

RESEARCH ARTICLE SUMMARY

HUMAN CELL ATLAS

Single-nucleus cross-tissue molecular reference maps toward understanding disease gene function

Gökçen Eraslan[†], Eugene Drokhyansky[†], Shankara Anand[†], Evgenij Fiskin[†], Ayshwarya Subramanian[†], Michal Slyper[†], Jiali Wang[†], Nicholas Van Wittenberghe, John M. Rouhana, Julia Waldman, Orr Ashenberg, Monkol Lek, Danielle Dionne, Thet Su Win, Michael S. Cuoco, Olena Kuksenok, Alexander M. Tsankov, Philip A. Branton, Jamie L. Marshall, Anna Greka, Gad Getz, Ayellet V. Segre^{*§}, François Aguet^{*§}, Orit Rozenblatt-Rosen^{*§}, Kristin G. Ardlie^{*§}, Aviv Regev^{*§}

INTRODUCTION: Understanding and treating disease requires deep, systematic characterization of different cells and their interactions across human tissues and organs, along with characterization of the genetic variants that causally contribute to disease risk. Recent studies have combined single-cell atlases of specific human tissues and organs with genes associated with human disease to relate risk variants to likely cells of action. However, it has been challenging to extend these studies to profile multiple tissues and organs across the body, conduct studies at population scale, and integrate cell atlases from multiple organs to yield unified insights.

RATIONALE: Because of the pleiotropy and specificity of disease-associated variants, systematically relating variants to cells and molecular processes requires analysis across multiple tis-

suess and individuals. Prior cell atlases primarily relied on fresh tissue samples from a single organ or tissue. Single-nucleus RNA sequencing (snRNA-seq) can be applied to frozen, archived tissue and captures cell types that do not survive dissociation across many tissues. Deep learning methods can integrate data across individuals and tissues by controlling for batch effects while preserving biological variation.

RESULTS: We established a framework for multi-tissue human cell atlases and generated an atlas of 209,126 snRNA-seq profiles from eight tissue types across 16 individuals, archived as frozen tissue as part of the Genotype-Tissue Expression (GTEx) project. We benchmarked four protocols and show how to apply them in a pooled setting to enable larger studies. We integrated the cross-tissue atlas using a conditional variational autoencoder, annotated it

with 43 broad and 74 fine categories, and demonstrated its use to decipher tissue residency, such as a macrophage dichotomy and lipid associations that are preserved across tissues, and tissue-specific fibroblast features, including lung alveolar fibroblasts with likely roles in mechanosensation. We relate cells to human disease biology and disease-risk genes for both rare and common diseases, including rare muscle disease gene groups enriched in distinct subsets of myonuclei and nonmyocytes, and cell type-specific enrichment of expression and splicing quantitative trait locus (QTL) target genes mapped to genome-wide association study loci.

CONCLUSION: Our framework will empower large, cross-tissue population and/or disease studies at single-cell resolution. These frameworks and the cross-tissue perspective provided here will form a basis for larger-scale future studies to improve our understanding of cross-tissue and cross-individual variation of cellular phenotypes in relation to disease-associated genetic variation. ■

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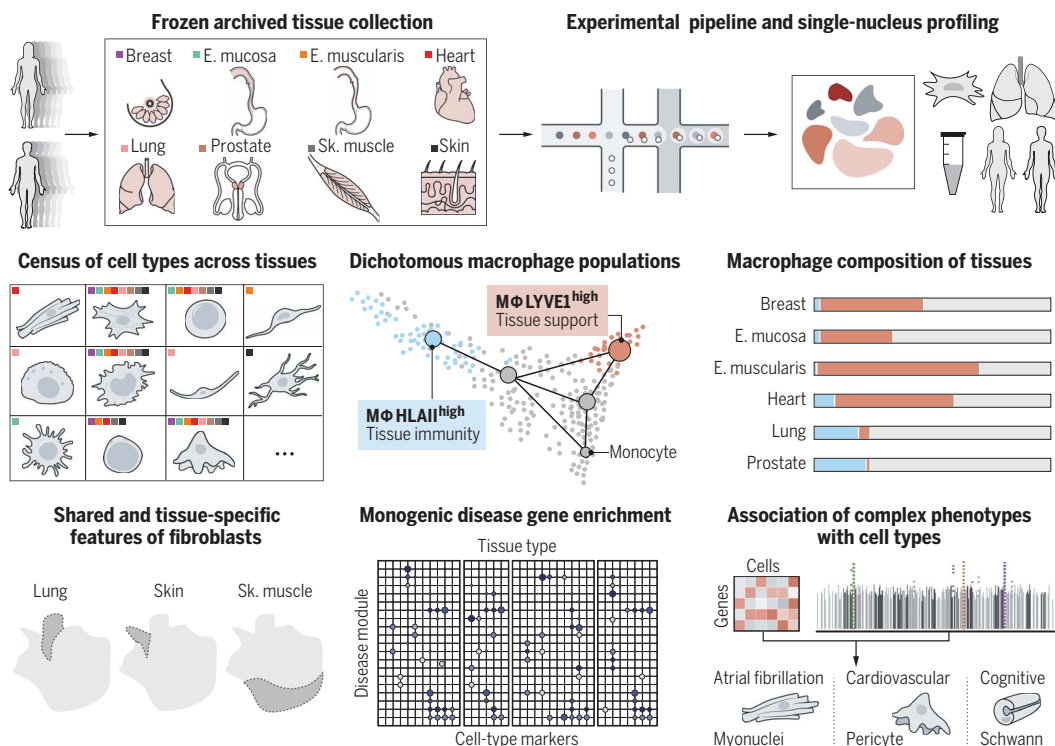
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Cross-tissue snRNA-seq atlas in eight frozen, archived adult human tissues.

Tissue sites and experimental pipeline (top row). The resulting atlas enables a cross-tissue census of tissue-specific and shared cell types (middle left). Differentiation trajectories and compositional analysis of dichotomous macrophage populations improve our understanding of tissue residency (middle center and right). Analyses of fibroblasts across tissues reveal tissue-specific and shared fibroblast features and their functional interpretation (bottom left). Robust and scalable computational methods enable comprehensive associations of monogenic and complex diseases to tissue-specific and shared cell populations (bottom center and right). E. mucosa, esophagus mucosa; E. muscularis, esophagus muscularis; Sk. muscle, skeletal muscle.



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Understanding gene function and regulation in homeostasis and disease requires knowledge of the cellular and tissue contexts in which genes are expressed. Here, we applied four single-nucleus RNA sequencing methods to eight diverse, archived, frozen tissue types from 16 donors and 25 samples, generating a cross-tissue atlas of 209,126 nuclei profiles, which we integrated across tissues, donors, and laboratory methods with a conditional variational autoencoder. Using the resulting cross-tissue atlas, we highlight shared and tissue-specific features of tissue-resident cell populations; identify cell types that might contribute to neuromuscular, metabolic, and immune components of monogenic diseases and the biological processes involved in their pathology; and determine cell types and gene modules that might underlie disease mechanisms for complex traits analyzed by genome-wide association studies.

Tissue homeostasis and pathology arise from an intricate interplay between different cell types, such that disease risk is influenced by variation in genes that affect the cells' functions and interactions. Human genetics studies, to date, have mapped tens of thousands of loci that either underlie rare monogenic disease or are associated with complex polygenic disease risk (1–3), including many in regulatory regions, whereas single-cell genomics has become instrumental in constructing cell atlases of both healthy organs and diseased tissues (4–6).

Coupling these advances in human genetics and single-cell genomics should enhance our understanding of cell type-specific changes in the function and regulation of disease genes. In particular, tissue (7), cell type (8–11), time point, and stimulation (12–14) all affect gene expression in disease-associated genetic loci. Recently, studies combining single-cell expression atlases and genetic signals have been able to associate risk genes with specific cell types and states in relevant tissues (15–18).

Because complex diseases often manifest in and implicate cells across multiple tissues, fully understanding the way in which genetic variation affects disease requires generating atlases from diverse tissues across the body and from many individuals, spanning different populations. This poses several challenges. First, collecting fresh tissue samples at scale is logistically challenging, and some tissues, such as brain, muscle, and adipose, are difficult to process into single-cell suspensions (19–23). As a result, large-scale single-cell profiling studies in human populations (24, 25) have focused on peripheral blood mononuclear cells, which can be frozen and thawed for multiplexed single-cell analysis. Single-nucleus RNA sequencing (snRNA-seq) offers a compelling alternative because it can be applied to archived, frozen tissues (26, 27) from multiple organs and captures diverse cell types. Second, annotation and classification of cell types and states require defining biological relationships between parenchymal, immune, and stromal cells across tissue types. Finally, data integration and interpretation require cross-tissue analytical frameworks to remove unwanted vari-

ation while preserving biological differences; identify cell types and states; and relate cell types and states to monogenic and complex trait genetics.

Results

A multitissue, multi-individual single-nucleus reference atlas from archived, frozen human tissues

We constructed a cross-tissue snRNA-seq atlas from 25 archived, frozen tissue samples, previously collected and banked by the Genotype-Tissue Expression (GTEx) project (7), that span three or four samples from each of eight tissue sites—breast, esophagus mucosa, esophagus muscularis, heart, lung, prostate, skeletal muscle, and skin—from 16 individuals (seven males and nine females) (Fig. 1A). We selected the samples by RNA quality, tissue autolysis score, and the availability of existing bulk RNA-seq and genome sequencing data [(28); table S1]. Histology slides corresponding to each tissue were reviewed by a pathologist to provide detailed annotations (table S1). Because different nucleus extraction protocols can be optimal for different tissues (26, 29), we isolated nuclei from each sample using four protocols that vary in detergents, salt, buffer, and mechanical preparation methodology [CST (0.49% CHAPS detergent, salts, and Tris buffer), NST (NP-40, salts, and Tris buffer), TST (0.03% Tween 20 detergent, salts, and Tris buffer), and the EZ nuclei isolation kit (proprietary composition; NUC101, Sigma-Aldrich); (26, 28, 29); table S1], followed by droplet-based single-cell RNA-seq (scRNA-seq) (28).

We processed the initial snRNA-seq profiles to retain high-quality nuclei profiles and remove the effects of contaminant transcripts from ambient RNA (28). In breast and skin, the majority of nuclei profiles were recovered from only one individual sample for each tissue (breast: 61.3%, skin: 93.1%; table S1). Some tissues and protocols had higher ambient RNA contamination, reflected as spurious expression of highly expressed transcripts from one cell type in nuclei profiles of other cell types. Such effects were more prominent in skeletal muscle and heart [false discovery rate (FDR) < 0.05], irrespective of protocol, but were also present in other tissues [(28); fig. S1]. We corrected for ambient RNA contamination with CellBender v2.1 (30) (fig. S1) and further applied standard quality control metrics (28), retaining 209,126 nuclei profiles across the eight tissues, with a mean of 918 genes and 1519 transcripts (unique molecular identifiers) detected per profile.

Cross-tissue atlas annotation recovers diverse cell types, including difficult-to-profile and rare cell subsets

We integrated data from all samples and methods using a conditional variational autoencoder

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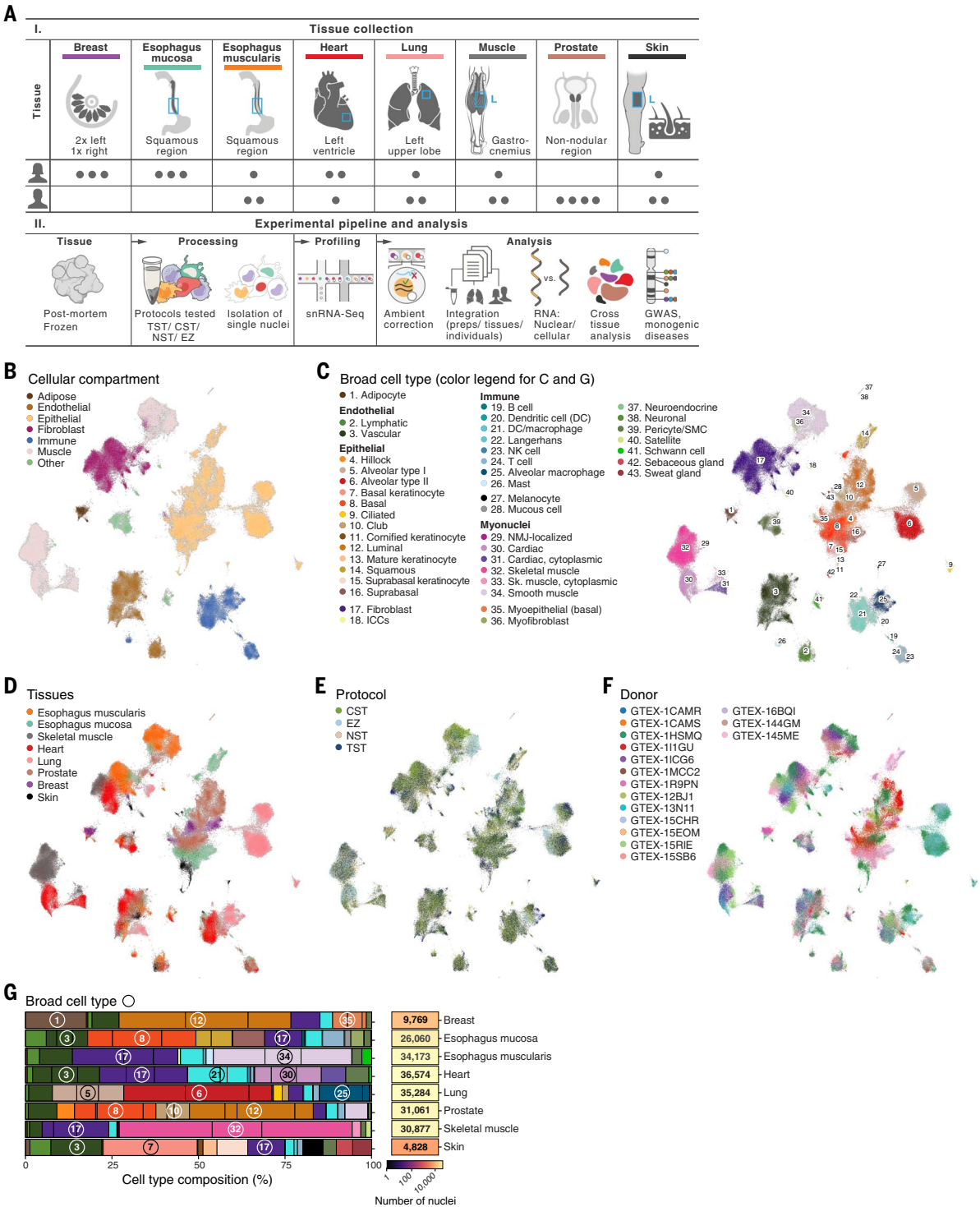


Fig. 1. Cross-tissue snRNA-seq atlas in eight archived, frozen adult human tissues. (A) Study design. **(B to F)** Cross-tissue single-nucleus atlas. Uniform manifold approximation and projection (UMAP) representation of single-nucleus profiles (dots) colored by main compartments (B), broad cell types (C), tissues (D), isolation protocol (E), and individual donors (F). **(G)** Cell-type composition across tissues. The overall proportion of cells (%) of each type and number of nuclei profiled in each tissue (rows) are shown. Numbers in circles indicate the corresponding broad cell type. Black vertical lines indicate the relative proportion of nuclei from each individual.

(cVAE), which is designed to correct for multiple sources of variation in expression, such as individual-, sex-, and protocol-specific effects, while preserving tissue- and cell type-specific variation [(28); Fig. 1, B to F, and figs. S2A and

S3]. We benchmarked the cVAE against several other data integration methods, obtaining comparable or improved results and providing guidelines for future integration efforts [fig. S4 and supplementary text note S1; (28)]. Cells

grouped first by cell type and then by tissue-specific subclusters (Fig. 1, B to D), suggesting that the variation between cell types is larger than the variation within a cell type across tissues.

We annotated cell types within each tissue after dimensionality reduction and graph-based clustering (28) by identifying genes that are differentially expressed between clusters and comparing them with literature-based marker genes [(28); tables S2 and S3]. We curated comprehensive lists of cell-type markers from the literature for each tissue (figs. S5 and S6 and table S3), including markers for relatively poorly characterized cells, such as interstitial cells of Cajal (ICCs). We defined cellular compartments shared across tissues (e.g., adipose, endothelial, epithelial, fibroblast, immune, muscle) (Fig. 1B), broad cell types (e.g., luminal epithelial cells, vascular endothelial cells) (Fig. 1C and fig. S5), and granular cell subsets (e.g., luminal epithelial cell 1 and 2) (fig. S6). The annotations were consistent across extraction protocols, tissues, and donors (figs. S2, B and D, and S7).

The atlas features 43 broad cell classes (Fig. 1C and tables S2 and S3), with both tissue-shared cell types and tissue-specific subsets (e.g., Fig. 1G and figs. S2, B and D, and S5). For example, tissue-specific cell types such as pneumocytes (alveolar type I and II) and keratinocytes were the predominant cell types in the lung and skin, respectively. Many shared broad cell types such as immune and stromal cells were detected across all tissues (fig. S2, D and E), but with tissue-specific specializations (discussed later in the text). For example, macrophages made up the largest immune population, with diverse subsets of tissue-resident cells.

The atlas captured profiles from cell classes that are difficult to profile by dissociation-based scRNA-seq (23, 31, 32), including 2350 adipocytes, 21,607 skeletal muscle myonuclei, and 9619 cardiac myonuclei. We detected adipocytes in five of the eight tissue types (breast, muscle, heart, esophagus muscularis, and skin), with 86% of adipocytes from breast tissue (fig. S2D), making up 18% of all breast nuclei profiles (Fig. 1G) (28). Skeletal and cardiac myonuclei included key subsets (33, 34). Cardiac myonuclei primarily included the previously distinguished classical myonuclei as well as the recently reported “cytoplasmic myonuclei” (33) (fig. S8 and supplementary text note S2). Other myonuclei subsets included neuromuscular junction (NMJ)-localized skeletal muscle myonuclei, which have also been observed in scRNA-seq and snRNA-seq studies in mice (21, 31, 35), and “fast-twitch” and “slow-twitch” subtypes (Fig. 1, B and C, and fig. S6G), which are characterized by differentially expressed genes (fig. S9, A and B) that are concordant with previously reported markers (36) (fig. S9, C and D).

Cross-tissue and cross-sample integration enhanced our ability to resolve multiple rare cell subsets (Fig. 1C and figs. S2, B and D, S5, and S6). For example, we detected Schwann

cells that support peripheral nerves (37) in multiple tissues (esophagus mucosa and muscularis, heart, prostate, and skeletal muscle), rare neuroendocrine cells in the prostate (38), and rare (26) ICCs and enteric neurons in the esophagus. Because these rare cells can contribute to various pathologies, their profiling in human tissues will enable disease studies (26, 37).

snRNA-seq protocols perform well across tissues and correspond to scRNA-seq

We benchmarked the performance of our nucleus extraction and profiling protocols (26, 29) relative to each other across all eight profiled tissues and to other snRNA-seq, scRNA-seq, and bulk RNA-seq datasets in relevant tissues. For each dataset, we compared standard quality control metrics per profiled cell or nucleus, as well as the diversity and proportions of captured cell types.

Of the four tested nucleus isolation protocols (CST, NST, TST, and EZ; table S1), the EZ protocol displayed lower performance in each of the eight profiled tissues by multiple quality metrics (28, 39) (Fig. 2A and fig. S10). These included the lowest total number of nuclei captured (fig. S10, A and B), higher levels of ambient RNA (FDR < 0.05; fig. S1, A and B), and separate grouping of EZ-profiled samples [fig. S2C; (28)].

The extraction protocols also varied in the proportion of nuclei recovered from each cell type (figs. S2, B and D, and S11A; supplementary text note S3), consistent with our previous observations in tumors (29). The TST, CST, and NST protocols had comparable cell-type diversity as measured by Shannon entropy [Fig. 2A; (28)], whereas the EZ protocol resulted in significantly lower diversity (Fig. 2A; linear mixed-effects model effect size of -1.08 , $P = 5 \times 10^{-11}$). Overall, TST yielded the highest cell-type diversity, on average, across tissues (Fig. 2A) and significantly higher proportions of T cells, fibroblasts, and vascular endothelial cells (FDR < 10%; fig. S11A). Because the protocols varied by their performance (most diverse, high capture of the desired cell types), users should choose protocols by matching protocol features to scientific goal, tissue type, and complexity; and further protocol optimization may still be required (29) (for further details and guidance, see supplementary text note S3).

We compared cell-type compositions between our four protocols and other snRNA-seq studies, focusing on our frozen heart left-ventricle samples, for which two recently published snRNA-seq studies evaluated similar samples (33, 34). We found agreement in broad cell types such as mast cells, adipocytes, “cytoplasmic” cardiac myonuclei, and Schwann cells (fig. S11, B and C, and table S4)—but differences in some of their proportions (fig. S11, D and E). Protocols used in the published

studies and the EZ protocol in our study captured a higher proportion of muscle nuclei, whereas CST, NST, and TST yielded a higher proportion of endothelial cells (fig. S11E). There was also high concordance between the expression profiles of bulk RNA-seq [from GTEx; (7)] and pseudobulk profiles derived from our snRNA-seq (accuracy 92.3%; fig. S12). A few samples showed lower agreement (heart-EZ, breast-EZ, and two breast-TST samples), suggesting that these particular tissue-protocol combinations may not reflect cellular composition as accurately.

We next compared snRNA-seq data to fresh-tissue scRNA-seq data from lung (40), skin [current study; (28)], and prostate (38). For cell composition (Fig. 2B), we recovered the same main cell groups across compartments. We confirmed the accuracy of our annotations by training a multiclass random forest classifier on snRNA-seq data and predicting cell types on scRNA-seq data [(28); Fig. 2, C to E], and vice versa (fig. S11, F to H). In addition, cell-type intrinsic (pseudobulk) profiles of protein-coding genes were overall similar between snRNA-seq and scRNA-seq [average Spearman ρ across cell types of 0.58 (skin), 0.69 (prostate), and 0.47 (lung); table S4]. Moreover, integrating cell and nuclei profiles from prostate, skin, and lung and annotating the cells with a random forest classifier that is trained on our nuclei profiles with our granular annotations yielded well-mixed groupings, similar marker genes, and high concordance between protocols (fig. S13).

Divergences observed include the greater expression in cells versus nuclei of a dissociation-induced stress signature (41, 42) (Wilcoxon rank sum test, Benjamini-Hochberg FDR < 10^{-16} ; Fig. 2F and fig. S14, A and B), as reported (29), and of ribosomal and nuclear-encoded mitochondrial protein genes [Fig. 2G; (28); linear model], consistent with their longer half-lives and higher cytoplasmic levels (43, 44). Conversely, nuclei profiles had higher levels of longer transcripts (fig. S14, H and I) and of transcripts with a larger number of adenine stretches (Fig. 2G and fig. S14, C to G), consistent with previous reports (45).

Notably, our snRNA-seq generally captured relatively lower proportions of lymphocytes. For example, for lung and skin, respectively, T cells represented 1.7 and 1.4% of all cells (aggregated) compared with 8.73 and 6.83% by scRNA-seq. We observed similar patterns for B cells in skin. Furthermore, these immune cell proportions varied across samples and protocols. A study comparing snRNA-seq and in situ measurements (46) suggested that scRNA-seq may oversample immune cells.

Myeloid populations across tissues

Our cross-tissue atlas allowed us to characterize tissue-specific and shared features of

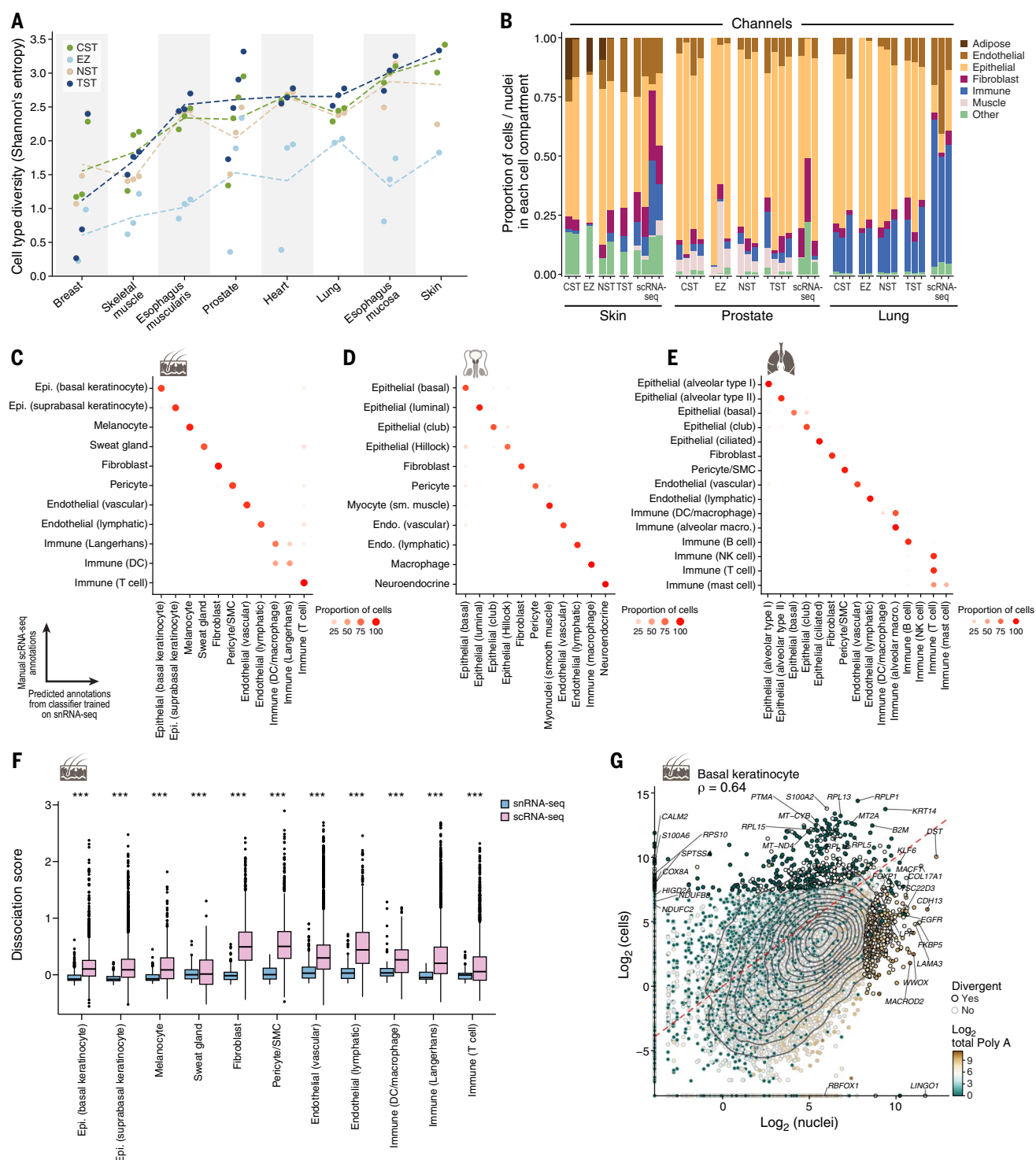


Fig. 2. Concordance of cell-type diversity and cell-intrinsic profiles between snRNA-seq and scRNA-seq. (A) Cell-type diversity (Shannon entropy, y axis) of each protocol (color) in each sample (dot) and tissue (x axis). Dashed lines indicate the average across samples. (B) Differences in cell proportions. The proportions (y axis) of cells from major categories (color) in each individual by tissue and protocol (x axis) are shown. (C to E) Concordance of cell-intrinsic programs. Proportions of cells (dot color and size) of a manually annotated group (rows) predicted to belong to a given nucleus profile annotation label (columns) by a random forest classifier trained on nuclei and applied to cells of the same tissue for skin (C), lung (D) or prostate (E) are shown. (F) Tissue dissociation expression

signatures in scRNA-seq. Scores [y axis, average background corrected log(TP10K+1)] of a dissociation-related stress signature (41) in scRNA-seq (pink) and snRNA-seq (blue) profiles in each major lung cell type (x axis) are shown (***Benjamini-Hochberg FDR < 10⁻¹⁶, Wilcoxon rank sum test). Box plots show median, quartiles, and whiskers at 1.5 times the interquartile range (IQR). (G) Divergent genes between cell and nucleus profiles. Averaged pseudobulk expression (28) of protein-coding genes (dots) in skin basal keratinocyte nuclei (x axis) and cells (y axis) is shown. Divergent genes are represented by a black dot outline. The color scale shows the total length of polyA stretches with at least 20 adenine bases in log₂ scale. Epi., epithelial; sm., smooth; SMC, smooth muscle cell.

tissue-resident immune cells, which play key roles in immune surveillance and tissue support (47, 48). Integration and annotation of 14,156 myeloid nuclei profiles (28) (60% of immune nuclei) revealed 14 distinct monocyte, macrophage, and dendritic cell (DC) subsets (Fig. 3A; fig. S15, A and B; and table S5). These included $CD16^+$ monocytes, $CD14^+$ monocytes, two transitional Mo/MΦ $FCGR3A^{low}$ and Mo/MΦ $FCGR3A^{high}$ populations with co-expression of both monocyte and macrophage markers (see next section), DC1s, DC2s (49), mature DCs, and Langerhans cells (Fig. 3B). Tissue macrophage states further included lung macrophages expressing an alveolar macrophage signature (50) (fig. S15C), proliferating macrophages, cytokine- and chemokine-expressing inflammatory macrophages, and two additional macrophage subsets: MΦ $LYVE1^{high}$ and MΦ $HLAI^{high}$ (Fig. 3B), where $HLAI$ is HLA class II. Finally, lipid-associated macrophage (LAM)-like nuclei highly expressed LAM signature (Fig. 3B and fig. S15C) (51) as well as lipid metabolism-related, myeloid cell immune activation, and macrophage migration genes (fig. S15D). Although most myeloid subsets were present in multiple tissues, notable exceptions included $PPARG^{high}$ lung macrophages, which were present only in lung, and $CD207^+$ Langerin⁺ Langerhans cells, which were present only (97%) in skin and esophagus mucosa, consistent with their role in antigen sampling within stratified epithelia (52, 53) (Fig. 3C and fig. S15, F and I).

Myeloid cell proportions were more highly correlated between samples within a tissue type (fig. S15, E and F) than between different tissues (fig. S15, E to H), confirming the reproducibility of tissue-specific myeloid state proportions. Moreover, related tissues—such as muscle (heart, esophagus muscularis, skeletal muscle) or epithelial barriers (esophagus mucosa, skin)—grouped by their myeloid composition profiles (fig. S15H). Macrophage types and proportions varied by tissue, with breast, esophagus mucosa, esophagus muscularis, heart, and skeletal muscle having significantly higher proportions of MΦ $LYVE1^{high}$ macrophages [$P < 10^{-6}$, Dirichlet regression likelihood ratio test (LRT); (28)] and lung and prostate having significantly higher proportions of MΦ $HLAI^{high}$ [$P < 10^{-8}$, Dirichlet regression LRT; (28)] (Fig. 3C and fig. S15, F and I).

A dichotomy between $LYVE1$ - and $HLAI$ -expressing macrophages is preserved across tissues

Two expression states of $LYVE1^{high}$ and $HLAI^{high}$ macrophage populations were dichotomous—either $LYVE1^{high}HLAI^{low}$ or $LYVE1^{low}HLAI^{high}$ —and represented the end points of two alternative branches. Specifically, low-dimensional representation of monocytes, macrophages, and Mo/MΦ populations as a continuum with

diffusion maps captures the $HLAI^{high}$ and $LYVE1^{high}$ cells as “terminal” points in two branches that emanate from $CD14^+$ monocytes at the root. Each of the two terminals is preceded by distinct earlier putative transitional states: a Mo/MΦ $FCGR3A^{low}$ state between $CD14^+$ monocytes and the $LYVE1^{high}HLAI^{low}$ population and a Mo/MΦ $FCGR3A^{high}$ state between the monocytes and the $LYVE1^{low}HLAI^{high}$ cells (Fig. 3D and fig. S16, A and B). (There is also a putative secondary path between $FCGR3A^{low}$ and $LYVE1^{low}HLAI^{high}$ cells through a $FCGR3A^{high}$ intermediate.) The position of the $FCGR3A^{high}$ transitional state is consistent with lung data from a humanized mouse model (54, 55). These key features are consistent overall in the map that is constructed only from macrophage nuclei from a single tissue (fig. S16A). Thus, $FCGR3A^{low}$ and $FCGR3A^{high}$ states might be Mo/MΦ populations that are less-differentiated or less-activated states of $LYVE1^{high}$ and $HLAI^{high}$ MΦs, respectively.

Each of the $LYVE1^{high}$ and $HLAI^{high}$ subsets expressed a combination of a common signature and tissue-specific markers (Fig. 3E, fig. S16C, and table S5) and was enriched for distinct functions, mirroring those of $Lyve1^{high}$ MHCII^{low} and $Lyve1^{low}$ MHCII^{high} resident macrophage populations in mouse tissues (47) (fig. S16, C and E). $HLAI^{high}$ cells were enriched for immune-related processes and expressed higher levels of complement components $APOE$ and $C1QA$, $C1QB$, and $C1QC$ (Fig. 3B and fig. S16, C and D). Genes differentially expressed between human $HLAI^{high}$ and $LYVE1^{high}$ subsets corresponded to those in murine counterparts (fig. S16E), but with higher expression in human $HLAI^{high}$ macrophages of $C1QB$ and $C1QC$ complement genes in lung and phagocytic receptors $MARCO$ and $CD36$ in heart (fig. S16E). $HLAI^{high}$ macrophages were also enriched for immune interactions with B cells, DCs, mast cells, natural killer (NK) cells, and T cells (fig. S16G). $LYVE1^{high}$ profiles were enriched for tissue-supporting modules and had putative receptor-ligand interactions (28) with lymphatic endothelial cells, fibroblasts, adipocytes, and myocytes (fig. S16F). In mice, MΦ $Lyve1^{high}$ cells were located near blood vessels (47) and regulated vascular tone (56), and cardiac $LYVE1^+$ macrophages have been implicated in regulating the lymphatic network (57). Thus, $LYVE1^{high}$ macrophages may have a homeostatic role in the human heart, lung, and esophagus.

LAM-like macrophages are prevalent across human tissues and share a regulatory program

In our atlas, we identified LAM-like cells as widely distributed across healthy human tissues, with the vast majority of LAMs (97%, 268 of 283) from the breast, heart, lung, and prostate (Fig. 3C). LAMs and LAM-like cells

have been previously reported in disease contexts in adipose tissue from obese humans and mice (51), injured and fibrotic liver (58–60), obese liver (61), fibrotic lung (50, 62), atherosclerotic aortic tissue (63, 64), leprosy (65), and the brains of individuals with Alzheimer's disease (66–68). However, an understanding of their distribution and heterogeneity across human tissues is still lacking.

To characterize LAMs across the body, we analyzed LAM-like cells in the expanded context of our study and 17 other published atlases spanning 14 tissues. We trained a linear classifier with published omental adipose scRNA-seq containing LAMs (28, 51) and classified each myeloid profile in our dataset and the published compendium as LAM-like macrophages, non-LAM macrophages, and non-macrophages. From this, we recovered 283 LAM-like cells in our study and 4285 LAM-like cells in the 17 published studies (Fig. 3, F to H; fig. S17, A to D; and table S6).

LAM-like cells were present among tissue-resident macrophages across a broad range of tissues and pathologies. These included adipose (51, 69) and atherosclerotic (70, 71) tissue, as reported, as well as healthy tissues [placenta (72), testis (73), kidney (74), pancreas (75), prostate (38), decidua (72), liver (76), ovary (76, 77), skeletal muscle (78), and intestine (26)] and other disease contexts [acne (79), leprosy (79) and atopic dermatitis (80, 81) (skin), and Crohn's disease (ileum) (82)]. Microglia from the central nervous system of epileptic patients (83) were also classified as LAM-like cells, indicating that microglia that express LAM signature genes extend beyond Alzheimer's disease (66–68). LAM signature genes were enriched for genome-wide association study (GWAS) genes associated with levels of high-density lipoprotein (HDL) and low-density lipoprotein (LDL) cholesterol, triglycerides, type 2 diabetes, and the tau to Aβ1-42 ratio in cerebrospinal fluid (fig. S17H and table S7), further supporting their role in lipid homeostasis.

Although a core set of signature LAM genes was expressed across most tissues and studies, many other genes varied across tissues. For example, both $CHIT1$ and $CTSK$ were highly expressed in LAMs from leprosy skin samples (but had very low expression in other skin LAMs) (Fig. 3H and fig. S17F), in line with higher serum chitotriosidase activity in leprosy patients (84). Select lipid pathway genes, including fatty-acid binding protein $FABP4$, lipoprotein lipase LPL , and phagocytic lipid receptor $CD36$, were highly expressed in LAMs from tissues with high adipose content, including adipose, atherosclerotic lesions, and intestine creeping fat samples, possibly reflecting increased lipid-induced transcriptional stimulation of these target genes under these conditions (85). LAMs in creeping fat from

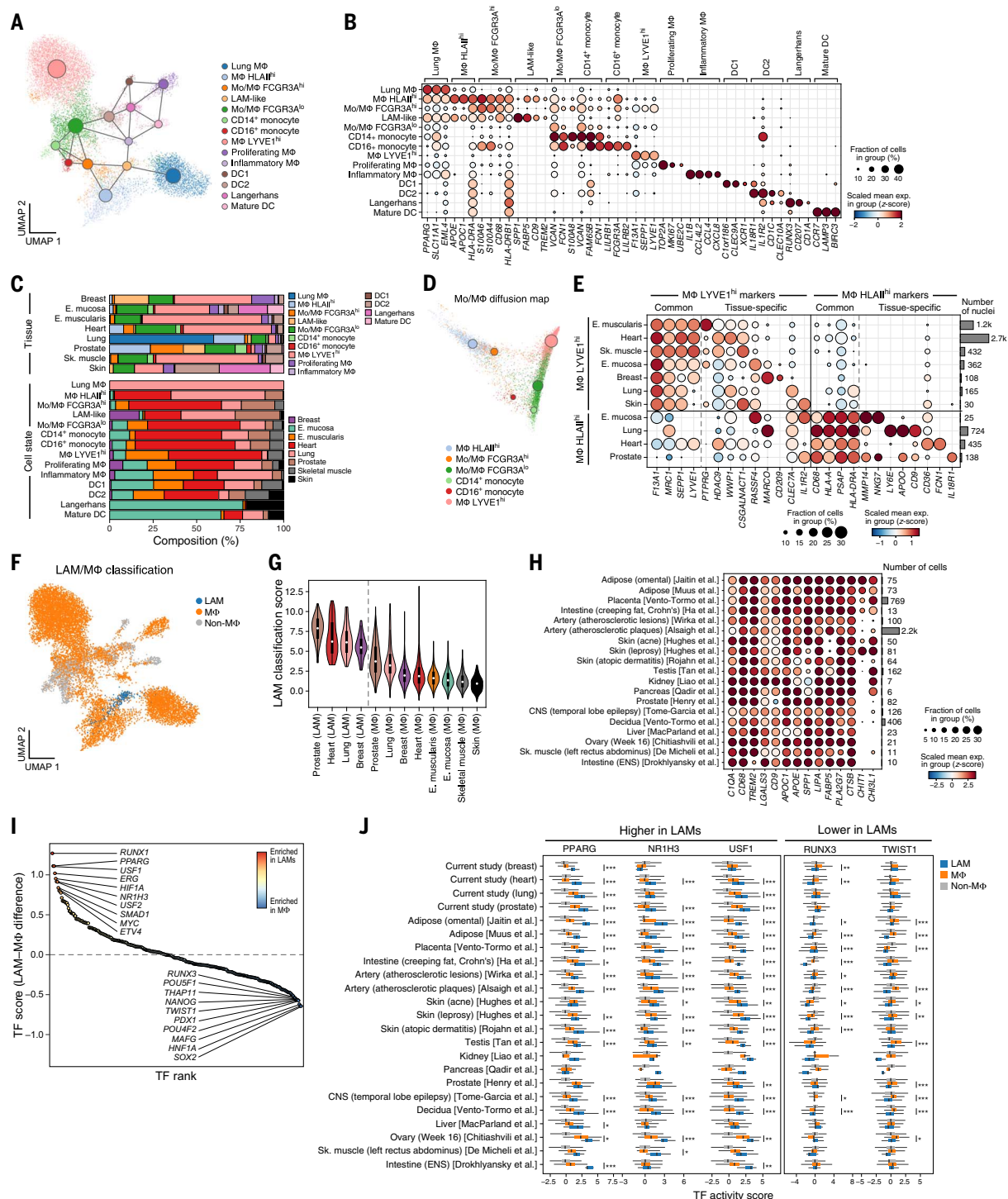


Fig. 3. A dichotomy between LYVE1- and HLAII-expressing macrophages and LAM-like populations across tissues. (A) Myeloid profiles (dots), colored by cell type and state and overlaid with a PAGA graph of myeloid states (large nodes). (B) Expression of marker genes (columns) associated with each subset (rows). (C) Myeloid cell distribution across tissues. The overall proportion of myeloid cell subsets (colors) in each tissue (bars) is shown at the top, and the overall proportion of cells from each tissue in each subset (bars) is shown at the bottom. (D) LYVE1^{high} and HLAII^{high} macrophages are end points of two differentiation trajectories. A diffusion map of monocytes, macrophages, and transitional subsets (colors) is shown. Large circles represent centroids (sizes are proportional to population size). (E) Cross-tissue and tissue-specific markers. Expression of marker genes (columns) associated with two myeloid subsets (left) in each tissue

(rows) is shown. The right bar plot shows the number of nuclei. (F to H) LAM-like cells across tissues. Myeloid cells (dots) colored by their classification [legend; (28)] are shown in (F). Classification scores (y axis) of LAM-like and other macrophages across tissues (x axis) are shown in (G). Expression of LAM marker genes (columns) in LAM-like profiles from other studies (rows) is shown in (H). (I and J) Inferred TFs regulating the LAM-like program. TF differential activity scores between LAMs and other macrophages (y axis) for each TF (dot) ranked by score (x axis) are shown in (I). TF differential activity scores (x axis) for three TFs with significantly high scores (two tailed *t* test; Benjamini-Hochberg *FDR < 0.05, **FDR < 0.01, ***FDR < 0.001) in LAMs or other macrophages are shown in (J). Box plots show median, quartiles, and whiskers at 1.5 times the IQR. E., esophagus; ENS, enteric nervous system; Sk., skeletal.

Crohn's intestine and atherosclerotic lesions were additionally characterized by higher expression of interleukin *IL1B* and multiple chemokines (*CXCL8*, *CXCL3*, *CCL4*), possibly reflecting the inflammatory environment under these conditions (fig. S17F).

We predicted transcription factors (TFs) that could mediate LAM-like gene expression by inferring TF activity from target expression (28) and ranking TFs by the mean difference between their activities in LAMs versus non-LAM macrophages [(28); Fig. 3I]. LAM-associated TFs inferred across all classified LAM-like cells included *PPARG*, *USF1*, and *NR1H3* (*LXR4*) (Fig. 3J and fig. S17G), suggesting a shared core regulatory mechanism. These are major regulators of lipid metabolism-related expression (86) and have been proposed to regulate *Trem2* expression in mice (87).

Shared and tissue-specific features of fibroblasts

To characterize fibroblast heterogeneity (88), we analyzed 32,421 fibroblast nuclei profiles across the eight profiled tissues (fig. S18A), identifying shared and tissue-specific signatures. The cross-tissue, shared fibroblast expression program consisted of markers that were significantly more highly expressed in fibroblasts than in nonfibroblast cell types within each tissue (FDR < 0.05, Welch's *t* test) and included multiple extracellular matrix (ECM) constituents (Fig. 4A and table S8). Conversely, the tissue-enriched fibroblast signatures were defined based on genes exclusive to or highly enriched in fibroblasts from a given tissue versus fibroblasts from all other tissues (Fig. 4, B to D, and fig. S18B).

Tissue-enriched fibroblast features were consistent with the specific functions and interactions required in the respective tissues. For example, the esophagus mucosa fibroblast signature (table S8) included genes involved in neuron and axon development (e.g., *NTN1*, *PLXNB1*, *FGF13*), suggesting interactions with the enteric nervous system, possibly through NTN1-DCC and NTN1-UNC5C interactions (fig. S18E). The cardiac fibroblast signature genes included TFs involved in cardiac development (e.g., *GATA4* and *GATA6*), revealing that expression of these developmental TFs is retained in the adult cardiac fibroblast compartment (89–92). The skeletal muscle signature showed increased expression of the *CXCL14* and *CXCL12* chemokines and of components of the renin-angiotensin-aldosterone system (*AGTR1* and *MME*), which regulate skeletal muscle mass (93, 94) [reminiscent of the expression of the same components by WNT2B⁺ fibroblasts in a local renin-angiotensin system in the colon (16)].

Lung fibroblast signatures were enriched for ECM, cation transport, and contractile functions (Fig. 4E), including multiple components

of the basement membrane (BM) that are required for epithelial-mesenchymal adhesion [nephronectin (*NPNT*), *FRAS1*, hemiscitin-1 (*HMCN1*), and integrin *ITGA8* (Fig. 4, F and G), which form a protein complex (95–98) that anchors NPNT to the BM (99–103)]. Mutations in *ITGA8*, *FRAS1*, and *HMCN1* have been linked to Fraser syndrome, a congenital disorder that affects cell adhesion and results in skin, kidney, lung, and craniofacial abnormalities (104–107), and common variants in *NPNT* are significantly associated with chronic obstructive pulmonary disease (COPD) and forced expiratory volume (FEV) in GWASs (Fig. 4H). Granular annotation of the lung fibroblast profiles (40, 108) (fig. S18C) suggests that the adhesion complex is expressed by alveolar fibroblasts (Fig. 4F and fig. S18D). Alveolar fibroblasts also specifically expressed *FGFR4* (Fig. 4B), which is essential for alveologenesis in mice (109) and is genetically linked with bronchopulmonary dysplasia that affects alveoli in humans (110).

Of all tissue fibroblasts, lung alveolar fibroblasts distinctively expressed a calcium signaling and actomyosin contractility program (Fig. 4, E and F, and fig. S18D), including the mechanosensitive calcium ion channel *PIEZO2* (Fig. 4G), which has been proposed to sense pulmonary stretch (111, 112); multiple calcium channels and adrenergic and purinergic receptors, which have been implicated in stimulating calcium release from intracellular stores (*CACNA1D*, *TRPC6*, *MCOLN2/TRPML2*); and myosin light chain kinase (*MYLK*), which is involved in mediating calcium-induced actomyosin contraction (113). This suggests that alveolar fibroblasts could integrate mechanical stretch and forces (through *PIEZO2*) and neuronal excitatory signals (40), which could affect their migratory or mechanical properties. Notably, although alveolar fibroblasts express myofibroblast markers (e.g., *ACTA2*; fig. S18D), they are distinct from previously reported myofibroblasts (40), and our data suggest that they constitute a distinctive contractile and excitable fibroblast state.

Intra- and cross-tissue cell-type associations with monogenic disorders

Human genetics has identified numerous rare monogenic disease genes, and many have been experimentally mapped to cell type(s) of action (114, 115). To characterize the expression of monogenic disease genes across cell types, we related disease genes associated with different phenotypes or disease categories in the Online Mendelian Inheritance in Man (OMIM) database (116) to the cell populations in which they are expressed in our cross-tissue atlas. Because OMIM entries are not organized by disease categories, we leveraged topic modeling to aggregate 5812 genotype-phenotype associations based on similarities in text

descriptions of clinical features, resulting in 229 distinct disease topics [(28); figs. S19 and S20]. We then related topics to a cell type based on the enriched expression of the topic genes in the cell type (Fig. 5A, figs. S21 and S22, and table S9).

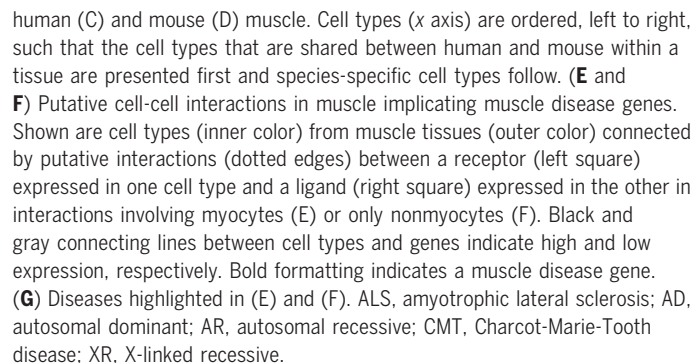
Many topics mapped to their expected cell populations. For example, cardiac disease topics (topics 65 and 66) mapped to cardiac myonuclei and/or endothelial cells, immune and infection topics (topics 132, 205, and 217) mapped to immune cells across tissues, and a diabetes and lipodystrophy topic (topic 222) mapped to adipocytes in skeletal muscle and skin (Fig. 5A and fig. S23A). Male infertility and sperm motility topic (topic 129) was associated with ciliated lung cells and stromal cells (fig. S23B). The link between sperm dysmotility and bronchitis is well established, with genes common to flagella and cilia perturbed across both sperm and lung (117); we further detect this specific link in topic 155, which associates ciliary dyskinesias and the lung ciliated epithelium (fig. S23C).

Genes from monogenic muscle disease groups are enriched in distinct subsets of myocyte and nonmyocyte nuclei in three muscle types

Among the monogenic disorders, muscle disease phenotypes are a well characterized subset and are known to arise from mutations in genes that are expressed in myocytes (e.g., structural genes involved in contraction) and/or other cells in the surrounding tissue (e.g., NMJ, ECM, and adipose tissue) (118–121). We leveraged the three muscle types represented in our atlas—cardiac, skeletal, and smooth muscle—to map 605 well-curated monogenic muscle disease genes (118) (table S10), recovering known biology and extending hypotheses beyond those obtained from bulk-tissue RNA-seq (122, 123). We tested disease groups (e.g., hereditary cardiomyopathies, motor neuron diseases) for their enrichment with cell type-specific markers across our muscle tissues (FDR < 0.1) [(28); Fig. 5B, fig. S24, and table S11].

As expected, different disease gene sets were associated with different myonuclei subsets (113 of 605 genes; table S11) in patterns that recapitulated known disease mechanisms (Fig. 5B). For example, skeletal muscle myonuclei were associated with congenital myopathy genes (FDR = 7.07×10^{-5}), and cardiac myonuclei were associated with hereditary cardiomyopathy genes (FDR = 4.24×10^{-12}). Some associations highlighted finer myonuclei subsets. For example, genes linked to congenital myasthenic syndrome, a disorder affecting neuromuscular transmission (124), were specifically expressed in NMJ-localized myonuclei but not in other skeletal myonuclei (tables S2 and S12). These included acetylcholine receptor subunits (*CHRNA1*, *CHRNA2*, *CHRND*), the NMJ-organizing receptor tyrosine kinase *MUSK*,





hypertrophic neuropathy (a subtype of Charcot-Marie-Tooth disease) was specific to skeletal muscle ($\text{FDR} = 2.28 \times 10^{-5}$), consistent with the need to maintain innervation to prevent muscle atrophy (125). Thus, distinctive cell expression patterns can provide insight into onset (Dejerine-Sottas is an early-onset disease), severity, and the predominant peripheral nerves that are affected (i.e., motor, sensory, or autonomic), where heart can possibly act as a proxy for autonomic nerves. In other examples, adipocytes from esophagus muscularis were associated with metabolic myopathies related to lipid metabolism ($\text{FDR} = 0.0003$); the congenital muscular dystrophies genes *COL6A1*, *COL6A2*, and *COL6A3* are expressed in fibroblasts, consistent with joint laxity and progressive contractures along with muscle weakness (126); and the recessive limb girdle muscular dystrophies gene *DYSF* is expressed in immune cells, consistent with the role of more aggressive monocytes in disease progression (127). Some of the enrichments in non-myocytes are also present in the same cell types in other nonmuscle tissues (e.g., breast adipocytes for metabolic myopathies or breast and skin pericytes for cardiomyopathies; fig. S25I), highlighting that tissue-specific pathology may arise from the relation between an accessory cell's broader function and specialized tissue demand.

There were differences in the expression of monogenic muscle disease genes between slow and fast myocyte subsets for both classical and cytoplasmic myonuclei (fig. S9). As expected, slow myocytes preferentially expressed type 1 fiber markers (36, 128) and disease genes *MYH7* ($\text{FDR} < 10^{-15}$ in cytoplasmic myonuclei) and *PPARGC1A* ($\text{FDR} < 10^{-15}$ in regular myonuclei). Fast myocytes specifically expressed the type 2 fiber marker *MYH2* ($\text{FDR} < 10^{-15}$ in cytoplasmic myonuclei), in line with respective clinical myosinopathy phenotypes (129). Congenital myopathy disease genes, which include genes known to be active in slow and fast myocytes, were enriched in both groups. The myotonic dystrophy (MD) type 2 gene *CNBP*, was enriched in regular myonuclei from fast myocytes [$\text{FDR} < 10^{-12}$; (28); fig. S9, E and F], which are preferentially affected in MD type 2 (130, 131). By contrast, MD type 1 disease gene *DMPK* was preferentially expressed in cytoplasmic myonuclei from slow myocytes ($P < 0.036$, Welch's *t* test), which are perturbed in MD type 1 (132). Regular myonuclei subsets from slow myocytes were also enriched in *TPM3* [$\text{FDR} < 10^{-15}$; (28); fig. S9, E and F], which has been linked to congenital fiber type disproportion, a condition characterized by smaller slow myocyte fibers (133). Metabolic myopathy disease genes (*GBE1*, *ETFDH*, and *SLC25A20*) were also enriched in slow myocyte markers ($P < 0.05$, Fisher's exact test; fig. S9, E and F).

Many of the cell-type associations for monogenic muscle disease genes were conserved between our human muscle atlas and corresponding mouse snRNA-seq data (26, 28) (fig. S25), but there were also notable differences. In both mouse and human, there were significant associations between skeletal muscle myonuclei and various dystrophies and myopathies ($\text{FDR} < 0.1$, Fisher's exact test), between cardiac myonuclei and cardiomyopathies ($\text{FDR} < 0.1$), between adipocytes (in skeletal muscle, esophagus, and heart) and metabolic myopathies ($\text{FDR} < 0.1$), and between Schwann cells in skeletal muscle and hereditary motor and sensory neuropathies ($\text{FDR} < 0.1$) (fig. S25H). However, dystrophin (*DMD*) expression across accessory cell types varied between human and mouse. In humans, high levels of *DMD* expression (comparable to that in myonuclei) were observed in adipocytes in all muscle types; pericytes and Schwann cells in skeletal muscle and esophagus muscularis; and enteric neurons in esophagus muscularis [mean expression $\log(\text{TP10K}+1) > 2.0$; Fig. 5C]. Lower, intermediate levels of *DMD* expression were observed in skeletal muscle fibroblasts and satellite cells, as well as esophagus muscularis ICCs [mean expression $\log(\text{TP10K}+1) < 2.0$; Fig. 5C]. *DMD* expression in adipocytes supports the possibility of local metabolic perturbation (134) (Fig. 5C), whereas its expression in the enteric nervous system and ICCs raises the possibility that perturbation of these cells contributes to the gastrointestinal dysfunction phenotype in Duchenne muscular dystrophy (135). In mouse, whereas *Dmd* expression is high in myonuclei, pericytes, satellite cells, and Schwann cells, it is low in adipocytes and fibroblasts across all three muscle tissue types [mean expression $\log(\text{TP10K}+1) < 1.0$; Fig. 5D]. These differences suggest that the effects of *DMD* mutation on accessory cells and their contribution to disease (135, 136) may differ between human patients and mouse disease models (137).

Cell-composition and cell-intrinsic secondary effects in muscle disease tissue

Muscle dystrophies commonly display a secondary shift in cellular composition, with fibrotic or adipogenic replacement of muscle tissue (138), which can introduce secondary pathologic processes that exacerbate muscle loss (139). To characterize the cellular expression patterns of disease genes whose bulk-tissue expression levels are altered in muscle diseases, we analyzed the cell-type specificity of genes up-regulated in bulk RNA-seq data from muscle tissues from 43 patients with rare muscle disorders (122) compared with healthy muscle tissues (140) (fig. S26). Some of the cell type-specific expression patterns reflected known biology, including up-regulation of hallmark fibroblast, adipocyte, and immune genes, likely because of changes in cell pro-

portions (e.g., *COL6A2*, *ADIPOQ*, and *HLA-A*, respectively). Other patterns suggest additional, cell-intrinsic regulatory events (beyond cell composition changes) that may modify disease progression. For example, dermatopontin (*DPT*), an adipokine that promotes ECM remodeling and inflammation (141), is expressed in adipocytes but is not correlated with *ADIPOQ* expression across adipocytes from different tissues. This suggests that its expression is regulated and is not merely reflecting changes in cell composition. Similarly, *ELK3*, which is expressed in endothelial cells, suppresses angiogenesis (142) and may contribute to functional muscle ischemia in muscle disease (143), which is otherwise typically attributed to nitric oxide signaling (136). Thus, the increased resolution of a single-cell atlas can help disentangle secondary effects related to cell-composition and cell-regulatory events in accessory cells during disease.

Disease genes may affect receptor-ligand interactions

Some disease genes encode receptors or ligands that participate in cell-cell interactions, such that loss-of-function mutations can affect tissue function through non-cell autonomous effects through these cell-cell interactions. We related cell types in muscle tissue to one another through receptor-ligand interactions (16, 26, 144) that included at least one monogenic disease gene (28) in every tissue in our atlas (table S13).

Our analysis suggests that mutations in some disease-causing genes may disrupt interactions between myocytes and other cell types. For example, mutations in *ERBB3* (the disease gene for lethal congenital contracture syndrome) may disrupt interactions between myocytes and Schwann cells (Fig. 5E and table S13) and contribute to joint contractures as an associated, but not primary, disease phenotype (145). Mutations in *DAG1* (the disease gene for congenital muscular dystrophy), although known to interact primarily with laminin produced by fibroblasts (146, 147), may additionally disrupt interactions with immune cells through *LGALS9* and alter their function (148). Putative cell-cell interactions involving only non-myocytes included the disease genes *LICAM* (MASA syndrome), *MET* (arthrogryposis and muscular dysplasia), and *NGF* (hereditary sensory and autonomic neuropathy), each potentially affecting multiple cell pairs, including neurons and Schwann, satellite, immune, and stromal cells (Fig. 5, F and G, and table S13).

Cell type-specific enrichment of QTL genes mapped to GWAS loci

Single-cell atlases can also provide insights into cell-type specificity and mechanisms of action of the genes in disease-associated loci identified by GWASs. Studies associating genetic variants to changes in gene expression or splicing

quantitative trait loci (eQTL or sQTL, respectively) showed tissue-specific colocalization with multiple loci from GWASs of human traits, including disease risk (7, 149–151), but lacked cellular resolution (17). To prioritize cell types of action and causal genes for complex diseases and traits in specific cells and tissues, we used ECLIPSER (28, 152) to test whether GWAS loci from 21 complex traits (table S14), with likely effects in at least one of the eight tissues analyzed, are enriched for genes with high cell type-specific expression in each tissue. We defined putative causal genes for each GWAS locus as the set of genes whose eQTLs and sQTLs (7) were in linkage disequilibrium [squared genotype correlation (r^2) > 0.8] with the lead GWAS variant(s) [Fig. 6A; (28)]. We further included genes prioritized by additional genomic data [e.g., Hi-C and protein QTLs (pQTLs)] and linkage to predicted deleterious protein-coding variants (153, 154). Because more than one gene typically maps to a GWAS locus using this approach (mean = 2, and maximum = 37 for selected traits and 170 for null traits), we scored loci by the fraction of cell type-specific genes in the locus [Fig. 6A; (28)]. We assessed enrichment for each GWAS locus set against a null distribution of GWAS loci associated with tissue-unrelated traits using a Fisher's exact test [Fig. 6A; (28)].

Seventeen of the traits were enriched in both expected and previously undescribed cell types at a tissue-wide FDR less than 0.05 (Benjamini-Hochberg), 16 of which were significant (FDR < 0.05) across tissues (Fig. 6B, figs. S27 and S28, and tables S15 and S16). Among the expected associations are skin pigmentation traits in melanocytes, autoimmune and inflammatory diseases in T and NK cells, COPD in lung fibroblasts, prostate cancer in luminal epithelial cells, atrial fibrillation and heart rate in myonuclei, and heart rate in lymphatic endothelial cells (155) (Fig. 6B). Type 2 diabetes loci were enriched in skeletal muscle adipocytes and in lymphatic endothelial cells in multiple tissues, which might contribute to the predisposition of type 2 diabetes to vascular disease (156, 157) (Fig. 6B). Less well-characterized cell type-trait associations included DCs (in almost all tissues) with non-melanoma skin cancer (158) and adipocytes (breast) with atrial fibrillation (table S15). GWAS loci enriched in a specific cell type from a known tissue of action frequently showed similar enrichment in the same cell type from other uninvolved tissues. For example, atrial fibrillation GWAS loci were enriched in myonuclei in heart, skeletal muscle, esophagus muscularis, and prostate (Fig. 6, B and C); coronary artery disease and heart rate loci were enriched in pericytes in five or six tissues in addition to heart; and prostate cancer loci were enriched in luminal epithelial cells in both prostate and breast (Fig. 6B and figs. S29 and S30).

Cell-type enrichment helped identify putative causal genes in GWAS loci with multiple QTL-mapped genes (tables S15 and S16). On average, about two-thirds of genes driving the cell type-specific enrichment for a given trait in a relevant tissue [mean = 66%, 60 to 71.4% (95% confidence interval)] were also driving the enrichment in the same cell type in other tissues. For example, in the case of atrial fibrillation and cardiac myonuclei, 26 out of 31 myonuclei-specific genes that drove the enrichment signal in myonuclei in heart and were shared with at least one other tissue (Fig. 6D) were enriched in muscle system-related processes, such as muscle contraction (FDR < 0.05; table S17). The five myonuclei-specific genes that were specific to heart only (pink vertical lines in Fig. 6D) were instead enriched in heart development processes, such as cardiac muscle tissue development (FDR < 0.05; table S18). Cardiac myonuclei were also found to be the most relevant cell type for atrial fibrillation in two separate snRNA-seq studies of the human heart (33, 34) and based on ECLIPSER analysis of these two studies [fig. S31 and table S19; (28)]. *CASQ2*, which encodes a cardiac muscle member of the calsequestrin family, and the myosin heavy chain 6 and 7 genes (*MYH6* and *MYH7*) were the top myonuclei-specific genes driving the enrichment signal for atrial fibrillation in all three studies (table S19).

Associating GWAS genes with gene programs across cell types reveals six main trait groups

To chart cellular programs and processes that may be affected by genetic variants, we associated a larger set of >2000 complex phenotypes with cell types and the covarying gene modules that these cell types express (28). We defined gene modules in the cells in our atlas by hierarchically clustering genes based on correlation across all cells, as well as within cell types. We then scored modules for their overlap with GWAS genes [defined by variant to gene mapping in Open Targets Genetics (153, 154)] that are also highly expressed by cell type [(28); fig. S32 and table S20]. Next, we grouped GWAS phenotypes into major groups by the similarity of their module enrichment across cell types (Fig. 7, A and B, and fig. S33). Finally, for each major group of traits, we identified the relevant cell types associated with the underlying modules (Fig. 7, A and C) and tested the GWAS genes that overlapped with gene modules for functional enrichments (Fig. 7, A, D, and E).

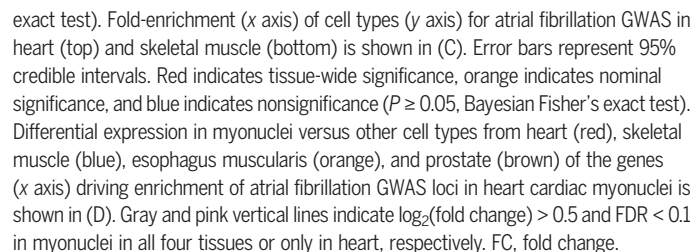
Traits and diseases partitioned into six major groups, spanning immune hypersensitivity, cardiovascular, calcium channel-related, cognitive and psychiatric, pigmentation, and HDL cholesterol-related based on their associations with cell types (Fig. 7B). The immune hypersensitivity disorders were associated with T cells, including the expected relation between lung T cells and hay fever, allergic rhinitis,

and respiratory disease [(159, 160); Fig. 7C]. GWAS genes in the modules associated with these traits were enriched for lymphocyte activation and differentiation and T cell receptor signaling and cell-cell adhesion (Fig. 7D), with interleukin-35 (IL-35) signaling genes enriched in hypothyroidism and inflammatory bowel disease (IBD), consistent with IL-35 up-regulation in Hashimoto's thyroiditis (161) and IBD (162) and down-regulation in Graves' disease (163). The cardiovascular traits group was associated with pericytes and smooth muscle cells and enriched with blood circulation, smooth muscle contraction, muscle structure development, and cardiocyte differentiation genes. The cardiovascular group overlapped with the calcium channel-related group—which included blood pressure medication, pulse rate, medication use of calcium-channel blockers, and vascular system traits, as well as schizophrenia and autism spectrum disorder, which are psychiatric disorders with known calcium channel associations (164)—and was enriched with membrane depolarization and calcium ion channel genes (Fig. 7D). Other cognitive and psychiatric phenotypes grouped separately and were enriched with neuronal synapse organization, structure, and activity genes (Fig. 7D). Finally, a group of HDL cholesterol traits was associated with adipocytes in all three muscle tissues and enriched for fatty acid, triglyceride, and lipid homeostasis and related metabolic processes (monocarboxylic and glycerol metabolism), including *ANGPTL8* and *PNPLA3* (Fig. 7E and table S20), which are genes involved in lipolysis regulation in adipocytes that might affect extrahepatic cholesterol transport via HDLs (165, 166).

Toward large-scale snRNA-seq of human tissues with pooling

To enable future studies at the population scale, we tested whether frozen samples from different individuals can be pooled for snRNA-seq, followed by computational demultiplexing, as a cost-effective and scalable approach (167), as previously applied to large-scale studies of human peripheral blood mononuclear cells (24). We processed lung or prostate samples jointly from three individuals using the CST and TST protocols. We pooled tissue samples from three individuals and processed the pool for single-nucleus extraction, thus minimizing technical batch effects and wet-lab time. After sequencing, we removed ambient RNA (30) and performed de novo genotype-based demultiplexing to assign nuclei to donors [using *souporcell* (168); fig. S34; (28)]. We validated the demultiplexing by comparing the genotype-based assignments to those from an expression-based multinomial logistic classifier that assigned donor identity to each nucleus profile after training with unpooled samples of the same donors, which showed high concordance

B



through lipid-bound receptors on macrophages, such as *TREM2* or *CD36*, up-regulates the expression of more lipid receptors, as well as of lipid-modifying enzymes through *PPARG* and *NRIH3*. Because we observed LAMs in healthy organs and in conditions linked to lipid accumulation, future studies may identify other tissue-specific signals or conditions that can trigger LAM-like states.

In lung alveolar fibroblasts, we identified an interconnected functional module that may enable an adequate response to alveolar distortion, through transduction of mechanical cues from the ECM to cytosolic calcium-induced cytoskeletal contraction (fig. S35, D and E). These include BM adhesion genes, including *ITGA8* and its interacting partner *NPNT* (95–100); a calcium transport module featuring neurotransmitter receptors and multiple calcium ion channels, including the mechanosensitive channel *PIEZO2*; and an actomyosin contractility program that features *MYLK* and mediates cytosolic calcium-induced cytoskeletal contraction, which is required for the transduction of mechanical cues from the ECM (169). *PIEZO2* and *ITGA8* are also coexpressed in intraglomerular mesangial cells (170, 171), where *PIEZO2* may sense mechanical forces resulting from changes in blood flow and *ITGA8* confers contractility and adhesion (172). Because the alveolus and glomerulus are stretch sensors of air pressure and blood pressure, respectively, *PIEZO2*⁺/*ITGA8*⁺ alveolar fibroblasts may contribute to mechanosensing of alveolar tension, which so far has been mainly attributed to *Piezo2*⁺ sensory neurons in mice (173), suggesting conserved features in mechanosensing across organs. Both lung alveolar fibroblasts and myofibroblasts have been localized to alveoli (40), suggesting a possible relationship, possibly through *FGFR4* in alveolar fibroblasts and its ligand *FGF18* in myofibroblasts (fig. S18D) (109, 173).

We further demonstrated the utility of our tissue atlas for monogenic and polygenic disease biology. Many monogenic disease gene modules that are defined by comorbidity [e.g., diabetes and lipodystrophy (174)] or similarity of clinical phenotypes (e.g., muscle diseases) were enriched in expected cell types (fig. S35F). Focusing on the pathobiology of monogenic muscle diseases (fig. S35G), we highlighted nonmyocyte cell populations with a potential role in muscle diseases, including nervous system, immune, and stromal cells (118), as well as specific myonuclei subsets that express muscle disease genes. Whether the multiple myonuclei subsets we observed are related to multinucleation and specialization of different nuclei in one syncytium (31) remains unclear. Some disease-risk genes may also disrupt cell-cell interactions in the muscle. Notably, we observed varying levels of *DMD* expression across cell types, as well as between human

and mouse. Variation in *DMD* isoform expression, which has critical implications in Duchenne muscular dystrophy (135), can be investigated in future studies. For common complex diseases, we found significant enrichment in specific cell groups for multiple traits (fig. S35H). For more than half of the traits, there was enrichment for the same cell type in different tissues driven by both common and tissue-specific genes. Future work will be needed to extend these analyses across a broad set of tissues and cell types and examine the role of disease-associated genes and cell types in a disease context in patient samples.

Advances in single-cell epigenomics (175) and multi-omics (176–178) should further enable linking GWAS variants to their target genes and the cell types and programs in which they act. Recent findings indicating that a large fraction of genetic regulatory effects linked to GWAS variants can only be detected at the cellular level (11, 25) suggest that cell-level eQTL maps will be essential. The experimental and computational methods we developed for a cross-tissue atlas, and the biological queries we defined, will provide a basis for scaling such efforts to hundreds of individuals and diverse populations.

Methods summary

Tissue samples were selected from among a subset of GTEx project samples that were flash frozen and banked. Nuclei from each sample were extracted using the EZ, CST, NST, and TST protocols described in (26). Libraries for snRNA-seq were generated using the Chromium Single Cell 3' v2 Reagent Kit (10x Genomics), and sequencing was performed with Illumina HiSeq X (96 samples) or NextSeq (three samples), according to the manufacturer's protocols. The resulting snRNA-seq data were aligned and quantified using Cell Ranger v2.1.0 (10x Genomics), ambient RNA correction was performed using CellBender (30), and low-quality nuclei were filtered out using standard criteria (28). The resulting snRNA-seq expression profiles were integrated across samples using a total correlation variational autoencoder (28). Detailed descriptions of all computational analyses are provided in (28).

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SUPPLEMENTARY MATERIALS

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Single-nucleus cross-tissue molecular reference maps toward understanding disease gene function

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Cartography of human cells

The function of disease genes active in different cell types is modulated to meet the needs of the different tissues and organs in which the cells reside. Resolving these differences is critical to understanding homeostasis and disease. However, single-cell atlases generated to date have largely focused on individual tissues. Eraslan *et al.* applied single-nucleus RNA sequencing to frozen, banked samples from eight healthy human organs from 16 donors and characterized cell populations across tissues, including tissue-resident myeloid and fibroblast populations, and their role in tissue support and immunity (see the Perspective by Liu and Zhang). Using this cross-tissue atlas, the authors linked specific cell populations to monogenic and polygenic diseases, suggesting cell- and tissue-specific programs. — LZ and DJ

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