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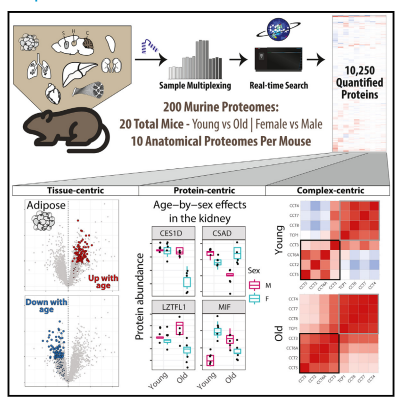
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Graphical abstract



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In brief

Organismal aging drives pleiotropic changes in tissue and cellular homeostasis. By quantifying the proteomes of 10 tissues, including three brain sections, in aging mice, Keele et al. explore the regulation of proteins and protein complexes with age. They observe cross-tissue and tissue-specific age- and age-by-sex-based remodeling of proteomes.

Highlights

- Quantified 10,250 total proteins across 20 mice ages 8 and 18 months
- Quantified 200 anatomical proteomes across 10 tissues and at two ages
- Immune, proteostatic, and metabolic protein abundances change with age
- Altered protein complex stoichiometry and substrates correlate with age







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Global and tissue-specific aging effects on murine proteomes

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SUMMARY

Maintenance of protein homeostasis degrades with age, contributing to aging-related decline and disease. Previous studies have primarily surveyed transcriptional aging changes. To define the effects of age directly at the protein level, we perform discovery-based proteomics in 10 tissues from 20 C57BL/6J mice, representing both sexes at adult and late midlife ages (8 and 18 months). Consistent with previous studies, age-related changes in protein abundance often have no corresponding transcriptional change. Aging results in increases in immune proteins across all tissues, consistent with a global pattern of immune infiltration with age. Our protein-centric data reveal tissue-specific aging changes with functional consequences, including altered endoplasmic reticulum and protein trafficking in the spleen. We further observe changes in the stoichiometry of protein complexes with important roles in protein homeostasis, including the CCT/TriC complex and large ribosomal subunit. These data provide a foundation for understanding how proteins contribute to systemic aging across tissues.

INTRODUCTION

Aging results in a progressive decline in physiological function with increased risks of disease and death. Gene expression studies have revealed age-related changes in transcripts that are shared across tissues and others that are tissue specific, as well as transcripts that contrast healthy and diseased aging in humans. However, these studies are unable to directly ascertain age-related changes in proteins. Untargeted, quantitative proteomics studies can reveal how proteins change with age, which can confirm findings from gene expression, but more importantly, they can identify molecular aging signatures that occur independent of gene expression changes.

Previous proteomics studies have investigated aging-related changes in rodent tissues, including rats⁶ and mice.^{7–9} We previously investigated the effects of age and sex on gene expression and protein abundance in kidneys¹⁰ and hearts¹¹ from genetically diverse mice. We found that differences in protein abundance between males and females in the kidney were largely mediated through their transcripts. In contrast, differences in protein abundance across ages were largely independent of their transcripts. A similar dynamic between sex and age differences was observed in heart, although fewer sex differences were present. From these studies, we concluded that many age-related changes in protein abundance are not due to corresponding changes in gene expression and that transcriptomics provides an incomplete picture of aging in kidney and

heart. Multiple mechanisms could result in the discordance of age-related changes in proteins compared with changes in their transcripts, including reduced proteasome activity¹² and reduced ribosome occupancy¹³ with age.

A key question is whether this discordance of age-related changes between proteins and their transcripts is seen in most tissues. Furthermore, we are interested in which age-related changes in proteins are shared across tissues and which are unique. Comparing the kidney and heart revealed common signatures of increased immune cell infiltration in both tissues for proteins and transcripts. We also observed tissue-specific changes, particularly among proteins. Changes in the kidney proteome correspond to functions specific to the substructures and cell subtypes of the nephron, including the podocytes and proximal tubule cells. In the heart, we observed age-related changes in fatty acid metabolism and autophagy. These tissue-specific changes relate to the unique biological functions and stressors of these tissues during aging.

To address these questions, we performed a survey of protein abundance across 10 tissues (kidney, liver, fat, spleen, lung, heart, skeletal muscle, striatum, cerebellum, and hippocampus) collected from female and male C57BL/6J (B6) mice at 8 and 18 months of age. We performed multiplexed, quantitative mass spectrometry on bulk tissue samples and analyzed differences in protein abundance between age and sex, as well as sex-specific changes with age. We compared age and sex differences in proteins in our study with transcript changes in





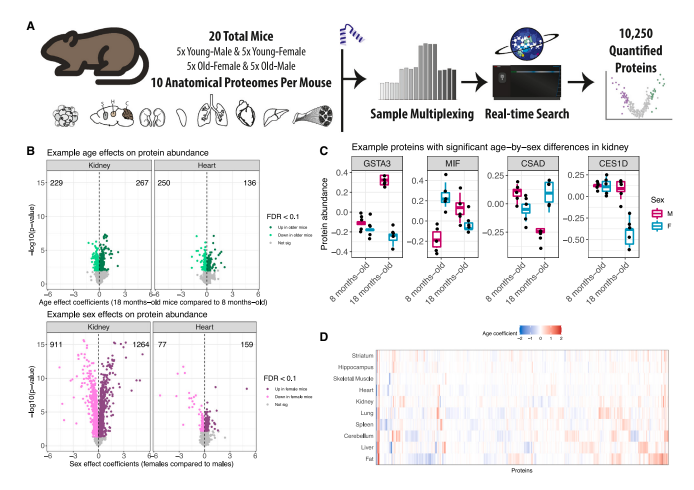


Figure 1. Quantitative proteomics study on the effects of age and sex on protein abundance across 10 tissues from B6 mice

(A) Using sample multiplexing, 10 anatomical proteomes (adipose tissue, striatum, hippocampus, cerebellum, kidney, spleen, lung, heart, liver, skeletal muscle) were profiled across age- and sex-matched mice (n = 20).

(B) Age (top) and sex (bottom) differences for protein abundance from kidney (left) and heart (right) tissues, depicted as volcano plots. Differences in protein abundance are summarized as regression coefficients (x axis) and corresponding -log10(p value) (y axis). Points are colored based on statistical significance (FDR < 0.1) and direction of effect. Counts of proteins with significantly higher abundance in 18- and 8-month-old mice are included. Dashed vertical lines at

- (C) Examples of proteins in kidney tissue with significant age-by-sex differences (FDR < 0.1).
- (D) Age differences detected across the 10 tissues (FDR < 0.1), represented as a heatmap. Differences are summarized as regression coefficients.

corresponding tissues that were reported by Schaum et al.² We used enrichment analyses to assess how aging affects biological processes, as reflected by coordinated changes in proteins across and within tissues. Finally, we characterized agingrelated changes in protein complexes, in terms of overall and relative abundance of member proteins. Our findings confirm that the discordance between age-related changes in proteins and gene expression occurs across multiple tissues. Our data survey a broad range of age-related changes in proteins that occur globally across tissues and others that are tissue specific.

RESULTS

We quantitatively profiled protein abundance across 10 tissues (Table S1) representing a range of organ systems (kidney, liver, fat, spleen, lung, heart, skeletal muscle, striatum, cerebellum, and hippocampus) from 20 B6 mice using tandem mass tags (TMTs) and real-time search (RTS) mass spectrometry. 14 Animals represented an equal balance across males, females, and two age groups (8 and 18 months), with five animals per ageby-sex group (Figure S1A). Outlying samples were identified using principal-component analysis (PCA), resulting in the removal of one sample from liver, fat, spleen, lung, and skeletal muscle and two samples from striatum (STAR Methods). Outlying samples across tissues were not from the same mouse. Cumulatively, we detected 11,853 proteins across the 10 tissues. We filtered the data to a high-confidence set of proteins (observed in both batches per tissue), resulting in 10,250 proteins used in further quantitative analysis (Figure 1 and S1B). Spleen had the highest number of analyzed proteins (6,556) and skeletal muscle had the lowest (2,353) (UpSet plot 15 in

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Figure S1C). Many proteins were detected across multiple tissues. We observed 676 different cross-tissue detection patterns for proteins, with detection in all tissues being the most common (1,229), followed by proteins detected in only spleen (1,003), and then in only the three brain tissues (425) (Figure S1C). Using gene set enrichment analysis¹⁶ (STAR Methods), the 425 proteins (compared with the overall set of 10,250 proteins observed across all tissues) were enriched for gene ontology (GO) terms related to synapses, which is consistent with brain biology and supports the quality of the protein data (Figure S1D).

To assess how different technical features of the experiment contributed to variation in the abundance of individual proteins, we jointly modeled data across tissues by fitting random effects models (STAR Methods) for each of the 1,229 proteins measured in all 10 tissues (Table S2). As expected, the protein abundance varies greatly with tissue/batch, which is confounded due to each tissue being measured in two separate runs, i.e., batches, of the mass spectrometer (Figure S1E). Our aim here is not to detect protein abundance differences between tissues, which would be invalid due to confounding, but rather to detect agerelated change patterns across many tissues. One unique finding was 13 proteins that had abundance patterns that were consistent across the 10 tissues and highly specific to individual mice, including IGHG2C and four other immunoglobulins (Figures S1E-S1H). The 13 proteins (compared with the overall set of 1,229 proteins observed in all tissues) were enriched for GO terms related to adaptive immune response, suggesting cross-tissue responses to infections that are unique to each mouse.

Effects of age and sex on the abundance of individual proteins

Protein abundance can vary with age 10,11,17 and between sexes. 18-20 For each tissue, we characterized age and sex effects (Table S3) and declared differences to be significant based on false discovery rate (FDR) < 0.1 (Figures 2 and S2A, respectively; STAR Methods). The number of proteins with age effects varied greatly across the tissues, ranging from lung with the most (866) to striatum with none meeting statistical significance (partially due to loss of power from the removal of two animals). The number of proteins with sex effects varied across the tissues, from kidney with 2,175 to the three brain tissues with 10 or fewer. Comparison of sex and age effects across the tissues revealed generally more differences between sexes than between ages, most notably in kidney, liver, fat, and spleen. However, proteins with significant age effects outnumbered those with sex effects in lung, heart, and cerebellum (Figure S2B). The three brain tissues were distinctly buffered from differences based on age and sex. In total, 2,356 proteins had a significant age effect in at least one tissue, 5,125 had a sex effect, and 43 had an age-by-sex interaction effect (FDR < 0.1). We also identified proteins for which sex differences in abundance depended on age by testing for age-by-sex interaction effects (STAR Methods), detecting 21 in kidney, 6 in liver, 5 in fat, and 11 in skeletal muscle (FDR < 0.1) (Figures S2C-S2G). The skeletal muscle proteins, for which males had distinctly higher abundance than females in older mice, were associated with the endoplasmic reticulum lumen.

We jointly modeled individual proteins across multiple tissues (STAR Methods) to test whether the age or sex effects on proteins were consistent across tissues or unique to specific tissues (Table S3), declaring significance based on FDR < 0.1. Among the 7,745 proteins detected in more than one tissue, 643 had consistent age differences across the tissues in which they were quantified. For example, IKGC is an immunoglobulin that has increased abundance in older mice for all tissues (cross-tissue age p = 1.1e-6; Figure 2B). We detected 1,028 proteins with age effects that differed between tissues, representing proteins that had age differences in only some of the tissues or even age effects with differing directions, such as BCAT1 (age-by-tissue p = 1.8e-10), CES1D (age-by-tissue p = 1.1e-10), FBLN1 (age-by-tissue p = 2.1e-8), and STAB1 (age-by-tissue p =2.6e-8) (Figure 2C). These age-related differences reflect tissue-specific features related to aging decline. For example, in kidney, there was reduced abundance of BCAT1, which promotes mitochondrial biogenesis and ATP production and has been shown to promote breast cancer formation when knocked down,²¹ and REN1, which has been shown to play a role in modulating vascular tone and tubular function in the kidney.²² For sex effects, we detected 1,006 proteins with consistent differences between sexes across tissues and 2,565 proteins with tissue specificity.

Most age-related changes in proteins show no corresponding change in their transcripts

To compare age effects on proteins to corresponding effects on transcripts, we obtained data from Schaum et al.,2 in which bulk RNA sequencing was performed across 17 tissues of mice from 10 age groups, ranging from 1 to 27 months. Each age group consisted of four to six C57BL/6JN (B6N) mice. Overlapping tissues between studies included kidney, liver, and heart. We selected the transcriptomics data from the 9 and 18 months age groups, which are closest to 8 and 18 months in this study's proteomics data, and characterized the age and sex effects on transcripts (STAR Methods). We contrasted the age effects between proteins and their transcripts, and for comparison, we also compared the sex effects on proteins and their transcripts.

The age effects on proteins and transcripts are generally not consistent. In contrast, sex differences are highly concordant between proteins and transcripts, most notably in kidney (Figures 3A and 3B). The consistency of strong sex effects correlated between protein and transcript supports the validity of comparing data across distinct but related mouse strains²³ with mice that were raised at different sites as part of different experiments. We observed a similar dynamic between age and sex differences in the kidneys of genetically diverse outbred mice. 10 Genes with consistent age effects on transcripts and proteins in kidney include Vcam1 and Keg1 (Figures 3C and 3D). VCAM1 is an immunoglobulin that facilitates interactions between vascular and immune cells, and its increase with age has been associated with age-related disease in humans.²⁴⁻²⁶ We observed increases in VCAM1 abundance with age across multiple tissues (kidney, liver, fat, and cerebellum), which resulted in a significant cross-tissue age effect (cross-tissue age p = 1.59e - 7). We note that the expression data suggest greater



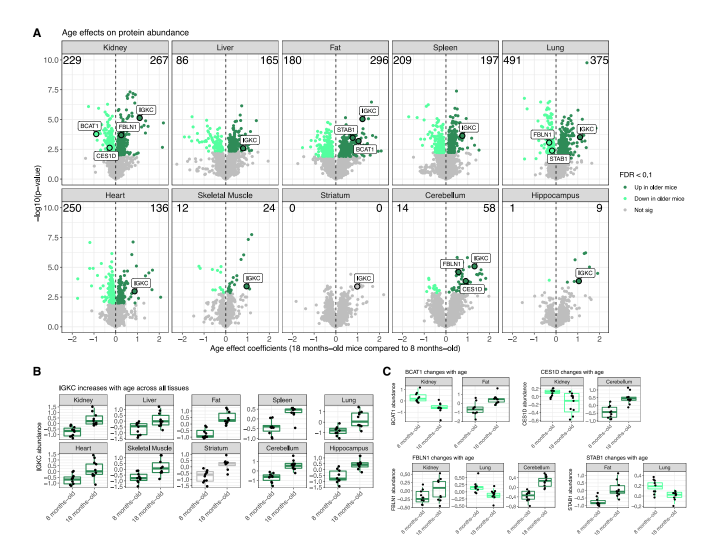


Figure 2. Proteins with age-related differences in abundance across 10 tissues

(A) Proteins with age differences in abundance, represented as volcano plots. Differences in protein abundance are summarized as regression coefficients (x axis) and corresponding -log₁₀(p value) (y axis). Points are colored based on statistical significance (FDR < 0.1) and direction of effect. Counts of proteins with significantly higher abundance in older mice and younger mice are included. Dashed vertical lines at 0 included for reference. Proteins with sex differences in abundance are shown in Figure S2.

- (B) The immunoglobulin IGKC has consistent increased abundance in older mice across all 10 tissues.
- (C) Proteins with significant age differences that vary between tissues: BCAT1, CES1D, FBLN1, and STAB1.

variation within ages and sexes than the protein data, which has more mice per age-by-sex group.

Heart had the least consistency between the age effects on proteins and on their transcripts (Figure 3E). It also had far fewer significant sex effects than kidney, consistent with our previous work, 11 although they were still more consistent between proteins and transcripts than age effects (Figure 3F). Notable genes with consistent sex effects on proteins and transcripts are Ddx37 and Eif2s3x, which are encoded on the Y and X chromosomes, respectively. These findings validate the quality of the protein data by demonstrating the consistency of sex effects with transcript data. They also highlight the importance of assessing aging-related changes among proteins, many of which are not observed at the transcript level.

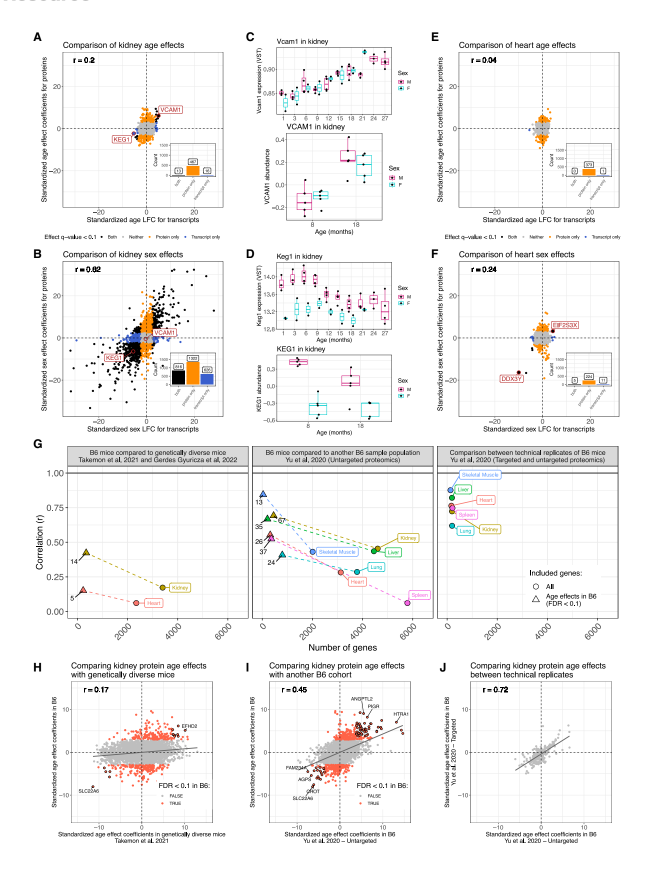
Age-related changes in proteins are consistent across

We examined the consistency of age effects across several proteomics studies of murine aging. One of the comparison studies also used B6 mice,8 but with a smaller sample size of 10 male mice that were evenly split between 4 and 20 months of age. This study included nine tissues, of which six were in common with ours. We also compared our results with those from a proteomics study of aging in genetically diverse outbred mice 10,11; this study looked at kidney and heart tissues from 188 mice, with approximately equal representation across sexes and 6, 12, and 18 month age groups.

We computed the correlation of age effects between studies across all shared proteins as well as across proteins with effects

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that met statistical significance (FDR < 0.1) in our study. The strength of correlations was greater in comparisons between independent studies of B6 mice than between B6 mice and genetically diverse mice (Figure 3G). All correlations are statistically significant, with the stringent comparison of heart between genetically diverse mice and B6 mice the weakest (r = 0.15; p = 0.022). In the genetically diverse mice, 27 higher levels of genetic variation compared with the isogenic B6 strain likely contribute to the reduced concordance. We also had an opportunity to examine technical variation in protein age effects estimated by either targeted or untargeted mass spectrometry on the same 10 B6 mice⁸; these comparisons show the strongest correlations, indicating that biological factors as well as experimental ones influence the concordance of age effects across studies. Comparisons of kidney protein age effects between these populations are highlighted in Figures 3H-3J, along with consistent strong age effects across datasets (Table S4). These results demonstrate concordance of age-related changes in proteins across studies and reveal the extent to which technical, biological, and genetically driven variation contribute to cross-study consistency.

Immune-associated proteins change with age across all tissues

We clustered the 2,356 proteins that had significant age effects in at least one tissue after setting effects to zero in tissues in which proteins were not observed. Looking broadly across the clustered age effects reveals consistent immune-related differences between age groups across all 10 tissues (Figure 4A). Components of the innate immune system, most notably members of the complement cascade, such as C8A and C8B, were less abundant in all tissues of older mice. Immunoglobulins and other proteins related to humoral immunity were distinctly more abundant in older mice. Proteins involved in proteolysis, including immunoproteasomes like PSMB8, were also more abundant in older mice to varying degrees across the 10 tissues.

The consistency of age effects across tissues for immunerelated proteins is striking, as highlighted by the adaptive im-

mune response GO set (GO: 0002250) (Figure 5A). Even tissues with few statistically significant age effects, most notably striatum with none, show differences in the same direction, such as increased abundance of immunoglobins in older mice. We note that the abundance of the immunoproteasomes (PSMB8, PSMB9, and PSMB10) is higher in older mice in the tissues with the most pronounced age-related increases in immunoglobins, fat and cerebellum (Figures 5B and 5C). The immunoproteasomes are inducible components that replace the constitutive components (PSMB5, PSMB6, and PSMB7) in the 20S catalytic core of the proteasome. The immunoproteasome is more efficient at degrading ubiquitin-labeled proteins as antigens for presentation on MHC class I molecules, a key process in distinguishing between self and non-self in adaptive immunity.²⁸ However, it is not a perfect marker of immune cells, as it is also expressed by non-immune cells during inflammation.²⁹

The co-regulation of the overall proteasome complex, encompassing the catalytic core of the 20S proteasome, its 19S regulator, and the 11S regulator, is multifaceted. We have previously shown that the abundance of individual components and subcomplexes of the proteasome is influenced by genetic variation¹⁹ and age¹¹ in genetically diverse mouse populations. In this B6 population, where genetic variation has been fixed (excluding spontaneous mutations specific to individuals), the subcomplex structure of the proteasome, most notably the 20S catalytic core and 19S regulator, is reflected in the correlations between complex members. The individual subcomplexes become more tightly correlated within themselves (and anti-correlated with each other) in the older mice in fat tissue (Figure 5D). The correlation among the immunoproteasome components (and anti-correlation with their constitutive analogs) becomes more pronounced in the older mice, likely due to increased levels of immunoproteasome from immune cells. Changing immune cell-related tissue composition with age is further supported by the reduction in complement activation proteins with age.

Figure 3. Age- and sex-related protein differences show varying levels of consistency with previously published transcriptomics and proteomics datasets

(A and B) Comparisons of (A) age- and (B) sex-related differences in proteins with transcripts in kidney. Points are colored based on statistical significance (FDR < 0.1) in proteins and transcripts. Correlation (r) between protein differences and transcript differences and dashed vertical and horizontal lines at 0 included for reference. Counts of genes with significant differences included as bar plots in the bottom right quadrant.

(C) Vcam1 significantly increases with age in terms of both transcripts (top) and proteins (bottom). Transcript data represent 10 age groups compared with 2 age groups for proteins.

(D) Keg1 expression had significant age and sex differences (top), whereas its protein had a matching sex effect (bottom). The age effect did not meet significance at FDR < 0.1, but the direction is consistent with transcripts. Transcript data represent 10 age groups compared with 2 age groups for proteins.

(E and F) Comparisons of (E) age- and (F) sex-related differences in proteins with transcripts in heart. Points are colored based on statistical significance (FDR < 0.1) in proteins and transcripts. Correlation (r) between protein differences and transcript differences and dashed vertical and horizontal lines at 0 included for reference. Counts of genes with significant differences included as bar plots in the bottom right quadrant.

(G) Correlations between protein age differences across tissues comparing three mouse sample populations. The number of genes being summarized by the correlation is on the x axis. Circle points represent correlations across all overlapping genes. Triangle points represent correlations across overlapping genes that had a significant age difference (FDR < 0.1) in this study's B6 mice. Dashed lines connect correlations from the same tissue and study comparison. Horizontal line at 0 included for reference as the upper limit of correlation. Numbers associated with each point indicate the number of proteins associated with each com-

(H–J) Comparisons of protein age-related differences in kidney between (H) this study's cohort of B6 mice and genetically diverse mice, (I) this study's cohort of B6 mice and another smaller cohort of male B6 mice, and (J) targeted and untargeted protein measurements from the smaller cohort of male B6 mice, representing a technical replication. Proteins with consistent strong age effects (same sign in both datasets and absolute Z scores within each population greater than 2) across two studies are outlined in black. Correlation (r) between protein differences and dashed vertical and horizontal lines at 0 included for reference. Black best fit lines also included for reference.

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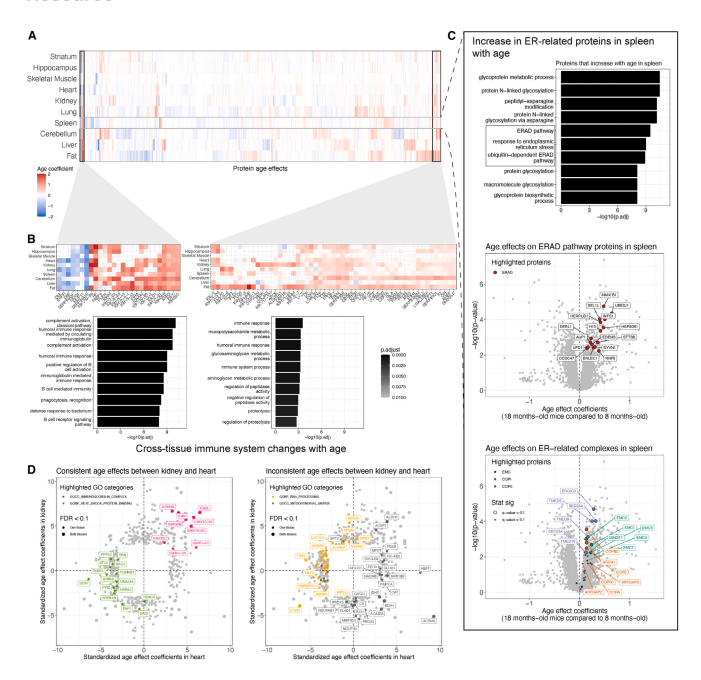


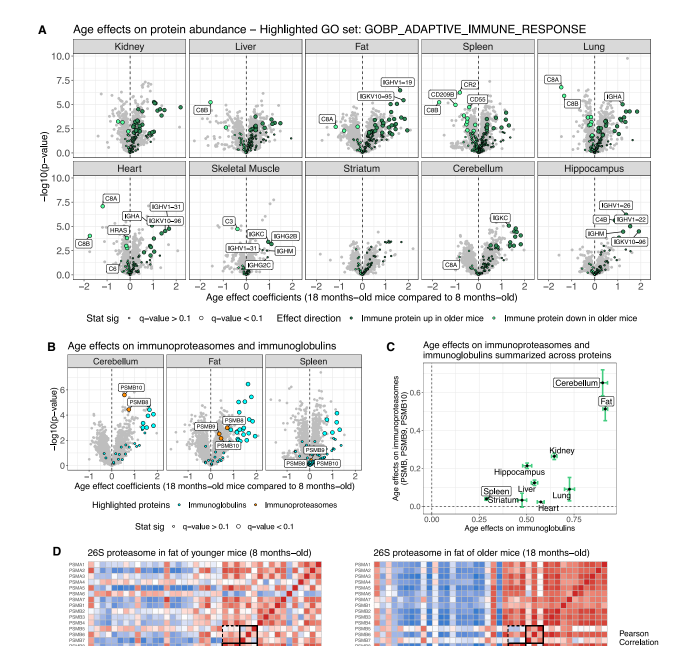
Figure 4. Cross-tissue and tissue-unique patterns of aging

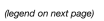
(A) Age-related differences detected across the 10 tissues (FDR < 0.1), represented as a heatmap. Differences are summarized as regression coefficients. Hierarchical clustering of the proteins (columns) reveals sets of proteins with age difference patterns across tissues and unique to specific tissues. (B) Proteins with age differences that are shared across tissues are enriched for immune-related GO categories. Additional tissue-unique patterns are highlighted in Figure S3.

(C) The proteins with age differences in a specific tissue can be enriched in GO categories, with spleen highlighted here for proteins with higher abundance in older mice. Abundance differences with age for proteins analyzed in spleen are represented as volcano plots. Differences in protein abundance are summarized as regression coefficients (x axis) and corresponding -log₁₀(p value) (y axis). The ERAD pathway (GO: 0036503), EMC, COPI, and COPII proteins are highlighted. Highlighted proteins with significant differences (FDR < 0.1) have larger point size. Proteins with age p < 0.05 are labeled.

(D) Comparison of age differences between kidney and heart with highlighted GO categories that are consistent (left) and inconsistent (right) between the tissues. Proteins with a significant age difference (FDR < 0.1) in kidney or heart are shown. Proteins with significant differences (FDR < 0.1) in both tissues have a larger point size.







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Tissue-specific signatures of aging

Age effects reveal unique patterns specific to tissues (Tables S5 and S6). Spleen exhibits unique increases in abundance for proteins related to the endoplasmic reticulum (ER), such as the ER-associated degradation (ERAD) pathway, ER membrane complex (EMC), and COPI and COPII complexes (Figure 4C), suggesting potential changes to protein quality control³⁰ in the spleen. The EMC enables the biogenesis of multipass transmembrane proteins and has been associated with pleiotropic phenotypes across organisms. 31,32 COPI and COPII are involved in trafficking proteins between the ER and the Golgi. 33 In addition to the immune signatures observed across all 10 tissues, we saw tissue-specific immune patterns, such as decreased abundance for proteins involved in leukocyte and lymphocyte activation in the spleens of older mice (Figure S3B) and increased abundance for proteins involved in a broad range of immune system-related GO categories in the fat of older mice (Figure S3C). We observed decreased abundance in older mice for proteins related to mitochondrial chain complex I and cellular respiration (Figure S3A) and increased abundance in older mice for proteins related to multiple metabolic processes (Figure S3D). Comparison of gene set enrichment results between tissues can highlight shared or distinct aging processes. For example, we compared kidney and heart, which revealed consistent GO categories like immunoglobulins (increasing with age) and heat shock proteins (decreasing with age) and inconsistent GO categories like the mitochondrial matrix and RNA processing (Figure 4D).

Age and sex influence the abundance and stoichiometric balance of protein complexes

Individual members of protein complexes are often co-regulated to maintain stoichiometric balance of components.34-38 Biological factors can affect this balance, including genetics, sex, and age. 6,11,18-20 More than one factor can influence a complex (or its subcomplexes or individual components), as demonstrated by the effects of both genetic variation and age on the proteasome. 11,19 To measure a protein complex's co-abundance, referred to here as cohesiveness, for each of 228 protein complexes³⁹⁻⁴¹ across the 10 tissues, we used the median correlation across all pairs of complex members (Table S7; STAR Methods). We note that the cohesiveness of a complex could reflect its stoichiometric balance as well as cell-type heterogeneity. We observed some conservation of protein complex cohesiveness between tissues, most notably among the three brain tissues (r > 0.68, p < 2.2e-16; Figure S4).

We tested whether age or sex had consistent effects on abundance across the members of a protein complex (Table S7; STAR Methods), as would be expected if the entire protein complex were changing with age or sex. Examples include the previously mentioned COPI and COPII complexes in the spleen (Figure 4C). The number of protein complexes with a multiprotein age effect varied across the tissues, ranging from lung with 29 to skeletal muscle and hippocampus with none at FDR < 0.1 (Figure S5A). More complexes had consistent sex effects across proteins, ranging from spleen with 76 to striatum and cerebellum with none at FDR < 0.1 (Figure S5B).

Changes due to age or sex on protein complexes may be more subtle than a change in mean abundance across complex members. Cohesiveness of a protein complex can vary with age or sex. We first calculated the correlations among protein complex members for each age group and then looked for changes in overall correlation patterns with a paired t test (Table S7; STAR Methods). The same approach was used for sex. We used a stricter threshold of significance (FDR < 0.01) to focus on only the most significant effects. The association of age with cohesiveness of complexes varied across the tissues, ranging from 35 complexes in spleen to 3 in skeletal muscle at FDR < 0.01 (Figure S5C). For sex, spleen had the largest number of complexes with changes in cohesiveness (26), and fat, hippocampus, and striatum had the fewest with 6 each, all at FDR < 0.01 (Figure S5D). Ribosomes have been shown to lose stoichiometric balance with age in the brains of killifish, 12 and we see similar signals across many of our tissues (Figure S5C), indicating that this previous finding generalizes across tissues and species.

For each complex, we counted the number of tissues for which a significant age and sex effect on abundance or cohesiveness was detected (Figures S5E-S5H). The co-distribution of age and sex effects differed between effects on abundance and effects on cohesiveness. Effects on abundance were more likely to be detected in smaller subsets of tissues, whereas effects on cohesiveness represent a distinct minority of complexes with both age and sex effects on cohesiveness across many tissues, such as the chaperonin-containing T complex (CCT complex) and cytoplasmic ribosomal large subunit (CRLS).

CCT complex is more cohesive in older B6 mouse

The CCT complex, also known as the tailless complex polypeptide 1 ring complex (TRiC), is required for folding proteins such as actin and tubulin. The CCT complex was significantly more cohesive in older mouse cerebellum than in younger (p = 2.0e-8; Figure 6). This signal is due to CCT6A, CCT2, CCT5, and, to a lesser extent, CCT3 being anti-correlated with other complex members in younger mice but more correlated in older mice (Figure 6B). The significance of the change in individual correlations was

Figure 5. Increased immunoglobulin abundance is a signature of aging detected in all 10 tissues

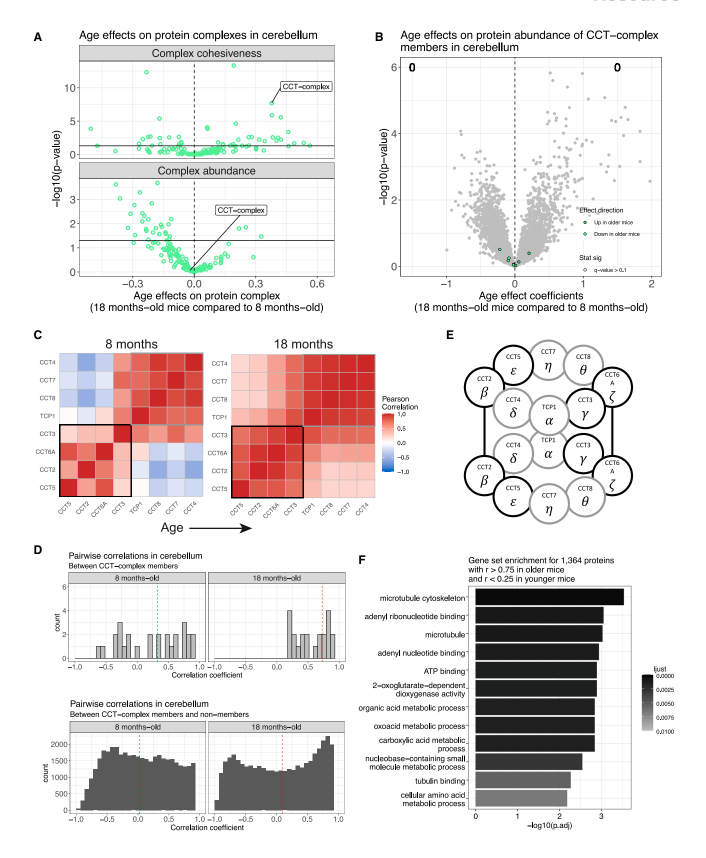
(A) Proteins with age differences in abundance, represented as volcano plots. Differences in protein abundance are summarized as regression coefficients (x axis) and corresponding -log₁₀(p value) (y axis). Points are colored based on being a member of the adaptive immune response GO category (GO: 0002250) and direction of effect. Highlighted proteins with significant differences (FDR < 0.1) have larger point size. Dashed vertical lines at 0 included for reference.

(B) Volcano plots for cerebellum, fat, and spleen, with immunoglobins and immunoproteasomes (PSMB8, PSMB9, and PSMB10) highlighted.

(C) Age differences summarized across the immunoproteasomes (y axis) and immunoglobins (x axis) for all 10 tissues. Points represent mean differences and bars represent standard errors. Horizontal and vertical dashed lines at 0 included for reference.

(D) Pearson correlations from the proteasome in younger (top) and older (bottom) mouse fat. Rows and columns are ordered to reflect key subcomplexes of the proteasome, which are labeled. The immunoproteasomes and their matching constitutive analogs are highlighted with black squares.





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determined using permutations (STAR Methods). The pattern of correlations reflects the physical structure of the CCT complex, which is composed of two octameric rings made from eight proteins (TCP1, CCT2, CCT3, CCT4, CCT5, CCT6A, CCT7, and CCT8)^{42,43} (Figure 6E). Notably, the CCT6A and CCT2 components from the upper and lower rings are in physical contact with their matching protein. CCT5 and CCT3 are immediately adjacent to CC2 and CCT6A, respectively, in both rings. We have previously shown that genetic variation at Cct6a regulates other members of the CCT complex in genetically diverse mice. 18,19 There is no complex-wide age effect on abundance (p = 0.87), and none of the proteins differ significantly in mean abundance between the two age groups (Figure 6B). In older mouse cerebellum, members of the CCT complex are correlated with more non-CCT-complex members (1,364 genes with r > 0.75 in older mice and r < 0.25 in younger mice), which enrich for many GO categories related to its function in folding cytoskeleton proteins, 44 such as the microtubule category (GO: 0005874) (Figures 6F and S6).

Ribosomal large subunit complex is more cohesive in young female liver tissue

The CRLS was found to be significantly more cohesive in livers of younger mice (p = 4.8e-57) and females (p = 2.7e-8) (Figures 7A and 7C). There was no complex-wide age effect on abundance (p = 0.86) or any significant age differences detected for individuals' CRLS proteins (FDR < 0.1), whereas female liver had lower complex-wide abundance (p = 2.9e-6) and significantly lower abundance for 19 proteins (Figures 7B and 7D). This replicates our previous finding of decreased ribosomal protein abundance in female liver tissue from genetically diverse mouse populations. 19,20 The co-occurrence of effects on individual protein abundance and complex-wide abundance and cohesiveness led us to examine the age-by-sex interaction effects on individual proteins, where 14 had age-by-sex differences (age-by-sex p < 0.05), with a consistent pattern of females having distinctly lower abundance in the younger mouse liver (Figure 7E). These differences contribute to the unique age-by-sex co-abundance patterns of the CRLS in the liver, which is more cohesive in young females compared with older females or males (Figures 7F and 7G). Notably, age and sex effects on the CRLS vary across the tissues; in lung there is a complex-wide greater abundance in older mice and little clear effect from sex (Figure S7).

DISCUSSION

We performed quantitative protein profiling across 10 tissues from the mouse reference strain, C57BL/6J. We characterized differences in protein abundance based on age, sex, and their interaction. We compared age-related changes observed in our study with transcriptomics and proteomics data from other studies of aged mice and observed broad consistency, but also evidence of protein-specific and population-specific aging changes. Looking within and across tissues, we identified unique functional patterns of proteins that vary with age, including global changes to components of the immune system, tissue-specific changes to cellular respiration and metabolism, and proteostasis. Furthermore, we examined how protein complexes differ based on age and sex in terms of complex-wide abundance and cohesiveness or correlation.

We detected a handful of proteins with age differences that varied between sexes (e.g., age-by-sex interaction effects). Tissues with prominent age-by-sex effects were kidney (CES1D, MIF, PSMC4), liver (GSTP1, GSTP3, NDRG1), fat (CHDH, CKA P4, RPL35), and skeletal muscle (HSPA5, CDNF, ADAMT S1). The age-by-sex effect patterns varied across proteins and tissues. In the kidney, macrophage migration inhibitory factor (MIF) had higher protein abundance in older males than in younger males, but an inverted response in females (younger females had more MIF protein). Other proteins like CES1D had no age-related abundance change in male kidneys, but a striking decrease in protein abundance in female kidneys. The age-bysex effects in skeletal muscle were highly consistent across the 11 proteins, with females showing no age-related abundance changes compared with males having an increased abundance with age. These findings highlight diverse cellular and tissue responses driven by age-by-sex effects and emphasize the need for studies to include both sexes to provide a complete picture of aging.

The primary aging pattern shared across all tissues is an increase in immunoglobulin proteins, implicating the immune system in the aging process. This parallels immune-aging signatures in transcriptomics,² and the strongest immune signal for both proteins and transcripts occurs in fat tissues. Even in tissues like striatum and hippocampus that had very few proteins with significant age differences, specific immune proteins increased in abundance with age, consistent with the other tissues. Given that the data represent bulk tissue samples, age

Figure 6. CCT complex is more stoichiometrically balanced in older mouse cerebellum

(A) Comparison of age-related differences and p values in complex cohesiveness (top) with complex-wide abundance (bottom) in cerebellum. CCT complex is highlighted. Dashed vertical lines at 0 included for reference. Horizontal line at $p = -\log_{10}(0.05)$ included to indicate statistical significance.

(B) Volcano plot for age differences in individual protein abundance for cerebellum with CCT-complex members highlighted with color based on direction of effect. Counts of proteins with significantly higher abundance in older mice and younger mice are included (FDR < 0.1). Dashed vertical lines at 0 included for reference.

- (C) Pearson correlations from the CCT complex in younger (left) and older (right) mouse cerebellum. Black and gray squares highlight patterns in the correlation matrix that mirror the structure of the CCT complex.
- (D) Histograms of pairwise correlation coefficients between CCT-complex members with each other (top) and other proteins (bottom). Vertical red dashed lines represent median correlations.
- (E) The CCT complex is composed of two identical octomeric rings. The CCT2 (β) and CCT6A (ζ) from each ring are in physical contact with their twin. Outline of proteins matches correlation structure previously highlighted.
- (F) GO categories enriched in proteins that are more correlated with CCT-complex members in older mouse cerebellum than in younger. The microtubule GO category (GO: 0005874) is explored further in Figure S6.



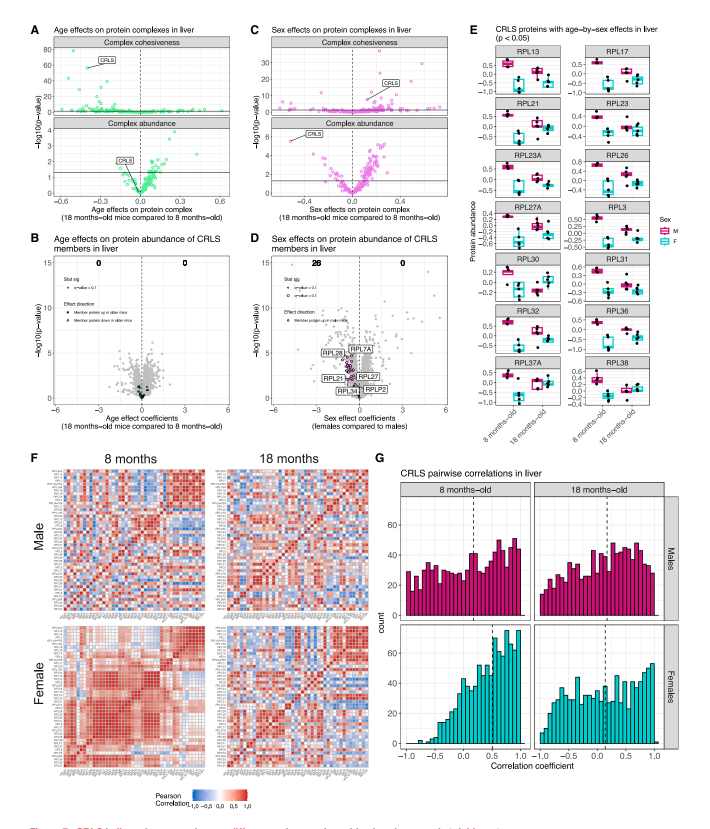


Figure 7. CRLS in liver shows age-by-sex differences in complex-wide abundance and stoichiometry

(A) Comparison of age-related differences and p values in complex cohesiveness (top) with complex-wide abundance (bottom) in liver. CRLS is highlighted. Dashed vertical lines at 0 included for reference. Horizontal line at $p = -\log_{10}(0.05)$ included to indicate statistical significance.

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effects within a tissue may be driven by changes in cellular composition with age. These patterns suggest that, with age, changes in the adaptive immune response occur, potentially due to increased presence of immunoglobulin-producing cells, to varying degrees across tissues. This is further supported by matching increases in the immunoproteasomes.

In addition, we observed immune-related aging patterns that were specific to tissues, which further highlights changes to the immune system as a signature of aging. In the spleen, which has a unique role in the immune system compared with the other tissues in this study, we saw reduced abundance with age for proteins involved in leukocyte activation, particularly T cells (including ITGB7, SLFN1, SATB1, FOXP1, FOXO1, SIT1, and THY1). We note that many of these proteins were primarily quantified in spleen. Fat also exhibited unique increases with age for immune-related proteins. Together these findings point to dynamic immune system changes across tissues.

Previous studies have demonstrated that protein complexes can be co-regulated by biological factors, such as sex, diet, and genetics. 18-20 We assessed how age and sex affect complex-wide abundance and cohesiveness, replicating our previous finding of reduced ribosomal protein abundance in female liver. 19 We note an excess of age differences across tissues for protein complexes, in terms of both cohesiveness and abundance, related to proteostasis, which is hypothesized to contribute to aging, 45-48 including ribosomes, 49 proteasomes, and the CCT complex. Similar changes to the cohesiveness of these protein complexes were observed in genetically diverse mice. 11 We also note that the direction of change was not necessarily consistent across tissues for a complex, nor was it possible to distinguish loss of stoichiometric balance with age from changes in tissue compositions, as is likely the case for the immunoproteasome. Nevertheless, our findings reveal changes to and potential disruption of proteostasis with age that vary across tissues.

Further studies are needed to understand the mechanisms underlying the large-scale aging dynamics of proteins discovered here, many of which are independent of age-related changes in transcription. A broader time series of age groups, similar to the gene expression study of Schaum et al.,2 would provide a higher-resolution picture of how protein abundance changes with age. Such a study could characterize non-linear trends of aging, which could then be compared between groups of ageco-regulated proteins or even with aging trends in gene transcripts. For example, we observed relatively few age and sex effects at the protein level across the murine brain sections profiled (hippocampus, striatum, and cerebellum), which is overall consistent with other studies based on two age groups.^{2,8} Therefore, future studies could improve the resolution of aging at the level of proteins by including B6 mice older than late midlife, which could reveal new temporal changes in brain proteins.

Lifespan has been shown to vary across mouse strains,50 which suggests that age-related protein changes may also vary. Our comparisons of age-related changes in proteins across different mouse populations are consistent with some population-specific differences. Age differences seen in B6 mice were more consistent with another study of B6 mice than with genetically diverse mice. Nevertheless, overall, tissue-specific changes in proteins during aging are significantly correlated between populations. There are key differences across these studies, including the numbers and ages of the mice, which can have an impact on how the age effect is estimated. Compared with sex differences in kidney, the most consistent effect type between transcripts and proteins across populations, age differences are small and thus more subject to error, particularly when comparing across studies.

Without corresponding gene expression data from the same samples, it is experimentally challenging to distinguish whether aging changes in proteins observed in bulk tissue reflect consistent changes across the cells that make up a tissue or changes in tissue composition. If gene expression data were available, tissue deconvolution⁵¹ of bulk tissue RNA sequencing would be possible using single-cell data from overlapping tissues from a resource, such as Tabula Muris,³ to estimate cell-type proportions per sample. The relationship between cell-type proportions and age could then be tested to identify proteins with age effects that correspond to changes in tissue composition.

Even in the absence of gene expression data, there are hints that some aging effects on proteins reflect changes to tissue composition with age. For example, increased levels of immunoproteasomes and decreased levels of complement cascade proteins across multiple tissues could be explained by a changing balance of immune cells with age. Estimation of cell-type identity through integration of reference single-cell RNA sequencing with bulk protein abundance rather than gene expression is challenging because fewer genes are measured at the protein level, likely resulting in less information distinguishing cell types. Furthermore, single-cell gene expression and mass spectrometry proteomics are less comparable. Single-cell proteomics data for samples could more directly elucidate age changes in proteins at a cellular level. However, these approaches are newly developing^{52–54} and pose both technical and analytical challenges to overcome, such as extreme data sparsity.

Based on our prior studies, we sought to evaluate the effect of aging on proteins across a wide range of tissues in the reference mouse strain to assess the concordance of age effects on

⁽B) Volcano plot for age differences in individual protein abundance for liver with CRLS members highlighted with color based on direction of effect. Counts of proteins with significantly higher abundance in older mice and younger mice are included (FDR < 0.1).

⁽C) Comparison of sex-related differences and p values in complex cohesiveness (top) with complex-wide abundance (bottom) in liver. CRLS is highlighted. Dashed vertical lines at 0 included for reference. Horizontal line at $p = -\log_{10}(0.05)$ included to indicate statistical significance.

⁽D) Volcano plot for sex differences in individual protein abundance for liver with CRLS members highlighted with color based on direction of effect. Counts of proteins with significantly higher abundance in females and males are included (FDR < 0.1).

⁽E) CRLS proteins with age-by-sex differences in abundance in liver (age-by-sex p < 0.05).

⁽F) Pearson correlations from the CRLS, stratified by age (left, younger; right, older) and sex (top. male; bottom, female) in mouse liver.

⁽G) Histograms of pairwise correlation coefficients from the CRLS, stratified by age (left, younger; right, older) and sex (top, male; bottom, female) in mouse liver. Vertical dashed lines represent median correlations.





proteins and their transcripts across tissues, as well as to identify global and tissue-specific patterns of aging at the protein level. This study functions as a quantitative protein resource for the aging-focused research community. We provide our data and corresponding processed results as an interactive Shiny application, available online at http://aging-b6-proteomics.jax.org, allowing proteins of interest to be easily queried. This tool can be used to confirm or replicate findings from previous studies in mice (BCAT1) or other models (VCAM1 in humans and REN1 in aging rat kidney) and generate hypotheses for future studies of aging.

Limitations of the study

Sample size (n = 20) is a key limitation of this study. We are primarily powered to detect fairly large differences between groups (age and sex); more subtle differences are likely to be undetected. The aging changes revealed by this study cover the span from adult to late midlife in mice and thus do not necessarily reflect biomedically relevant changes that occur only at late stages of life. All tissue samples were exhausted in generating these proteomics data, and thus further relevant -omics data (e.g., transcripts) cannot be collected for these specific mice.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. celrep.2023.112715.

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AUTHOR CONTRIBUTIONS

R.K., S.P.G., G.A.C., and D.K.S. conceptualized the project and designed the experiment. J.S. and D.K.S. performed the experiments. G.R.K and G.A.C. designed the methodology. G.R.K. performed the analysis. G.R.K. and J.-G.Z. built the interactive webtool. G.R.K., R.K., G.A.C., and D.K.S. wrote the manuscript. All authors reviewed the manuscript.

DECLARATION OF INTERESTS

G.R.K., R.K., J.-G.Z., and G.A.C. are employees of The Jackson Laboratory. S.P.G. serves on the scientific advisory board for Thermo Fisher Scientific.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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REFERENCES

- 1. López-Otín, C., Blasco, M.A., Partridge, L., Serrano, M., and Kroemer, G. (2013). The Hallmarks of Aging. Cell 153, 1194-1217.
- 2. Schaum, N., Lehallier, B., Hahn, O., Pálovics, R., Hosseinzadeh, S., Lee, S.E., Sit, R., Lee, D.P., Losada, P.M., Zardeneta, M.E., et al. (2020). Ageing hallmarks exhibit organ-specific temporal signatures. Nature 583, 596-602
- 3. Tabula Muris Consortium; Almanzar, N., Antony, J., Baghel, A.S., Bakerman, I., Bansal, I., Barres, B.A., Beachy, P.A., Berdnik, D., Bilen, B., et al. (2020). A single-cell transcriptomic atlas characterizes ageing tissues in the mouse. Nature 583, 590-595.
- 4. Shokhirev, M.N., and Johnson, A.A. (2021). Modeling the human aging transcriptome across tissues, health status, and sex. Aging Cell 20, e13280.
- 5. Zeng, L., Yang, J., Peng, S., Zhu, J., Zhang, B., Suh, Y., and Tu, Z. (2020). Transcriptome analysis reveals the difference between "healthy" and "common" aging and their connection with age-related diseases. Aging Cell 19, e13121.
- 6. Ori, A., Toyama, B.H., Harris, M.S., Bock, T., Iskar, M., Bork, P., Ingolia, N.T., Hetzer, M.W., and Beck, M. (2015). Integrated Transcriptome and Proteome Analyses Reveal Organ-Specific Proteome Deterioration in Old Rats. Cell Syst. 1, 224-237.

Resource



- 7. Walther, D.M., and Mann, M. (2011). Accurate Quantification of More Than 4000 Mouse Tissue Proteins Reveals Minimal Proteome Changes During Aging. Mol. Cell. Proteomics 10. M110.004523-S7.
- 8. Yu, Q., Xiao, H., Jedrychowski, M.P., Schweppe, D.K., Navarrete-Perea, J., Knott, J., Rogers, J., Chouchani, E.T., and Gygi, S.P. (2020). Sample multiplexing for targeted pathway proteomics in aging mice. Proc. Natl. Acad. Sci. USA 117, 9723-9732.
- 9. Li, Y., Yu, H., Chen, C., Li, S., Zhang, Z., Xu, H., Zhu, F., Liu, J., Spencer, P.S., Dai, Z., and Yang, X. (2020). Proteomic Profile of Mouse Brain Aging Contributions to Mitochondrial Dysfunction, DNA Oxidative Damage, Loss of Neurotrophic Factor, and Synaptic and Ribosomal Proteins. Oxid. Med. Cell. Longev. 2020, 5408452-5408521.
- 10. Takemon, Y., Chick, J.M., Gerdes Gyuricza, I., Skelly, D.A., Devuyst, O., Gygi, S.P., Churchill, G.A., and Korstanje, R. (2021). Proteomic and transcriptomic profiling reveal different aspects of aging in the kidney. Elife 10, e62585.
- 11. Gerdes Gyuricza, I., Chick, J.M., Keele, G.R., Deighan, A.G., Munger, S.C., Korstanje, R., Gygi, S.P., and Churchill, G.A. (2022). Genome-wide transcript and protein analysis highlights the role of protein homeostasis in the aging mouse heart. Genome Res. 32, 838-852.
- 12. Kelmer Sacramento, E., Kirkpatrick, J.M., Mazzetto, M., Baumgart, M., Bartolome, A., Di Sanzo, S., Caterino, C., Sanguanini, M., Papaevgeniou, N., Lefaki, M., et al. (2020). Reduced proteasome activity in the aging brain results in ribosome stoichiometry loss and aggregation. Mol. Syst. Biol. 16,
- 13. Anisimova, A.S., Meerson, M.B., Gerashchenko, M.V., Kulakovskiy, I.V., Dmitriev, S.E., and Gladyshev, V.N. (2020). Multifaceted deregulation of gene expression and protein synthesis with age. Proc. Natl. Acad. Sci. USA. 117, 15581-15590.
- 14. Schweppe, D.K., Eng, J.K., Yu, Q., Bailey, D., Rad, R., Navarrete-Perea, J., Huttlin, E.L., Erickson, B.K., Paulo, J.A., and Gygi, S.P. (2020). Full-Featured, Real-Time Database Searching Platform Enables Fast and Accurate Multiplexed Quantitative Proteomics. J. Proteome Res. 19, 2026-2034
- 15. Conway, J.R., Lex, A., and Gehlenborg, N. (2017). UpSetR: an R package for the visualization of intersecting sets and their properties. Bioinformatics 33, 2938-2940
- 16. Yu, G., Wang, L.-G., Han, Y., and He, Q.-Y. (2012). clusterProfiler: an R Package for Comparing Biological Themes Among Gene Clusters. OMICS A J. Integr. Biol. 16, 284-287.
- 17. Williams, E.G., Pfister, N., Roy, S., Statzer, C., Haverty, J., Ingels, J., Bohl, C., Hasan, M., Čuklina, J., Bühlmann, P., et al. (2022). Multiomic profiling of the liver across diets and age in a diverse mouse population. Cell Syst. 13, 43-57.e6.
- 18. Chick, J.M., Munger, S.C., Simecek, P., Huttlin, E.L., Choi, K., Gatti, D.M., Raghupathy, N., Svenson, K.L., Churchill, G.A., and Gygi, S.P. (2016). Defining the consequences of genetic variation on a proteome-wide scale. Nature 534, 500-505.
- 19. Keele, G.R., Zhang, T., Pham, D.T., Vincent, M., Bell, T.A., Hock, P., Shaw, G.D., Paulo, J.A., Munger, S.C., de Villena, F.P.M., et al. (2021). Regulation of protein abundance in genetically diverse mouse populations. Cell Genom. 1, 100003.
- 20. Romanov, N., Kuhn, M., Aebersold, R., Ori, A., Beck, M., and Bork, P. (2019). Disentangling Genetic and Environmental Effects on the Proteotypes of Individuals. Cell 177, 1308-1318.e10.
- 21. Zhang, L., and Han, J. (2017). Branched-chain amino acid transaminase 1 (BCAT1) promotes the growth of breast cancer cells through improving mTOR-mediated mitochondrial biogenesis and function. Biochem. Biophys. Res. Commun. 486, 224-231.
- 22. Jung, F.F., Kennefick, T.M., Ingelfinger, J.R., Vora, J.P., and Anderson, S. Down-Regulation of the Intrarenal Renin-Angiotensin System in the Aging Rat. J. Am. Soc. Nephrol., 8.

- 23. Mekada, K., and Yoshiki, A. (2021). Substrains matter in phenotyping of C57BL/6 mice. Exp. Anim. 70, 145-160.
- 24. Timmers, P.R.H.J., Tiys, E.S., Sakaue, S., Akiyama, M., Kiiskinen, T.T.J., Zhou, W., Hwang, S.-J., Yao, C., et al.; Biobank Japan Project; FinnGen (2022). Mendelian randomization of genetically independent aging phenotypes identifies LPA and VCAM1 as biological targets for human aging. Nat. Aging 2, 19-30.
- 25. Yousef, H., Czupalla, C.J., Lee, D., Chen, M.B., Burke, A.N., Zera, K.A., Zandstra, J., Berber, E., Lehallier, B., Mathur, V., et al. (2019). Aged blood impairs hippocampal neural precursor activity and activates microglia via brain endothelial cell VCAM1. Nat. Med. 25, 988-1000.
- 26. Perner, C., Perner, F., Gaur, N., Zimmermann, S., Witte, O.W., Heidel, F.H., Grosskreutz, J., and Prell, T. (2019). Plasma VCAM1 levels correlate with disease severity in Parkinson's disease. J. Neuroinflammation 16, 94.
- 27. Churchill, G.A., Gatti, D.M., Munger, S.C., and Svenson, K.L. (2012). The diversity outbred mouse population. Mamm. Genome 23, 713-718.
- 28. Murata, S., Takahama, Y., Kasahara, M., and Tanaka, K. (2018). The immunoproteasome and thymoproteasome: functions, evolution and human disease. Nat. Immunol. 19, 923-931.
- 29. Kimura, H., Caturegli, P., Takahashi, M., and Suzuki, K. (2015). New Insights into the Function of the Immunoproteasome in Immune and Nonimmune Cells. J. Immunol. Res. 2015, e541984.
- 30. Phillips, B.P., Gomez-Navarro, N., and Miller, E.A. (2020). Protein quality control in the endoplasmic reticulum. Curr. Opin. Cell Biol. 65, 96-102.
- 31. Guna, A., Volkmar, N., Christianson, J.C., and Hegde, R.S. (2018). The ER membrane protein complex is a transmembrane domain insertase. Science 359, 470-473.
- 32. Shurtleff M.J. Itzhak D.N. Hussmann J.A. Schirle Oakdale, N.T. Costa E.A., Jonikas, M., Weibezahn, J., Popova, K.D., Jan, C.H., Sinitcyn, P., et al. (2018). The ER membrane protein complex interacts cotranslationally to enable biogenesis of multipass membrane proteins. Elife 7, e37018.
- 33. Szul, T., and Sztul, E. (2011). COPII and COPI Traffic at the ER-Golgi Interface. Physiology 26, 348-364.
- 34. Huttlin, E.L., Bruckner, R.J., Navarrete-Perea, J., Cannon, J.R., Baltier, K., Gebreab, F., Gygi, M.P., Thornock, A., Zarraga, G., Tam, S., et al. (2021). Dual proteome-scale networks reveal cell-specific remodeling of the human interactome. Cell 184, 3022-3040.e28.
- 35. Huttlin, E.L., Ting, L., Bruckner, R.J., Gebreab, F., Gygi, M.P., Szpyt, J., Tam, S., Zarraga, G., Colby, G., Baltier, K., et al. (2015). The BioPlex Network: A Systematic Exploration of the Human Interactome. Cell 162, 425-440.
- 36. Szklarczyk, D., Gable, A.L., Lyon, D., Junge, A., Wyder, S., Huerta-Cepas, J., Simonovic, M., Doncheva, N.T., Morris, J.H., Bork, P., et al. (2019). STRING v11: protein-protein association networks with increased coverage, supporting functional discovery in genome-wide experimental datasets. Nucleic Acids Res. 47, D607-D613.
- 37. Lapek, J.D., Greninger, P., Morris, R., Amzallag, A., Pruteanu-Malinici, I., Benes, C.H., and Haas, W. (2017). Detection of dysregulated protein-association networks by high-throughput proteomics predicts cancer vulnerabilities. Nat. Biotechnol. 35, 983-989.
- 38. Taggart, J.C., Zauber, H., Selbach, M., Li, G.-W., and McShane, E. (2020). Keeping the Proportions of Protein Complex Components in Check. Cell Syst. 10, 125-132.
- 39. Giurgiu, M., Reinhard, J., Brauner, B., Dunger-Kaltenbach, I., Fobo, G., Frishman, G., Montrone, C., and Ruepp, A. (2019). CORUM: the comprehensive resource of mammalian protein complexes - 2019. Nucleic Acids Res. 47, D559-D563.
- 40. Ori, A., Iskar, M., Buczak, K., Kastritis, P., Parca, L., Andrés-Pons, A., Singer, S., Bork, P., and Beck, M. (2016). Spatiotemporal variation of mammalian protein complex stoichiometries. Genome Biol. 17, 47.
- 41. Vinayagam, A., Hu, Y., Kulkarni, M., Roesel, C., Sopko, R., Mohr, S.E., and Perrimon, N. (2013). Protein Complex-Based Analysis Framework for High-Throughput Data Sets. Sci. Signal. 6, rs5.



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- 42. Leitner, A., Joachimiak, L.A., Bracher, A., Mönkemeyer, L., Walzthoeni, T., Chen, B., Pechmann, S., Holmes, S., Cong, Y., Ma, B., et al. (2012). The Molecular Architecture of the Eukaryotic Chaperonin TRiC/CCT. Structure 20, 814-825.
- 43. Kalisman, N., Adams, C.M., and Levitt, M. (2012). Subunit order of eukaryotic TRiC/CCT chaperonin by cross-linking, mass spectrometry, and combinatorial homology modeling. Proc. Natl. Acad. Sci. USA 109, 2884-2889.
- 44. Grantham, J. (2020). The Molecular Chaperone CCT/TRiC: An Essential Component of Proteostasis and a Potential Modulator of Protein Aggregation. Front. Genet. 11, 172.
- 45. Toyama, B.H., and Hetzer, M.W. (2013). Protein homeostasis: live long, won't prosper. Nat. Rev. Mol. Cell Biol. 14, 55-61. https://doi.org/10. 1038/nrm3496
- 46. Cheng, J., North, B.J., Zhang, T., Dai, X., Tao, K., Guo, J., and Wei, W. (2018). The emerging roles of protein homeostasis-governing pathways in Alzheimer's disease. Aging Cell 17, e12801.
- 47. Morimoto, R.I., and Cuervo, A.M. (2009). Protein Homeostasis and Aging: Taking Care of Proteins From the Cradle to the Grave. J. Gerontol. A Biol. Sci. Med. Sci. 64, 167-170.
- 48. Morimoto, R.I., and Cuervo, A.M. (2014). Proteostasis and the Aging Proteome in Health and Disease. J. Gerontol. A Biol. Sci. Med. Sci. 69,
- 49. Stein, K.C., Morales-Polanco, F., van der Lienden, J., Rainbolt, T.K., and Frydman, J. (2022). Ageing exacerbates ribosome pausing to disrupt cotranslational proteostasis. Nature 601, 637-642.
- 50. Yuan, R., Musters, C.J.M., Zhu, Y., Evans, T.R., Sun, Y., Chesler, E.J., Peters, L.L., Harrison, D.E., and Bartke, A. (2020). Genetic differences and longevity-related phenotypes influence lifespan and lifespan variation in a sex-specific manner in mice. Aging Cell 19, e13263.
- 51. Newman, A.M., Steen, C.B., Liu, C.L., Gentles, A.J., Chaudhuri, A.A., Scherer, F., Khodadoust, M.S., Esfahani, M.S., Luca, B.A., Steiner, D., et al. (2019). Determining cell type abundance and expression from bulk tissues with digital cytometry. Nat. Biotechnol. 37, 773-782.
- 52. Kelly, R.T. (2020). Single-cell Proteomics: Progress and Prospects. Mol. Cell. Proteomics 19, 1739-1748.
- 53. Labib, M., and Kelley, S.O. (2020). Single-cell analysis targeting the proteome. Nat. Rev. Chem 4, 143-158.
- 54. Pham, T., Tyagi, A., Wang, Y.-S., and Guo, J. (2021). Single-cell proteomic analysis. WIREs Mech. Dis. 13, e1503.
- 55. Bates, D.W., Mächler, M., Bolker, B., and Walker, S. (2015). Fitting Linear Mixed-Effects Models Using Ime4. BMJ Qual. Saf. 24, 1-3.

- 56. Kuznetsova, A., Brockhoff, P.B., and Christensen, R.H.B. (2017). ImerTest Package: Tests in Linear Mixed Effects Models. J. Stat. Software 82.
- 57. Love, M.I., Huber, W., and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15, 550.
- 58. Korotkevich, G., Sukhov, V., Budin, N., Shpak, B., Artyomov, M.N., and Sergushichev, A. (2021). Fast gene set enrichment analysisPrecheck. bio-Rxiv. https://doi.org/10.1101/060012.
- 59. R Core Team (2020). R: A Language and Environment for Statistical Computing (R Foundation for Statistical Computing).
- 60. Navarrete-Perea, J., Yu, Q., Gygi, S.P., and Paulo, J.A. (2018). Streamlined Tandem Mass Tag (SL-TMT) Protocol: An Efficient Strategy for Quantitative (Phospho)proteome Profiling Using Tandem Mass Tag-Synchronous Precursor Selection-MS3. J. Proteome Res. 17, 2226-2236.
- 61. Eng, J.K., Jahan, T.A., and Hoopmann, M.R. (2013). Comet: An opensource MS/MS sequence database search tool. Proteomics 13, 22-24.
- 62. Rad, R., Li, J., Mintseris, J., O'Connell, J., Gygi, S.P., and Schweppe, D.K. (2021). Improved Monoisotopic Mass Estimation for Deeper Proteome Coverage. J. Proteome Res. 20, 591-598.
- 63. Cunningham, F., Allen, J.E., Allen, J., Alvarez-Jarreta, J., Amode, M., Armean, I., Austine-Orimoloye, O., Azov, A., Barnes, I., Bennett, R., et al. (2021). Ensembl 2022. Nucleic Acids Res. 50, D988-D995.
- 64. Huttlin, E.L., Jedrychowski, M.P., Elias, J.E., Goswami, T., Rad, R., Beausoleil, S.A., Villén, J., Haas, W., Sowa, M.E., and Gygi, S.P. (2010). A Tissue-Specific Atlas of Mouse Protein Phosphorylation and Expression. Cell 143, 1174-1189.
- 65. Stacklies, W., Redestig, H., Scholz, M., Walther, D., and Selbig, J. (2007). pcaMethods-a bioconductor package providing PCA methods for incomplete data. Bioinformatics 23, 1164-1167.
- 66. Benjamini, Y., and Hochberg, Y. (1995). Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. J. Roy. Stat. Soc. B 57, 289-300.
- 67. Giesbrecht, F.G., and Burns, J.C. (1985). Two-Stage Analysis Based on a Mixed Model: Large-Sample Asymptotic Theory and Small-Sample Simulation Results. Biometrics 41, 477-486.
- 68. Hrong-Tai Fai, A., Cornelius, P.L., and Cornelius, P.L. (1996). Approximate F-tests of multiple degree of freedom hypotheses in generalized least squares analyses of unbalanced split-plot experiments. null 54, 363–378.
- 69. Anders, S., and Huber, W. (2010). Differential Expression Analysis for Sequence Count Data12.
- 70. Dolgalev, I. (2021). Msigdbr: MSigDB Gene Sets for Multiple Organisms in a Tidy Data Format.

Cell ReportsResource



STAR***METHODS**

KEY RESOURCES TABLE

SOURCE	IDENTIFIER
Thermo Fisher	A34808
Wako Chemicals	Barcode No. 4987481427648
	V5113
Roche	4693116001
Millipore Sigma	I1149-25G
, ,	D0632-10G
Thermo Fisher	23227
ProteomeXchange (http://www.proteomexchange.org)	PXD034029
Schaum et al.; Gene Expression Omnibus (https://www.ncbi.nlm. nih.gov/geo/)	GSE132040
Yu et al.; ProteomeXchange (http://www.proteomexchange.org)	PXD017385
https://doi.org/10.6084/m9.figshare. 19765849	N/A
http://aging-b6-proteomics.jax.org; https://github.com/gkeele/Aging_ B6_Proteomics_RShiny	N/A
The Jackson Laboratory	JAX: 000664
Charles River Laboratories	N/A
The Jackson Laboratory	JAX: 009376
Bates et al. ⁵⁵	https://cran.r-project.org/web/ packages/lme4/index.html; RRID: SCR_015654
Kuznetsova et al. ⁵⁶	https://cran.r-project.org/web/ packages/lmerTest/index.html; RRID: SCR_015656
Love et al. ⁵⁷	https://bioconductor.org/ packages/release/bioc/html/ DESeq2.html; RRID: SCR_015687
Yu et al. ¹⁶	https://bioconductor.org/ packages/release/bioc/html/ clusterProfiler.html; RRID: SCR_016884
Korotkevich et al. ⁵⁸	https://bioconductor.org/ packages/release/bioc/ html/fgsea.html; RRID: SCR_020938
	Thermo Fisher Wako Chemicals Promega Roche Millipore Sigma Millipore Sigma Thermo Fisher ProteomeXchange (http://www.proteomexchange.org) Schaum et al.; Gene Expression Omnibus (https://www.ncbi.nlm. nih.gov/geo/) Yu et al.; ProteomeXchange (http://www.proteomexchange.org) https://doi.org/10.6084/m9.figshare. 19765849 http://aging-b6-proteomics.jax.org; https://github.com/gkeele/Aging_ B6_Proteomics_RShiny The Jackson Laboratory Charles River Laboratories The Jackson Laboratory Bates et al. 55 Kuznetsova et al. 56 Love et al. 57 Yu et al. 16

(Continued on next page)





Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
R	The R Project	https://www.r-project.org; RRID: SCR_001905
Shiny		https://cran.r-project.org/ web/packages/shiny/ index.html; RRID: SCR_001626
Other		
Protein complex annotations	Ori et al. ⁴⁰	http://doi.org/10.1186/s13059-016-0912-5
Waters 100mg Sep-Pak	Waters	WAT036820
Orbitrap Fusion Lumos	Thermo Fisher	IQLAAEGAAPFADBMBHQ
Zorbax 300 Extend C18 column	Agilent	770995-902
1260 Infinity II LC System	Agilent	

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Devin Schweppe (dkschwep@uw.edu).

Materials availability

None generated, see Date and code availability for more information on proteomics data.

Data and code availability

- The mass-spec proteomics data for all samples reported here have been deposited in ProteomeXchange (http://www. proteomexchange.org/) via the PRIDE partner repository (ProteomeXchange: PXD034029). All statistical analyses were performed using the R statistical programming language (v4.0.3)⁵⁹.
- All starting data, key forms of processed data, and the analysis pipeline to process the data, run analysis, and produce the reported findings have been made publicly available at figshare (https://doi.org/10.6084/m9.figshare.19765849). The processed data are also interactively viewable through an Shiny application, which is available online (http://aging-b6-proteomics.jax.org) or can be downloaded from GitHub (https://github.com/gkeele/Aging_B6_Proteomics_RShiny) and run locally through RStudio (https://posit.co).
- Any additional information required to reanalyze the data reported in this work paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Mice

Female and male C57BL/6J mice (stock JR#000664) were obtained from The Jackson Laboratory. Animals were maintained on pine shavings and given a standard rodent diet (LabDiet 5KOG) and acidified water in a pathogen free room (health report included in figshare repository). The room was maintained at 21°C with a 12-hour light/dark cycle (6am to 6pm). At the time of tissue collection (at 8 and 18 months of age) animals were euthanized by cervical dislocation. Kidney, liver, fat (inquinal adipose), spleen, lung, heart, skeletal muscle (quadriceps), striatum, cerebellum, and hippocampus tissues were collected from each animal. Whole organs were used for kidney, liver, spleen, lung, and heart. All animal experiments were performed in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals (National Research Council) and were approved by The Jackson Laboratory's Animal Care and Use Committee.

METHOD DETAILS

Sample preparation for proteomics analysis

Tissue samples were dounce homogenized and resuspended in lysis buffer (8M urea, 150 mM NaCl, Roche protease inhibitor tablets) and cells were lysed by sonication (procedure). Lysates were cleared by centrifugation (15 min at 20,000xg) and protein concentrations were measured using Pierce BCA assay kits. Proteins were then reduced with dithiothreitol (5mM for 30 minutes at room

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temperature) and alkylated with iodoacetamide (15mM for 60 minutes in the dark). The alkylation reaction was quenched by adding an additional aliquot of DTT. For each sample, 100ug of protein was aliquoted and diluted to a final concentration of 1mg/mL. The bridge channel included equal amounts protein from each tissue for all 20 mice as 20µg per mouse split into two bridge samples (100µg each) and added to each of two plexes per tissue. Proteins were digested using LysC (Wako, overnight, room temperature, moderate agitation) followed by trypsin (6 hr, 37C, 200rpm). The resulting peptides were then labeled with individual TMT (Thermo) reporters (1.5 hours at room temperature) and the reaction was quenched with hydroxylamine (5% in water for 5 minutes). Labeled peptides were mixed into a set of two plexed for each tissue analysis. After labeling and mixing, peptide mixtures were desalted using C18 seppak cartridges (1mg, Waters). Desalted peptides were then fractionated using basic-pH reverse phase chromatography⁶⁰. Briefly, peptides were resuspended in Buffer A (10mM ammonium bicarbonate, 5% acetonitrile [ACN], pH 8) and separated on a linear gradient from 13% to 42% Buffer B (10mM ammonium bicarbonate, 90% acetonitrile [ACN], pH 8) over an Agilent 300Extend C18 column using an Agilent 1260 HPLC equipped with single wavelength detection at 220nm). Fractionated peptides were desalted using Stage-tips⁶⁰ prior to LC-MS/MS analysis.

Mass spectra data analysis

Peptides were separated prior to MS/MS analysis using an Easy-nLC (Thermo) equipped with an in-house pulled fused silica capillary column with integrated emitter packed with Accucore C18 media (Thermo). Separation was carried out with 90-minute gradients from 96% Buffer A (5% ACN, 0.125% formic acid) to 30% Buffer B (90% ACN, 0.125% formic acid). Mass spectrometric analysis was carried out on an Orbitrap Fusion Lumos (Thermo). Multiplexed analysis of samples was done using real-time search data acquisition 4, based on canonical SPS-MS3 acquisition. Briefly, a survey MS1 scan was used to identify potential peptide precursors (R = 120000, Mass range: 400-2000 m/z, max Inject time: 50ms, AGC: 200000, RF lens: 30%). The top 10 precursors were selected for fragmentation and analysis in the ion trap (Dynamic exclusion: 120s at 10ppm, CID collision energy: 35%, max inject time: 120ms, AGC: 20000, scan rate: rapid, isolation width: 0.5 m/z). Real-time spectral matching was carried out using the Comet search algorithm⁶¹. If, and only if, a peptide was matched with high confidence, the instrument would then acquire an additional SPS-MS3 scan for quantification of relative abundances (R = 50000, HCD NCE: 65, max inject time: 200ms).

Raw spectral information was converted to mzXML format using Monocle⁶², and spectra were matched using the Comet search algorithm comparing against the ENSEMBL_GRCm39 database^{61,63}. Peptides and proteins were filtered to a 1% using rules of protein parsimony⁶¹.

QUANTIFICATION AND STATISTICAL ANALYSIS

Protein abundance estimation from peptides

Samples from each tissue were run across two tissue-specific batches. For each tissue, the abundance level for proteins was estimated as a scaled sum of their component peptides. For protein j from tissue k of mouse i, the abundance is calculated as y_{ijk} $\frac{\sum_{M} y'_{imk} 1_{imk}}{\theta_{ik}}$ where M is the set of peptides that map to protein j, y'_{imk} is the intensity of peptide m from tissue k of mouse i, 1_{imk} is

the indicator function that peptide m was observed in tissue k of mouse i, and $\theta_{ik} = \frac{\sum_{p} y'_{ip}}{\max_{l \in Bijk!} (\sum_{p} y'_{l})}$ is the within-batch scaling factor⁶⁴

for mouse i in tissue k, representing the ratio of the sum of all peptide intensities for mouse i to the maximum sum total for the 11 samples in the batch of mouse i for tissue k, B[ik]. To standardize quantities across the two batches, abundances were ratio normalized to a pooled sample that was included in both batches: $\tilde{y}_{ijk} = log_2\left(\frac{y_{ijk}+1}{y_{b_ikijk}+1}\right)$ where b[ik] is the bridge sample from the batch of mouse i for tissue k.

Filtering out low quality proteins and samples

We filtered out proteins that were only observed in one of the two batches for a tissue because we found single batch proteins could be influential in downstream analysis. After protein abundance estimation and removal of single batch proteins, we performed PCA 65 to identify tissue samples that were clear outliers across many proteins. We removed one sample from liver (young male), fat (young male), spleen (old male), lung (young male), and skeletal muscle (old female). Two samples were removed from striatum, both old female mice.

Testing for age, sex, and age-by-sex interaction effects on proteins within tissues

We tested for significant differences based on age, sex, and age-by-sex using ordinary least squares (OLS) regression. For each protein j observed in tissue k, we fit the following model:

$$\tilde{y}_i = \mu + \beta_{old} X_i^{old} + \beta_{female} X_i^{female} + \beta_{batch} X_i^{batch} + \varepsilon_i$$
 Equation

where μ is the intercept, β_{old} represents the effect of being in the old group, β_{female} represents the effect of being female, β_{batch} represents the effect of being in second batch, x_i^{old} , x_i^{female} , and x_i^{batch} are indicator variables that mouse i is old, female, or in batch2, respectively, and ε_i is the error for mouse i, modeled as $\varepsilon_i \sim N(0, \sigma^2)$ and σ^2 is the variance of the noise. To test for an age effect,



we used analysis of variance (ANOVA), comparing the model in Equation 1 to a model excluding the age term and recorded the age effect coefficients, standard errors, and p-values. Similarly, we tested for a sex effect by comparing the Equation 1 model to a model excluding the sex term and again recorded sex effect coefficients, standard errors, and p-values. Finally, we assessed age-by-sex differences by adding an age-by-sex interaction term to the model which was then compared to the Equation 1 model with ANOVA and recorded the age-by-sex p-value. We estimated the FDR using the Benjamini-Hochberg (BH) method⁶⁶ for each effect type, producing age, sex, and age-by-sex q-values. This process was used each of the 10 tissues.

When plotting data (not effect parameters), we first regressed out the effect of batch to make the signal from age or sex clearer. We fit the model in Equation 1 and then calculated $\tilde{\hat{y}}_i = \tilde{y}_i - \hat{\beta}_{batch} x_i^{batch}$, where $\hat{\beta}_{batch}$ is the estimated coefficient for the second batch, for all proteins *j* across all tissues *k*.

Testing for consistent and unique age and sex effects on proteins across tissues

We tested whether age and sex effects on a protein were similar or distinct across tissues for all proteins detected in more than one tissue using linear mixed effects models (LMM) fit in the Ime4 R package⁵⁵. For each protein j, we fit the following model:

$$\tilde{y}_{ik} = \mu + \beta_{old} X_i^{old} + \beta_{female} X_i^{female} + tissue[k] + u_i + \varepsilon_{ik}$$
 Equation 2

where tissue [k] represents the effect of tissue k, u_i is a random term specific to mouse i, modeled as $u_i \sim N(0,\tau^2)$, τ^2 is the variance component underlying the mouse-specific effect, and all other terms as previously defined. A batch effect was not included because tissue and batch are highly confounded (batch pairs specific to each tissue), and we are not interested in the marginal effect of either tissue or batch. We tested for a consistent age effect across tissues by comparing the model in Equation 2 to reduced model excluding the age effect with ANOVA using Satterthwaite's approximation for an LMM^{67,68}, implemented in the ImerTest R package⁵⁶. We next looked for age effects that were unique to tissues or were even flipped by testing an age-by-tissue term by comparing the model in Equation 2 to an expanded model that included the interaction term. The same approach was used to test for consistent sex effects across tissues and tissue-unique and flipped sex effects. To account for multiple testing across proteins, we again estimated q-values using the BH method⁶⁶.

Modeling how factors contribute to variation in proteins measured in all 10 tissues

To evaluate how various technical and biological factors contribute to variation in the abundance of individual proteins, we fit an LMM with variance components for each factor for all proteins measured in all 10 tissues. For each protein j, we fit the following model:

$$\tilde{y}_{ik} = \mu + \beta_{old} X_i^{old} + \beta_{female} X_i^{female} + u_i^{Mouse} + u_{t[i]}^{Tag} + u_{b[i]}^{Batch} + \varepsilon_{ik}$$
 Equation 3

where u_i^{Mouse} is a random term specific to mouse i (20 levels), $u_{t[i]}^{Tag}$ is a random term for TMT tag t of mouse i (10 levels), $u_{b[i]}^{Batch}$ is a random term for TMT batch b of mouse i (20 levels), and all other terms as previously defined. Each random effect is modeled as $u_{[i]}^{factor} \sim N(0, \tau_{factor}^2)$. Proportion of variation explained by each factor was calculated as $VarP_{factor} = \frac{\hat{\tau}_{factor}^2}{\hat{\tau}_{Mouse}^2 + \hat{\tau}_{Tag}^2 + \hat{\tau}_{Batch}^2 + \hat{\sigma}^2}$. For point and interval estimates of the random terms, best linear unbiased predictors (BLUPs) and 95% predictive intervals were used.

Testing for age and sex effects on transcripts within tissues

We obtained transcriptomics data reported in Schaum et al 2020², which represent 17 tissues and 10 age groups (1 to 27 months) from the closely related C57BL/6JN (B6N) mice. Each age group consisted of samples from four males and two females, except for 24 and 27 months, which only had the four males. We filtered the data to the 9 and 18 months-old age groups, which most closely match our age groups of 8 and 18 months. We then tested for and characterized age and sex effects (log fold change) on gene expression within each tissue. We used the DESeq2 R package⁵⁷ to fit models similar to Equation 1 (excluding the batch covariate), but now using a negative binomial generalized linear model (GLM) to accommodate that the data are gene counts. Age and sex effects on proteins and transcripts were compared based on aligning Ensembl gene IDs. When plotting the data to illustrate effects, we first used DESeq2's variance stabilizing transformation⁶⁹ across samples from all tissues and age groups.

Age and sex effects on proteins from previously published murine proteomics data

We obtained proteomics data from hearts¹¹ and kidneys¹⁰ from genetically diverse DO mice. These studies included three age groups (6, 12, and 18 months-old). Rather than re-estimating age and sex effects from the individual-level data, we used the publicly available effects summaries from the studies. For age, the effect represents a regression coefficient corresponding to age fit as a continuous variable.

We also obtained proteomics data from nine tissues from 10 B6 males8. These data included both targeted (Tomahto) and untargeted mass-spec proteomics. We estimated age effects from the individual-level data (for both targeted and untargeted) use ANOVA, using a model similar to Equation 1 but without a sex or batch term.

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Protein complex annotations

To define the set of protein complexes to assess, we used the annotations from Ori et al 2016⁴⁰, which were manually curated from their in-house data along with resources like the CORUM database³⁹ and COMPLEAT protein complex resource⁴¹. We first filtered out proteins that did not have an orthologous ENSEMBL protein ID in mouse, and then filtered out protein complexes that did not have four or more proteins observed in one of the tissues. This resulted in 228 protein complexes across the 10 tissues.

Summarizing protein complex co-abundance

We quantified how co-abundant, i.e., cohesive, a protein was with its complex⁴⁰ as the median Pearson correlation coefficient between it and other complex members. An overall summary for the complex was then taken as the median across all the individual medians, an approach we used previously 19. We calculated cohesiveness only for protein complexes with more than three member proteins observed for a given tissue.

Testing for consistent age and sex effects on abundance across a protein complex

We tested for consistent age and sex effects on protein complexes with more than three observed members by jointly modeling all proteins. For each protein complex c observed in tissue k, we fit the following LMM for proteins $j \in J_c$:

$$\tilde{y}_{ij} = \mu + \beta_{old} X_i^{old} + \beta_{female} X_i^{female} + U_i^{Mouse} + U_j^{Protein} + \varepsilon_{ij}$$
 Equation 4

where J_c is the set of proteins in complex c, $u_i^{Protein}$ is a random term for protein j, modeled as $u_j^{Protein} \sim N(0, \tau_{Protein}^2)$, $\tau_{Protein}^2$ is the variance component underlying proteins, and all other terms as previously defined. We tested for an age or sex effect by comparing the model in Equation 4 to either a model excluding the age or sex term, respectively, through ANOVA, again with Satterthwaite's approximation ^{67,68}, producing a *p*-value. FDR was again estimated using the BH method ⁶⁶ across protein complexes and tissues, for age and sex separately.

Testing for age and sex effects on protein complex cohesiveness

Age or sex could affect how tightly co-abundant a protein complex is. We evaluated whether age or sex had a consistent effect on the correlation structure of the complex using a paired t-test. For example, with age, we calculated all the pairwise correlations between members for both age groups for a given complex within a tissue: (r_{Old}, r_{Young}) , resulting in a t-test p-value and effect. This process was repeated based on sex. The BH method 66 was again used to estimate FDR and produce t-test q-values for both age and sex. We used a more stringent significance threshold of FDR < 0.01 because correlation coefficients are non-standard quantities to model with a t-test.

Testing changes in correlation between pairs of proteins with permutations

To test whether individual pairwise correlations between protein complex members differed based on age or sex, we used permutations. When testing for an age difference, we swapped mouse identifiers among males and among females, thus maintaining the effect of sex and potentially avoiding anti-conservative permutation p-values. When testing for a sex difference, we swapped labels while maintaining the age groups. We estimated empirical p-values as $\frac{1}{P}\sum_{P}J(|r_{mn}|>\left|r_{mn}^{(p)}\right|)$ where P is the number of permutations, I(.)is an indicator function, $|r_{mn}|$ is the observed absolute Pearson correlation coefficient between proteins m and n, and $r_{mn}^{(p)}$ is the absolute Pearson correlation coefficient for permutation p. We set P to 100,000.

Gene set enrichment analyses

We used both set-based and score-based gene set enrichment analyses and confirmed that they often conferred with each other. We used the clusterProfiler R package 16 for set-based analysis, in which we defined gene sets based on various criteria, such as all genes from a given tissue that have significant age or sex effects on protein abundance compared to all genes analyzed in the tissue. We also evaluated tissue-specific gene sets defined by the direction of the significant age or sex effect for given tissues or based on how genes clustered according to age effects across the tissues. For the score-based analysis, we used the fgsea R package⁵⁸ paired with GO pathways from the msigdbr R package⁷⁰. For each tissue, we input all analyzed genes with scores as the age or sex coefficients from Equation 1, standardized by their standard errors. We compared GO findings between pairs of tissues by intersecting tissue-specific results based on pathway ID and gene ID.