Aging increases cell-to-cell transcriptional variability upon immune stimulation

Celia Pilar Martinez-Jimenez#1,2, Nils Eling#1,3, Hung-Chang Chen1, Catalina A. Vallejos3,4, Aleksandra A. Kolodziejczyk2,3, Frances Connor1, Lovorka Stojic1, Timothy F. Rayner1, Michael J.T. Stubbington3, Sarah A. Teichmann2,3, Maike de la Roche1, John C. Marioni1,2,3, and Duncan T. Odom1,2

1University of Cambridge, Cancer Research UK Cambridge Institute, Robinson Way, Cambridge, CB2 0RE, UK
2Wellcome Trust Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, UK
3European Molecular Biology Laboratory, European Bioinformatics Institute (EMBL-EBI), Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SD, UK
4MRC Biostatistics Unit, Cambridge Institute of Public Health, Forvie Site, Robinson Way, Cambridge Biomedical Campus, Cambridge CB2 0SR, UK

# These authors contributed equally to this work.

Abstract

Aging is characterized by progressive loss of physiological and cellular functions, but the molecular basis of this decline remains unclear. We explored how aging impacts transcriptional dynamics using single-cell RNA-sequencing of unstimulated and stimulated naive and effector memory CD4+ T cells from young and old mice from two divergent species. In young animals, immunological activation drives a conserved transcriptomic switch resulting in tightly regulated gene expression, characterized by a strong up-regulation of a core activation program, coupled with a decrease in cell-to-cell variability. Aging perturbed the activation of this core program, and increased expression heterogeneity across populations of cells in both species. These discoveries suggest that increased cell-to-cell transcriptional variability will be a hallmark feature of aging across most, if not all, mammalian tissues.

The progressive decline of physiological and cellular functions (1) caused by aging is associated with complex effects on tissue-specific and species-specific gene expression levels (2, 3). For instance, aging affects distinct functional pathways even in closely related...
CD4⁺ and CD8⁺ T cells (4). Approaches analyzing the expression of sets of genes in single-cells have suggested that aging alters cell-to-cell transcriptional variability (5), though this may not be a universal attribute of aging (6). Whether cell-to-cell gene expression variability increases during aging on a genome-wide basis, particularly for dynamic activation programs, remains largely unexplored.

Naive CD4⁺ T cells are an excellent model system to evaluate the impact of aging on gene expression levels and cell-to-cell transcriptional variability. They are readily isolated as single, phenotypically homogeneous cells and can be stimulated into a physiologically-relevant activated transcriptional state in vitro. Naive CD4⁺ T cells are maintained in a quiescent state, and are essential for lifelong maintenance of adaptive immune function against infection and cancer (7). Comparing gene expression levels in matched tissues from different mammalian species is a powerful strategy to reveal conserved cell-type-specific regulatory programs (8) (9, 10), and has been successfully employed to identify a conserved set of immune activation genes in CD4⁺ T cells (11). However, it is not known whether conservation of gene expression levels is also reflected in cell-to-cell variability.

Here, we dissect the activation dynamics of naive CD4⁺ T cells at the single cell level during aging in two inbred mouse sub-species separated by 1 million years of divergence: the reference C57BL/6J, Mus musculus domesticus (B6), and Mus musculus castaneus (CAST). These mice have similar lifespans and aging phenotypes (12). We isolated naive CD4⁺ T cells and characterized their gene expression programs by single-cell RNA-sequencing (scRNA-seq) during aging in young (~3 months) and old (~21 months) individuals of each strain (Fig. 1A, (13)).

Purified naive CD4⁺ T cells were loaded directly into the Fluidigm C1, or were loaded three hours after stimulation in vitro with plate-bound anti-CD3ε/anti-CD28 antibodies (13). After extensive quality control and filtering (Fig. S1 and S2, (13)), a total of 1514 high-quality CD4⁺ T cell transcriptomes were analyzed with BASiCS (14) to identify differences in levels of gene expression and variability of gene expression. Naive CD4⁺ T cells cluster by species (Figs. 1B; S3A) with 15% of genes differentially transcribed between the two species (Figs. 1C; S3A, table S1). Species-specific genes were not enriched for any functional ontology and were not the result of errors in genome assembly or read mapping (Fig. S3A-D). Species-specific transcribed genes are more variable on a cell-to-cell basis than genes expressed in both species, consistent with neutral drift (Fig. 1D, Fig. S3E).

Functional CD4⁺ T cell transcriptional responses start with an early, targeted activation of translational machinery and cytokine networks, followed by large-scale transcriptional changes associated with lineage commitment (11, 15). We stimulated naive CD4⁺ T cells for three hours with plate-bound anti-CD3ε/anti-CD28 antibodies (Figs. 2A, S4A).

The resulting transcriptional response was highly coordinated, with 2063 genes (log₂FC>2) differentially expressed in CD4⁺ T cells upon activation in vitro (Figs. 2B, S4B). We observed a reduction in cell-to-cell transcriptional variability among genes whose transcription levels do not change after activation (Mann-Whitney-Wilcoxon test, p<10⁻¹⁰) (Fig. 2C). Transcriptionally variable genes in the unstimulated condition showed no
population substructure (Fig. S4C, (13)). Genes up-regulated after stimulation were expressed in more cells than downregulated genes (Figs. 2D; S4D).

The transcriptional switch driven by TCR engagement and co-stimulation included activation markers and translational machinery (11, 15) (Figs. 2E; S4E; table S2). Even genes not up-regulated after three hours of immune stimulation showed a marked decrease in their cell-to-cell variability, consistent with increased regulatory coordination (Fig. 2F). As expected, we observed the coordinated suppression of Sell (Cd62l) (Figs. 2G; S3D; table S2). CD4+ T cells isolated from CAST showed similar molecular phenotypes (Fig. S5).

We identified functional targets activated upon immune stimulation by searching for gene expression levels conserved between species and expressed in most cells (11). Therefore, we stimulated naive CD4+ T cells isolated from young CAST males (13). The 225 up-regulated genes shared between B6 and CAST were enriched for cellular processes activated by immune stimulation (Fig. 3, table S3) and were expressed across most CD4+ T cells. Species-specific genes (96 for B6, 75 for CAST) showed no enrichment for biological function and were more sporadically expressed. Thus, target genes involved in translational control and immune function represent the conserved early activation response signature.

Aging can cause a shift in the functional balance between self-renewal and differentiation in hematopoietic stem cells (16). We considered whether aging might similarly perturb the transcriptional response of CD4+ T cells to immune stimulation.

The global expression profiles of naive or activated CD4+ T cells did not change due to aging (Fig. S6A, B). Nevertheless, approximately 10% of genes are differentially expressed between cells from young and old mice (Fig. S6C, D). These genes are only expressed in a small subset of cells, and are not conserved between B6 and CAST (Fig. S6E, F). Most genes in the core-conserved program responded even after three hours of stimulation, irrespective of age (Fig. S7). However, the magnitude of the response was consistently reduced in older mice (Fig 4A).

The cell-to-cell transcriptional variability of the core activation program was higher in older animals (Mann-Whitney-Wilcoxon test; p<0.05, Fig. 4B, Fig. S7), independent of changes in gene expression levels. In both B6 and CAST mice, genes involved in the activation process are typically expressed in fewer cells in old mice (Fig. 4C).

Finally, we asked whether additional cellular components of the immune system also show increased transcriptional variability upon aging. We used a stringent gating strategy to isolate effector memory CD4+ T cells in young and old B6 mice (17) (Fig. S8, (13)). scRNA-seq experiments revealed that immune activation genes showed an increase in variability in effector CD4+ T cells from older animals, similar to naïve CD4+ T cells (Fig. S9). These results indicated that aging reduces the fraction of cells in which immune activation genes are up-regulated, thus increasing cell-to-cell heterogeneity and attenuating the response to stimulation across multiple CD4+ T cell subsets.

How cell-type-specific gene expression programs change during organismal lifespan has long been debated (5, 6) and what cell-to-cell transcriptome-wide differences accumulate
during aging are relatively unexplored (16). By activating naive CD4+ T cells and quantifying the transcriptional responses of hundreds of single-cells using scRNA-seq, we confirmed that translation processes and immune response genes are rapidly up-regulated (11) and discovered a reduction in cell-to-cell variability across thousands of transcripts. Thus, immune activation rapidly reduces transcriptional heterogeneity across the population of CD4+ T cells, revealing a regulatory strategy comparable to induced pluripotent stem cell reprogramming (18, 19).

Many attempts have been made to identify transcriptional signatures associated with aging (16, 20). Aging appears to have modest effects on mean expression levels in unstimulated and stimulated CD4+ T cells. However, regardless of species or analyzed cell-type, older mice showed substantially greater transcriptional heterogeneity; in simpler terms, older mice have more sporadic transcriptional responses. The discovery that CD4+ T cells from aged mice are less able to robustly up-regulate a core activation program may in part explain the aging associated decrease of immune function observed across mammals (21).

Our results indicate that in addition to transcriptional dysregulation and chromatin destabilization (1), increased cell-to-cell transcriptional variability is a major hallmark of aging.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

Thanks to R Miragaia, A Lun, A Scialdone and A Richard for input; A Mowbray, N Jacobs, M Clayton, M Mitchell for animal husbandry; S Lorenz and the Single Cell Genomics Core Facility at WTSI and the Genomics, Flow Cytometry, and BRU Cores at CRUK CI for technical support. Funded by the European Research Council (FC, TFR, DTO, SAT, MJTS), EMBO Young Investigators Programme (DTO), Cancer Research UK (HCC, MdLR, DTO, JCM), Janet Thornton Fellowship (WT098051) (CPJM), Sir Henry Dale Fellowship (WT107609, MdLR), EMBL (NE, AAK, MJTS, SAT, JCM), MRC Biostatistics Unit (CAV), Wellcome Trust Sanger Institute (CPMJ, SAT, JCM, DTO), BBSRC CASE Studentship with Abcam plc (AAK).

References

13. Materials and methods are available as supplementary materials on Science Online.


Single-cell sequencing of unstimulated and stimulated CD4+ T cells reveals how aging destabilizes a conserved transcriptional activation program.
Fig. 1. Divergence in gene expression levels and cell-to-cell variability among naive CD4+ T cells isolated from two species of mice.
(A) Single cells were isolated from spleens of young (~3 month) and old (~21 month) individuals of two mouse sub-species, and sequenced before or after activation (13); (B) Species-specific tSNE clustering of naive CD4+ T cells from young animals; (C) Representative heatmap of 30 genes and 30 cells randomly selected from young animals from both species shows species-specific variation; (D) Violin plots show distribution of expression levels of selected, species-specifically transcribed genes (grey background).
Fig. 2. Activation of naive CD4⁺ T cells induces a transcriptomic switch.

(A) Activation of CD4⁺ T cells from young B6 mice induces large-scale transcriptional changes (see S4 for CAST CD4⁺ T cells); (B) Genes up-regulated (red) and down-regulated (blue) by immune stimulation in young B6 mice (13); (C) Genes with no overall gene expression differences (black inset, (B)) during activation show decreased cell-to-cell variability in transcription (Mann-Whitney-Wilcoxon test, ***: p< 10⁻¹⁰); (D) Up-regulated genes were expressed in a large fraction of activated CD4⁺ T cells after stimulation (median 36%); down-regulated genes were expressed in a smaller fraction of naive CD4⁺ T cells.
(median 18%); (E-G) Example genes showing transcriptional changes in unstimulated and activated CD4+ T cells: (E) Il2ra (Cd25); (F) Eif1; (G) Sell (Cd62l).
Fig. 3. Interspecies comparison of CD4+ T cell activation.
(A) CD4+ T cells isolated from B6 (black) and CAST (gold) show similar large-scale transcriptional changes upon immune stimulation; (B) Immune activation of CD4+ T cells triggers up-regulation of conserved and species-specific transcriptional programs. For visualization purposes, genes in shared and species-specific categories were proportionately and randomly selected (Material and Methods). Thirty cells were randomly selected for each condition/species (13); (C) Only genes up-regulated in both B6 and CAST highly enrich for known T cell functionality (Bonferroni multiple testing corrected p-values, red line is 0.1); (D) Fractions of cells in which a gene is detected are displayed as histograms; 70 genes were randomly selected from each gene set.
Fig. 4. Aging increases the cell-to-cell variability during early activation of CD4+ T cells. 
(A) Upon activation, changes in expression of the core immune activation program are greater in younger mice than in older mice; (B) Among the genes from (A) that do not change their averaged gene expression, transcriptional variability increases during aging (13); (C) The fraction of cells in which genes of the shared activation process are expressed is reduced in CD4+ T cells from old animals. The distribution of fraction values is plotted on each corresponding axis (medians of fraction values are indicated in red); changes in fraction values were tested using a binomial test (significant changes are shaded in red or blue where multiple testing corrected p-values <0.1).