

HHS Public Access

Author manuscript *Nature*. Author manuscript; available in PMC 2021 June 29.

Published in final edited form as:

Nature. 2020 July ; 583(7817): 590-595. doi:10.1038/s41586-020-2496-1.

A Single Cell Transcriptomic Atlas Characterizes Aging Tissues in the Mouse

The Tabula Muris Consortium^{*}

Abstract

Aging is characterized by a progressive loss of physiological integrity, leading to impaired function and increased vulnerability to death¹. Despite rapid advances over recent years, many of the molecular and cellular processes which underlie progressive loss of healthy physiology are poorly understood². To gain a better insight into these processes we have created a single cell transcriptomic atlas across the life span of Mus musculus which includes data from 23 tissues and organs. We discovered cell-specific changes occurring across multiple cell types and organs, as well as age related changes in the cellular composition of different organs. Using single-cell transcriptomic data we were able to assess cell type specific manifestations of different hallmarks of aging, such as senescence³, genomic instability⁴ and changes in the organism's immune system². This Tabula Muris Senis provides a wealth of new molecular information about how the most significant hallmarks of aging are reflected in a broad range of tissues and cell types.

We performed single cell RNA sequencing on more than 350,000 cells from male and female C57BL/6JN mice belonging to six age groups ranging from one month (human early childhood equivalent) to thirty months (human centenarian equivalent) (Figure 1a). We prepared single cell suspensions of the bladder, bone marrow, brain (cerebellum, cortex, hippocampus and striatum), fat (brown, gonadal, mesenteric and subcutaneous), heart and aorta, kidney, large intestine, limb muscle and diaphragm, liver, lung, mammary gland, pancreas, skin, spleen, thymus, tongue and trachea for all mice. Data were collected for all six age groups using microfluidic droplets (droplet), while the 3m, 18m and 24m time points were also analyzed using single cells sorted in microtiter well plates (FACS) (Extended Data

Reporting summary

- Ethics declarations
- Competing interests

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Data and code availability

The entire dataset can be explored interactively at tabula-muris-senis.ds.czbiohub.org. Gene counts and metadata are available from figshare (https://doi.org/10.6084/m9.figshare.8273102.v2) and GEO (GSE132042), the code used for the analysis is available from GitHub (https://github.com/czbiohub/tabula-muris-senis) and the raw data are available from a public AWS S3 bucket (https://s3.console.aws.amazon.com/s3/buckets/czb-tabula-muris-senis/).

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

The authors declare no competing interests.

Supplementary Information is available in the online version of the paper.

Figures 1-3; Supplementary Tables 1&2). Due to technical constraints, not every tissue was analyzed at all timepoints; a complete list is in Extended Data Figure 4a. The droplet data allow large numbers of cells to be analyzed using 3' end counting, while the FACS data allow for higher sensitivity measurements over smaller numbers of cells as well as sequence information across the entire transcript length. Analyzing multiple organs from the same animal enables data controlled for age, environment, and epigenetic effects.

The previously published 3m time point, referred to as the *Tabula Muris*⁵, represents $\sim 20\%$ of the cells in the entire dataset and was used as a basis to perform semi-automated cell type annotation of the additional time points (Figure 1b, Extended Data Figure 4b). Using this approach, we were able to automatically annotate over 70% of the cells. All the automated cell annotations were reviewed and approved by human experts, and the remaining cells were annotated by hand, creating one of the largest manually curated single cell transcriptomic resources in existence. Many of these cell types have not previously been obtained in pure populations, and these data provide a wealth of new information on their characteristic gene-expression profiles. Out of 529,823 total cells sequenced, 110,824 cells for FACS and 245,389 cells for droplet passed our strict filtering criteria (Extended Data Figure 4b) and were annotated (Extended Data Figure 2a,b), separately for each tissue and method; the remaining cells are also included in the on-line dataset but were not used for further analysis here. To investigate whether cell annotations were consistent across the entire organism, we used bbknn⁶ to correct for method-associated batch effects (Supplementary Table 3). Following batch correction, we clustered all cells using an unbiased, graph-based clustering approach^{7,8} (Figure 1c,d) and assessed the co-occurrence of similarly annotated cells in the same clusters. For example, cells annotated as B cells or endothelial cells tend to occupy the same clusters irrespectively of their tissue of origin or method with which they were processed (Figure 1e,f; Extended Data Figure 1g-l).

Tabula Muris Senis enables discovery of aging related changes in specific cell types. Single cell data enables one to resolve whether gene expression changes observed in bulk experiments are due to changes in gene expression in each cell of the population, or whether the gene expression in each cell stays constant but the number of cells of that type changes, or both. In a global analysis of gene expression changes using the Tabula Muris Senis and bulk RNAseq from tissues⁹, we observed that in many cases changes in gene expression are due to both changes in the numbers of cells in a population and to changes in the gene expression levels in each cell (Extended Data Figure 5a,b). As one specific example, we investigated how the fraction of cells expressing of Cdkn2a changes with age. Cdkn2a/p16 is one of the most commonly used markers of senescence¹⁰ and an important hallmark of aging¹¹, and the proportion of cells expressing the gene more than doubled in older animals in both FACS (Figure 2a) and droplet (Figure 2b), accompanied by a 2-fold increase in the actual expression level of p16 by those cells that did express it (Figure 2c,d). Interestingly, the fraction of cells expressing p16 in the 30m mice is smaller than at 24m, perhaps because long-living animals have a slower rate of senescence. Using a list of previously characterized senescence markers^{12–15} we plotted the fraction of cells expressing each marker across all age groups (Supplementary Table 4). Cdkn2a has the highest correlation between aging and fraction, and other genes with positive correlation include E2f2¹⁶, Lmnb1^{17,18} and Tnf and Itgax¹⁹. For some genes the fraction of cells expressing decreased with age, including

The cellular composition of each tissue tends to vary with age, and we investigated changes for tissues with at least three time points (Supplementary Table 5). Since dissociation does not affect all cell types in a tissue equally, changes in the relative composition of a given cell type with age are more meaningful than comparing proportions of different cell types at a single age^{22-24} . The bladder has pronounced changes in cell type composition with age (Figure 2e). While the mesenchymal compartment of this tissue decreases by a factor of three over the lifetime of the mouse (Figure 2e left), the urothelial compartment increases by a similar amount (Figure 2e right). The observation that the bladder urothelial cells increase with age is concordant with known age-related urothelial changes²⁵. Differential gene expression (DGE) analysis of overall tissue changes with age revealed that stromalassociated genes (Col1a1, Col1a2, Col3a1, Dcn) are downregulated while epithelialassociated genes (Krt15, Krt18, Sfn) are upregulated, supporting the compositional observations (Figure 2f; Supplementary Table 6). The decline of the endothelial population suggests that bladder aging in mice may be associated with lower organ vascularization, consistent with recent findings^{26,27} and with the observed downregulation of vasculature associated genes Htra1 and Fos (Figure 2f; Supplementary Table 6). The increase in the leukocyte population could indicate an inflammatory tissue microenvironment, a common hallmark of aging which is consistent with literature on overactive bladders²⁸ and supported by a significant overexpression of Lgals3, Igfbp2 and Ly6d across the tissue (Figure 2f; Supplementary Table 6) and by the overexpression of immune response associate genes such as Tnfrsf12a and Cdkn1a, by both bladder (mesenchymal) cells and bladder urothelial cells (Supplementary Table 6). Moreover, when comparing across ages, we observed that old leukocytes show increased expression of pro-inflammatory markers, such as Cd14, Lgals3 and Tnfrsf12a, and decreased expression of anti-inflammatory ones, such as Cd9 and Cd81 (Supplementary Table 6).

Age-dependent changes in the kidney include a decrease in the relative abundance of mesangial cells, capillary endothelial cells, loop of Henle ascending limb epithelial cells and loop of Henle thick ascending limb epithelial cells (Figure 2g). Both mesangial cells and capillary endothelial cells are core glomerular cells and their relative abundances reduction (Figure 2g top panels), together with a tissue-wide reduction of Egf and Atp1a1 expression (Figure 2h; Supplementary Table 6) suggest impaired glomerular filtration rate 29,30 . Interestingly, local Atp1a1 expression actually increases with age in both capillary endothelial cells and mesangial cells, suggesting that a compensation mechanism counteracts the effects of the cell proportion declining with age. This finding is reinforced by differential gene expression results indicating that uromodulin (Umod), the most abundant protein in urine³¹, is also reduced in expression across the tissue. Umod is produced by the epithelial cells that line the thick ascending limb, and therefore given the relative decrease in the proportion of epithelial cells in the ascending and thick ascending limb, our results suggest that normal kidney functions are impaired³² (Figure 2g bottom panels, Figure 2h; Supplementary Table 6). As with Atp1a1, we see that Umod expression increases in a cell type whose abundance decreases with age, leading to an overall reduction of Umod expression in the organ.

In spleen the proportion of T cells decreases with age while the relative amount of plasma cells increases (Figure 2i). This is supported by upregulation of B/plasma cell markers (Cd79a, Igj; Figure 2j; Supplementary Table 6) and downregulation of Cd3d (Figure 2j; Supplementary Table 6). Similarly, in mammary gland we observed a decline of the T cell population (Extended Data Figure 5c). Age-related decline of T cell populations has been associated with increased risk of infectious disease and cancer³³ and our results suggest this may also happen in spleen and mammary gland. Moreover, AP1 transcription factors³⁴ (Junb, Jund and Fos) were upregulated with age (Extended Data Figure 5d; Supplementary Table 6), consistent with the observation that normal involution of the mammary gland is accompanied by increased expression of this gene family³⁵.

The liver also displays changing tissue compositions with age, as the relative number of hepatocytes decreases with age (Extended Data Figure 6a-d), which is supported by the reduction in the expression of albumin (Alb; Extended Data Figure 6e; Supplementary Table 6). DGE showed an increased immune signature, as illustrated by overexpression of H2-Aa, H2-Ab1, H2-D1, H2-Eb1, Cd74, Lyz2 and others (Extended Data Figure 6e). Previous findings suggested that pro-inflammatory macrophages drive cellular senescence and identified II1b as a gene whose liver expression was remarkably different with age¹² (Extended Data Figure 6f). We stained liver Kupffer cells (Extended Data Figure 6g) with Clec4f and found the number of Clec4f+ cells does not change with age, consistent with the results of the tissue composition analysis (Supplementary Table 7; Extended Data Figure 6h). However, when co-staining with Il1b, we found an increase with age in the number of cells expressing Clec4f and Il1b (Extended Data Figure 6h-j). Il1b has low expression in normal physiological conditions³⁶. Specific blocking of IL1-RI in hepatocytes has been shown to attenuate cell death upon injury, supporting the idea that increased expression of Il1b in Kupffer cells is typically a poor prognostic³⁷. Regarding immune defense within the liver, sinusoidal endothelial cells (LSECs) play a unique role, being the main carriers of the mannose receptor (Mrc1) in the liver³⁸ (Extended Data Figure 6k). Our findings identified increased Mrc1 age-related expression in Kupffer cells, while the overall expression of Mrc1 in liver endothelial cells was reduced with age (Supplementary Table 6). By performing in situ RNA staining for Mrc1 alongside classical LSEC marker Pecam1 (Supplementary Table 7; Extended Data Figure 61), we found that the number of Mrc1 expressing LSECs increased with age (Extended Data Figure 6m-o). While the Mrc1 expression did not increase with age in LSECs (Supplementary Table 6), the overall number of cells expressing Mrc1 did increase significantly with age (Extended Figure 6n). LSECs have been found to have a reduced endocytic capacity in aged livers, while it has been suggested that LSECs proliferate after injury or that bone-marrow derived LSECs progenitors are recruited to the liver. This suggests that changes in LSEC gene signatures with age are linked closely with their function in immune response.

Genomic instability is among the most widely studied aging hallmarks¹ and the full-length transcript data enables analysis of somatic mutation accumulation with age. We used the Genome Analysis ToolKit (GATK)³⁹ to perform SNP discovery across all FACS samples simultaneously (Supplementary Table 8)^{40,41}. We focused on genes expressed in at least 75% of cells for each age group within a particular tissue and observed an age-related increase in the number of mutations across all of the organs we analyzed (Figure 3;

genome stability.

Extended Data Figure 7a,c,e), with tongue and bladder being the most affected. We controlled for sequencing coverage and gene expression levels (Extended Data Figure 8a,c,e), and verified that the number of mutations exceeded technical errors due to amplification and sequencing errors, which can be estimated using ERCC controls that were spiked into each well⁴² (Figure 3; Extended Data Figure 7b,d,f; Extended Data Figure 8b,d,f). Despite the fact that it is difficult to infer absolute genome-wide mutation rates from the transcriptome, which is known to inflate apparent mutational rates for a variety of reasons⁴², the observed trend is a useful indirect estimate of mutational frequency and

Aging also affects the immune system², and we analyzed clonal relationships between Bcells and T-cells throughout the organism. We computationally reconstructed the sequence of the B-cell receptor (BCR) and T-cell receptor (TCR) for B and T cells present in the FACS data using singlecell-ige and TraCeR, respectively^{43,44}. BCRs were assembled for 6,050 cells (Figure 4a, Extended Data Figure 9a) and TCRs for 6,000 cells (Figure 4b, Extended Data Figure 9a). The number of cells with assembled BCRs was 1,818 for 3m, 1,356 for 18m and 2,876 for 24m old mice. We parsed the singlecell-ige⁴³ output to define B-cell clonotypes based on the sequence of the assembled BCR (Supplementary Table 9) and found that while most of the cells at 3m were not part of a clone (9% were part of a clonal family), the number of B-cells belonging to a clonotype doubled at 18m (20%) when compared to 3m and doubled again from 18m to 24m (~38%). The number of cells with assembled TCRs were roughly equal between 3m, 18m and 24m (2,076, 2,056 and 1,868 cells, respectively). Clonotype assignment is part of the output obtained by TraCeR⁴⁴ (Supplementary Table 9). Interestingly, only ~3% (55 out of 1,895) of the cells at 3m were part of a clone. For 18m and 24m, ~23% (479 out of 2,056) and ~20% (348 out of 1,780) of the cells, respectively, were part of a clone, indicating again an increase in clonality of the T-cell repertoire at later ages. These changes in clonality for both B and T cell repertoires are noteworthy because they suggest that the immune system of a 24m mouse is less likely to respond to new pathogens, corroborating literature suggesting that older individuals have higher vulnerability to new infections and lower benefits from vacination^{45,46}.

Finally, we computed an overall diversity score to identify which cell types were more susceptible to changes with age (Extended Data Figure 10). The diversity score is computed as the Shannon entropy of the cluster assignment and then regressed against age to provide a p-value (see Methods). We observed significant changes in diversity affecting cells of the immune system originating from the brain and the kidney (Figure 4c, Extended Data Figure 11a,b). These results were not confounded by the number of genes expressed per cell (Extended Data Figure 11c,d). In brain myeloid microglial cells, the majority of young (3m) microglia occupy clusters 1 and 6, while old (18m, 24m) microglia constitute the vast majority of cells in clusters 10, 12 and 14 (Figure 4d). Trajectory analysis suggests that young microglia go through an intermediate state, represented by the clusters mostly occupied by 18m microglial cells before acquiring the signature of old microglia (Extended Data Figure 11e). Clusters 10, 12 and 14 are mainly comprised of 18- and 24-month old microglia. These cells up-regulate MHC class I genes (H2-D1, H2-K1, B2m), along with genes associated with degenerative disease (e.g. Fth1)^{47,48}. When contrasting with clusters 1 and 6, which contain mostly 3m microglia, clusters 10, 12 and 14 gene expression is

enriched with interferon responsive or regulatory genes (e.g. Oasl2, Oasla, Ifit3, Rtp4, Bst2, Stat1, Irf7, Ifitm3, Usp18, Ifi204, Ifit2), suggesting an expansion of this small proinflammatory subset of microglia in the aging brain⁴⁹. Moreover, the list of differentially expressed genes between "young" and "old" clusters resembled the Alzheimer's disease specific microglial signature previously reported⁴⁷, with 55 out of the top 200 differential expressed genes being shared between the two differential gene expression lists (Figure 4e; Supplementary Table 10). Regarding kidney macrophages, we found two clusters that remarkably changed their composition with age. Cluster 10 is primarily composed of cells of 1m- and 3-month old mice while cluster 13 is mostly composed of cells of 18-, 21-, 24- and 30-month old mice (Figure 4f). Differential gene expression revealed that cluster 10 is enriched for an M2-macrophage gene signature (e.g. II10, H2-Eb1, H2-Ab1, H2-Aa, Cd74, C1qa, Cxcl16, Hexb, Cd81, C1qb, Cd72) while cluster 13 resembles a M1-proinflammatory macrophage state⁵⁰ (e.g. Hp, Itgal, Spex1, Gngt2) (Extended Data Figure 11f; Supplementary Table 10).

The Tabula Muris Senis is a comprehensive resource for the cell biology community which offers a detailed molecular and cell-type specific portrait of aging. We view such cell atlas as an essential companion to the genome: the genome provides a blueprint for the organism but does not explain how genes are used in a cell type specific manner or how the usage of genes changes over the lifetime of the organism. The cell atlas provides a deep characterization of phenotype and physiology and serves as a reference for understanding many aspects of the cell biological changes that mammals undergo during their lifespan.

Methods

All data, protocols, analysis scripts and an interactive data browser are publicly available.

Experimental Procedures

Mice and organ collection

Male and virgin female C57BL/6JN mice were shipped from the National Institute on Aging colony at Charles River (housed at 67-73 °F) to the Veterinary Medical Unit (VMU; housed at 68–76 °F)) at the VA Palo Alto (VA). At both locations, mice were housed on a 12-h light/ dark cycle and provided food and water ad libitum. The diet at Charles River was NIH-31, and Teklad 2918 at the VA VMU. Littermates were not recorded or tracked, and mice were housed at the VA VMU for no longer than 2 weeks before euthanasia, with the exception of mice older than 18 months, which were housed at the VA VMU beginning at 18 months of age. Before tissue collection, mice were placed in sterile collection chambers at 8 am for 15 min to collect fresh fecal pellets. After anaesthetization with 2.5% v/v Avertin, mice were weighed, shaved, and blood was drawn via cardiac puncture before transcardial perfusion with 20 ml PBS. Mesenteric adipose tissue was then immediately collected to avoid exposure to the liver and pancreas perfusate, which negatively affects cell sorting. Isolating viable single cells from both the pancreas and the liver of the same mouse was not possible; therefore, two males and two females were used for each. Whole organs were then dissected in the following order: large intestine, spleen, thymus, trachea, tongue, brain, heart, lung, kidney, gonadal adipose tissue, bladder, diaphragm, limb muscle (tibialis anterior), skin

(dorsal), subcutaneous adipose tissue (inguinal pad), mammary glands (fat pads 2, 3 and 4), brown adipose tissue (interscapular pad), aorta and bone marrow (spine and limb bones). Organ collection concluded by 10 am. After single-cell dissociation as described below, cell suspensions were either used for FACS of individual cells into 384-well plates, or for preparation of the microfluidic droplet library. All animal care and procedures were carried out in accordance with institutional guidelines approved by the VA Palo Alto Committee on Animal Research.

Tissue dissociation and sample preparation

All tissues were processed as previously described⁵.

Sample size, randomization and blinding

No sample size choice was performed before the study. Randomization and blinding were not performed: the authors were aware of all data and metadata-related variables during the entire course of the study.

Single-cell methods

All protocols used in this study are described in detail elsewhere⁵. Those include: i) preparation of lysis plates, ii) FACS sorting, iii) cDNA synthesis using the Smart-seq2 protocol^{51,52}, iv) library preparation using an in-house version of Tn5^{53,54},v) library pooling and Quality control and vi) sequencing. For further details please refer to http://dx.doi.org/ 10.17504/protocols.io.2uwgexe

Microfluidic droplet single-cell analysis

Single cells were captured in droplet emulsions using the GemCode Single-Cell Instrument (10x Genomics) and scRNA-seq libraries were constructed as per the 10x Genomics protocol using GemCode Single-Cell 3' Gel Bead and Library V2 Kit. In brief, single cell suspensions were examined using an inverted microscope, and if sample quality was deemed satisfactory, the sample was diluted in PBS with 2% FBS to a concentration of 1000 cells per µl. If cell suspensions contained cell aggregates or debris, two additional washes in PBS with 2% FBS at 300g for 5 min at 4 °C were performed. Cell concentration was measured either with a Moxi GO II (Orflo Technologies) or a haemocytometer. Cells were loaded in each channel with a target output of 5,000 cells per sample. All reactions were performed in the Biorad C1000 Touch Thermal cycler with 96-Deep Well Reaction Module. 12 cycles were used for cDNA amplification and sample index PCR. Amplified cDNA and final libraries were evaluated on a Fragment Analyzer using a High Sensitivity NGS Analysis Kit (Advanced Analytical). The average fragment length of 10x cDNA libraries was quantitated on a Fragment Analyzer (AATI), and by qPCR with the Kapa Library Quantification kit for Illumina. Each library was diluted to 2 nM, and equal volumes of 16 libraries were pooled for each NovaSeq sequencing run. Pools were sequenced with 100 cycle run kits with 26 bases for Read 1, 8 bases for Index 1, and 90 bases for Read 2 (Illumina 20012862). A PhiX control library was spiked in at 0.2 to 1%. Libraries were sequenced on the NovaSeq 6000 Sequencing System (Illumina).

Computational methods

Data extraction

Sequences from the NovaSeq were de-multiplexed using bcl2fastq version 2.19.0.316. Reads were aligned using to the mm10plus genome using STAR version 2.5.2b with parameters TK. Gene counts were produced using HTSEQ version 0.6.1p1 with default parameters, except 'stranded' was set to 'false', and 'mode' was set to 'intersection-nonempty'. Sequences from the microfluidic droplet platform were de-multiplexed and aligned using CellRanger version 2.0.1, available from 10x Genomics with default parameters.

Data pre-processing

Gene count tables were combined with the metadata variables using the Scanpy⁵⁵ Python package version 1.4.2. We removed genes not expressed in at least 3 cells and then cells that did not have at least 250 detected genes. For FACS we removed cells with less than 5000 counts and for droplet cells with less than 2500 UMIs. The data was then normalized using size factor normalization such that every cell has 10,000 counts and log transformed. We computed highly variable genes using default parameters and then scaled the data to a maximum value of 10. After we computed PCA, neighborhood graph and clustered the data using Louvain⁷ and Leiden⁸ methods. The data was visualized using UMAP projection. When performing batch correction to remove the technical artifacts introduced by the technologies, we replaced the neighborhood graph computation with **bbknn**⁶. Step-by-step instructions to reproduce the pre-processing of the data are available from GitHub.

Cell type annotation

To define cell types we analyzed each organ independently but combining all ages. In a nutshell, we performed principal component analysis on the most variable genes between cells, followed by Louvain and Leiden graph-based clustering. Next we subset the data for 3m (Tabula Muris⁵) and compute how many cell types map to each individual cluster. For the clusters that we had a single 1:1 mapping (cluster:cell type) we propagate the annotations for all ages; in case there is a 1:many mapping we flagged that cluster for manual validation. Step-by-step instructions to reproduce this method are available from GitHub. For each cluster, we provide annotations in the controlled vocabulary of the cell ontology⁵⁶ to facilitate inter-experiment comparisons. Using this method, we were able to annotate automatically (~1min per tissue) over 70% of the dataset. The automatic annotations were then reviewed by each of the tissue experts leading to a fully curated dataset for all the cell types in Tabula Muris Senis.

Tissue cell composition analysis

For each tissue and age, we computed the relative proportion of each cell type. Next we used scipy.stats linregress to regress the relative tissue-cell type changes against age and considered significant the changes with p-value<0.05 for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic and a $r^2>0.5$.

Differential gene expression

We performed differential gene expression analysis on each tissue with a well-powered sample size (>100 cells in both young (1m and 3m) and old age group (18m, 21m, 24m and 30m)). We used a linear model⁵⁷ treating age as a numerical variable while controlling for sex and technology. We applied a false-discovery rate (FDR) threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change).

In Situ RNA Hybridization and quantification.

In situ RNA hybridization was performed using the Advanced Cell Diagnostics RNAscope® Multiplex Fluorescent Detection kit v2 (323110, Bio-techne) according to the manufacturer's instructions. Staining of mouse liver specimens was performed using 5µm paraffin-embedded thick sessions. Mouse livers were fixed in 10% formalin buffer saline (HT501128, Sigma Aldrich) for 24h at room temperature before paraffin embedding. For multiplex staining the following probes were used; *Clec4f* (Mm-Clec4f 480421, *II1b* (Mm-II1b 316891-C2), *Pecam1* (Mm-Pecam-1 316721), *Mrc1* (Mm-Mrc1 437511-C3). Slides were counter stained with Prolong gold antifade reagent with DAPI (P36931, Life technologies). Mounted slides were imaged on a Leica DM6 B fluorescent microscope (Leica Biosystems). Image quantification was performed using the starfish open source image-based transcriptomics pipeline (please refer to Starfish: Open Source Image Based Transcriptomics and Proteomics Tools available from http://github.com/spacetx/starfish and ⁵⁸)

Comparison between bulk and single-cell datasets

The differential gene analysis was defined on a per tissue basis. First, we investigated genes based on the single-cell data. We only considered cells from male animals and perform our analysis on the log (1 + CPM) transformed single-cell count matrices. Note that normalization of the single-cell data was done on a per cell basis. We defined two groups of cells based on age: young cells with age ≤ 3 months (Y) and old cells with age > 3 months (O). For each gene we compute the log2 fold-change of cell and read counts between O and Y. We defined cell count as the fraction of cells that express the gene. Similarly, we defined read count as the mean read count of the gene in the cells that express it. The calculated log2 fold-changes of a gene reflect its expression changes with aging within the single-cell data. Next we analyze each gene based on the bulk data. We computed the Spearman (Sp) correlation of bulk DESeq2 normalized gene expression with aging. We defined two groups of genes based on the bulk data, increasing with age Sp > 0.7 (U) and decreasing with age Sp < -0.7 (D). Finally, we compared the single-cell data based log2 fold-changes between the bulk data defined groups U and D. Specifically, we run Wilcoxon-Mann-Whitney test in order to understand if log2 fold-changes of cell or read counts could distinguish between the two groups. We used the U statistic for effect size.

T-Cell processing

We used TraCeR⁴⁴ version 0.5 to identify T-Cell clonal populations. We ran tracer assemble with --species Mmus set. We then ran tracer summarise with –species Mmus to create the final results. We used the following versions for TraCeR dependencies: igblast version 1.7.0,

kallisto version v0.43.1, Salmon version 0.8.2, Trinity version v2.4.0, GRCm38 reference genome. Step-by-step instructions to reproduce the processing of the data are available from GitHub.

B-Cell processing

We used singlecell-ige⁴³ version eafb6d126cc2d6511faae3efbd442abd7c6dc8ef (https://github.com/dcroote/singlecell-ige) to identify B-Cell clonal populations. We used the default configuration settings except we set the species to mouse. Step-by-step instructions to reproduce the processing of the data are available from GitHub.

Mutation analysis

We used samtools⁵⁹ version 1.9 and GATK³⁹ version v4.1.1.0 for mutation analysis. We used samtools faidx to create our index file. Then we used GATK CreateSequenceDictionary and GRCm38, as the reference, to create our sequence dictionary. Next we used GATK AddOrReplaceReadGroups to create a single read group using parameters -RGID 4 -RGLB lib1 -RGPL illumina -RGPU unit1 -RGSM 20. Finally we used GATK HaplotypeCaller to call the mutations. We disabled the following read filters: MappingQualityReadFilter, GoodCigarReadFilter, NotSecondaryAlignmentReadFilter, MappedReadFilter, MappingQualityAvailableReadFilter, NonZeroReferenceLengthAlignmentReadFilter, NotDuplicateReadFilter, PassesVendorQualityCheckReadFilter, and WellformedReadFilter, but kept all other default settings. The results were summarized per gene in the form of a mutation count per cell table. We started by removing genes mutated in over 60% of cells, to eliminate the possible bias of germline mutations. Then for each tissue we selected genes expressed in at least 75% of the cells for all the time points to avoid confounding the mutation results with differential gene expression associated with age. Next we computed the average number of mutations in the gene set (or ERCC spike-in controls) per cell and also the average number of raw counts (Supplementary Table 8) and plotted the different distributions. Step-by-step instructions to reproduce the processing of the data are available from GitHub.

Trajectory analysis

We used partition-based graph abstraction (PAGA⁶⁰) to reconstruct the aging trajectory in brain microglial cells. Step-by-step instructions to reproduce the processing of the data are available from GitHub.

Diversity score

The raw FACS or droplet dataset were used as the input. We filtered genes expressed in fewer than 5 cells, filtered cells if expressing fewer than 500 genes and discarded cells with total number of counts less than 5000. Next we performed size factor normalization such that every cell had 1e4 counts and performed a log1p transformation. This was followed by clustering, where we clustered every tissue and every tissue-cell type for every mouse separately using 6 different configurations: resolution parameters (0.3, 0.5, 0.7) * clustering method (Louvain, Leiden). This is to provide a robust clustering result. For each combination (each tissue-mouse and each tissue-cell_type-mouse), we computed the

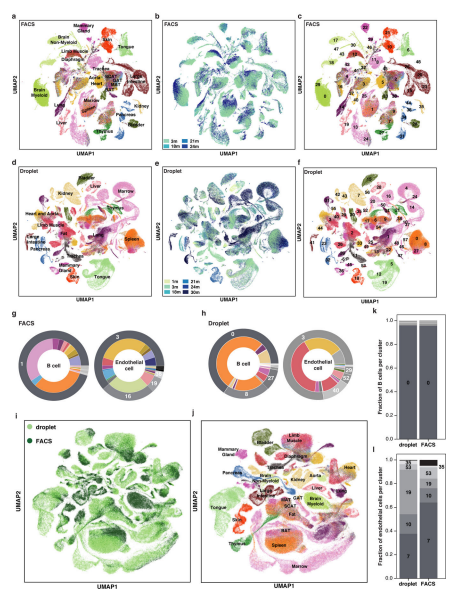
clustering diversity score as the Shannon entropy of the cluster assignment. We then regressed the diversity score against age to detect the systematic increase/decrease of clustering diversity with respect to age. FDR was used to correct for multiple comparisons. A tissue or a tissue-cell type was selected if the slope was consistent (having the same sign) in all 6 clustering configurations and at least 2 out of 6 clustering configurations had FDR<0.3. For each selected tissue or tissue-cell type, a separate UMAP was computed using cells from all mice for visualization using Leiden clustering with resolution parameter 0.7.

Interactive Data Browsers

http://tabula-muris-senis.ds.czbiohub.org/

https://tabula-maris-senis.cells.ucsc.edu

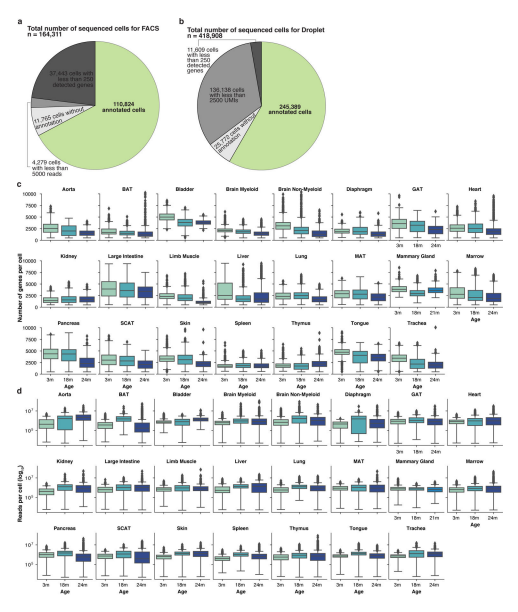
Extended Data

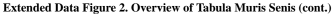


Extended Data Figure 1. Overview of Tabula Muris Senis (cont.)

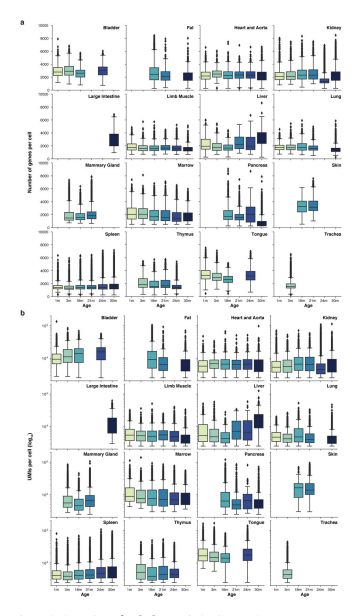
a,b, UMAP plot of all cells collected for FACS colored by tissue (**a**) or age (**b**). **c**, UMAP plot of all cells collected by FACS, colored by organ (Extended Data Figure 4c), overlaid with the Louvain cluster numbers. n = 110,824 individual cells for FACS. **d,e**, UMAP plot of all cells collected for droplet colored by tissue (**d**) or age (**e**). **f**, UMAP plot of all cells collected by droplet, colored by organ (Extended Data Figure 4c), overlaid with the Louvain cluster numbers. n = 245,389 individual cells for droplet. **g**, B cells (top) and endothelial cells (bottom) in FACS independently annotated for each organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ they originate from. **h**, B cells (and endothelial cells) in droplet independently annotated for each organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ set organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ set organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering clustering, irrespectively of the organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ where they were found. **i**, **j**, UMAP plot of all cells collected colored by method (**i**) or tissue

(j). n = 356,213 individual cells for FACS and droplet combined. k,l, B cells (k) and endothelial cells (l) cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the technology with which they were found.



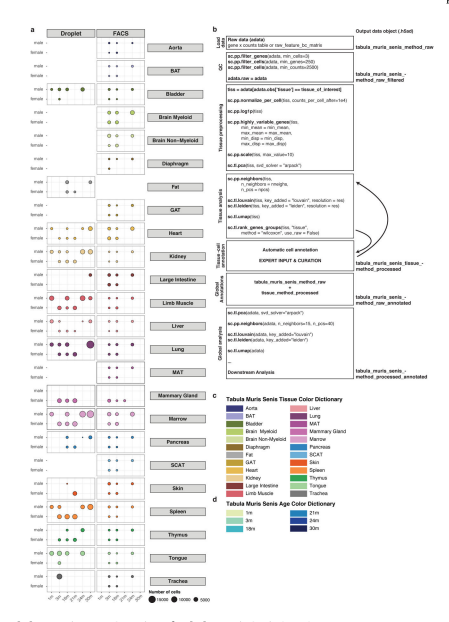


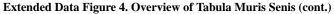
a, Pie chart with the summary statistics for FACS. **b**, Pie chart with the summary statistics for droplet. **c**, Box plot of the number of genes detected per cell for each organ and age for FACS **d**, Box plot of the number of reads per cell (log-scale) for each organ and age for FACS. For **c** and **d**, all data are expressed as mean \pm s.d. The sample size (number of cells for each tissue and age) is available in Supplementary Table 1.



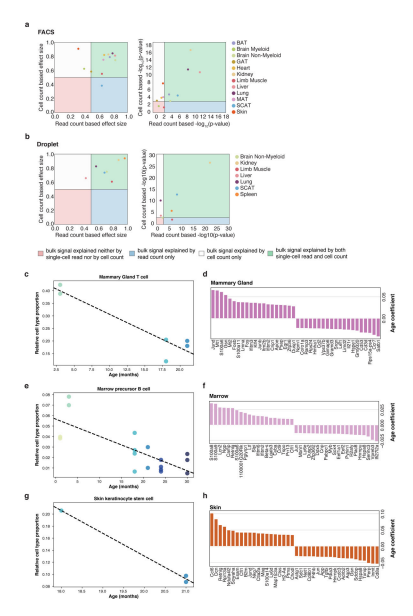
Extended Data Figure 3. Overview of Tabula Muris Senis (cont.)

a, Box plot of the number of genes detected per cell for each organ and age for droplet. **b**, Box plot of the number of UMIs per cell (log-scale) for each organ and age for droplet. All data are expressed as mean \pm s.d. The sample size (number of cells for each tissue and age) is available in Supplementary Table 2.





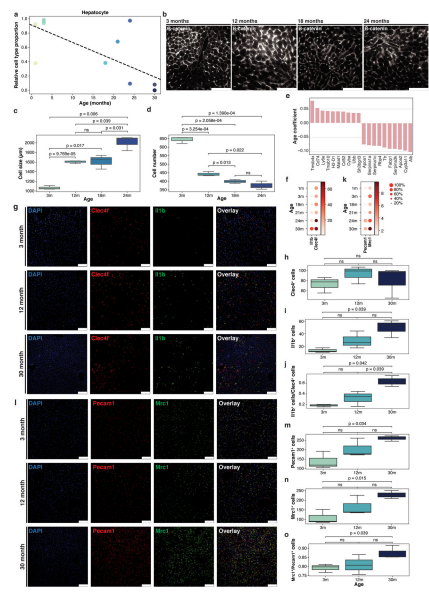
a, Balloon plot showing the number of sequenced cells per sequencing method per organ per sex per age. **b**, Schematic analysis workflow. **c**,**d**, Tabula Muris Senis color dictionary for organs and tissues (**c**) and ages (**d**).



Extended Data Figure 5. Comparison of bulk and single-cell datasets and Tissue cell compositions.

a,b, Aging patterns from bulk and single-cell data are consistent. Strong changes in bulk gene expression with aging can be either explained by cell or read count-based changes in single-cell data FACS (**a**) and droplet (**b**). Two-sided Wilcoxon–Mann–Whitney indicates that single-cell data based log2 fold-changes of cell or read counts distinguish between up and down regulated genes in bulk data. n = 110,824 individual cells for FACS and n = 245,389 individual cells for droplet. **c**, Mammary gland T cell relative abundances change significantly with age (p-value<0.05 and r²>0.7 for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). **d**, Top 20 upregulated and downregulated genes in mammary gland computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=6,393; 3,635; and 5,549 individual cells for

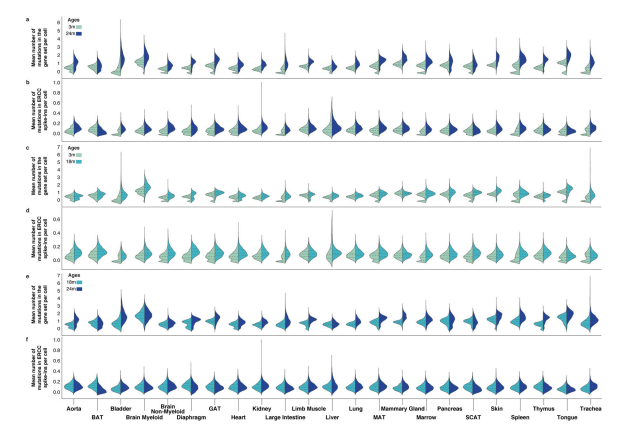
mammary gland 3m; 18m and 21m, respectively. e, Marrow precursor B cell relative abundances change significantly with age (p-value<0.05 and $r^2>0.7$ for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). f, Top 20 upregulated and downregulated genes in marrow computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=3,027; 8,559; 11,496; 5,216; 12,943 and 13,496 individual cells for marrow 1m; 3m; 18m; 21m; 24m and 30m, respectively. g, Skin keratinocyte stem cell relative abundances change significantly with age (p-value<0.05 and r^2 >0.7 for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). h, Top 20 upregulated and downregulated genes in skin computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=2,346; 1,494; 4,352= and 1,122 individual cells for skin 3m; 18m; 21m and 24m, respectively. The p-values for the cell type compositional changes are shown in Supplementary Table 5.



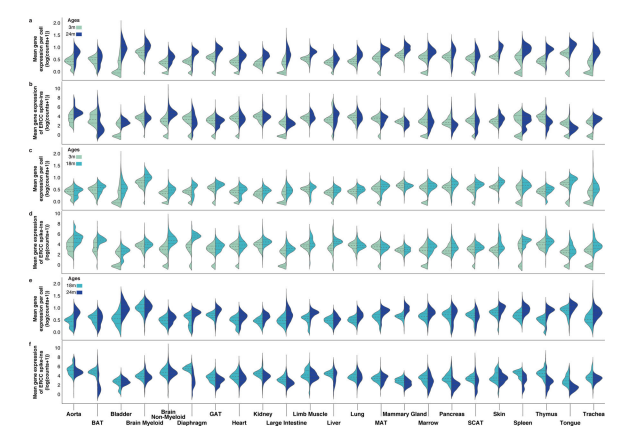
Extended Data Figure 6. Cellular changes during aging in the liver.

a, Liver hepatocyte relative abundances change significantly with age (p-value<0.05 and $r^2>0.7$ for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). n=2,791; 2,832; 3,806; 2,257; 6,384 and 5,713 individual cells for liver 1m; 3m; 18m; 21m; 24m and 30m, respectively. The p-values for the cell type compositional changes are shown in Supplementary Table 5. **b-d**, Brightfield imaging of hepatocytes across age (**b**) and respective quantification (**c-d**). **e**, Top 10 upregulated and downregulated genes in liver computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). The sample size is the same as for panel **a**. **f,k**, Gene expression of Il1b and Clec4f (**f**) and Pecam1 and Mrc1 (**k**) in the liver droplet dataset for the six ages. **g-j**, Staining of Kupffer cells across age (**g**) and respective quantification (**h-j**).

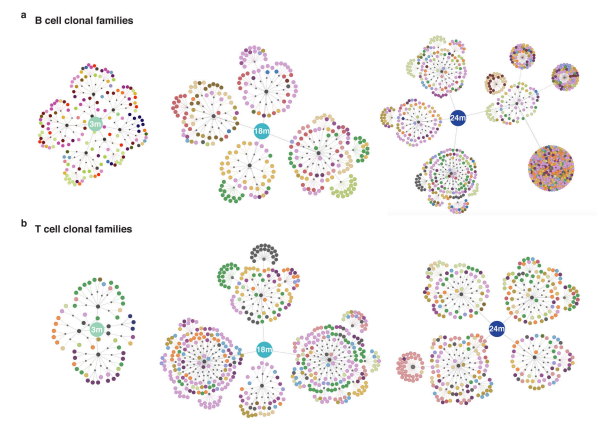
l-o, Staining of liver endothelial cells across ages (l) and respective quantification (**m-o**). The white scale bar corresponds to 100 μ m. For panels **c-d**, **h-j** and **m-o**, all data are expressed as mean \pm s.d. and p-values were obtained using a Welch's test. The sample size for each group is available in Supplementary Table 7.



Extended Data Figure 7. Mutational burden across tissues in the aging mice (cont.). a,b, Mean number of somatic mutations in genes and ERCC spike-in controls across all tissues per age group (3m and 24m (**a**), 3m and 18m (**b**), 18m and 24 (**c**)). Mutations are presented as the mean number of mutations per gene or ERCC spike-inn per cell.

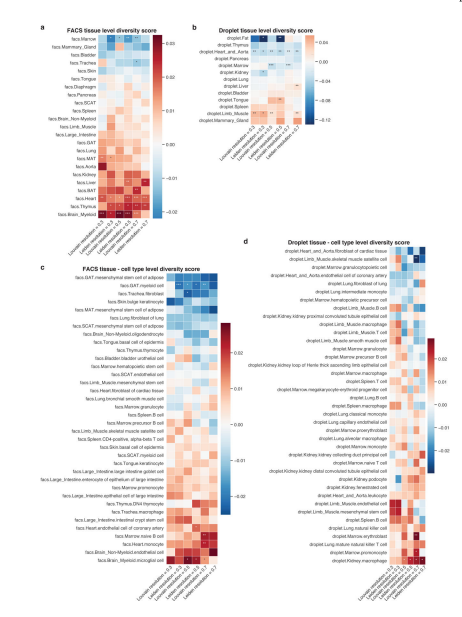


Extended Data Figure 8. Mutational burden across tissues in the aging mice (cont.). a,b,c, Gene raw expression and ERCC spike-inn control raw expression across all tissues per age group (3m and 24m (**a**), 3m and 18m (**b**), 18m and 24 (**c**)). Raw expression are presented as the mean number of counts per gene or ERCC spike-inn control per cell.



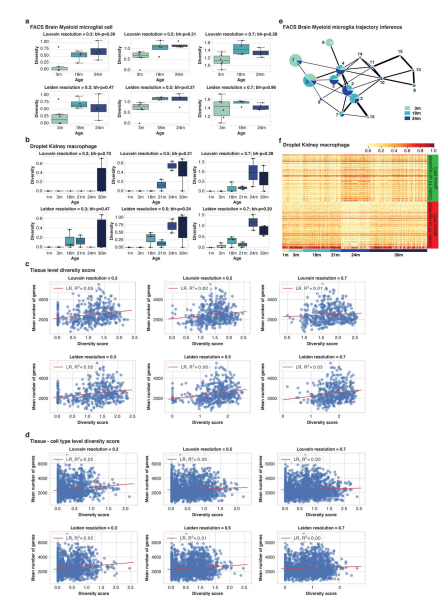
Extended Data Figure 9. Immune repertoire clonality analysis.

a, B-cell clonal families. For each time point, the clonal families are represented in a tree structure for which the central node is age. Connected to the age node there is an additional node (dark gray) that represents each animal and the clonal families are depicted for each animal. For each clonal family, cells that are part of that family are colored by the organ of origin. **b**, T-cell clonal families. For each time point, clonal families are represented in a tree structure for which the central node is age. Connected to the age node there is an additional node (dark gray) that represents each animal and the clonal families are depicted for each animal. For each clonal family, cells that are part of that family are colored by the organ of origin.



Extended Data Figure 10. Diversity score summary.

a,b, Heatmap summary of the overall tissue diversity score for FACS (a) and droplet (b).c,d, Heatmap summary of the tissue cell-type diversity score for FACS (c) and droplet (d).



Extended Data Figure 11. The aging immune system (cont.)

a,b, Diversity score at different cluster resolutions for FACS brain myeloid microglia cell (**a**) and droplet kidney macrophage (**b**). n = 14 mice for **a** and n = 16 mice for **b**. All data are expressed as quantiles. The p-values were obtained using a linear regression and two-sided F-test, adjusted for multiple comparison using the Benjamini-Hochberg procedure (i.e., bh-p value). **c,d**, Diversity score correlation with the number of genes expressed per tissue (**c**) or tissue cell-type (**d**). The red line corresponds to the linear regression curve. **e**, Trajectory analysis for brain myeloid microglia cell. **f**, Heatmap showing differential gene expression analysis of cluster 10 (mostly young macrophages) versus clusters 13 (mostly old macrophages). For the complete gene list please refer to Supplementary Table 10.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

We thank Sony Biotechnology for making an SH800S instrument available for this project. Some cell sorting/flow cytometry analysis for this project was done on a Sony SH800S instrument in the Stanford Shared FACS Facility. Some fluorescence activated cell sorting (FACS) was done with instruments in the VA Flow Cytometry Core, which is supported by the US Department of Veterans Affairs (VA), Palo Alto Veterans Institute for Research (PAVIR), and the National Institutes of Health (NIH). This work was supported by the Chan Zuckerberg Biohub, Department of Veterans Affairs grant IK6 BX004599 (TWC) and NIH/NIA DP1 AG053015 grant (TWC). We would like to thank Bruno Tojo for the artwork. We thank Chenling Xu and Joshua Batson for helpful discussions.

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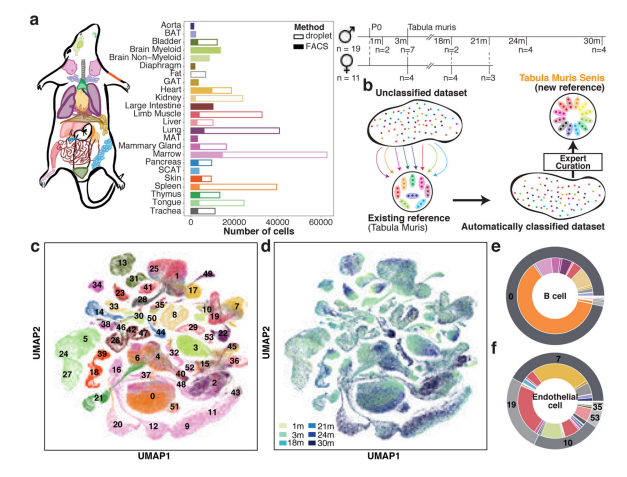


Figure 1. Overview of Tabula Muris Senis.

a, 23 organs from 19 male and 11 female mice were analyzed at 6 different time points. The bar plot shows the number of sequenced cells per organ prepared by FACS (n=23 organs) and microfluidic droplets (n=16 organs). For the droplet dataset the Fat sub-tissues were processed together (Fat = BAT+GAT+MAT+SCAT). BAT, Brown Adipose Tissue; GAT, Gonadal Adipose Tissue; MAT, Mesenteric Adipose Tissue; SCAT, Subcutaneous Adipose Tissue. **b**, Annotation workflow. Data were clustered together across all time points. We used the Tabula Muris (3m time point) as a reference for the automated pipeline and the annotations were manually curated by tissue experts. **c**,**d**, UMAP plot of all cells, colored by organ and overlaid with the Louvain cluster numbers (**c**) and age (**d**); n = 356,213 individual cells. For the color dictionaries please refer to Extended Data Figure 2c. **e**, B cells (top) and endothelial cells (bottom) independently annotated for each organ cluster together by unbiased whole-transcriptome Louvain clustering, irrespectively of the organ they were found.

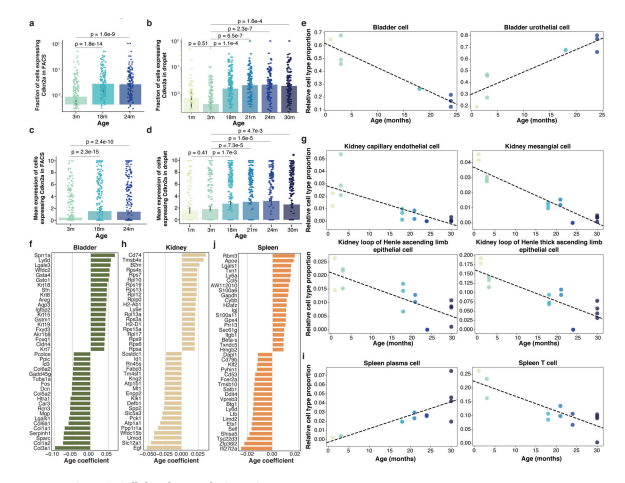


Figure 2. Cellular changes during aging.

a,b, Bar plot showing the fractions of cells expressing Cdkn2a at each age group for FACS (a) and droplet (b). c,d, Bar plot of the median expression of Cdkn2a for the cells that do express the gene at each age group for FACS (c) and droplet (d). The y-axis corresponds to log-transformed and scaled values. All data are expressed as mean \pm s.d. with individual data points shown. p-values were obtained using a Mann-Whitney-Wilcoxon rank-sum two-sided test. n=44,518; 34,027 and 31,551 individual cells for FACS 3m; 18m and 24m, respectively. n=25,980; 45,602; 44,645; 35,828; 37,660 and 55,674 individual cells for droplet 1m; 3m; 18m; 21m; 24m and 30m, respectively. e, Bladder cell (left) and bladder urothelial cell (**right**) relative abundances change significantly with age (p-value<0.05 and r^2 >0.7 for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). f, Top 20 upregulated and downregulated genes in bladder computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=970; 3,804; 2,739 and 3,864 individual cells for bladder 1m; 3m; 18m and 24m, respectively. g, Kidney capillary endothelial cell (top-left), mesangial cell (top-right), loop of Henle ascending limb epithelial cell (bottom-left) and loop of Henle thick ascending limb epithelial cell (bottom-right) relative abundances change significantly with age (pvalue<0.05 and r²>0.7 for a hypothesis test whose null hypothesis is that the slope is zero,

using two-sided Wald Test with t-distribution of the test statistic). **h**, Top 20 upregulated and downregulated genes in kidney computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=2,488; 2,832; 3,806; 2,257; 6,384 and 5,713 individual cells for kidney 1m; 3m; 18m; 21m; 24m and 30m, respectively. **i**, Spleen plasma cell (**left**) and T cell (**right**) relative abundances change significantly with age (p-value<0.05 and r²>0.7 for a hypothesis test whose null hypothesis is that the slope is zero, using two-sided Wald Test with t-distribution of the test statistic). **j**, Top 20 upregulated and downregulated genes in spleen computed using MAST⁵⁷, treating age as a continuous covariate while controlling for sex and technology. Genes were classified as significant under an FDR threshold of 0.01 and an age coefficient threshold of 0.005 (corresponding to ~10% fold change). n=2,986; 8,839; 7,141; 6,395; 5,245 and 8,946 individual cells for spleen 1m; 3m; 18m; 21m; 24m and 30m, respectively. The p-values for the cell type compositional changes are shown in Supplementary Table 5.

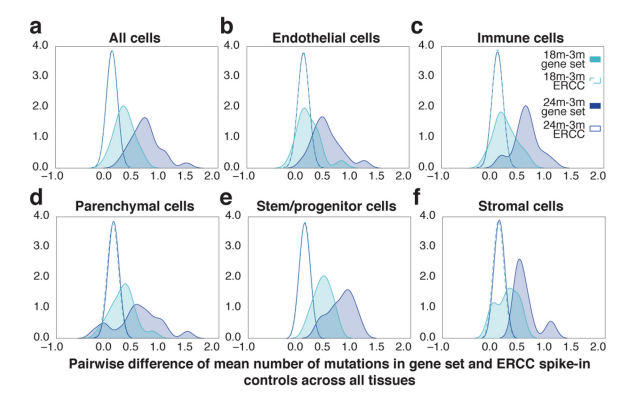


Figure 3. Mutational burden across tissues in the aging mice.

Distribution of the difference of the mean mutation in the gene set (and ERCC spike-in controls) per cell between 24m and 3m and 18m and 3m for all tissues and cells (**a**) and with the cell types split in five functional groups, endothelial (**b**), immune (**c**), parenchymal (**d**), stem/progenitor cell (**e**) and stromal (**f**). Filled and solid line distributions correspond to the mean mutation difference in gene set. White and dashed line distributions correspond to the mean mutation difference in ERCC spike-in controls. Please note that the mean mutation difference in ERCC spike-in controls overlaps for both age groups.

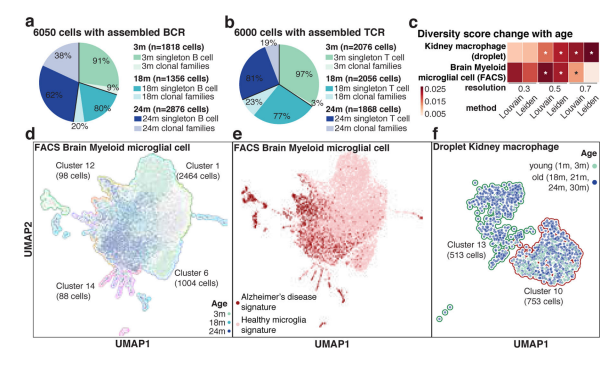


Figure 4. The aging immune system.

a, B-cell clonal families. The pie chart shows the proportion of singleton B cells and B cells that are part of clonal families at 3m, 18m and 24m. Please refer to Extended Data Figure 9 for the clonal networks. **b**, T-cell clonal families. The pie chart shows the proportion of singleton T cells and T cells that are part of clonal families at 3m, 18m and 24m. Please refer to Extended Data Figure 9 for the clonal networks. **c**, Diversity score for the two cell types that significantly change with age. **d**, UMAP plot of the brain myeloid microglial cell Leiden clusters (numbers) colored by age. Faded clusters do not change their relative age cell composition; colored clusters change their relative cell composition. **e**, UMAP plot of the brain myeloid microglial cells when scored using the microglia Alzheimer's disease signature (Supplementary Table 10). n = 4,532; 4,461 and 4,424 individual microglia cells for brain myeloid 3m, 18m and 24m, respectively. **f**, UMAP plot of the kidney macrophage Leiden clusters (numbers) colored by age group. n = 62; 139; 264; 105; 284 and 553 individual macrophage cells for kidney 1m, 3m, 18m, 21m, 24m and 30m, respectively.