Immune ageing at single-cell resolution

Denis A. Mogilenko, Irina Shchukina and Maxim N. Artyomov

Abstract | Ageing leads to profound alterations in the immune system and increases susceptibility to some chronic, infectious and autoimmune diseases. In recent years, widespread application of single-cell techniques has enabled substantial progress in our understanding of the ageing immune system. These comprehensive approaches have expanded and detailed the current views of ageing and immunity. Here we review a body of recent studies that explored how the immune system ages using unbiased profiling techniques at single-cell resolution. Specifically, we discuss an emergent understanding of age-related alterations in innate and adaptive immune cell populations, antigen receptor repertoires and immune cell-supporting microenvironments of the peripheral tissues. Focusing on the results obtained in mice and humans, we describe the multidimensional data that align with established concepts of immune ageing as well as novel insights emerging from these studies. We further discuss outstanding questions in the field and highlight techniques that will advance our understanding of immune ageing in the future.

Inflammageing

Chronic low-grade systemic inflammation occurring in the absence of a pathogen or exogenous inflammatory agent that develops with advanced age owing to endogenous pro-inflammatory processes.

Ageing is a multimodal process that occurs at several levels, with organs, tissues and cells undergoing substantial and simultaneous remodelling, thus limiting the power of conventional molecular biology approaches to resolve this complexity. Loss of physiological integrity, disturbed tissue homeostasis and increasing deterioration of various biological systems, including the immune system, are among the key features of ageing1. Together, these changes reshape physiological and immune landscapes as well as increase vulnerability to some diseases, frailty and multimorbidity2. Interestingly, not all immune processes are uniformly sensitive to ageing, suggesting that various immune cell populations are heterogeneously impaired in older organisms (for example, ≥65 years in humans and ≥1.5 years in mice)3. The current development of multi-omics and single-cell techniques4 (BOX 1) and progress in high-throughput biology and systems immunology5 make it possible to comprehensively characterize cellular and functional heterogeneity and dynamic changes in the immune system during ageing. Novel techniques, including single-cell RNA sequencing (scRNA-seq) (BOX 2), have confirmed previous findings and expanded existing paradigms in ageing and immunity built on evidence from prior targeted studies. In this analysis identified two distinct gene clusters related to immune responses that are rapidly upregulated on transition from middle to advanced age in mice3. These transcriptional alterations likely reflect an active infiltration of immune cells into aged tissues and consequent systemic inflammation. Intriguingly, the breadth of expression changes in these clusters is greatest in adipose tissues, especially in visceral fat deposits6, suggesting that fat-associated macrophages and adipocytes might be among the key regulators of inflammageing. In addition to immune cell infiltration, inflammageing might result from alterations in basic tissue architecture owing to age-associated remodelling. Broadly, there is enhanced formation of fibrotic structures and senescent cell accumulation in many tissues7–11, but some tissues are also characterized by a distinct set of immune alterations. For example, in mice, type I interferon signalling has been...
A comprehensive assessment of a set of molecules, such as DNAs, RNAs, proteins and metabolites, is collectively termed ‘multi-omics’. Flow cytometry is historically the first single-cell-based technique capable of simultaneously identifying 5–20 characteristics in individual cells. Typically, flow cytometry experiments use fluorescently conjugated antibodies to detect surface and intracellular epitopes and/or fluorescent probes that measure cellular characteristics such as an abundance of DNA, mitochondria, lysosomes and lipid droplets. Mass cytometry, also known as cytometry by time of flight, is a variation of flow cytometry that relies on antibodies labelled with heavy metal ions tags, which allows the measurement of up to 40 individual epitopes in individual cells. During the past decade, unbiased single-cell–based applications of omics have been extensively developed. Single-cell RNA sequencing (scRNA-seq) is a rapidly growing technology that can detect and quantitatively measure 500–5,000 of the most highly expressed transcripts per cell (Fig. 1). Computational analysis of transcriptomes of individual cells can identify novel cell populations, activation states and differentiation programmes in immune cells. The variation of the scRNA-seq technique that identifies sequences of variable regions in the T cell receptor or B cell receptor in addition to RNA transcripts in individual cells (termed ‘single-cell T cell receptor sequencing’ and ‘single-cell B cell receptor sequencing’) has become instrumental in studying antigen receptor repertoires at single-cell resolution (Box 1). Cellular indexing of transcriptomes and epitopes by sequencing is a method in which oligonucleotide–labelled antibodies are used to integrate cellular protein and transcriptome measurements into a single-cell readout (Fig. 1). It exceeds the limitations of mass cytometry and, theoretically, may extend to measuring thousands of proteins and transcripts simultaneously. Single-cell assay for transposable-accessible chromatin using sequencing was recently developed, and it enables the analysis of epigenetic landscapes of individual cells. Moreover, single-cell techniques that combine transcriptional information with epigenetic characteristics in individual cells have been developed. An important limitation of these single-cell methods is that they use cell suspensions or isolated nuclei and require disaggregation of an organ, which leads to loss of spatial information relating to the anatomical localization of cells. Adding the positional information to single-cell experiments is the next frontier in the field of multi-omic immunology. Thus, spatial applications of mass cytometry (imaging mass cytometry) and spatial transcriptomics are rapidly growing fields that will help to integrate transcriptional and proteomic information in tissue sections to combine multimodal characteristics of individual cells or groups of cells with the complexity of tissue architecture.

Box 1 | Multi-omics and single-cell technologies in immunology

A comprehensive assessment of a set of molecules, such as DNAs, RNAs, proteins and metabolites, is collectively termed ‘multi-omics’. Flow cytometry is historically the first single-cell–based technique capable of simultaneously identifying 5–20 characteristics in individual cells. Typically, flow cytometry experiments use fluorescently conjugated antibodies to detect surface and intracellular epitopes and/or fluorescent probes that measure cellular characteristics such as an abundance of DNA, mitochondria, lysosomes and lipid droplets. Mass cytometry, also known as cytometry by time of flight, is a variation of flow cytometry that relies on antibodies labelled with heavy metal ions tags, which allows the measurement of up to 40 individual epitopes in individual cells. During the past decade, unbiased single-cell–based applications of omics have been extensively developed. Single-cell RNA sequencing (scRNA-seq) is a rapidly growing technology that can detect and quantitatively measure 500–5,000 of the most highly expressed transcripts per cell. Computational analysis of transcriptomes of individual cells can identify novel cell populations, activation states and differentiation programmes in immune cells. The variation of the scRNA-seq technique that identifies sequences of variable regions in the T cell receptor or B cell receptor in addition to RNA transcripts in individual cells (termed ‘single-cell T cell receptor sequencing’ and ‘single-cell B cell receptor sequencing’) has become instrumental in studying antigen receptor repertoires at single-cell resolution (Box 1). Cellular indexing of transcriptomes and epitopes by sequencing is a method in which oligonucleotide–labelled antibodies are used to integrate cellular protein and transcriptome measurements into a single-cell readout. It exceeds the limitations of mass cytometry and, theoretically, may extend to measuring thousands of proteins and transcripts simultaneously. Single-cell assay for transposable-accessible chromatin using sequencing was recently developed, and it enables the analysis of epigenetic landscapes of individual cells. Moreover, single-cell techniques that combine transcriptional information with epigenetic characteristics in individual cells have been developed. An important limitation of these single-cell methods is that they use cell suspensions or isolated nuclei and require disaggregation of an organ, which leads to loss of spatial information relating to the anatomical localization of cells. Adding the positional information to single-cell experiments is the next frontier in the field of multi-omic immunology. Thus, spatial applications of mass cytometry (imaging mass cytometry) and spatial transcriptomics are rapidly growing fields that will help to integrate transcriptional and proteomic information in tissue sections to combine multimodal characteristics of individual cells or groups of cells with the complexity of tissue architecture.

Effects of aging on myeloid cells

Ageing is associated with a gradual expansion of circulating myeloid cell populations relative to lymphoid cell populations (Table 1). In part, this shift is due to a decline in thymic activity required for the generation of nascent T cells. At the same time, there is a differential effect of age on early myelopoiesis and lymphopoiesis in the bone marrow. Changes in the bone marrow of aged mice lead to clonal skewing of haematopoietic stem cells (HSCs) towards the myeloid lineage. RNA sequencing analysis of the bone marrow stromal microenvironment in mice revealed an age-driven progressive upregulation of pro-inflammatory genes (including Il6 and Il1b), which can shift the balance towards HSCs with myeloid potential. Indeed, analysis of the young bone marrow microenvironment at single-cell resolution demonstrated that stress-induced changes in molecular pathways (such as downregulation of the Notch ligands Delta-like proteins encoded by Dil1 and Dil4) also initiate the myeloid transcriptional programme in mouse HSCs. It is plausible that these and other similar stress-related molecular circuits underlie the accumulation of myeloid-biased HSCs in haematopoietic niches during ageing. Nevertheless, these developmental shifts do not imply the emergence of inflamaging directly. For instance, ageing of humans does not appear to be associated with an active shift of circulating monocytes to a pro-inflammatory phenotype, given that these cells show similar transcriptional profiles in young and old healthy individuals. Further studies of human HSC ageing at single-cell resolution are needed to better understand how myelopoiesis is regulated in aged individuals.

Ageing is also associated with alterations in subpopulations of tissue-resident myeloid cells (Fig. 2). scRNA-seq analyses revealed substantial heterogeneity among tissue-resident myeloid cells, especially in response to inflammation, and showed increased proportions of interstitial macrophages expressing chemokine receptor genes Ccr2 and Cx3cr1 but reduced proportions of alveolar macrophages in aged mice (Table 1). Moreover, alveolar macrophages in aged mice highly express genes that contribute to inflamaging, such as Spp1 (which encodes osteopontin) and Cd68 (Ref. 29). In line with this, the proportions of fetal-derived tissue-resident macrophages markedly decrease in aged mouse mammary tissues. By contrast, the proportions of macrophages that arise from bone marrow–derived myeloid precursors are not changed, but these cells show increased expression of pro-inflammatory genes (such as Ccl5 and Gdf15) in the peritoneal cavity, scRNA-seq identified markedly reduced numbers of large peritoneal macrophages (which originate from embryonic precursors and express Icam2) in old mice versus young mice, as previously observed by means of flow cytometry. Moreover, single-cell analysis of ageing mice demonstrated increased inflammatory function (including expression of the cytokine gene Il1b) of liver-resident macrophages (also known as Kupffer cells), which likely contributes to hepatic inflammation and damage. Thus, the tissue environment in aged mice is generally associated with a substitution of tissue-resident subsets of macrophages originating from embryonic precursors for monocye-derived populations. Their transcriptional programmes shift towards pro-inflammatory phenotypes, contributing to inflamaging. It is crucial to further understand the fate of these macrophage lineages in various organs during human ageing, and one important resource is a recently published dataset dissecting the development of human tissue-resident macrophages at single-cell resolution.

To a certain degree, pro-inflammatory shifts of myeloid cell populations in old organisms might depend on effects of ageing on physiological processes and might be ameliorated by systemic interventions, such as dietary restrictions, including caloric restriction. Recent single-cell transcriptomics analysis of the effects of caloric restriction on multiple tissues in ageing rats showed that caloric restriction reverses age-related inflammation. Specifically, that study demonstrated that caloric restriction reverses the pro-inflammatory polarization of aged M1 macrophages, increases the proportions of anti-inflammatory macrophages with high phagocytic activity and modifies

Senescent cell
A terminally differentiated cell characterized by irreversible cell-cycle arrest and a specific secretory profile that develops in response to various stress triggers and accumulates in aged organisms.

Caloric restriction
A dietary intervention that reduces average daily caloric intake below typical or habitual levels, while containing essential nutrients and not leading to malnutrition.
ligand–receptor-dependent communication between macrophages and stromal cells\(^4\). Further studies are needed to translate the effect of caloric restriction and similar interventions to myeloid cell populations in ageing individuals.

Neutrophils are also important myeloid cells, which although generally short-lived may also be affected by ageing. Neutrophil development and numbers seem to be unaltered systemically by ageing, yet neutrophils show a decline in phagocytic activity and killing of bacteria in elderly humans\(^6\). Furthermore, recent studies showed that age-related changes in the tissue environment could reprogramme neutrophils. Thus, in mice, ageing interferes with transmigrational migration of neutrophils in injured tissues via a process regulated by CXC-chemokine ligand 1 (CXCL1), resulting in aberrant neutrophil trafficking and subsequent remote organ damage\(^3\). CXCL1 also recruits neutrophils to the liver in aged mice, where they induce tissue senescence and inflammation via the production of reactive oxygen species\(^6\). This crosstalk between the inflamminge-associated chemokine CXCL1, tissue damage and aberrant neutrophil functions possibly contributes to the development of age-related chronic inflammation and can be further studied in the context of exacerbated pulmonary and hepatic chronic diseases in ageing individuals. Furthermore, a recent multi-omics profiling study demonstrated that age-associated neutrophil remodelling is evident in bone marrow-residing neutrophils\(^7\), suggesting a role for the aged bone marrow microenvironment in age-associated neutrophil dysregulation.

Among other cellular phenotypes linked to ageing, disturbed phagocytic activities of neutrophils, macrophages and dendritic cells (DCs) and their reduced capacity to engulf apoptotic cells via efferocytosis have been described in mice and humans\(^8,9\). In old mice, conventional type 1 dendritic cells, but not conventional type 2 dendritic cells, are decreased in abundance in the spleen\(^10\). However, it remains controversial whether aged dendritic cell subtypes demonstrate specific alterations in antigen presentation capacity\(^11\).

**Effects of ageing on innate lymphocytes**

Innate lymphoid cells (ILCs) include natural killer (NK) cells, group 1 ILCs (ILC1s), group 2 ILCs (ILC2s) and group 3 ILCs\(^12\). ILCs play a unique role in tissue immunity via cytokine secretion and interaction with other immune and stromal cells. However, surprisingly little is known about the role of ILCs in the homeostasis of aged tissues. Age-related changes in NK cells in older humans (≥60 years) include a shift towards a mature CD14\(^+\)CD56\(^-\) subset, coupled with diminished cytokine secretion, cytotoxicity and especially the release of perforin\(^13,14\). At least some of the age-associated alterations in NK cell functions are programmed by the aged environment in mice. Indeed, NK cell transfer experiments show that the aged environment interferes with NK cell maturation and proliferation\(^15\). In line with these observations, scRNA-seq analysis of immune cells from the spleen, lungs and liver demonstrated a decrease in NK cell proportions in old mice\(^16\). These data suggest a general ability of an aged environment to suppress NK cell homeostasis. It is possible that the decreased NK cell numbers contribute to impaired antiviral immunity in older organisms\(^17\). Intriguingly, tissue-specific proliferation and accumulation of NK cells was shown to occur in distinct aged niches, such as the dentate gyrus neurogenic niche of human and mouse aged brains. scRNA-seq analysis of NK cells in the aged dentate gyrus revealed features of increased activation and cytotoxicity\(^18\). Moreover, mouse NK cells eliminate senescent neuroblasts and mediate age-related decline in cognitive function\(^19\), providing a striking example of a deleterious effect of NK cells in old organisms.

ILC1s are non-cytotoxic cells that produce interferon-γ (IFNγ) and function as the first line of defence against viruses and certain bacteria\(^20\). Similarly to NK cells, hepatic ILC1 numbers decrease in old mice\(^21\). Tissue-resident liver ILC1s develop locally from SCA1\(^+\)MAC1\(^-\) HSCs via a mechanism driven by IFNγ\(^22\). It remains unclear which signals from the aged environment disturb ILC1 development and whether ILC1s in other organs are affected in old mice and humans. ILC2s are involved in innate immune responses against parasites and regulate tissue homeostasis\(^23\). Single-cell analyses in mice showed that ILC2s are less abundant in old lungs\(^24\) and adipose tissue\(^25\). However, the decrease in ILC2 populations is not a universal hallmark of ageing; for example, the numbers of ILC2s increase in aged mouse brains\(^26\). Ageing increases early bone marrow development of ILC2s, and the decline in tissue ILC2

---

**Box 2 | Single-cell RNA sequencing**

Single-cell RNA sequencing (scRNA-seq) approaches recently evolved owing to rapid progress in next-generation sequencing technologies. Single-cell isolation is the first step of scRNA-seq. Historically, fluorescence-activated cell sorting of hundreds of individual cells was one of the early strategies that rely on the availability of monoclonal antibodies that recognize proteins expressed on the cells of interest. As a more efficient variant, microfluidic technologies have recently become popular for the low-cost isolation of individual cells. One of the microfluidic technologies, Fluidigm C1, provides single-cell isolation, RNA extraction and cDNA synthesis for up to 800 cells on a single chip\(^27\). A popular alternative for single-cell isolation is microdroplet-based microfluidics\(^28\). This approach allows the dispersion of liquid droplets in an oil phase, requires lower volumes than standard microfluidic chips and can screen thousands to millions of cells. An example of a commercial microdroplet-based microfluidics system is the Chromium system from 10X Genomics, which can take a suspension of cells or nuclei as an input. Consequent steps include cell lysis, first-strand cDNA synthesis by reverse transcription of mRNA, second-strand synthesis and cDNA amplification by PCR to generate scRNA-seq libraries. Some scRNA-seq methods produce full-length coverage across transcripts (such as Smart-Seq2\(^29\)), whereas other methods limit sequencing coverage to 3′ or 5′ regions of mRNAs (such as 10X Genomics). The most popular approaches to make scRNA-seq libraries are also limited by capturing only mRNA with poly(A) tails. Although bulk RNA-seq allows almost genome-wide detection of mRNA expression in pooled cell samples, scRNA-seq cannot measure the expression levels of most genes in individual cells. Indeed, the sequencing depth usually allows reliable identification of limited numbers of highly expressed transcripts per single cell (usually 1,000–5,000 individual transcripts). Because of the ongoing development of scRNA-seq techniques, different generations can differ in sequencing depth and create technical limitations to compare results from early versions versus late versions of scRNA-seq. In the latest versions, incorporation of unique molecular identifiers (random 10–16-bp sequences, also known as barcodes) in the reverse transcription step removes bias from PCR amplification steps and increases sequencing accuracy. A more detailed analysis of applications of scRNA-seq to analyse the immune system is presented in recent excellent reviews\(^30,31\).
Fig. 1 | Cellular and molecular components of inflammainge. Low-grade chronic systemic inflammation, or inflammainge, comprises multiple inflammatory factors that originate from various cellular sources in aged organisms. Aged pro-inflammatory tissue macrophages produce cytokines and chemokines, and senescent cells secrete a plethora of pro-inflammatory components and matrix metalloproteinases (MMPs) as components of the senescence-associated secretory phenotype (SASP). Moreover, non-immune cells, such as adipocytes, preadipocytes and stromal cells (including fibroblasts and endothelial cells), contribute to inflammainge by secreting soluble inflammatory factors and altering the tissue microarchitecture of aged organs. The pro-inflammatory chemokines drive excessive immune cell infiltration in tissues, where the cytokines (such as IL-6 and IL-1β) and immunoregulatory factors (such as prostaglandin E2, PGE2) reprogramme immune cell subsets to a more pro-inflammatory and dysfucntional state. In turn, the dysfunctional immune cells, including macrophages and T cells, amplify the inflammatory and destructive processes in ageing tissues. This crosstalk between cellular and molecular components of inflammainge ultimately results in a progressive functional decline in various organs and leads to age-associated diseases.

**Effects of ageing on T cells**

**Age-associated CD8**+ **memory T cells.** Essential for antiviral and antitumour immunity, CD8+ T cells are substantially affected by ageing in both mice and humans. In general, older age has been associated with a loss of naive CD8+ T cells coupled with an increase in memory subsets and an overall increase in clonally expanded CD8+ T cells in multiple tissues in old mice, whereas in the ageing brain, ILCs2 function to counteract cognitive decline. ILCs2 have been found in human adipose tissue, but it remains unclear whether they contribute to age-associated metabolic dysfunction in ageing individuals.

**T cell exhaustion**
A specific state of T cell dysfunction driven by a persistent antigen stimulation and chiefly defined by reduced cytotoxicity and inability to produce pro-inflammatory cytokines (such as interferon-γ (IFNγ)) and tumour necrosis factor (TNF) and proliferation on T cell receptor stimulation.

**Cellular indexing of transcriptomes and epigenomes by sequencing (CITE-seq).** A method that uses oligonucleotide-labelled antibodies to integrate cellular protein and transcriptome measurements into an integrated single-cell readout.

populations might be due to dysfunctional migratory cues. Functionally, ILC2s appear to be associated with healthier ageing phenotypes. In old adipose tissues, a decrease in the numbers of ILC2s is linked to a metabolic impairment and cold-induced lethality exacerbated in old mice, whereas in the ageing brain, ILCs2 function to counteract cognitive decline. ILC2s have been found in human adipose tissue, but it remains unclear whether they contribute to age-associated metabolic dysfunction in ageing individuals.

**Ageing-associated morbidities**
- Frailty
- Obesity
- Vulnerability to infections
- Cognitive function decline

**Immunoregulatory factors:**
- Prostaglandin E2 (PGE2)
- TGFβ
- IL-1β
- CXCL1, CXCL8
- MMPs
- GDF15
- CCL2, CCL5

**Chemokines:**
- CCL2, CCL5
- CXCL1, CXCL8

**Cytokines:**
- Interferon-γ (IFNγ)
- Pro-inflammatory tissue macrophages produce cytokines and chemokines, and senescent cells secrete a plethora of pro-inflammatory components and matrix metalloproteinases (MMPs) as components of the senescence-associated secretory phenotype (SASP). Moreover, non-immune cells, such as adipocytes, preadipocytes and stromal cells (including fibroblasts and endothelial cells), contribute to inflammainge by secreting soluble inflammatory factors and altering the tissue microarchitecture of aged organs. The pro-inflammatory chemokines drive excessive immune cell infiltration in tissues, where the cytokines (such as IL-6 and IL-1β) and immunoregulatory factors (such as prostaglandin E2, PGE2) reprogramme immune cell subsets to a more pro-inflammatory and dysfunctional state. In turn, the dysfunctional immune cells, including macrophages and T cells, amplify the inflammatory and destructive processes in ageing tissues. This crosstalk between cellular and molecular components of inflammainge ultimately results in a progressive functional decline in various organs and leads to age-associated diseases. CCL, CC-chemokine ligand; CXCL, CXC-chemokine ligand; GDF15, growth/differentiation factor 15; IFNγ, interferon-γ; PD1, programmed cell death 1; PD1, programmed cell death 1; IFNγ, interferon-γ; PGE2, prostaglandin E2. Aged cells include macrophages, senescent cells, preadipocytes, adipocytes, structural cells, Fibrosis, Loss of cell identity, Impaired cec-cells communication.

**Structural cells**
- adipocytes
- preadipocytes
- senescent cells
- macrophages
- structural cells

**Adipocytes**
- preadipocytes
- senescent cells
- macrophages
- structural cells

**Preadipocytes**
- adipocytes
- senescent cells
- macrophages
- structural cells

**Senescent cells**
- adipocytes
- preadipocytes
- senescent cells
- macrophages
- structural cells

**Adipocytes**
- preadipocytes
- senescent cells
- macrophages
- structural cells

**Aged cells**
- macrophages
- senescent cells
- preadipocytes
- adipocytes
- structural cells

**Immunologic and Inflammatory factors**
- TGFβ
- IL-1β
- IFNγ
- PGE2
- MMPs
- GDF15

**Organ system dysfunction**
- Cognitive function decline
- Frailty
- Obesity
- Vulnerability to infections

**Inflammageing**
- Low-grade systemic inflammation
- Cell reprogramming
- Immune cell infiltration
- Ageing-associated morbidities

**Immunoregulatory factors**
- TGFβ
- IL-1β
- CXCL1, CXCL8
- MMPs
- GDF15, PGE2

**Cytokines**
- IL-1β
- IFNγ
- PGE2

**Chemokines**
- CCL2, CCL5
- CXCL1, CXCL8

**Inflammageing**
- Low-grade systemic inflammation
- Cell reprogramming
- Immune cell infiltration
- Ageing-associated morbidities

**Effects of ageing on T cells**

**Age-associated CD8**+ **memory T cells.** Essential for anti-viral and antitumour immunity, CD8+ T cells are substantially affected by ageing in both mice and humans. In general, older age has been associated with a loss of naive CD8+ T cells coupled with an increase in memory subsets and an overall increase in clonally expanded CD8+ T cells. Most recently, age-associated remodelling of the mouse immune T cell compartment was comprehensively characterized by scRNA-seq and single-cell T cell receptor (TCR) sequencing of four tissues (spleen, peritoneum, lungs and liver). These data confirmed a decrease in the abundance of naive CD8+ T cells across the four tissues and defined a distinct age-associated CD8+ T cell population (referred to by the study authors as “Taa cells”) uniquely characterized by co-expression of the checkpoint receptor PD1 and the transcription factor TOX. Although absent in young mice, these age-associated PD1+PD1- TOX+CD8+ T cells accumulate in multiple tissues in old animals, including the spleen, liver, lungs and fat. They constitute up to 60% of all CD8+ T cells in these tissues and are a striking example of systematic changes in the ageing immune system that affect multiple tissues. This observation is consistent with the organ-wide single-cell resolution profiling data produced by the Tabula Muris Senis Consortium and Calico Life Sciences, although these studies did not perform a detailed analysis of CD8+ T cell ageing.

The age-associated CD8+ T cells exhibit a phenotype of T cell exhaustion characterized by high expression levels of TOX, PD1, lymphocyte activation gene 3 protein (LAG3) and T cell immunoreceptor with immunoglobulin and ITIM domains (TIGIT), which generally suggests a loss of classical effector functions. However, upon TCR stimulation, they produce a distinct set of pro-inflammatory molecules such as granzyme K (GZMK) and CCL5. This observation suggests a functional role for these cells in murine ageing as a source of pro-inflammatory signals that might be particularly important in the context of old tissue remodelling or under conditions of an acute infection.

In humans, most of the studies focus on the ageing of circulating CD8+ T cells and use mass cytometry and scRNA-seq to analyse peripheral blood mononuclear cells (TABLE 1). For example, recent scRNA-seq studies of healthy donors reported a twofold decrease in the naive CD8+ T cell fraction of peripheral blood mononuclear cells from old individuals compared with young individuals. Another study used cellular indexing of transcriptomes and epigenomes by sequencing (CITE-seq) to characterize old and young peripheral blood mononuclear cells and also demonstrated an age-associated decrease in the abundance of naive CD8+ T cells and significant remodelling of the T cell memory compartment. Specifically, in healthy lean individuals, GZMK-expressing, but not granzyme B (GZMB)-expressing, CD8+ effector memory T cells (TEM cells) proportionally increase in abundance with age relative to overall CD8+ T cells. A recent mass cytometry study of a large cross-sectional...
## Changes in main immune cell populations in ageing

| Immune cell population | Age-associated changes | Methods used for analysis |
|------------------------|------------------------|---------------------------|
| **Mice**               |                        |                           |
| Monocytes              | Increased abundance in the spleen and blood\(^{155,156}\) | Flow cytometry            |
|                       | Decreased abundance of alveolar macrophages with reduced proliferative capacity and phagocytosis\(^{5,150,151}\) |                           |
|                       | Increased abundance of Fn1\(^-\) Ccr2\(^-\) interstitial macrophages\(^{5,27}\) |                           |
|                       | Decreased abundance of peritoneal macrophages\(^{5,96}\) |                           |
|                       | Increased abundance of CX, CR1\(^-\) macrophages in the liver\(^{9}\) and kidney\(^{152}\) |                           |
|                       | Increased abundance of fetal-derived macrophages in mammary tissue\(^{38}\) |                           |
| Macrophages           | Decreased abundance of cDC1s in the spleen\(^{25}\) | scRNA-seq, flow cytometry, microscopy |
|                       | Decreased abundance of cDC2s in the peritoneum\(^{25}\) |                           |
|                       | Decreased abundance of pDCs in the spleen\(^{5,153}\) and liver\(^{25}\) |                           |
| DCs                   | Increased abundance of age-associated B cells in the spleen and meninges\(^{50,51,90}\) | scRNA-seq, flow cytometry |
|                       | Increased abundance of plasma B cells in the kidney, adipose tissue and bone marrow\(^{55}\) |                           |
|                       | Accumulation of Apoe\(^-\) age-associated B cells in the spleen\(^{57}\) |                           |
|                       | Accumulation of Zbtb32\(^-\) age-associated B cells in the spleen\(^{25}\) |                           |
|                       | Increased abundance of Zcwpw1\(^-\) B1 cells in the peritoneum\(^{25}\) |                           |
| Neutrophils           | Increased abundance of Vγ6\(^\dagger\) cells in lymph nodes\(^{52}\) | scRNA-seq, flow cytometry |
|                       | Decreased abundance of Sox4\(^\dagger\) cells in the spleen\(^{25}\) |                           |
|                       | Increased abundance in the lungs and liver\(^{25}\) |                           |
| ILCs                  | Decreased abundance of NK cells in the spleen, peritoneum, lungs and liver\(^{25}\) | scRNA-seq, flow cytometry, microscopy |
|                       | Increased abundance of NK cells in neurogenic niches\(^{47}\) |                           |
|                       | Decreased abundance of ILC1s in the liver\(^{25}\) |                           |
|                       | Decreased abundance of ILC2s in the lungs\(^{25}\) |                           |
|                       | Increased abundance of ILC2s in the brain\(^{51}\) |                           |
| B cells               | Increased abundance of age-associated B cells in the spleen and meninges\(^{50,51,90}\) | scRNA-seq, flow cytometry |
|                       | Increased abundance of plasma B cells in the kidney, adipose tissue and bone marrow\(^{55}\) |                           |
|                       | Accumulation of Apoe\(^-\) age-associated B cells in the spleen\(^{57}\) |                           |
|                       | Accumulation of Zbtb32\(^-\) age-associated B cells in the spleen\(^{25}\) |                           |
|                       | Increased abundance of Zcwpw1\(^-\) B1 cells in the peritoneum\(^{25}\) |                           |
| γδ T cells            | Increased abundance of Vγ6\(^\dagger\) cells in lymph nodes\(^{52}\) | scRNA-seq, flow cytometry |
|                       | Decreased abundance of Sox4\(^\dagger\) cells in the spleen\(^{25}\) |                           |
|                       | Increased abundance in the lungs and liver\(^{25}\) |                           |
| CD4\(^+\) T cells     | Decreased abundance of naive cells in the spleen, peritoneum and lungs\(^{55,154}\) | scRNA-seq, flow cytometry |
|                       | Increased abundance of exhausted-like PD1\(^\dagger\) cells in the spleen, peritoneum, lungs and liver\(^{55,58}\) |                           |
|                       | Increased abundance of cytotoxic cells in the spleen\(^{69}\) |                           |
|                       | Increased abundance of activated T\(^\text{reg}\) cells in the spleen\(^{55,98,154}\) and adipose tissue\(^{55}\) |                           |
| CD8\(^+\) T cells     | Decreased abundance of naive cells in the spleen, peritoneum and lungs\(^{55,156}\) | scRNA-seq, flow cytometry |
|                       | Increased abundance in neurogenic niches\(^{157}\) |                           |
|                       | Increased abundance of virtual memory cells in the spleen, blood and lymph nodes\(^{158,158}\) |                           |
|                       | Increased abundance of a CD49d\(^\dagger\) subset in the spleen\(^{159}\) |                           |
|                       | Accumulation of exhausted-like PD1\(^\dagger\) cells in the spleen, peritoneum, lungs, liver, kidney, adipose tissue, meninges and blood\(^{55,90}\) |                           |
| Humans                | Increased abundance in the blood\(^{55,151}\) | Flow cytometry            |
|                       | Increased abundance of non-classical CD14\(^-\)CD16\(^-\) cells in the blood\(^{55,152}\) |                           |
| DCs                   | Decreased abundance of pDCs in the blood\(^{152}\) | Flow cytometry            |
| B cells               | Decreased abundance of B1 cells (CD19\(^-\)CD20\(^-\)CD27\(^-\)CD38\(^\text{low/mid}\)CD43\(^+\)) in the blood\(^{155}\) | scRNA-seq, flow cytometry |
|                       | Decreased abundance of the ZBTB32\(^-\) (CD27\(^-\)CD38\(^\text{low/mid}\)) subset in the blood\(^{25}\) |                           |
| MAIT cells            | Increased abundance in the blood\(^{53,154}\) | scRNA-seq, flow cytometry |
| γδ T cells            | Decreased abundance in the blood\(^{155}\) | scRNA-seq, flow cytometry |
|                       | Decreased abundance of Vδ2-Vγ9\(^\dagger\) cells in the blood\(^{55,156,157}\) |                           |
|                       | Decreased abundance of SOX4\(^\dagger\) cells in the blood\(^{55}\) |                           |
Table 1 (cont.) | Changes in main immune cell populations in ageing

| Immune cell population | Age-associated changes | Methods used for analysis |
|------------------------|------------------------|---------------------------|
| **Humans (cont.)**     |                        |                           |
| CD4+ T cells           | Decreased abundance of naive cells in the blood\(^{25,61,137,138}\) | scRNA-seq, flow cytometry |
|                        | Increased abundance of T\(_{EM}\) cells in the blood\(^{118}\) |                           |
|                        | Decreased abundance of recent thymic emigrants in the blood\(^{17}\) |                           |
|                        | Decreased abundance of an interferon-activated subset in the blood\(^{17}\) |                           |
|                        | Increased abundance of the cytotoxic subset in the blood (supercentenarians)\(^{37}\) |                           |
| CD8+ T cells           | Decreased abundance of naive cells in the blood\(^{12,61,138}\) | scRNA-seq, flow cytometry, mass cytometry |
|                        | Increased abundance of T\(_{EM}\) cells in the blood\(^{75,118}\) |                           |
|                        | Increased abundance of the GZMK\(^{+}\)(CD28\(^{-}\)CD57\(^{-}\)) subset of T\(_{EM}\) cells in the blood\(^{35}\) |                           |
|                        | Increased abundance of the CD57\(^{+}\) subset in the blood\(^{40}\) |                           |

\(cDC1\), conventional type 1 dendritic cell; \(cDC2\), conventional type 2 dendritic cell; CX\(_C\)R1, CX\(_C\)C chemokine receptor 1; DC, dendritic cell; GZMK, granzyme K; ILC, innate lymphoid cell; ILC1, group 1 innate lymphoid cell; ILC2, group 2 innate lymphoid cell; MAIT cells, mucosal associated invariant T cells; NK, natural killer; pDC, plasmacytoid dendritic cell; scRNA-seq, single-cell RNA sequencing; T\(_{EM}\) cell, effector memory T cell; T\(_{reg}\) cell, regulatory T cell.

Ageing cohort of 156 healthy individuals confirmed these observations and found that GZMK-expressing CD8\(^+\) T cells gradually increase in abundance in the circulation with age along with CD8\(^+\) central memory T cells\(^{25}\). Of note, GZMK\(^{+}\)CD8\(^+\) T cells express CD28 but lack CD57 and are distinct from CD45RA\(^{+}\) TEM\(_{CD8} \) cells — the age-associated expansion of which was reported in cytokemalovirus-exposed older individuals (range 21–96 years)\(^{25}\) but was not significant in a cohort of healthy donors (range 25–80 years)\(^{61}\).

Although GZMK-expressing human CD8\(^+\) TEM\(_{CD8} \) cells are similar to the mouse age-associated PD1\(^{+}\)TOX\(^{+}\) GZMK\(^{+}\)CD8\(^+\) T cells in some aspects (such as high expression of EOMES and GZMK), in humans this cell subpopulation cannot be identified solely on the basis of expression of PD1 (REFS\(^{25,61}\)). This difference between mouse and human CD8\(^+\) T cell ageing landscapes might arise because, unlike laboratory mice, humans experience multiple immune challenges during their lifespan, which forms a broader and more phenotypically diverse pool of antigen-experienced memory T cells.

**Effect of ageing on CD4\(^+\) T cells.** Similarly to CD8\(^+\) T cells, the proportion of naive CD4\(^+\) T cells declines and the memory population expands with age in mice and humans\(^{25,61,64}\). This switch to a memory phenotype seems to be driven by a combination of reduced thymic output and cumulative effect of lifelong antigen exposure as CD4\(^+\) T cells with transgenic TCRs tend to maintain a naive phenotype even in aged mice\(^{65}\). Another long-known feature of age-associated CD4\(^+\) T cell remodelling is increased abundance of regulatory T cells (T\(_{reg}\) cells) in the spleen and lymph nodes\(^{61–68}\). However, this increase in abundance of T\(_{reg}\) cells was not observed in ageing lungs or liver of mice\(^{69}\). The accumulation of splenic T\(_{reg}\) cells has been suggested to play a role in restricting immune responses in aged animals (such as compromising control of *Leishmania major* infection)\(^{69,70}\). A recent study used scRNA-seq to dissect the ageing of splenic CD4\(^+\) T cells in mice and reported that the age-associated increase is not a feature of all T\(_{reg}\) cells but is instead attributed to a subpopulation of activated T\(_{reg}\) cells\(^{69}\). These activated T\(_{reg}\) cells are defined by over-expression of Foxp3, Cd81, Cd74 and genes encoding activation markers (such as **Tnfrsf9**\(^{70}\)). Activated T\(_{reg}\) cells are approximately ten times more abundant in old mice and are superior in suppressive activities in comparison with resting T\(_{reg}\) cells\(^{69}\).

The accumulation of anti-inflammatory activated T\(_{reg}\) cells is paralleled by a marked expansion of so-called cytotoxic CD4\(^+\) T cells. These cells are characterized by the expression of Eomes and Gzmk and can produce not only IFN\(_y\) and tumour necrosis factor (TNF) but also cytotoxic GzmB and perforin upon stimulation\(^{69}\). Similarly to activated T\(_{reg}\) cells, the cytotoxic CD4\(^+\) T cell subpopulation was not associated with ageing before the single-cell study but was known in other physiological contexts (reviewed by Takeuchi and Saito\(^{70}\)). Strikingly, scRNA-seq of ageing human CD4\(^+\) T cells identified a similar subset of cytotoxic CD4\(^+\) T cells that are profoundly expanded at the extreme end of the longevity spectrum in individuals who reached the age of 110 years — supercentenarians\(^{72}\). The matching single-cell TCR sequencing data suggested that clonal expansion of CD4\(^+\) T cells with age is restricted primarily to the subpopulation of cytotoxic CD4\(^+\) T cells. By contrast, other T helper cell subsets remain clonally diverse, even in supercentenarians\(^{72}\). Interestingly, cytotoxic CD4\(^+\) T cell subpopulations are also more abundant in older patients with COVID-19 (range 32–91 years) with moderate symptoms compared with individuals with severe disease\(^{71}\), which might indicate that these cells can be beneficial during viral infections.

Finally, ageing is also characterized by increased expression of exhaustion markers by splenic CD4\(^+\) T cells in mice. Although the concept of T cell exhaustion is less well established for CD4\(^+\) T cells than for CD8\(^+\) T cells, an age-associated increase in the abundance of CD4\(^+\) T cells with high levels of inhibitory receptors (such as PD1 and LAG3) has been observed\(^{72,74}\). PD1\(^{+}\)-CD4\(^+\) T cells have a memory phenotype and are hyporesponsive to TCR stimulation\(^{71}\), which matches the canonical
features of exhausted CD8+ T cells. Exhausted-like CD4+ T cells constitute about 30% of all CD4+ T cells in the spleen of aged mice. Of note, age-associated changes in CD4+ T cell subpopulations are not universal across the tissues. scRNA-seq profiles of immune cells across four mouse tissues corroborated the data on splenic CD4+ T cell remodelling and revealed that while accumulation of PD1+ cells was observed in all tissues considered, the expansion of Treg cells was evident only in the spleen.

Age-related cell-intrinsic deficiencies of naive CD4+ T cells have also been described in studies that used aged mice with a transgenic TCR to investigate CD4+ T cell ageing with experimentally controlled antigen exposure.

**Expanded cell subsets**

- Oligoclonal PD1+CD8+ T cells (all tissues)
- PD1+CD4+ T cells (spleen, liver, fat)
- Activated Treg cells (PD1+CCR8+)
- Age-associated B cells (Zbtb32+Apoe+Tbx21+)
- CD38+ macrophages (ICAM2
- Interstitial macrophages (FN1+CCR2+ profibrotic?)

**Reduced cell subsets**

- B1 cells (Zcwpw1+Cd54+)
- Alveolar macrophages (inhibited proliferation)
- NK cells
- NK cells
- NK cells
- ILC2s

**Fig. 2** Altered immune cell populations in ageing. Single-cell techniques identified expanded and reduced immune cell populations with distinct phenotypes in multiple organs of old mice. Exhausted PD1+TOX+CD8+ T cells and activated PD1+CD4+ T cells accumulate across multiple tissues in ageing, whereas natural killer (NK) cells and innate lymphoid cells (ILCs) are among the cell subsets that are reduced in abundance in various organs. Functionally and phenotypically similar CD38+ macrophages expand in metabolically active organs — liver and fat — of old mice. Unlike these cell subsets commonly affected by age in many tissues, alterations of other immune cell populations show organ-specific patterns in ageing. For example, activated regulatory T cells (Treg cells) increase in abundance in the spleen of old mice. Tissue-resident alveolar macrophages decrease in abundance and fibronectin 1-positive (FN1+), CC-chemokine receptor 2-positive (CCR2+) interstitial macrophages increase in abundance in the lungs of old mice. Distinct subsets of age-associated B cells accumulate in the spleen (B cells expressing Zbtbb32, Apoe and Tbx21) and peritoneal cavity (B1 cells expressing Zcwpw1 and Cita4) of aged mice. cDC1, conventional type 1 dendritic cell; ICAM1, intercellular adhesion molecule 1; ILC1s, group 1 innate lymphoid cells; ILC2s, group 2 innate lymphoid cells.

These studies demonstrated that aged naive CD4+ T cells are less responsive to activation in vivo and in vitro than their young counterparts; they have decreased proliferation rates and produce less IL-2. The defect in TCR responsiveness seems to be specific for CD4+ T cells that have spent an extended time in circulation because CD4+ T cells from young thymectomized mice exhibit this phenotype, but recent immigrants from either young or aged bone marrow do not. Furthermore, a recent study used scRNA-seq to demonstrate that the responses of naive CD4+ T cells isolated from the spleen of old mice to in vitro stimulation are more heterogeneous and variable on the transcriptional level than those of naive CD4+ T cells from young mice, suggesting that activation programmes of aged CD4+ T cells are less tightly regulated.

An improper activation has been suggested to drive an incomplete differentiation of aged CD4+ T cells into Thelper 1 (T1) and T2 effector cells. At the same time, the ability of aged CD4+ T cells to acquire a T17-like phenotype is not compromised, creating an imbalance and a skew towards antimicrobial responses. In bulk, splenic CD4+ T cells from aged mice and humans produce much more IL-17 than young controls, which might reflect an age-associated accumulation of T17-type memory cells. As T17 cells have been implicated in the development of multiple autoimmune conditions, including inflammatory bowel diseases, one can speculate that the second peak of inflammatory bowel disease prevalence that occurs in older adults might be partially ascribed to this skewing of T cell populations. Indeed, transfer of aged CD4+ T cells to Rag1-/- recipients induced severer colitis than transfer of cells from a young donor. However, further research is required to prove a causal relationship between these events. IL-17-producing cells are also over-represented among γδ T cells in lymph nodes of aged mice, substituting IFNγ-producing effector γδ T cells, which are more abundant in young animals. Numbers of γδ T cells change with age in a tissue-dependent manner, as demonstrated by a recent single-cell profiling study of multiple mouse tissues. For example, γδ T cells are less frequent in the aged liver and lungs but not in the spleen. However, under conditions of acute lung inflammation caused by a murine betacoronavirus, old mice show lower numbers of lung γδ T cells, which could be restored by a ketogenic diet.

Altogether, the described age-associated differences in T cell populations appear to be partly responsible for altered humoral responses in old animals. It has been shown that CD4+ T cell help is required to launch a germinal centre reaction and produce high-affinity antibodies. Eaton et al. used an adoptive transfer model to compare germinal centre reactions facilitated by naive CD4+ T cells from either young mice or old mice with transgenic TCRs. In that study, cell-intrinsic changes in naive CD4+ T cells from old donors led to fewer antigen-specific B cells in the germinal centre and reduced production of class-switched antibodies. By contrast, naive CD4+ T cells from young mice were sufficient to elicit robust responses in the aged environment. These observations highlight the importance of
CD4+ T cell ageing biology for developing vaccination strategies customized to the state of the aged population.

**TCR repertoire.** Early studies of TCR repertoire changes during ageing — first using TCRβ chain-specific antibodies92 then by bulk TCR sequencing — suggested that TCR diversity loss is associated with the increased prevalence of memory T cells that undergo clonal expansion87,93. However, it can also be affected by a modest decrease in TCR diversity of naive T cells88–90. Overall, CD8+ T cells show a greater increase in clonality than CD4+ T cells, which is consistent with the more considerable reduction in naive CD8+ T cell subpopulations with ageing. scRNA-seq technology provides a unique opportunity to investigate TCR repertoire complexity via direct sequencing of paired α-chains and β-chains at single-cell resolution, which unequivocally defines TCR clones — a feature not available in bulk level analysis, which often uses only β-chain sequences. In mice, it was recently demonstrated that the accumulation of highly clonal age-associated PD1+TOX+CD8+ T cells is responsible for the clonality increase in the CD8+ T cell compartment91. The clonal expansion of these cells is ‘idiiosyncratic’91, meaning that individual mice differ in the specific TCRαβ chains used by the expanded T cell clones. Therefore, there might not be a shared ‘ageing antigen’ sensed by T cells in all ageing mice. Further studies are needed to decipher single-cell clonality landscapes in older individuals and identify specific age-related antigens that drive the accumulation of distinct memory T cell subsets.

**Effects of ageing on B cells.** Remodelling of adaptive immunity during ageing also includes altered B cell composition and function. A phenotypically and functionally separate subset of B cells has been shown to expand in the spleen and bone marrow of aged mice91–93. These cells are distinct from the conventional naive and memory B cells and have been termed ‘age-associated B cells’. Hao et al. defined age-associated B cells as CD43+CD21/CD35+CD23+ B cells92, whereas Rubtsov et al. identified age-associated B cells as CD11b+CD11c+ B cells93. Interestingly, age-associated B cells express Tbx21 (which encodes T-bet), which makes them similar to the B cells that have been shown to contribute to lupus-like autoimmunity in mice94. Although the age-related factors that prime the expansion of these B cell subsets are not entirely understood, a body of evidence suggests that damage-associated molecular patterns, such as debris and chromatin from apoptotic cells, initiate age-associated B cells via the Toll-like receptor 7 (TLR7) or TLR9 axis95. Once age-associated B cells are generated, their survival in old mice requires IFNγ signalling96. Age-associated B cells contribute to inflammaging by altering immune homeostasis. For example, age-associated B cells secrete IL-4 and IL-10 on activation97 and can present antigens to T cells98.

Over the past 3 years, several scRNA-seq studies reported age-associated dynamics of B cells across multiple tissues in mice99,100,101. In a study profiling both immune and non-immune cells from three tissues of young and old mice, Kimmel et al. characterized a distinct cluster of B cells that increases in abundance with age in the spleen7. This cluster is identified by the expression of the apolipoprotein gene *ApoE*, yet the precise identity of this B cell subset in terms of immunological nomenclature has not been specified. The aforementioned scRNA-seq profiling study of mouse immune cells in four tissues102 defined an age-associated population of conventional splenic B cells that express the transcription factor genes *Tbx21* and *Zbtb32* and genes encoding other markers of age-associated B cells (such as *Ccxr3* [REF93]). This *Zbtb32*+ B cell subset also highly expresses *ApoE*, suggesting a similarity with the age-associated B cell population reported by Kimmel et al.102. Single-cell analysis of B cells in the meninges revealed *ApoE*-expressing age-associated B cells that accumulate at the central nervous system borders in old mice102. Analysis of single-cell B cell receptor (BCR) sequencing data provided additional insights into this population. Indeed, splenic age-associated B cells have been recognized as cells displaying characteristics of antigen-experienced B cells with increased somatic hypermutation93. Consistently, the cluster of splenic *Tbx21*-expressing age-associated B cells demonstrates increased BCR clonality while still preserving BCR repertoire diversity7.

Another study that used a combination of scRNA-seq and single-cell BCR sequencing also reported a progressive increase in B cell clonality in ageing mice93. This increase was attributed to a cluster of cells expressing markers of plasma B cells *Jchain*, *Derl3* and *Deri3* found in the spleen, bone marrow, kidney and fat of aged mice, albeit in tiny proportions (~0.1% in these tissues)7. Moreover, tracing the cells from this cluster across the mouse lifespan showed that, similar to age-associated B cells, age-associated plasma B cells initially arise in the bone marrow and spleen and subsequently increase in abundance in peripheral tissues, such as adipose tissue and kidneys, later in life7. Functionally, plasma B cells can produce pro-inflammatory cytokines and enhance myelopoiesis in the ageing bone marrow7. Interestingly, a phenotypically similar population of plasma B cells increases in abundance in adipose tissue of older humans (>70 years)7.

A focused scRNA-seq analysis of immune cells in the peritoneal cavity (a location not included in *Tabula Muris Senis*) identified B1-like cells as heavily accumulating with age and characterized by a very limited BCR repertoire7. Highly enriched in *V H4*, 11-2 chains, these BCRs recognize the cell membrane component phosphatidylcholine95. Similarly to splenic age-associated B cells, peritoneal B1-like cells express the transcription factor gene *Zbtb32* yet represent a distinct transcriptional cluster characterized by high levels of an exclusive set of genes, including *Zcwpw1* and *Clta4* [REF5,101]. The Biragyn group showed an age-associated accumulation of B1-like cells, which were identified in old mice and humans by surface expression of the TNF ligand superfamily member 4–1BBL29,102,103. Functionally, B1-like cells become more activated with age, serve as potent inducers of cytotoxic CD8+ T cells and respond to commensal bacteria29,102,103. The insights from single-cell studies expand our understanding of age-associated B cell heterogeneity and point to potential antigen specificity of age-associated B cell subsets and their tissue-specific inflammatory functions in old organisms.
Role of non-immune cells in immune ageing

**Immune functions of stromal cells in ageing.** Immune function is not limited to cells of haematopoietic origin, and multiple types of stromal and epithelial cells are equipped to respond to pathogens and partake in immune homeostasis. For instance, fibroblasts are critical structural components of most tissues and a stromal cell type extensively studied in pathological states and ageing. Transcriptional analysis showed a loss of cell identity by dermal populations of old fibroblasts accompanied by upregulation of inflammatory genes in mice. Similarly, an analysis of transcriptomic and epigenomic changes in aged fibroblasts identified an increased abundance of ‘activated fibroblasts’ (for example, fibroblasts involved in tissue repair), including pathways related to cytokine signalling and inflammation. In line with these observations in mice, scRNA-seq analysis of human skin fibroblasts demonstrated a higher expression of genes contributing to low-grade inflammation in distinct populations of aged fibroblasts and predicted a decline in cell–cell communications between fibroblasts and other cell types in aged skin.

Endothelial cells line vascular beds and regulate immune trafficking. Ageing appears to substantially modify the inflammatory transcriptional signatures of endothelial cells in various tissues. For instance, single-cell transcriptional profiling of endothelial cells in aged mouse brain prominently implicated altered inflammation and cytokine signalling pathways associated with neurodegenerative diseases. Moreover, in mice, a single-cell analysis revealed transcriptional alterations in aged liver sinusoidal endothelial cells, which are essential regulators of immune defence. Finally, communication between endothelial cells and various immune cell types (including macrophages and T cells) is markedly altered in old tissues, including liver and fat, as was revealed by a single-cell transcriptional analysis in rats.

**Senescent cell contribution to inflamaging.** Ageing is characterized by a gradual accumulation of senescent cells throughout the body. Senescence is a terminal stage of cell differentiation, often acquired by damaged or stressed cells, that leads to permanent withdrawal from the cell cycle and a switch to a distinct secretome. This secretome, termed the ‘senescence-associated secretory phenotype’ (SASP), involves the secretion of multiple chemokines, cytokines and growth factors and contributes to low-grade systemic inflammation in ageing. Although a core transcriptional signature of senescent cells in vitro has been described, no robust approach to determine the transcriptional landscape of senescent cells in vivo has been developed to date. Moreover, senescent cells are profoundly variable in their expression of regulatory and SASP genes even in vitro. Nonetheless, recent scRNA-seq analyses provided some insights into the development of senescent cells in ageing mice. The scRNA-seq atlas of ageing mouse tissues (Tabula Muris Senis) showed that the fraction of cells expressing Cdkn2a (which encodes p16INK4A, a commonly used marker of senescent cells) increases in different tissues in old mice. However, another single-cell analysis found no difference in Cdkn2a expression between young mice and old mice, emphasizing current conceptual and technical limitations precluding identification of bona fide senescent cells in living organisms. A recent article reported scRNA-seq of p16INK4A+ senescent cells with heterogeneous senescence-associated phenotypes in a reporter mouse model. Interestingly, senescent cells include various cell types that are enriched in hepatic endothelium and renal proximal and distal tubule epithelia. Elimination of these p16INK4A+ senescent cells decreases immune infiltration in a model of metabolic liver disease. Further studies are needed to better understand how heterogeneous senescent cells contribute to low-grade chronic inflammation and reshape immunity in ageing.

**Interplay between immune and senescent cells.** Recognition and elimination of senescent cells by specialized immune cells is a crucial step in maintaining tissue homeostasis and preventing maladaptive inflammation (Fig. 5). Components of the SASP, such as CCL2 and CCL7, attract immune cells, including macrophages, NK cells and T cells. Senescent cells also recruit neutrophils, tissue infiltration of which is one of the hallmarks of immune ageing. Macrophages can physically interact with and eliminate senescent cells in developmental and pathological processes. Whether the impaired phagocytosis by aged macrophages is causative for senescent cell accumulation in old tissues remains to be clarified. NK cells recognize and kill senescent cells to coordinate tissue remodelling and limit fibrogenic responses to acute tissue damage. Intriguingly, the cytotoxic functions of NK cells and T cells appear to control the accumulation of senescent cells in ageing because mice lacking perforin.

---

**Box 3 | Cellular senescence and senescent cells**

Cellular senescence is a state of irreversible cell-cycle arrest due to cellular replicative limit, tumour suppression, inflammation, embryogenesis, wound healing or age-associated loss of regenerative capacity. Cellular senescence has been defined as a stress response to developmental and environmental stimuli, including genotoxic agents, hypoxia, mitochondrial dysfunction, oncogene activation, epigenetic modifiers and cytokines. Senescent cells are a product of cellular senescence and feature a combination of molecular and morphological characteristics. These cells undergo permanent cell-cycle arrest that is distinct from terminal differentiation and quiescence. They have rewired cell-cycle regulatory pathways with a specific phosphorylation profile of cyclin-dependent kinases (CDKs) and upregulated expression of the CDK2 inhibitor p21CIP1 (also known as CDKN1A) and the CDK4/CDK6 inhibitor p16INK4A (also known as CDKN2A). Although increased expression of p21CIP1 and p16INK4A is not unique to senescent cells and can be found in, for example, non-senescent macrophages, p21CIP1 and p16INK4A are often used both as markers of senescent cells and as targets to develop genetic models allowing selective elimination of senescent cells. Morphological characteristics of senescent cells include an enlarged cell size due to a dilated cytoplasmic domain and increased activity of lysosomal senescence-associated β-galactosidase — the most widely used histological biomarker of senescent cells. Other features of senescent cells include DNA, protein and lipid damage, dysfunctional mitochondria and lysosomes, and an altered chromatin landscape. Importantly, senescent cells have a specific secretory profile involving excessive secretion of pro-inflammatory cytokines and chemokines, growth modulators, angiogenic factors and matrix metalloproteinases, collectively named the ‘senescence-associated secretory phenotype’. Different components of the senescence-associated secretory phenotype mediate the communication of senescent cells with immune cells, affect the functions of stromal cells in tissues and contribute to the low-grade systemic inflammatory phenotype in ageing.
(the crucial effector of cytotoxicity) exhibit both higher senescent-cell tissue burden and chronic inflammation while they age. Consistent with this is the observation of decreased frequencies of NK cells and replacement of tissue-resident CD8+ T cells with exhausted-like PD1+ T cells in various tissues of aged mice. In addition to cytotoxic CD8+ T cells, at least in the context of cellular senescence in cancer, CD4+ T cells also provide immune surveillance of premalignant senescent cells and require macrophages to eliminate them. In mice, elimination of senescent cells via genetic manipulation can prevent or delay tissue dysfunction and extend 'healthspan' (the period of time an organism lives free of chronic diseases of ageing), suggesting that immune surveillance and elimination of senescent cells contribute to delay of age-related diseases. However, the immune cell-mediated elimination of senescent cells is context dependent and tightly regulated because a transitory accumulation of senescent cells is critical for tumour suppression and certain regenerative processes such as wound healing.

**Immune ageing in mice versus humans**

Despite tremendous progress in single-cell techniques and their application to mouse immune ageing in various tissues, current single-cell data in humans are...
Single-cell technologies have already significantly expanded the current paradigm of immune ageing. For example, they have made it possible to better understand the heterogeneity of human CD8+ T cell populations and distinguish between GZMK+ and GZMB+ T cell populations, revealing the GZMK+ cells as the crucial age-associated CD8+ T cell subset in humans. Single-cell analyses also defined cytotoxic CD4+ T cells as a key feature of immune ageing conserved between mice and humans. In addition, combined single-cell sequencing of RNA and antigen receptors expanded our capability to study TCR and BCR repertoires to an unprecedented level. One of the recent discoveries stemming from this expanded toolbox is the age-associated loss of TCR repertoire diversity in mouse CD8+ T cells appears to be driven primarily by the replacement of polyclonal naive T cells and T EM cells by oligoclonal PD1-TOX-CD8+ T cells. Finally, unbiased single-cell profiling of immune cells across multiple tissues led to the demonstration that a dietary intervention with a beneficial effect on healthspan not only reshapes age-disturbed immune landscapes but also specifically reprograms distinct pro-inflammatory macrophage subsets in rodents. These results suggest an intriguing possibility that some immune cell populations might be a future target for anti-ageing therapies.

The next-generation profiling techniques will allow further exploration of the many remaining questions in the field of immune ageing. For instance, the interaction between immune cells and the ageing environment is an important avenue for investigation, including the need to understand which processes are driven by immune cells and which reflect adaptation of immune cells to the aged environment. Spatial transcriptomics and imaging cytometry can reveal spatial relationships between immune cells and non-immune cells and promise to solve this puzzle in an unbiased manner. Also, tissue-resident immune cell populations are not sufficiently studied in ageing humans. Thus, spatial techniques will be an essential foundation for human immune ageing research. This approach is especially important for a better understanding of immune responses to pathogens in older individuals, in which the severity of some (but not all) viral and bacterial infections — such as pneumococcal disease, influenza virus and COVID-19 — is profoundly increased compared with that in younger adults. Single-cell analyses across individuals of different ages can distinguish between the effects of ageing and the effects of the infectious disease itself, which will help to distinguish between age-associated changes and the immune mechanisms of diseases such as COVID-19.

Another frontier in single-cell immunology is multiplexing of orthogonal approaches to characterize the ageing immune system. For instance, a simultaneous characterization of hundreds and thousands of genes and proteins in individual immune cells (for example, using CITE-seq) will deepen the immune profiling and facilitate prediction and validation of immune cell functions in ageing. The same approach is valid for a combination of transcriptomics and epigenomics of ageing immune cells. Parallel application of scRNA-seq and single-cell assay for transposase-accessible chromatin using sequencing (scATAC-seq) has already demonstrated encouraging potential to decipher new regulatory programmes in age-associated immune cell populations of CD8+ T cells in mice and humans. The same approach is valid for a combination of transcriptomics and epigenomics of ageing immune cells. Parallel application of scRNA-seq and single-cell assay for transposase-accessible chromatin using sequencing (scATAC-seq) has already demonstrated encouraging potential to decipher new regulatory programmes in age-associated immune cell populations of CD8+ T cells in mice and humans. The same approach is valid for a combination of transcriptomics and epigenomics of ageing immune cells.
Table 2 | Single-cell transcriptomic and epigenetic datasets of immune ageing

| Dataset*                                                                 | Species and organs                                                                 | Details                                                                 | Ref. |
|-------------------------------------------------------------------------|------------------------------------------------------------------------------------|-------------------------------------------------------------------------|------|
| Profiling of single cells across the lifespan in mice (Tabula Muris Sensis) | Mouse immune and non-immune cells from the bladder, bone marrow, brain, fat, heart and aorta, kidney, large intestine, limb muscle, diaphragm, liver, lung, mammary gland, pancreas, skin, spleen, thymus, tongue and trachea | scRNA-seq of total cells from male and female C57BL/6J mice from six age groups: 1 month (the equivalent of early human childhood) to 30 months (the equivalent of a human centenarian). This dataset contains scTCR-seq and scBCR-seq data. | 26   |
| Profiling of immune cells in young and old mice                         | Sorted mouse immune cells from the spleen, liver, lung and peritoneal cavity       | scRNA-seq of CD45+ cells from young (3–4 months) and aged (17–18 months) male C57BL/6J mice. This dataset contains scTCR-seq and scBCR-seq data. Because of selective enrichment for CD45+ cells, this dataset captures rare immune cell populations in tissues. | 25   |
| Mouse ageing cell atlas in three organs                                 | Mouse immune and non-immune cells from the spleen, lung and kidney                 | scRNA-seq of total cells from young (7 months) and aged (22–23 months) male C57BL/6J mice. This dataset contains scTCR-seq and scBCR-seq data. | 57   |
| Mouse ageing lung atlas                                                 | Mouse single-cell suspensions of whole lungs                                       | scRNA-seq of total cells from young (3 months) and aged (24 months) male C57BL/6N mice. | 27   |
| Immune cells in young and old mice                                      | Mouse cells from the meninges, bone marrow and blood                               | scRNA-seq of cells from young (3 months) and aged (25 months) female C57BL/6J mice. This dataset contains scBCR-seq data. | 96   |
| Analysis of immune cells in young and aged mice                         | Mouse immune cells from the spleen and dentate gyrus                               | scRNA-seq of CD45+ cells from young (3 months) and old (18 months) C57BL/6J mice. | 47   |
| Ageing-associated alterations in mammary cells in mice                  | Epithelial and stromal cells from mouse mammary tissues                            | scRNA-seq of cells from young (3–4 months) and aged (13–14 months) virgin female C57BL/6J mice. | 28   |
| Ageing-associated alterations in adipose tissue in mice                 | Tissue-resident immune cells from young and old gonadal adipose tissue             | scRNA-seq of CD45+ cells from young (2–3 months) and aged (18–24 months) male C57BL/6J mice. | 50   |
| Effect of ageing on CD4+ T cells in mice                               | Mouse CD4+ T cells from the spleen                                                 | scRNA-seq of CD4+ T cells from young (2–3 months) and aged (22–24 months) C57BL/6J mice. | 9    |
| Single-cell atlas of ageing in rats                                    | Rat immune and non-immune cells from the liver, fat, kidney, aorta, skin and bone marrow | scRNA-seq of total cells from young (5 months) and old (27 months) male and female Rattus norvegicus. One group of old rats was calorie restricted (70% of calories) starting at 18 months. | 35   |
| Cells from wounds of old mice                                          | Mouse cells from entire wounds                                                     | scRNA-seq of cells from skin wounds aged (24 months) C57BL/6J mice with fast-healing and slow-healing trajectories. | 109  |
| Cell composition in old neurogenic niches                              | Mouse cells from subventricular zone neurogenic niche                               | scRNA-seq of cells from subventricular zone young (3 months) and aged (28–29 months) male C57BL/6J mice. | 57   |
| PBMCs of healthy young and older individuals                           | Human PBMCs from healthy non-obese individuals                                     | scRNA-seq and CITE-seq (39 antibodies) of PBMCs from 11 young (25–29 years) and 10 older (62–70 years) male donors. This dataset contains scTCR-seq data. | 25   |
| PBMCs of young and older healthy adults with COVID-19                   | Human PBMCs                                                                        | scRNA-seq and scATAC-seq of PBMCs from young (20–45 years) and older (60–80 years) male and female donors — healthy or with COVID-19. This dataset contains scTCR-seq and scBCR-seq data. | 60   |
| PBMCs of supercentenarians and control adult individuals               | Human PBMCs from older individuals (controls) and supercentenarians               | scRNA-seq of PBMCs from five controls (50–80 years) and seven supercentenarians (110 years). This dataset contains scTCR-seq data. | 72   |
| Epigenetic landscapes of aged immune cells in mice and humans          | Mouse splenocytes and human PBMCs                                                  | scATAC-seq of mouse CD45+ cells from the spleen of young (3–4 months) and aged (17–18 months) male C57BL/6J mice and of human PBMCs from three young (25–29 years) and three older (62–70 years) healthy male donors. | 25   |

CITE-seq, cellular indexing of transcriptomes and epigenomes by sequencing; PBMCs, peripheral blood mononuclear cells; scATAC-seq, single-cell assay for transposase-accessible chromatin using sequencing; scBCR-seq, single-cell B cell receptor sequencing; scRNA-seq, single-cell RNA sequencing; scTCR-seq, single-cell T cell receptor sequencing. *Datasets can be explored online through an Atyomyx laboratory webpage.

of ageing depend on a combination of genotype, medical history and environment, it is crucial to understand how these factors impact the immune system during ageing. Furthermore, immune ageing can be studied in the context of various risk factors that affect healthspan. For example, genetic factors driving the variability of functional decline in ageing are recognized in multiple genetic association studies[148]. Future single-cell research can deepen our understanding of how these and other factors affect distinct populations of immune cells at different ages. In mouse models of immune ageing, this should prompt researchers to strictly control effects of ageing that are derivatives from associated physiological processes (for example, decreased activity and accumulation of adipose tissue). Similarly, single-cell analysis of the immune system from phenotypically and genetically
dive... different stages of the immune response to an infection in young hosts versus older hosts is a crucial future goal.

In conclusion, adding space, time, depth and diversity to single-cell research of immune ageing is the next goal of the field.

Published online 23 November 2021
This study reveals...
152. Baker, D. J. et al. Clearance of p16Ink4a-positive senescent cells delays ageing-associated disorders. Nature 479, 252–256 (2011).
153. Baker, D. J. et al. Naturally occurring p16Ink4a-positive cells in healthy lifespan. Nature 530, 184–189 (2016).
154. Demaria, M. et al. An essential role for senescent cells in the induction of immune resistance to cancer. J. Exp. Med. 204, 15–22 (2007).
155. Puchta, P. et al. TNF drives monocytic dysfunction with age and resembles p16Ink4a pneumococcal immunity. PLoS Pathog. 12, e1005658 (2016).
156. Loukov, D., Naidoo, A., Puchta, A., Marin, J. L. A. & Bowdish, D. M. E. Tumor necrosis factor drives increased splenic monocytes in old mice. J. Leuk. Biol. 100, 121–129 (2016).
157. Tong, Q.-F. et al. Human thymic involution and aging in humanized mice. Front. Immunol. 11, 1599 (2020).
158. Li, et al. Age-related human T cell subset evolution and senescence. Immunity. Aging 16, 24 (2019).
159. Jergovic, M., Contreras, N. A. & Nikolich-Zugich, J. Impact of CMV upon immune aging. facts and fiction. Med. Microbiol. Immunol. 208, 265–269 (2019).
160. Gordon, A. & Reingold, A. The burden of influenza: a complex problem. Curr. Epidemiol. Rep. 5, 1–9 (2018).
161. Johnson, R. W. et al. The impact of herpes zoster and immune age derived from high-dimensional longitudinal microenvironment shapes a dysfunctional response of alveolar macrophages in aging. J. Clin. Invest. 135, 291–300 (2016).
162. Shenoy, A. T. & Mizgerd, J. P. Seedy CD8 T cells. J. Immunol. 193, 1589–1592 (2014).
163. Kang, C. et al. The DNA damage response induces immune age. J. Exp. Med. 215, 3471–3479 (2018).
164. Colonna-Romano, G. et al. Impairment of gamma/δ T lymphocytes in elderly: implications for immunosenescence. Exp. Gerontol. 39, 1439–1446 (2004).
165. Tan, C. T. Y. et al. Vδ2-positive T cells show divergent trajectories during human aging. OncotARGET 7, 14906–14916 (2016).
166. Michishita, Y. et al. Age-associated alteration of γδ T cell repertoire and different profiles of activation-induced death of Vδ1 and Vδ2 T cells. Int. J. Hematol. 94, 240–250 (2011).
167. Oishi, Y. & Manabe, I. Macrophages in age-related chronic inflammatory diseases. NPJ Aging Mech. Dis. 2, 1–8 (2016).
168. Minhas, P. S. et al. Restoring metabolism of myeloid cells reverses cognitive decline in aging. Nature 590, 122–128 (2021).
169. Ferrucci, L. & Fabbri, E. Inflammaging: chronic inflammation in aging, cardiovascular disease, and frailty. Nat. Rev. Cardiol. 15, 505–522 (2018).
170. Desdin-Micó, G. et al. T cells with dysfunctional mitochondria induce multimorbidity and premature senescence. Science 368, 1571–1576 (2020).
171. This article shows that TFAM-deficient T cells with dysfunctional mitochondria increase inflammationaging, accelerate senescence and regulate organismal fitness and lifespan in mice.
172. Shenoy, A. T. & Mizgerd, J. P. Seedy CD8 T cells. J. Immunol. 193, 1589–1592 (2014).
173. Kang, C. et al. The DNA damage response induces immune age. J. Exp. Med. 215, 3471–3479 (2018).
174. Colonna-Romano, G. et al. Impairment of gamma/δ T lymphocytes in elderly: implications for immunosenescence. Exp. Gerontol. 39, 1439–1446 (2004).
175. Tan, C. T. Y. et al. Vδ2-positive T cells show divergent trajectories during human aging. Oncotarget 7, 14906–14916 (2016).
176. Michishita, Y. et al. Age-associated alteration of γδ T cell repertoire and different profiles of activation-induced death of Vδ1 and Vδ2 T cells. Int. J. Hematol. 94, 240–250 (2011).
177. Oishi, Y. & Manabe, I. Macrophages in age-related chronic inflammatory diseases. NPJ Aging Mech. Dis. 2, 1–8 (2016).
178. Minhas, P. S. et al. Restoring metabolism of myeloid cells reverses cognitive decline in aging. Nature 590, 122–128 (2021).
179. Ferrucci, L. & Fabbri, E. Inflammaging: chronic inflammation in aging, cardiovascular disease, and frailty. Nat. Rev. Cardiol. 15, 505–522 (2018).
180. Desdin-Micó, G. et al. T cells with dysfunctional mitochondria induce multimorbidity and premature senescence. Science 368, 1571–1576 (2020).
181. This article shows that TFAM-deficient T cells with dysfunctional mitochondria increase inflammationaging, accelerate senescence and regulate organismal fitness and lifespan in mice.
182. Shneyo, A. T. & Mizgerd, J. P. Seedy CD8 T cells. J. Immunol. 193, 1589–1592 (2014).
183. Kang, C. et al. The DNA damage response induces immune age. J. Exp. Med. 215, 3471–3479 (2018).
184. Colonna-Romano, G. et al. Impairment of gamma/δ T lymphocytes in elderly: implications for immunosenescence. Exp. Gerontol. 39, 1439–1446 (2004).
185. Tan, C. T. Y. et al. Vδ2-positive T cells show divergent trajectories during human aging. Oncotarget 7, 14906–14916 (2016).
186. Michishita, Y. et al. Age-associated alteration of γδ T cell repertoire and different profiles of activation-induced death of Vδ1 and Vδ2 T cells. Int. J. Hematol. 94, 240–250 (2011).
187. Oishi, Y. & Manabe, I. Macrophages in age-related chronic inflammatory diseases. NPJ Aging Mech. Dis. 2, 1–8 (2016).
188. Minhas, P. S. et al. Restoring metabolism of myeloid cells reverses cognitive decline in aging. Nature 590, 122–128 (2021).
189. Ferrucci, L. & Fabbri, E. Inflammaging: chronic inflammation in aging, cardiovascular disease, and frailty. Nat. Rev. Cardiol. 15, 505–522 (2018).
190. Desdin-Micó, G. et al. T cells with dysfunctional mitochondria induce multimorbidity and premature senescence. Science 368, 1571–1576 (2020).
191. This article shows that TFAM-deficient T cells with dysfunctional mitochondria increase inflammationaging, accelerate senescence and regulate organismal fitness and lifespan in mice.
192. Shneyo, A. T. & Mizgerd, J. P. Seedy CD8 T RM cells in aging lungs drive susceptibility to pneumonia and sequestration. Cell. Mol. Immunol. 18, 787–789 (2021).
193. Covarrubias, A. J. et al. Senescent cells promote tissue NAD+ decline during aging via the activation of CD38 macrophages. Nat. Metab. 2, 1265–1280 (2020).
194. Chiri, C. S. S. et al. CD38 ecto-enzyme in immune cells is induced during aging and regulates NAD+ and NNN levels. Nat. Metab. 2, 1265–1304 (2020).
195. Freund, A., Patil, C. K. & Campisi, J. p38MAPK is a novel DNA damage response-independent regulator of the senescence-associated secretory phenotype. EMBO J. 30, 1556–1564 (2011).
196. Acosta, J. C. et al. Chemokine signaling via the CXCR4 receptor rescues senescent cells. Cell 153, 1006–1018 (2008).
197. Kang, C. et al. The DNA damage response induces inflammationaging by inhibiting autophagy of GATA4. Science 349, eaa6512 (2016).
198. Gluck, S. et al. Intranuclear sensing of cytosolic chromatin fragments through cGAS promotes senescence. Nat. Cell Biol. 19, 1061–1070 (2017).
199. Dou, Z. et al. Cytoplasmic chromatin triggers inflammationaging and cancer. Nature 550, 402–406 (2017).
200. Sagiv, A. et al. NKG2D ligands mediate immunosurveillance of senescent cells. Aging 8, 528–534 (2016).