocrinology 144, 3799-3810 (2003).

45. Acosta-Rodríguez, V.A., Rijo-Ferreira, F., Green, C.B. & Takahashi, J.S. Importance of circadian timing for aging and longevity. Nature Communications 12, 2862 (2021).

46. Little, T.J. et al. Methylation-Based Age Estimation in a Wild Mouse. bioRxiv, 2020.07.16.203687 (2020).

47. Cossette, M.-L. et al. Differential methylation, epigenetic clocks, and island-mainland divergence in an insectivorous small mammal. bioRxiv, 2022.04.14.488253 (2022).

48. Horvath, S. et al. Epigenetic clock and methylation studies in marsupials: opossums, Tasmanian devils, kangaroos, and wallabies. Geroscience In Press(2022).

49. Hogg, C.J., Lee, A.V. & Hibbard, C.J. Managing a metapopulation: intensive to wild and all the places in between. in Saving the Tasmanian Devil: recovery through science based management 169-182 (CSIRO Publishing Melbourne, 2019).

50. Hogg, C. & Hockley, J. DPIPWE/ZAA husbandry guidelines for Tasmanian devil, Sarcophilus harrisii. Australia: Zoo and Aquarium Association (2013).

51. Sambrook, J. & Russell, D.W. Purification of nucleic acids by extraction with phenol: chloroform. Cold Spring Harbor Protocols 2006, pdb. prot4455 (2006).

52. Villar, D. et al. Enhancer evolution across 20 mammalian species. Cell 160, 554-66 (2015).

53. Berthelot, C., Villar, D., Horvath, J.E., Odom, D.T. & Flicek, P. Complexity and conservation of regulatory landscapes underlie evolutionary resilience of mammalian gene expression. Nat Ecol Evol 2, 152-163 (2018).

54. Roller, M. et al. LINE retrotransposons characterize mammalian tissue-specific and evolutionarily dynamic regulatory regions. Genome Biol 22, 62 (2021).

55. Yan, L. et al. OSAT: a tool for sample-to-batch allocations in genomics experiments. BMC Genomics 13, 689 (2012).

56. Seluanov, A. et al. Hypersensitivity to contact inhibition provides a clue to cancer resistance of naked mole-rat. Proceedings of the National Academy of Sciences 106, 19352-19357 (2009).

57. Seluanov, A. et al. Telomerase activity coevolves with body mass not lifespan. Aging Cell 6, 45-52 (2007).

Sampling strategy

We sampled all mammalian species for which existing tissues were available. These fresh frozen tissue samples were contributed by a large network of investigators from our Mammalian Methylation Consortium.

To guide the quality control (QC) of the study samples, we generated two variables ; the first being a variable indicating the confidence (0 to 100%) in the chronological age estimate of the sample. For example, a low confidence was assigned to samples from wild animals whose ages were estimated based on body length measurements. The epigenetic clocks were trained and evaluated in tissue samples whose confidence exceeded 90% (>=90%). The second quality control variable was an indicator variable (yes/no) that flagged technical outliers or malignant (cancer) tissue. Since we were interested in "normal" aging patterns we excluded tissues from preclinical studies surrounding anti-aging or pro-aging interventions.

Data collection

DNA for methylation profiling was extracted from the tissue samples collected from different species as described in Supplementary Information, Note 1. After bisulfite conversion and labeling of the DNA, methylation profiles were obtained by hybridizing labeled

	Neuroscience Genomics Core.
Timing and spatial scale	The tissue samples were collected over the last 30 years. The data come from many labs all over the world: US, Canada, Europe, Australia, New Zealand, South America.
Data exclusions	We excluded about 1900 samples that had insufficient DNA to provide reliable methylation values, low confidence in the chronological age estimate or unknown age. We discovered that samples with concentrations below 6 ng/µl could not be accurately scored at all sites on the array.
Reproducibility	 We used calibration data (synthetic DNA) to evaluate the accuracy of the methylation measurements (A mammalian methylation array for profiling methylation levels at conserved sequences by A. Arneson 2021 Nat Comm) We performed EWAS meta-analysis of age using Stouffer's method estimates from Metal algorithm. In addition, we verified the Stouffer's statistics in our in-house R code. The universal clocks were established via elastic net regression models. To assess the accuracies of our clocks, we used 3 approaches: leave-one-fraction-out (LOFO), leave one-species-out (LOSO) cross validation, and data splitting. In LOFO, we randomly split the entire dataset into 10 fractions each of which had the same distribution in species and tissue types. Each penalized regression model was trained in 9 fractions but evaluated in the 10th left out fraction. After circling through the 10 fractions, we arrived at LOFO predictions which were subsequently related to the actual values. The LOSO cross validation approach trained each model on all but one species. The left out species was used a test set. The LOSO approach was used to assess how well the penalized regression models generalize to species that were not part of the training data. To ensure unbiased estimates of accuracy, all aspects of the model fitting were only conducted in the training data in both LOFO and LOSO analysis. The reported EWAS p values are significant even after using the most stringent multiple comparison correction (Bonferroni)=0.05/37K based on 37K CpGs on the mammalian array. In GREAT enrichment analysis, we performed two different sensitivity analyses that were inspired by our GREAT enrichment analysis of the top 1 thousand age related CpGs (EWAS of age). The results are listed in Supplementary Info, Note 2. Our first sensitivity analysis involved a random set of 1000 CpG mammalian CpGs. Second, we evaluated the enrichment of the top 1087 most highly conserved CpGs across
Randomization	Not applicable since this is an observational study.

DNA to a custom Illumina methylation array (HorvathMammalMethylChip40) and scanning with an Illumina iScan at the UCLA

Nanuomization			
Blinding	Blinding was not relevant to our study, because this is an observational study and all available data were used		
Did the study involve fiel	d work? Yes XNo		

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems		thods	
n/a Involved in the study	n/a	Involved in the study	
Antibodies	\boxtimes	ChIP-seq	
Eukaryotic cell lines	\boxtimes	Flow cytometry	
Palaeontology and archaeology	\square	MRI-based neuroimaging	
Animals and other organisms	,		
🛛 🗌 Clinical data			
Dual use research of concern			

Animals and other research organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research, and <u>Sex and Gender in</u> Research

Laboratory animals

Details in Supplementary Data S1.2. This study leveraged existing tissue samples or data that had been collected as part of other studies. We profiled tissues from lab animals or animals kept in captivity for research. This includes mouse, rat, opossum, naked mole rat, deer mouse colonies, rhesus macaque, marmosets, vervet monkey. Companion pets: dogs, cats. Agricultural animals: horses, pigs, sheep.

Wild animals	Details in Supplementary Data S1.2. Some samples from zoo animals (e.g. elephants). Samples from zoo-based animals were opportunistically collected and banked during routine health exams. This study also includes samples from wild animals, which were collected in the field: bats, deer.
Reporting on sex	As the contributors are engaged in long-term field studies, care was taken to minimize disturbance during all the sample collections for different species. The details are described in species-specific papers.
Field-collected samples	As the contributors are engaged in long-term field studies, care was taken to minimize disturbance during all the sample collections for different species. The details are described in species-specific papers.
Ethics oversight	Institutional animal care and use protocols, or equivalent information from non-US contributors, is provided in Supplementary Information describing the data and also in the underlying species-specific papers published by Mammalian Methylation Consortium. Non university organizations are certified either by the Associated Zoos and Aquariums (Lubee Bat Conservancy) or by the Global Federation of Animal Sanctuaries (Bat World Sanctuary) or Elephant Taxon Advisory Group and Species Survival Plan. Details in appendix.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Ethics Oversight for Mammalian Methylation Consortium Studies

M1. Primates ¹ Ethics

This research complied with all relevant ethical regulations overseen by seven ethics review boards. The human skin samples were acquired with informed consent prior to collection of human skin samples with approved by the Oxford Research Ethics Committee in the UK; reference 10/H0605/1. Participants were not compensated. The secondary use of the other de-identified/coded human tissue samples (blood, postmortem tissues) is not interpreted as human subjects research under U.S. Department of Health & Human Services 45 CFR 46. Therefore, the need to obtain written, informed consent from human study participants was waived (secondary use of de-identified tissues). Human samples were covered by University of California Los Angeles IRB#18-000315. All procedures related to non-human primates were approved by different committees: baboons (UTHSCSA Animal Care and Use Committee), strepsirrhini (Duke Institutional Animal Care and Use Committee and the DLC Research Committee), rhesus macaques (Animal Care and Use Committee of the NIA Intramural Program)², vervet monkey (UCLA and VA Institutional Animal Care and Use Committees)³, marmosets (IACUC of UTHSA)⁴.

M2. Prairie voles ⁵ Ethics

All experimental procedures were conducted and approved by the Institutional Animal Care and Use Committee (IACUC) of Cornell University (2013-0102) and were in accordance with the guidelines set forth by the National Institutes of Health.

M3. Horses ⁶ Ethics

This collection protocol was approved by the UC Davis Institutional Animal Care and Use Committee (Protocol#19037). All collection protocols were approved by the UC Davis Institutional Animal Care and Use Committee (Protocols #20751 and 21455, respectively).

M4. Naked mole-rat ⁷

The NMR tissue samples were provided by two different labs: (i) Vera Gorbunova and Andrei Seluanov from the University of Rochester and (ii) Chris Faulkes from the Queen Mary, University of London.

Ethics for (i)

All animal experiments were approved and performed in accordance with guidelines set up by the University of Rochester Committee on Animal Resources with protocol number 2009-054 (naked mole-rat).

Ethics for (ii)

Naked mole-rats were maintained in the Biological Services Unit at Queen Mary University of London in accordance with UK Government Animal Testing and Research

Guidance.

M5. Sheep ⁸

Sheep DNA samples for this study were derived from two distinct tissues from two strains: ear tissue from New Zealand Merino, and blood from South Australian Merino.

Ethics for ear samples

Ear tissue was obtained from females and both intact and castrated male Merino sheep during routine on-farm ear tagging procedures in Central Otago, New Zealand. As a small piece of tissue is removed during the ear tagging process that is usually discarded by the farmer, we were able to source tissue and record the year of birth without altering animal experience, in accordance with the New Zealand Animal Welfare Act (1999) and the National Animal Ethics Advisory Committee (NAEAC) Occasional Paper No 2 [26].

Ethics for blood samples

All protocols involving OVT73 sheep were approved by the Primary Industries and Regions South Australia (PIRSA, Approval number 19/02) Animal Ethics Committee with oversight from the University of Auckland Animal Ethics Committee.

M6. Pig ⁹ <u>Ethics</u>

All animal procedures were approved by the University of Illinois and University of

Wisconsin Institutional Animal Care and Use Committee, and all animals received

humane care according to the criteria outlined in the Guide for the Care and Use of

Laboratory Animals.

M7. Odontocete species ¹⁰⁻¹² <u>Ethics</u>

The study was authorized by the management of each institution and was reviewed by their respective zoo research and animal use committees.

M8. Beluga whales ¹² <u>Ethics</u>

Skin tissue samples were collected from carcasses of beluga whales that were beachcast, stranded dead, or taken during subsistence hunting from 1992 to 2015 in Cook Inlet, Alaska, USA (NMFS Research Permit 932-1905-00/MA-009526 through the Marine Mammal Health and Stranding Response Program).

M9. Killer whales and bowhead whales <u>Ethics</u>

For bowhead (*Balaena mysticetus*) subsistence hunts, indigenous hunters had the authorization to conduct hunts and collected samples on behalf of Fisheries and Oceans Canada. Bowhead whale biopsy samples were collected in 2019 under Fisheries and Oceans Canada (DFO) license to Fish for Scientific Purposes (LFSP) S-19/20-1007-NU and Animal Care approval (AUP) FWI-ACC-2019-14. Skin samples from eastern North Pacific killer whales (*Orcinus orca*) were collected as previously described (Ford et al. 2018b) under NMFS General Authorization No. 781–1725, and scientific research permits 781-1824-01, 16163, 532- 1822-00, 532– 1822, 10045, 18786-03, 545-1488, 545-1761, and 15616.

M10. Humpback whales <u>Ethics</u>

Skin samples were collected by the Center for Coastal Studies under research permits issued by the U.S., National Marine Fisheries Service (21485, 16325, 20465, 14245, 633-1483, 633-1778, 932-1905), the Canadian Department of Fisheries and Oceans and IACUC #NWAK-18-02.

M11. Cats ¹³ Ethics

Sample collection was approved by the Clinical Research Ethical Review Board of the

RVC (URN: 2019 1947-2).

M12. Elephants ¹⁴ Ethics

This study was authorized by the management of each participating zoo and, where applicable, was reviewed and approved by zoo research committees. In addition, the

study received IACUC approval (#18-29) at the NZP; and endorsement from the elephant Taxon Advisory Group and Species Survival Plan.

M13. Yellow-bellied marmots ¹⁵ <u>Ethics</u>

Data and samples were collected under the UCLA Institutional Animal Care and Use protocol (2001-191-01, renewed annually) and with permission from the Colorado Parks and Wildlife (TR917, renewed annually).

M14. Roe deer ¹⁶ Ethics

The protocol of capture and blood sampling under the authority of the Office Français de la Biodiversité (OFB) was approved by the Director of Food, Agriculture and Forest (Prefectoral order 2009–14 from Paris). The land manager of both sites, the Office National des Forêts (ONF), permitted the study of the populations (Partnership Convention ONCFS-ONF dated 2005-12-23). All experiments were performed in accordance with guidelines and regulations of the Ethical Committee of Lyon 1 University (project DR2014-09, June 5, 2014)

M15. Zebras ¹⁷ Ethics

Plains zebra samples were collected under a protocol approved by the Research Safety and Animal Welfare Administration, University of California Los Angeles: ARC # 2009-090-31, originally approved in 2009.

M16 Rat 18

The rat tissues came from 4 different labs across three countries:(i) India: Nugenics Research in collaboration with School of Pharmacy SVKM's NMIMS University (K. Singh), (ii) United States: University of Tennessee Health Science Center (H. Chen) and Medical College of Wisconsin (L.C. Solberg Woods), and (iii) Argentina: University of La Plata (R. Goya).

Ethics for (i)

The experimental protocols received approval from the Institutional Animal Ethics Committee under two distinct approval numbers. The first is CPCSEA/IAEC/P-75/2018, and the second is CPCSEA/IAEC/P-6/2018. The second approval was granted in accordance with the norms of the Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA), Government of India, complying with the standard guidelines for handling experimental animals. It specifically pertains to the use of male Sprague Dawley rats, aged 8 weeks (200–250 g) and 20 months (400-450g), procured from the National Institute of Bioscience, Pune, India.

Ethics for (ii)

All procedures were approved by the Institutional Animal Care and Use Committee of the University of Tennessee Health Science Center or the Medical College of Wisconsin and followed the NIH Guide for the Care and Use of Laboratory Animals.

Ethics for (iii)

All experiments with animals were performed in accordance with the Animal Welfare Guidelines of NIH (INIBIOLP's Animal Welfare Assurance No A5647-01) and approved by our Institutional IACUC (Protocol # P05-02-2017). Ethics committee approval number - CPCSEA/IAEC/P-6/2018.

M17. Dog 19

For this study, DNA samples were collected from a total of 742 blood samples taken from dogs across 93 different breeds. These samples were generously provided by researchers at the National Human Genome Research Institute (NHGRI). Unfortunately, we did not have access to individual weight data for these dogs.

Ethics

The collection of these samples was conducted in compliance with ethical guidelines and was officially approved by the Animal Care and Use Committee of the Intramural Program of NHGRI at the National Institutes of Health (Protocol #8329254).

M18. Bats ²⁰ Ethics

The study was approved by the University of Maryland Institutional Animal Care and Use Committee (FR-APR-18-16).

M19. Cattle ²¹ Ethics

All animal procedures were carried out in accordance with the relevant guidelines at each institution. Specifically, procedures related to sample collection in Poland followed the EU Directive of the European Parliament and the Council on the protection of animals used for scientific purposes (22 September 2010; No 2010/63/EU), Polish Parliament Act on Animal Protection (21 August 1997, Dz.U. 1997 nr 111 poz. 724) with further novelization - Polish Parliament Act on the protection of animals used for scientific or educational purposes (15 January 2015, Dz.U. 2015 poz. 266). Blood and oocyte collection were approved by the Local Ethics Committee for Experiments on Animals, University of Warmia and Mazury in Olsztyn, Poland (Agreement No. LKE.065.27.2019). For animal procedures in the USA, approval from the University of Nebraska Institutional Animal Care and Use Committee was obtained (approval number is 1560).

M20. Mouse data ²²

The mouse data were sourced from various institutions or studies, as outlined below: (i) UCLA Lab. Animal breeding and husbandry ²³, (ii) BXD mice ²² from University of Tennessee Health Science Center, (iii) Growth hormone receptor knockout from the University of Michigan, (iv) Calorie restricted mice from the University of Texas Southwestern Medical Center ²⁴, (v) South African species, (vi) Apodemus mice²⁵, and (vii) Spiny mouse.

Ethics for (i)

All mice were maintained and bred under standard conditions consistent with National Institutes of Health guidelines and approved by the University of California, Los Angeles Institutional Animal Care and Use Committees.

Ethics for (ii)

All animal procedures were in accordance with the protocol approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Tennessee Health Science Center.

Ethics for (iv)

The Institutional Animal Care and Use Committee (IACUC) of the University of Texas Southwestern Medical Center approved the animal protocol (APN 2015-100925), which has been subsequently renewed every 3 years (2018 and 2021).

Ethics for (v)

The Animal Use and Care Committee of the University of Pretoria evaluated and approved the experimental protocol and collection of all samples (ethics clearance number: NAS022/2021, NAS209/2021, NAS021/2020), with DAFF section 20 approval (SDAH-Epi-21051907211, SDAH-Epi-12/11/1/1/8 (2002 LH), SDAH-Epi-20072707050).

In addition, permission to capture the various species was obtained from all landowners, and a collecting permit was obtained from the relevant nature conservation authorities (Permit number: Western Cape- CN44-87-13780, CN44-31-2285, Gauteng- CPF6-0124, Kwa-Zulu Natal- OP1545/2021). Necessary TOPS permits were also acquired for threatened species (Permit number: 68103).

Ethics for (vi)

All animal work was conducted in accordance with the UK Home Office in compliance with the Animals (Scientific Procedures) Act 1986, was approved by the University of Edinburgh Ethical Review Committee and was carried out under the approved UK Home Office Project License PP4913586.

Ethics for (vii)

Animal protocols were approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Kentucky (2019-3254).

M21. Nova Scotia masked shrews (*Sorex cinereus*) ²⁶ Shrews and small animals from Museum of Biological Diversity at The Ohio State

<u>University</u>

All wild-caught animals were collected and sacrificed in accordance with protocols approved by The Ohio State University IACUC (Institutional Animal Care and Use Committee) under protocol number 2017A00000036. All wild-caught animals were collected with scientific collecting permits issued from Ohio and Washington and according to guidelines established by the American Society of Mammalogy for the use of wild animals in research (Sikes & Animal Care and Use Committee of the American Society of Mammalogists 2016).

M22. Marsupials and mice ²⁷ Ethics

These procedures are in accordance with the AVMA Guidelines for the Euthanasia of Animals 2013: https://www.avma.org/KB/Policies/Documents/euthanasia.pdf, and all animal procedures were approved by the UCLA IACUC.

M23. Mammalian liver samples (Diego Villar Lozano and Duncan Odom) <u>Ethics</u>

The use of all animals in this study was approved by the Animal Welfare and Ethics Review Board under reference number NRWF-DO-02vs and followed the Cancer Research UK Cambridge Institute guidelines for the use of animals in experimental studies. Tissue samples from humans were obtained from Addenbrooke's Hospital at the University of Cambridge under license number 08-H0308-117, specifically for the study "Liver specific transcriptional regulation."

M24. Mammalian liver samples from the University of Rochester <u>Ethics</u>

All experiments were performed according to procedures approved by the University of Rochester Committee on Animal Resources (UCAR), under animal protocol #101939 / UCAR-2017-033. The tissues used in the study were obtained from the Gorbunova and Seluanov tissue bank at the University of Rochester.^{28,29}

Murine anti-aging studies <u>Ethics</u>

Experiments involving Snell, growth hormone receptor knockout (GHRKO), and liverspecific GHRKO mouse strains (PI: Richard Miller) were conducted at the University of Michigan. These experiments were approved by the University of Michigan's Institutional Animal Care and Use Committee.

Reference:

- 1. Horvath, S. *et al.* Pan-primate DNA methylation clocks. *bioRxiv*, 2020.11.29.402891 (2021).
- 2. Horvath, S. *et al.* Epigenetic clock and methylation studies in the rhesus macaque. *GeroScience* (2021).
- 3. Jasinska, A.J. *et al.* Epigenetic clock and methylation studies in vervet monkeys. *GeroScience* (2021).
- 4. Horvath, S. *et al.* DNA methylation age analysis of rapamycin in common marmosets. *GeroScience* (2021).
- 5. Sailer, L.L. *et al.* Pair bonding slows epigenetic aging and alters methylation in brains of prairie voles. *bioRxiv*, 2020.09.25.313775 (2020).
- 6. Horvath, S. *et al.* DNA methylation aging and transcriptomic studies in horses. *Nat Commun* **13**, 40 (2022).
- 7. Horvath, S. *et al.* DNA methylation clocks tick in naked mole rats but queens age more slowly than nonbreeders. *Nature Aging* **2**, 46-59 (2022).
- 8. Sugrue, V.J. *et al.* Castration delays epigenetic aging and feminizes DNA methylation at androgen-regulated loci. *eLife* **10**, e64932 (2021).
- 9. Schachtschneider, K.M. *et al.* Epigenetic clock and DNA methylation analysis of porcine models of aging and obesity. *GeroScience* (2021).
- 10. Robeck, T.R. *et al.* Multi-Tissue Methylation Clocks for Age and Sex Estimation in the Common Bottlenose Dolphin. *Frontiers in Marine Science* **8**(2021).
- 11. Robeck, T.R. *et al.* Multi-species and multi-tissue methylation clocks for age estimation in toothed whales and dolphins. *Commun Biol* **4**, 642 (2021).
- 12. Bors, E.K. *et al.* An epigenetic clock to estimate the age of living beluga whales. *bioRxiv*, 2020.09.28.317610 (2020).
- 13. Raj, K. *et al.* Epigenetic clock and methylation studies in cats. *GeroScience* (2021).
- 14. Prado, N.A. *et al.* Epigenetic clock and methylation studies in elephants. *Aging Cell* **20**, e13414 (2021).
- 15. Pinho, G.M. *et al.* Hibernation slows epigenetic ageing in yellow-bellied marmots. *Nature Ecology & Evolution* **6**, 418-426 (2022).
- 16. Lemaître, J.-F. *et al.* DNA methylation as a tool to explore ageing in wild roe deer populations. *Molecular Ecology Resources* **n/a**(2021).
- 17. Larison, B. *et al.* Epigenetic models developed for plains zebras predict age in domestic horses and endangered equids. *Communications Biology* **4**, 1412 (2021).
- 18. Horvath, S. *et al.* Reversing age: dual species measurement of epigenetic age with a single clock. *bioRxiv*, 2020.05.07.082917 (2020).
- 19. Horvath, S. *et al.* DNA methylation clocks for dogs and humans. *Proceedings of the National Academy of Sciences* **119**, e2120887119 (2022).
- 20. Wilkinson, G.S. *et al.* DNA methylation predicts age and provides insight into exceptional longevity of bats. *Nature Communications* **12**, 1615 (2021).
- 21. Kordowitzki, P. *et al.* Epigenetic clock and methylation study of oocytes from a bovine model of reproductive aging. *Aging Cell* **20**, e13349 (2021).
- 22. Mozhui, K. *et al.* Genetic loci and metabolic states associated with murine epigenetic aging. *eLife* **11**, e75244 (2022).

- 23. Lu, A.T. *et al.* DNA methylation study of Huntington's disease and motor progression in patients and in animal models. *Nature communications* **11**, 1-15 (2020).
- 24. Acosta-Rodríguez, V.A., Rijo-Ferreira, F., Green, C.B. & Takahashi, J.S. Importance of circadian timing for aging and longevity. *Nature Communications* **12**, 2862 (2021).
- 25. Little, T.J. *et al.* Methylation-Based Age Estimation in a Wild Mouse. *bioRxiv*, 2020.07.16.203687 (2020).
- 26. Cossette, M.-L. *et al.* Differential methylation, epigenetic clocks, and islandmainland divergence in an insectivorous small mammal. *bioRxiv*, 2022.04.14.488253 (2022).
- 27. Horvath, S. *et al.* Epigenetic clock and methylation studies in marsupials: opossums, Tasmanian devils, kangaroos, and wallabies. *Geroscience* In **Press**(2022).
- 28. Seluanov, A. *et al.* Hypersensitivity to contact inhibition provides a clue to cancer resistance of naked mole-rat. *Proceedings of the National Academy of Sciences* **106**, 19352-19357 (2009).
- 29. Seluanov, A. *et al.* Telomerase activity coevolves with body mass not lifespan. *Aging Cell* **6**, 45-52 (2007).