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The immune cell landscape in kidneys of patients with lupus nephritis

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COMPETING INTERESTS

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A.A., D.A.R, A.D., P.J.H., A.H.J. and D.J.L. analyzed the data; D.A.R., C.C.B., T.M.E., E.P.B., J.A.L. and D.A.H. developed the sample collection and processing protocols; D.A.R., Y.L., P.J.H., A.C., A.N., D.S. and S.S. processed the samples; S.L., D.J.L., A.N., D.S. and S.S. developed the scRNA-seq library preparation protocol; F.Z. and K.S. developed the web-based browser of the data; D.E.S., P.T., E.M., M.D.E., M.P., D.L.K., R.A.F., F.P.S., W.F.P., E.A.M., J.P.B., M.A.P., C.P., K.C.K., E.S.W., D.A.H., D.W. and J.H.A. acquired samples; A.A., D.A.R., C.C.B., A.D., W.A., J.A.L., D.A.H., C.N., D.W., M.K., J.H.A., M.B.B., N.H. and B.D. designed the study; A.D., W.A., D.A.H., C.N., S.R., D.W., M.K., J.H.A., M.B.B., N.H. and B.D. supervised the work; A.A., D.A.R., C.C.B., A.D., P.J.H., N.H. and B.D. wrote the manuscript.

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Abstract

Lupus nephritis is a potentially fatal autoimmune disease for which the current treatment is ineffective and often toxic. To develop mechanistic hypotheses of disease, we analyzed kidney samples from patients with lupus nephritis and from healthy control subjects using single-cell RNA sequencing. Our analysis revealed 21 subsets of leukocytes active in disease, including multiple populations of myeloid cells, T cells, natural killer cells and B cells that demonstrated both pro-inflammatory responses and inflammation-resolving responses. We found evidence of local activation of B cells correlated with an age-associated B-cell signature and evidence of progressive stages of monocyte differentiation within the kidney. A clear interferon response was

observed in most cells. Two chemokine receptors, *CXCR4* and *CX3CR1*, were broadly expressed, implying a potentially central role in cell trafficking. Gene expression of immune cells in urine and kidney was highly correlated, which would suggest that urine might serve as a surrogate for kidney biopsies.

Lupus nephritis (LN) is a frequent complication of systemic lupus erythematosus (SLE)^{1, 2}, for which current therapies are both toxic and insufficiently effective^{2, 3}. Despite the rapid pace of immunologic discovery, most clinical trials of rationally designed therapies have failed in both general SLE and LN, with only one new drug approved for the treatment of SLE in the last five decades^{2, 4}. Thus, there is a pressing need to decipher the immune mechanisms that drive LN.

Current knowledge of the molecular pathways dysregulated in SLE comes mainly from the unbiased analysis of blood cells⁵; however, the extent to which blood reflects the inflamed tissue is unclear. Immunohistochemistry and flow cytometry studies of kidney biopsies have indicated the presence of infiltrating subpopulations of immune cells^{6, 7, 8} but cannot reveal previously-unidentified cell types or activation states. Mouse models of LN provide detailed knowledge of the molecular pathways and cell types active in their kidney⁹ but vary in key aspects such as the degree of immune cell infiltration, the role of interferon and Fc receptors and responses to therapeutic interventions.

It is thus clear that the study of LN can greatly benefit from a resource allowing the generation and preliminary testing of new hypotheses. Single-cell transcriptomics is a powerful tool capable of producing a complete catalogue of cell types and states present in a given sample. Here, we employed it in the analysis of kidney, urine and blood samples from patients with LN and healthy individuals, while utilizing a standardized protocol to process patient samples acquired across a distributed clinical and research network. Our findings delineate the complex array of leukocytes active in human LN kidneys. Analysis of blood reveals both similarities to and differences from the molecular signatures detected in kidneys, highlighting the limitations of blood samples for deciphering renal disease processes. We further show that urine cells have the potential to serve as surrogates for kidney biopsies in assessing the molecular activation state of subsets of infiltrating leukocytes.

RESULTS

Isolation and processing of kidney cells for single-cell transcriptomics

To establish a uniform pipeline to analyze kidney biopsy samples from multiple institutions, we evaluated several strategies for their preservation and transport. Cryopreservation of intact kidney tissue immediately after acquisition, followed by batched processing at a central site, offered robust leukocyte yields, intact staining for lineage markers and high quality transcriptomes (Supplementary Fig. 1a-e). We employed this pipeline to analyze kidney biopsies from 24 patients with LN and 10 control samples, acquired from living donor kidney biopsies (Fig. 1a and Supplementary Table 1). Approximately half of the LN samples, independent of histologic classification, provided leukocyte yields well above those

obtained from control samples (Supplementary Fig. 2a,b), including B cells, T cells, macrophages and other leukocytes based on flow cytometry (Supplementary Fig. 2c,d).

Viable cells were sorted into 384-well plates for single-cell RNA sequencing (scRNA-seq) using a modified CEL-Seq2 protocol¹⁰. Since our focus was on characterizing the immune cells within LN kidneys, 90% of the cells sequenced from each sample were CD45⁺ cells, and the rest were CD45⁻CD10⁺ cells. The quality of the collected sequencing data was comparable across plates, and higher in leukocytes compared with epithelial cells, reflecting the lower viability of the latter in the processed samples (Supplementary Fig. 1f,g). Principal component analysis performed on the gene expression data from 2,736 leukocytes and 145 epithelial cells indicated that the main sources of variability in the data corresponded to cell types, rather than batch or technical factors (Supplementary Fig. 2e,f).

Stepwise cell clustering identifies cells of the myeloid, T, natural killer (NK), B and epithelial lineages

To identify the lineage and activation state of the cells extracted from kidney samples, we clustered them based on their gene expression data, taking a stepwise approach (Fig. 1b). Low-resolution clustering of all kidney cells identified 10 clusters (Supplementary Fig. 2g), which we labeled as myeloid cells (clusters C4 and C6), T/NK cells (C0, C1, C2 and C5), B cells (C3, C8), dividing cells (C9) and kidney epithelial cells (C7), based on the expression of canonical lineage markers and other genes specifically upregulated in each cluster. Sensitivity analysis demonstrated that this labeling of cells was highly robust (Supplementary Table 2).

We next clustered the cells of each lineage separately, and identified 21 immune cell clusters and a single epithelial cell cluster (Fig. 2a), each containing cells from multiple patients and plates (Supplementary Tables 3 and 4). Saturation analysis indicated that the size of the present cohort is adequate to reveal most of the major clusters in LN kidneys (Supplementary Fig. 2h). Only three clusters were substantially represented in living donor control samples (Supplementary Fig. 2i and Supplementary Table 3), as verified by analyzing two additional living donor control samples, using a droplet-based approach to maximize the number of processed cells; this increased the number of living donor high-quality cells from 183 to 305, yielding largely the same results (Supplementary Table 3). For clusters that were present in sufficient numbers in both patients with LN and living donor controls, we performed a cluster-based differential expression analysis, comparing the two patient populations; the results of this analysis (Supplementary Table 5) are reported below.

The renal and systemic interferon responses are highly correlated

Type I interferons are elevated in the peripheral blood of patients with lupus ¹¹. To assess the extent of this phenomenon in kidney, we calculated for each cell an interferon response score, defined as the average expression of several known interferon-stimulated genes (ISGs; Supplementary Table 6). We found that in all patients but one there was a significant upregulation of this score in kidney cells compared with living donor controls (Fig. 2b). Furthermore, this upregulation was observed in all clusters, although it was less pronounced in the kidney epithelial cells (Fig. 2c). Two clusters, one containing B cells and the other

CD4⁺ T cells (CB3 and CT6, respectively; see below), demonstrated particularly high values of the interferon response score; the majority of these cells were extracted from two patients (patient IDs 200–0841 and 200–0874; Supplementary Table 3). These two patients also featured B cells and CD4⁺ T cells with a substantially lower interferon score, suggesting that the secretion of this cytokine may be spatially localized, either in the kidney or outside of it. When we compared the interferon response score in matched blood and kidney samples from 10 patients with LN we found a significant correlation (Spearman's $\rho = 0.733$, P = 0.016; Fig. 2d), indicating that the interferon response is mainly an extrarenal process.

Classification and annotation of myeloid cell clusters reveal resident and infiltrating populations

Focused analysis of the 466 cells in myeloid clusters C4 and C6 revealed 5 finer clusters (clusters CM0-CM4, Fig. 3a and Supplementary Fig. 3a-c). We determined their putative identity by comparing their global gene expression patterns with those of published reference monocyte/dendritic cell (DC) clusters identified in blood samples of healthy individuals using scRNA-seq¹² (Fig. 3b,c), and by the expression of canonical lineage markers. Cluster CM3 was closest to CD1C+ DCs (reference clusters DC2 and DC3) or CLEC9A⁺ DCs (reference cluster DC1), in accordance with the expression of the canonical DC markers CD1C and FLT3 (Supplementary Fig. 3a), and the lack of expression of monocyte markers CD14 and CD16. Cluster CM0 cells were most similar to CD16+ patrolling monocytes (reference clusters Mono2 and DC4), with very high expression of CD16 (FCGR3A) and CX3CR1 and low expression of CD14 and CCR2. Similar results were found for clusters CM1 and CM4, though their correlation scores with the reference clusters were notably lower and, in the case of CM4, below a data-derived "assignability threshold". CM1 cells expressed lower levels of CX3CR1 and CD16 than CM0, while CM4 cells expressed even lower levels of these two genes and higher levels of CD14 and CD64 (FCGR1A), despite being dissimilar to classical CD14⁺ monocytes. These three clusters likely represent infiltrating kidney monocyte/macrophage subsets as they constitute a minority of myeloid cells in normal kidneys (Supplementary Table 3).

We next determined whether the pattern of gene expression in each cluster could indicate functional capabilities (Supplementary Fig. 3a). Cluster CM1 expressed upregulated levels of phagocytic receptors *CD36* (*SCARB3*), *SCARB2*, *CD68*, *CD163*, *NR1H3* (*LXR*) and *GPNMB*, and cluster CM4 expressed *VSIG4*, *MSR1*, *CD163*, *MERTK*, *STAB1* and *CD209*. Cells in CM1 and especially CM4 had upregulated expression of *C1Q*, which acts as an opsonin for phagocytosis and promotes apoptotic cell clearance by enhancing the expression of *MERTK* and its soluble ligand *GASo*¹³, ¹⁴. They also demonstrated upregulated levels of *CD169* (*SIGLEC1*), an endocytic receptor that is associated with a phagocytic and reparative phenotype ¹⁵. Cluster CM0 had the highest level of expression of inflammatory genes including *TNF*, *S100A8*, *S100A9*, *NFKB1* and the WNT pathway activator *TCF7L2*. By contrast, CM4 expressed many genes associated with alternatively activated macrophages, including *CD163* and *SLC40A1* (ferroportin), which control iron homeostasis ¹⁶; *IGF1* and *DAB2*, both drivers of the alternatively activated phenotype ¹⁷, ¹⁸; and folate receptor beta (*FOLR2*), a receptor expressed on alternatively activated CD14⁺ macrophages that are found in inflammatory and malignant tissues ¹⁹.

Finally, since CM2 was the main cluster found in normal kidneys (Supplementary Table 3), it probably corresponds to steady-state kidney macrophages. This cluster demonstrated low expression of *CD14*, *CD16*, *CX3CR1* and *CCR2*, and no clear similarity to the published reference clusters of peripheral myeloid subsets (Fig. 3b,c). In comparison to the other macrophage subsets, CM2 upregulated several genes associated with tissue remodeling including *MMP2*, *ADAMTS10* and *HTRA1*. These cells also upregulated *BHLHE41*, a gene expressed in microglia and lung resident macrophage populations²⁰, consistent with CM2 representing resident cells. Compared with CM2 cells from living donor controls, lupus CM2 cells expressed higher levels of ISGs, as well as anti-inflammatory genes (*GRN*, *TMSB4X*, *CREB5*) and inhibitors of TLR signaling (*GIT2*, *TNFAIP8L2*), and lower levels of pro-inflammatory genes (*ALOX15B*, *WNT5A*) (Supplementary Table 5)²⁰.

Trajectory analysis identifies a continuum of intermediate states spanning patrolling, phagocytic and alternatively activated monocytes

Dimensionality reduction using either diffusion maps²¹ (Fig. 3d) or t-Distributed Stochastic Neighbor Embedding²² (tSNE; Fig. 2a) indicated possible transitions between the three clusters of infiltrating monocytes/macrophages, with CM1 linking CM0 and CM4. Furthermore, since the cells in cluster CM0 tended to be the most similar to peripheral blood CD16⁺ monocytes, while the cells in cluster CM4 were the least similar (Fig. 3c), the suggested progression is from an inflammatory blood monocyte (CM0) to a phagocytic (CM1) and then an alternatively activated (CM4) phenotype. Indeed, we found a gradual reduction along the trajectory from CM0 to CM4 in the expression of *NFKB1*, an inflammatory gene; a transient increase in *CD36*, an important phagocytic receptor; and a continuous increase in *MERTK*, a key signaling receptor induced by *CD36* (Supplementary Fig. 3d-f)²³. Overall, a general downregulation of inflammatory genes and a concurrent upregulation of genes associated with phagocytosis (Supplementary Table 6) was observed along this trajectory (Fig. 3e,f).

To further investigate this hypothesized within-kidney transition, we analyzed blood samples from two of the patients who had high numbers of CM1 and CM4 cells in their kidneys (patient IDs 200–0873 and 200–0874; Supplementary Table 3). We used droplet-based scRNA-seq, yielding 1,411 sorted high-quality myeloid blood cells that included a subpopulation of CD16⁺ monocytes (Supplementary Fig. 3g). We next compared the gene expression data of each cell in this subpopulation with that of the myeloid kidney clusters. As expected, the vast majority of peripheral blood CD16⁺ cells were most similar to the CM0 cluster, with a few cells mapped to either CM1 or CM3 and no cell mapped to CM4 or CM2 (Supplementary Fig. 3h). This held true when considering all sorted blood myeloid cells, not just those identified as CD16⁺ monocytes.

To determine whether the hypothesized differentiation begins before entering the kidney, we examined the relative upregulation of phagocytosis-associated genes in cluster CM1 compared with CM0, in both blood and kidney (Supplementary Fig. 3i-j). We found that while there was a significant increase in these genes in kidneys (P < 0.001; Mann-Whitney U-test), no such increase could be observed in blood.

These analyses are consistent with differentiation of CD16⁺ monocytes into CM1 and CM4 cells within the kidney, but do not rule out differentiation of a small number of blood cells coupled with selective migration into the kidney. Furthermore, other schemes of transitions (or their absence) between these clusters are possible, and further investigation is required.

LN kidneys contain two clusters of NK cells and three clusters of CD8+ T cells

Clusters C0, C1, C2 and C5, comprising 1,764 cells, contained T cells and NK cells. A focused clustering of these cells separated them into seven finer clusters of NK, CD8⁺ T and CD4⁺ T cells (clusters CT0-CT6, Fig. 4a and Supplementary Fig. 4a). Cluster CT1 contained NK cells, which could be identified by the lack of CD3E and CD3D combined with expression of CD56 (NCAMI) and DAP12 (TYROBP), as well as high expression of cytotoxic genes including PRF1, GZMB, and GNLY. A similar cytotoxic program was observed in the CD8⁺ T cell cluster CT2, pointing to a cytotoxic T lymphocyte (CTL) identity. A second population of CD8+ T cells, demonstrating high levels of the granzyme GZMK²⁴ rather than GZMB and GNLY, populated cluster CT4. These cells expressed relatively low levels of PRF1 compared with cluster CT2 and also showed high expression of HLA-DR/DP/DQ molecules and CCR5, consistent with an earlier report²⁵. Cluster CT5 could be further split into two subclusters (Fig. 4b and Supplementary Fig. 4b): a third CD8⁺ T cell population (cluster CT5a), and a small population of NK cells (CT5b). The cells in cluster CT5a had features of resident memory cells, including expression of ZNF683 (HOBIT), ITGAE, ITGA1 and XCL1, and lack of KLF2^{26, 27}, and accordingly were relatively abundant in normal kidney biopsies (Supplementary Table 3). Cluster CT5b cells expressed TYROBP and CD56, suggesting an NK cell identity, but differed from CT1 NK cells by higher expression of KIT, TCF7, IL7R, and RUNX2, and lower expression of PRF1, GZMB, FCGR3A, TBX21, and S1PR5, consistent with the identification of these cells as tissue-resident CD56brightCD16- NK cells, in contrast to the CD56dimCD16+ NK cell features observed in CT1²⁸.

It was previously reported that an exhaustion signature in peripheral blood CD8⁺ T cells of patients with SLE associates with lower flare rates²⁹ and that such exhausted T cells are also detected in the kidneys of lupus mice³⁰. In our data, however, all three CD8⁺ T cell clusters (CT2, CT4, CT5a) expressed only low levels of canonical exhaustion markers (Supplementary Fig. 4c); this was probably not due to technical limitations, as *PD-1* (*PDCD1*), *CTLA4* and *BTLA* were highly expressed in the T follicular helper (TFH)-like cells (cluster CT3b – see the following section). We sorted CD8⁺ T cells from matched blood samples obtained at the time of kidney biopsy along with ten healthy control samples, and used data from bulk RNA-seq to measure an "exhaustion score" defined as the average expression of a comprehensive, published list of exhaustion markers³¹ (Supplementary Table 6). We found that this exhaustion score was significantly higher in blood CD8⁺ T cells of patients with LN compared with those of healthy controls (P < 0.01, Mann-Whitney U-test), but not in kidney CD8⁺ T cells (Supplementary Fig. 4d-f), suggesting that the CD8⁺ T cell exhaustion seen in blood does not occur in the affected organ.

Analysis of CD4+ T-cell subsets identifies five clusters, including TFH-like cells

Clusters CT0, CT3 and CT6 contained CD4⁺ T cells. CT3 could be divided into two subclusters, with one (CT3a) containing cells expressing genes associated with T regulatory (T_{reg}) cells, including *FOXP3* and *IKZF2* (*HELIOS*), and the other (CT3b) consisting of cells with low *FOXP3* and features consistent with TFH cells, including the expression of *CXCL13*, *CXCR5*, *PDCD1*, *MAF*, and *CD200* (Fig. 4c and Supplementary Fig. 4g).

Cluster CT0 could be further split into two subclusters, the first containing primarily effector memory CD4⁺ T cells (CT0a), with more frequent expression of *PRDM1*, *CCL5* and *CXCR6*, and the second consisting of mostly CCR7⁺SELL⁺TCF7⁺ central memory T cells (CT0b; Fig 4d and Supplementary Fig. 4h). The similar expression of *CD69* in both clusters suggests that CT0b cells are more likely to be central memory than naive CD4⁺ T cells. Expression of *TCF7*, *KLF2* and *LEF1* may indicate an early central memory T-cell (Tcm) phenotype of CT0b cells, in contrast to the late effector phenotype of CT0a cells³². Of note, CT0a was the only CD4⁺ T cell cluster found with substantial frequency in living donor samples (Supplementary Table 3). Differential expression analysis of this cluster indicated a dysregulation of ISGs in the LN samples (Supplementary Table 5).

While some LN kidney T cells have been previously annotated as T_H1 and T_H17 cells, in our data CD4⁺ T cells did not segregate into distinct clusters with characteristic effector lineage features. *IFNG* and *CXCR3* could be identified in few CT0 cells, primarily within CT0a (Supplementary Fig. 4h). In contrast, *IL17A*, *IL17F* and *CCR6* were very rarely detected, and no *IL4*, *IL5* or *IL13* expression was observed. *TBX21* and *RORC* were found in a minority of CT0 cells, with *TBX21* much more frequently expressed in CD56^{dim}CD16⁺ NK cells (cluster CT1) and CTLs (CT2; Supplementary Fig. 4a).

Finally, cluster CT6 contained CD4⁺ T cells demonstrating exceptionally higher levels of ISGs, including *ISG15, MX1, RSAD2, OAS3, IFIT1* and *IFIT2*, compared with other T cells (Supplementary Fig. 4a).

Analysis of B-cell clusters reveals age-associated B cells (ABCs)

Analysis of the 435 cells mapped to clusters C3 and C8 identified 4 different B-cell clusters in LN samples, but almost no B cells in healthy kidneys (clusters CB0–CB3; Fig. 5a, Supplementary Figs. 5a and 2d and Supplementary Table 3). Cluster CB1 contained plasmablasts/plasma cells, expressing high levels of *XBP1* and *MZB1*, as well as immunoglobulin genes. The cells in cluster CB3 demonstrated high levels of several ISGs, including *IFIT1*, *IFIT2*, *IFIT3*, *ISG15*, *OAS3* and *RSAD2*. Expression of these genes was also detected in the other B-cell clusters, but at substantially lower levels.

Cluster CB0 cells had upregulated expression of activation markers such as *CD27*, *CD86*, *IGJ* and *IGHG1*, and low levels of *IGHD* and *IGHM*, suggesting an activated B-cell identity. Furthermore, we could detect in this cluster a gene expression signature consistent with ABCs (Fig. 5b) that are implicated in both aging and autoimmunity³³. Based on a panel of genes reported to be differentially expressed in ABCs³⁴, we computed for each cell in cluster CB0 a score representing the extent to which its gene expression pattern matched that expected by an ABC ("ABC score"; Supplementary Table 6). A continuous range of values

of this score could be observed in cluster CB0, without a clear separation into distinct subpopulations (Fig. 5b). The ABC score per patient, calculated as the average of the ABC score over the CB0 cells of each patient, did not positively correlate with age (Spearman's ρ = -0.255), suggesting that the presence of these cells indeed reflected disease rather than age.

The tSNE plot for the B cells suggested that cluster CB2 may contain multiple subsets (Supplementary Fig. 5b). Accordingly, we were able to split the cells in cluster CB2 into two subclusters (Fig. 5c and Supplementary Fig. 5c): the first of these (CB2a), expressing the B cell markers CD19 and CD20 (MS4A1), demonstrated upregulation of genes typical of naive B cells, including high levels of IGHD, IGHM, TCL1A and IL4R, and had nearly undetectable expression of CD27; the other cluster (CB2b) expressed genes known to be upregulated in plasmacytoid DCs (pDCs) (Fig. 5c and Supplementary Fig. 5c), including PTPRS, GZMB, CLEC4C, CD123 (IL3RA) and CD317 (BST2). To further validate this hypothesized identification, we calculated the Pearson correlation in gene expression between each cell in cluster CB2 and 3 independent sets of reference samples: FANTOM5^{35,36}, containing bulk RNA-seq data from 360 cell types, 17 of which are immune cell subsets; bulk RNA-seq data from 13 immune cell populations sorted from healthy individuals (Browne et al., manuscript in preparation); and an scRNA-seq data set, which includes data from 10 different clusters of DCs and monocytes from healthy blood 12. This analysis classified all CB2b cells as pDCs, using any of the three reference data sets (Supplementary Fig. 5d-f). Furthermore, as predicted almost all of the CB2a cells were classified as naive B cells, when compared with the data from Browne et al. (the only data set of the three tested that contained multiple B-cell populations).

Trajectory analysis reveals intermediate states between naive B cells and ABCs

Projecting the gene expression data of the B cells onto two dimensions using diffusion maps²¹, we found that the naive (CB2a) and activated (CB0) B cells formed a continuum of states, demonstrating a gradual increase in *CD27* expression and a parallel decrease in *IGHD* expression, reflecting activation (Fig. 5d-f). Furthermore, traversing the trajectory from CB2a to CB0 coincided with a continuous increase in the ABC score (Fig. 5g), indicating that activation and differentiation into ABCs are highly correlated processes in our data. In contrast, very few cells occupied intermediate states between plasma and naive or activated B cells, consistent with a lack of differentiation into plasma cells in the inflamed kidney. However, as it was previously suggested³⁷ that ABCs are preplasma cells, this question requires additional investigation, in particular employing B-cell receptor (BCR) repertoire analysis.

The dividing cell cluster includes T and NK cells

Cluster C9 contained three subclusters. Two of them (CD1 and CD2) demonstrated upregulated levels of mitochondrial genes and genes associated with a stress response (Supplementary Fig. 6a,b), indicating lower viability and/or quality, and were excluded from subsequent analyses. Cluster CD0 demonstrated elevated levels of genes participating in cell division. Classification of its cells by comparison with FANTOM5 indicated CD8⁺ T cells, NK cells and CD4⁺ T_{reg} cells (Supplementary Fig. 6c).

Cluster-specific expression of genes associated with disease risk

Genome-wide association studies (GWASs) have identified numerous risk alleles and their susceptibility genes in SLE and LN^{38, 39}. We analyzed the expression of these genes across the 22 clusters identified in kidneys, and found both expected and surprising cluster-specific expression patterns (Fig. 6). For example, *TLR7*, whose role in nucleic acid sensing, B cell activation and differentiation is well established^{40, 41} is expressed here in pDCs, myeloid cells and B cells. We found *HIP1*, suggested to regulate DC activity⁴², to be expressed in resident memory CD8⁺ T cells (cluster CT5a), CD56^{bright}CD16⁻ NK cells (CT5b), conventional dendritic cells (cDCs) (CM3) and pDCs (CB2b). *LBH*, implied to modulate synovial hyperplasia⁴³, was expressed here in T- and B-cell subsets, raising the possibility that the LBH risk locus impacts both fibroblasts and lymphocyte subsets. We also observed cluster-specific expression of several poorly annotated SLE susceptibility genes, including *WDFY4*, *CXorf21* and *TMEM39A*. Finally, our analysis identified both innate and adaptive immune cell subsets expressing several transcription factors associated with SLE, including *ARID5B*, *CIITA*, *ETS*, *IKZF1*, *IKZF2*, *IRF7*, *IRF8* and *PRDM1*.

Expression patterns of chemokines and cytokines

We next analyzed the expression patterns of chemokine and cytokine receptors (Fig. 7a), focusing on receptors that were expressed by a relatively large fraction (> 30%) of the cells in at least 1 cluster (this threshold was set based on the observed distribution of expression frequency, considering all receptors; Supplementary Fig. 7a). We found that a single chemokine receptor, *CXCR4*, was expressed in the majority of infiltrating cells in nearly all clusters (Supplementary Fig. 7b). A second chemokine receptor, *CX3CR1*, was expressed in most myeloid cells, as well as CD56^{dim}CD16⁺ NK cells (cluster CT1) and CTLs (CT2) (Supplementary Fig. 7c). Of note, the expression frequency of other chemokine receptors previously implicated in LN, such as *CCR5*, *CXCR3* and *CCR2*, was found to be much lower (Supplementary Fig. 7d-f). For cytokine receptors, we observed that *IL2RG*, encoding the common gamma chain and important for signaling of several cytokines, was frequently expressed in almost all clusters. *TGFBR2*, a subunit of the receptor for the cytokine TGFβ, was also expressed on the majority of cells. *IL10RA*, *IL27RA*, *IL17RA* and *TNFRSF1B* were expressed by a large fraction of cells in all clusters, with the exception of the B cells.

To identify potential interactions between the cells acting in the inflamed kidney, we analyzed the expression patterns of the corresponding ligands. We found that the *CXCR4* ligand, *CXCL12*, was expressed mainly in the cells in cluster CM4, as well as in the epithelial cells (Fig. 7b and Supplementary Fig. 7g). The latter were also the main source of the *CX3CR1* ligand, *CX3CL1* (Fig. 7c and Supplementary Fig. 7h). Of note, CM4 cells were in addition the top producers of *CCL2* and *CCL8* (Supplementary Fig. 7i); these are the ligands of *CCR2*, which is expressed in a large fraction of plasma cells (cluster CB1) and pDCs (cluster CB2b). These findings imply that kidney epithelial cells and M2-like macrophages may be coordinating traffic of immune cells infiltrating the kidney. It should be noted though that this analysis does not cover other cell types, such as endothelial cells, which were not profiled here.

Comparison of urine and kidney leukocytes

Leukocytes isolated from urine samples of patients with LN were processed in the same way as kidney cells (Fig. 1a and Supplementary Fig. 1h). Following filtering, 577 high-quality cells, collected from 8 patients, were included in subsequent analyses.

We first assigned each urine cell to the kidney cluster most similar in its gene expression data. Urine samples had a higher frequency of myeloid cells (in particular cluster CM1) and fewer T cells than kidneys (Fig. 8a). We next compared gene expression across corresponding urine and kidney clusters, restricting the comparison to clusters with at least five urine cells. High correlations were observed, typically ranging from 0.85 to 0.95 (Fig. 8b and Supplementary Fig. 8), suggesting that urine cells can serve to estimate gene expression in their kidney counterparts.

DISCUSSION

Using single-cell transcriptomics to study kidney samples obtained from patients with LN and living donor controls, we reveal the complexity of immune populations in LN kidneys, identifying multiple disease-specific subsets of myeloid, NK, T and B cells, and giving rise to several observations. We found: (1) evidence for within-tissue differentiation of inflammatory CD16⁺ macrophages into M2-like cells, which may orchestrate the renal infiltration and retention of other leukocyte subsets; (2) an abundance of dividing CTLs and NK cells, indicated to be a major source of IFNγ and cytolytic molecules, and lack of expression of exhaustion markers in CD8⁺ T cells, suggesting a role for cytotoxic activity in LN; (3) two additional populations of CD8⁺ T cells, which could not be easily identified by cell surface markers; (4) a range of B-cell activation states from naive cells to ABCs in the kidney; (5) an interferon response signature in infiltrating leukocytes, correlated with the same signature in blood; (6) frequent expression by kidney immune cells of the chemokine receptors *CXCR4* and *CX3CR1*, suggesting they may serve as potential therapeutic targets; (7) cell subset-specific expression of genes associated with lupus in GWAS; and (8) a high correlation of gene expression in urine immune cells and corresponding kidney leukocytes.

Our transcriptomic analysis offered a detailed view of the T-cell populations in LN kidneys. The co-clustering of CD4⁺ TFH cells and FoxP3⁺Helios⁺ regulatory T cells raises the possibility that T follicular regulatory cells are also present^{44, 45}. CD4⁺ T cell clusters were not clearly associated with T_H1 or T_H17 signatures, suggesting that T cell polarization may not be a major feature in LN. The identification of three distinct CD8⁺ T-cell subsets raises the question whether the previously reported localization pattern of CD8⁺ T cells in the kidney⁴⁶ is subset-specific.

B cells are found in more than half of lupus biopsies but not in healthy samples⁴⁷. Our finding of B cells spanning a spectrum of states between naive and activated cells, together with the presence of TFH-like cells, is consistent with the view that immune responses to tissue damage are being driven in situ⁴⁸. Of note, activation was correlated with an ABC signature previously suggested to be driven by BCR/TLR ligands⁴⁹. Understanding whether these ABCs are clonally expanded or enhance inflammation locally in patients with LN, and

determining the clonal relatedness of naive, activated and antibody-secreting B cells will require a larger data set, combined with analysis of BCR sequences.

Peripheral monocytes can enter injured tissues and differentiate into inflammatory and reparative/resolving macrophages⁵⁰. If the cells are chronically exposed to damage-associated molecular pattern molecules (DAMPs) and endosomal TLR ligands, resolution may fail, and macrophages with mixed functions may emerge⁵¹. Here, the three subpopulations of CD16⁺ macrophages are suggested to transition through an inflammatory to a resolution phase; such a functional switch was previously identified in a mouse model of acute inflammation⁵². Of note, we did not identify a cluster of infiltrating cells with high similarity to CD14⁺ monocytes. It is still unclear why some types of tissue injury recruit CD14⁺ macrophages while others recruit CD16⁺ macrophages; influences may include the types of expressed DAMPS and/or other microenvironmental cues such as cytokines and chemokines.

Our study demonstrates the feasibility of profiling kidney samples using single-cell transcriptomics, employing a freezing strategy to minimize batch effects that could mask subtle gene expression signatures. While this strategy may result in the loss of neutrophils and can alter the relative frequency of other cell subsets, we did not observe a major effect on gene expression due to freezing.

Despite our cohort's relatively high diversity with respect to histologic appearance and intercurrent therapies, we observed a surprising number of commonalities. Furthermore, saturation analysis indicated that increasing the cohort size is not expected to drastically change the presented cell subset catalogue. Rather, such an increase can enable investigating how the presence and transcription profiles of particular cell infiltrates are related to disease manifestations and treatment responses. A study currently in progress, performed as part of the AMP RA/SLE consortium, will utilize the sample processing strategy developed here to analyze a much larger cohort, addressing these questions. Furthermore, profiling stromal renal cells together with leukocytes will elucidate their interactions. The results discovered in such studies will be further validated using tissue staining, functional studies in cell lines or primary human cells, and animal models of disease.

METHODS

Human kidney tissue, urine and blood acquisition

Renal tissue, urine and blood specimens from patients with LN were acquired at ten clinical sites in the United States. Institutional review board approval was received at each site. Research biopsy cores were collected from consented subjects either as an additional biopsy pass obtained specifically for research during a clinically indicated biopsy procedure (nine sites), or as a portion of a biopsy specimen acquired for diagnostic pathology during a clinically-indicated biopsy procedure (one site). Control kidney samples were obtained at a single site by biopsy of a living donor kidney after removal from the donor and before implantation in the recipient.

After acquisition, kidney biopsy samples were placed into HypoThermosol FRS preservation solution for 10–30min on ice and then transferred to a cryovial containing 1ml CryoStor CS10 cryopreservation medium (BioLife Solutions). The cryovial was incubated on ice for 20–30min and was then placed in a Mr. Frosty freezing container (Nalgene, catalog no. 5100–0001) and transferred to a –80°C freezer overnight. Cryopreserved samples were then stored in liquid nitrogen and shipped on dry ice to the central processing site, where they were stored in liquid nitrogen until processing.

Kidney tissue thawing and dissociation into a single-cell suspension

Kidney samples were thawed and processed in batches of four samples, with most batches containing both LN and control kidney samples. The cryovial containing the kidney tissue was rapidly warmed in a 37°C water bath until almost thawed. The sample was then poured into a well of a 24-well dish and rinsed in a second well containing warmed RPMI/10%FBS. The tissue was incubated for 10min at room temperature. Specimens were cut into 2–3 pieces and placed into a 1.5 ml centrifuge tube containing 445ul of Advanced DMEM/F-12 (ThermoFisher Scientific, catalog no. 12634–028) and 5ul of DNase I (Roche, catalog no. 04536282001, 100U/ml final concentration). 50ul of Liberase TL (Roche, catalog no. 05401020001, 250ug/mL final concentration) was added, and the tube was placed on an orbital shaker (300-500 r.p.m.) at 37°C for 12min. At 6min into the digestion, the mixture was gently pipetted up and down several times using a cut 1 mL pipette tip. After 12min, 500ul of RPMI/10% FBS was added to stop the digestion. The resulting cell suspension was filtered through a 70-µm filter into a new 1.7 ml microfuge tube. The cells were washed with RPMI/10% FBS, centrifuged at 300g at 4°C for 10min, and resuspended in cold PBS for downstream analyses. Quantification of cell yields was performed by hemocytometer with trypan blue exclusion and by flow cytometry with propidium iodide-exclusion. Yields of cell subsets (leukocytes, epithelial cells) were quantified by acquiring the entire sample through the flow sorter and plotting the number of intact, PI-negative cell events with the appropriate surface markers. Cell yields were normalized to input tissue mass.

Urine cell pellet collection and cryopreservation protocol

Midstream urine samples were collected from patients with LN before kidney biopsy. The total urine volume (15–90 mL) was split into two 50 mL Falcon tubes. Urine cells were pelleted by centrifugation at 200g for 10min, and then resuspended in 1 ml cold X-VIVO10 medium (Lonza BE04–743Q). Cells were transferred to a microcentrifuge tube, washed once in 1ml X-VIVO10 medium, and then resuspended in 0.5 ml cold CryoStor CS10. Cells were transferred into a 1.8 mL cryovial, placed in a Mr. Frosty freezing container, stored in at –80°C overnight, and then transferred to liquid nitrogen. For downstream analyses, cryopreserved urine cells were rapidly thawed by vigorous shaking in a 37°C water bath, transferred into warm RPMI/10%FBS, centrifuged at 300g for 10min, and resuspended in cold HBSS/1%BSA.

Flow cytometric cell sorting of kidney and urine samples

An 11-color flow cytometry panel was developed to identify epithelial cells and leukocyte populations within dissociated kidney cells. Antibodies include anti-CD45-FITC (HI30), anti-CD19-PE (HIB19), anti-CD11c-PerCP/Cy5.5 (Bu15), anti-CD10-BV421 (HI10A), anti-CD10-BV421

CD14-BV510 (M5E2), anti-CD3-BV605 (UCHT1), anti-CD4-BV650 (RPA-T4), anti-CD8-BV711 (SK1), anti-CD31-AlexaFluor700 (WM59), anti-PD-1-APC (EH12.2H7), and propidium iodide (all from BioLegend). Kidney or urine cells were incubated with antibodies in HBSS/1%BSA for 30min. Cells were washed once in HBSS/1%BSA, centrifuged, and passed through a 70 µm filter. Cells were sorted on a 3-laser BD FACSAria Fusion cell sorter. Intact cells were gated according to FSC-A and SSC-A. Doublets were excluded by serial FSC-H/FSC-W and SSC-H/SSC-W gates. Non-viable cells were excluded based on propidium iodide uptake. Cells were sorted through a 100 micron nozzle at 20 psi. For each sample, 10% of the sample was allocated to sort CD10+CD45- epithelial cells as single cells, and the remaining 90% of the sample was used to sort CD45+ leukocytes as single cells. Single cells were sorted into 384-well plates containing 6µl 1% NP-40 with index sorting, and plates were immediately frozen and stored at -80 °C. Flow cytometric quantification of cell populations was performed using FlowJo 10.0.7.

Library preparation and RNA sequencing of kidney and urine samples

scRNA-seq was performed using the CEL-Seq2 method¹⁰ with the following modifications. Single cells were sorted into 384-well plates containing 0.6µl 1% NP-40 buffer in each well. Then, 0.6µl dNTPs (10mM each; NEB) and 5nl of barcoded reverse transcription primer (1μg/μl) were added to each well along with 20nL of ERCC spike-in (diluted 1:800,000). Reactions were incubated at 65°C for 5min, and then moved immediately to ice. Reverse transcription reaction and second-strand complementary DNA (cDNA) synthesis were carried out as previously described¹⁰ and double-stranded c-DNA was purified using 0.8X volumes of AMPure XP beads (Beckman Coulter). In vitro transcription reactions were performed as described followed by treatment with ExoSAP-IT PCR Product Cleanup Reagent (ThermoFisher Scientific, catalog no. 78201.1.ML). Amplified RNA (aRNA) was fragmented at 80°C for 3min and purified using RNAClean XP beads (Beckman Coulter). The purified aRNA was converted to cDNA using an anchored random primer and Illumina adaptor sequences were added by PCR. The final cDNA library was purified using AMPure XP beads (Beckman Coulter). Paired-end sequencing of ~1 million paired-end reads per cell was performed on the HiSeq 2500 in Rapid Run Mode with a 5% PhiX spike-in using 15 bases for Read1, 6 bases for the Illumina index and 36 bases for Read2.

Frozen needle core biopsies obtained from two additional healthy donor kidneys before to reperfusion were processed as above to produce single cell suspensions. Unsorted cells in 0.04% BSA (Sigma) were used to generate single cell libraries with the Chromium Single Cell Gene Expression system using 3′ Library & Gel Bead Kit v2 (10X Genomics) and paired-end sequencing was performed on a HiSeq X.

Processing of blood samples

For profiling blood cells, blood was collected from 10 patients with LN before kidney biopsy and peripheral blood mononuclear cells (PBMCs) were isolated using Ficoll-Paque PLUS (GE Healthcare) density gradient centrifugation in 15ml SepMate tubes (Stemcell) according to manufacturer instructions and cryopreserved in CryoStor CS10 Freezing Media (STEMCELL Technologies). Sex-matched PBMCs from healthy donors isolated and cryopreserved at the University of North Carolina Kidney Center were used as controls.

For bulk RNA-seq experiments, thawed cells were stained with antibodies against CD45-PE (HI30), CD3-PE/Cy7 (UCHT1), and CD8a-APC (HIT8a). 5,000 viable (DAPI⁻) PBMC (CD45⁺) or 1,000–5,000 CD8⁺ T cells (CD45⁺CD3⁺CD8⁺) were sorted into microcentrifuge tubes containing 20µl TCL buffer (Qiagen) supplemented with 1% b-mercaptoethanol (Sigma) using a Sony SH800S cell sorter, and the lysate was frozen and stored at –80 degrees. Of ten LN patient, 8 had enough CD8⁺ T cells to allow sequencing. RNA was isolated with Agencourt RNAClean XP beads (Beckman Coulter) and converted to sequencing libraries using the Smart-seq2 method⁵³. 38bp paired-end reads were generated on a Nextseq500 (Illumina).

For droplet-based scRNA-seq, thawed cells were stained with antibodies from Biolegend: CD3-FITC (HIT3a), CD19-FITC (HIB19), CD20-FITC (2H7), CD56-FITC (HCD56), HLADR-PE/Dazzle (L243), CD16-PerCP/Cy5.5 (3G8), and from BD Biosciences: CD14-APC/Cy7 (M ϕ P9). Viable (DAPI⁻) monocytes (CD3⁻, CD19⁻, CD20⁻, CD56⁻, HLADR⁺, CD14⁺ or CD16⁺) were sorted into RPMI (Life Technologies) + 0.04% BSA (Sigma) and single cell libraries were generated using the Chromium Single Cell Gene Expression system using 3' Library & Gel Bead Kit v2 (10X Genomics). Paired-end sequencing was performed on a Nextseq500.

RNA-seq data processing

For the cells processed using CEL-Seq2, we used a modified version of the Drop-seq pipeline developed by the McCarroll lab⁵⁴, to perform all steps necessary to produce gene by cell expression matrices of reads as well as unique molecular identifiers (UMIs). These steps include demultiplexing, quality filtering, polyA and adapter trimming, aligning, and collapsing reads with unique combinations of cell+gene+UMI. We used STAR-2.5.1b to align reads to the Hg19 human genome reference. Only uniquely mapped reads were counted. UMIs with fewer than 10 reads were filtered out before creating the final expression matrices, to minimize read cross-contamination across cells. For each cell, the computed gene expression counts were then normalized for read depth and log transformed. For cells processed using 10x, sequencing output was aligned using the 10x standard pipeline.

Cell filtering and quality control

For kidney cells processed using CEL-Seq2, high-quality cells were defined as having at least 1,000 detected genes (that is, with positive count values); for urine cells, which tended to have fewer detectable genes, this threshold was set to 500 genes; for cells processed using 10x, the threshold used was 250 genes. We further required the percentage of reads mapped to mitochondrial genes per cell to be lower than 25% (8% for blood cells processed using 10x). To remove wells that were suspected to contain messenger RNA from multiple cells, we required the number of genes per cell to be smaller than 5,000 for the kidney cells processed using CEL-Seq; 4,000 for urine cells; 1,700 for blood cells processed using 10x; 3,500 for the kidney cells processed using 10x (all thresholds were set based on empirical distributions).

To minimize the effect of technical factors, we tested different regression models, taking into account such variables as the plate identifier, number of UMIs per cell, and the percentage of mitochondrial genes per cell. We found that using such models had a negligible effect on the gene by cell expression matrix, as well as the overall results of clustering. We therefore decided to avoid employing them in cleaning the data for subsequent analyses.

In the analysis of myeloid blood cells, initial comparison to the gene expression data of immune cell subsets in FANTOM5 was performed to further validate their myeloid cell identify, in addition to filtering based on flow cytometry cell sorting.

Cell clustering

Clustering of kidney cells was done using Seurat (v1.4.0.8)⁵⁵, in a stepwise manner. We initially performed low-resolution clustering, analyzing all cells together, then labeled each of the resulting clusters as myeloid cells, T/NK cells, B cells, dividing cells or epithelial cells. The cells of each such general class were then analyzed separately, to identify finer clusters. In some cases, as described in the main text, the resulting clusters were further split into subclusters. In each case, clustering was done following principal component analysis, based on context-specific variable genes that were identified independently for each set of analyzed cells.

Sensitivity analysis was performed in each clustering step, with a particular focus on the low-resolution clustering stage. Briefly, all parameters in the clustering process, including the number of variable genes and principal components considered, were varied, and the robustness of the results was determined. To assess this robustness, we estimated in each case the Rand index: looking at a large number (1,000) of random pairs of cells, we counted how many pairs were either included in the same cluster in both of the compared clustering runs, or not included in the same cluster, and referred to these as consistent pairs; we then calculated the fraction of consistent pairs of all random cell pairs considered. We repeated this procedure 100 times, to calculate the mean of the Rand index estimate.

Classification by correlation

In determining their putative identity, we compared the gene expression of individual cells with external gene expression data sets of reference samples. In each comparison, we computed the Pearson correlation between the log-transformed gene expression data of the cell and the reference sample, and chose the reference sample that produced the maximal correlation value (using Spearman correlation instead of Pearson correlation did not drastically change the classification results). To assess the confidence in this classification, we computed in each case an "assignability threshold": we generated 1,000 "random cells", by averaging for each gene the raw counts across the classified cells using random weights, such that the sum of weights for each gene was 1; we then normalized the sum of counts to 10,000. For each random cell, we identified the most similar reference sample, and recorded the corresponding Pearson correlation. The assignability threshold was set to the 95th percentile of the distribution of these Pearson correlations. We note that this approach preserves the main aspects of the original data; in particular, highly expressed genes (such as house-keeping genes) remain high in the generated random cells.

Myeloid cells were compared with the scRNA-seq data published in Villani *et al.*¹², such that each cluster in that study was represented by the average expression over the cells included in it, taking into account only genes showing high variability in that data set. Similar results were found if all genes, or only cluster markers, were considered. Comparison to FANTOM5 and the data from Browne *et al.* was done based on the median of reference sample replicates, considering all genes.

Differential expression analysis

Identification of genes differentially expressed between patients with LN and living donor controls was done using the framework proposed by McDavid *et al.*⁵⁶, as implemented by Seurat. P values were corrected for multiple comparisons using the Benjamini-Hochberg method⁵⁷. For each gene with a corrected P value smaller than 0.05 ("candidate differentially expressed genes"), a further correction for the number of patients was performed: 1,000 random permutations of patients across the two groups were generated, while keeping the number of patients in each group fixed. For each candidate differentially expressed gene and each random permutation, the McDavid test statistic was computed as above. We then calculated a new P value for each candidate differentially expressed gene, defined as the fraction of random permutations in which the value of the test statistic was more extreme than its value in the original partition of patients between groups (while adding 1 to both numerator and denominator). Finally, the new P values were corrected for multiple comparisons using the Benjamini-Hochberg method. This analysis was performed separately for each cluster with at least 20 cells in both patient groups.

Trajectory analysis

Trajectory analysis was performed based on dimensionality reduction using diffusion maps, as implemented in the Destiny software package (v2.6.2)²¹. In each case, only the cells relevant to the question at hand were analyzed.

Calculation of gene set-based scores

Scores based on specific gene sets (for example, interferon response score, ABC score etc.) were calculated for each cell as the average of the scaled (Z-normalized) expression of the genes in the list. To control for the variable quality and complexity of the data of different cells, the score of each cell was corrected by subtracting the average of a large set of similarly-expressed genes, as proposed by Tirosh $et\ a^{58}$. When the list contained genes that are expected to be upregulated in a particular condition and genes that are expected to be downregulated in it (as was the case for the ABC score), the average of scaled expression was calculated separately for each set of genes, and the difference between the scores of the upregulated genes and the downregulated genes was taken as the overall score. A similar approach was used when calculating gene-set based scores in bulk RNA-seq data. The gene sets used for particular scores can be found in Supplementary Table 6.

Analysis of interferon response score

To assess the statistical significance of ISGs upregulation per patient, an interferon response score was calculated for each cell based on a given list of ISGs, as explained above. For each

patient with LN, the distribution of the calculated scores was compared with that of cells collected from living donor controls, using the two-tailed Mann-Whitney U-test. The derived P values were then corrected for multiple comparisons, using the Benjamini-Hochberg method⁵⁷. A similar approach was used when comparing the distribution of ISG scores per cluster in the patient with LN, as compared with the LD cells taken as a whole.

Analysis of GWAS gene expression

We analyzed the expression patterns of 180 genes previously reported in GWASs of either SLE or LN. For each such gene, we calculated its average scaled (Z-normalized) expression in each cell cluster, taking into account only cells coming from LN samples. For biclustering of GWAS genes and cell clusters, we kept only genes that had an average scaled expression value of more than 1 or less than -1 in at least one cell cluster, such that biclustering was based only on the GWAS genes that were relatively variable in our data. Biclustering was then performed, based on the average scaled expression in each cell cluster and using a Euclidean distance metric. cDCs, conventional dendritic cells.

Analysis of chemokine/cytokine receptors

Analysis of chemokine/cytokine receptors was based on a receptor-ligand pairs list downloaded from the International Union of Basic and Clinical Pharmacology (IUPHAR) and British Pharmacological Society (BPS) database⁵⁹ and extended manually to incorporate a number of missing, previously published pairs. For each receptor, we calculated the percentage of cells expressing it in each cell cluster, where a receptor was said to be expressed by a cell if it had at least one mapped read (the results reported here were found to be robust to changes in this threshold). For biclustering of receptors and cell clusters, we kept only receptors that appeared in at least 30% of the cells in at least one cluster.

Assignment of urine cells to kidney clusters

For each urine cell, we computed its Pearson correlation with each kidney cluster, taking the average over the kidney cells included in the cluster. The urine cell was then assigned to the cluster that produced the highest correlation value.

CODE AVAILABILITY

All R scripts used to analyze the data reported in this publication are available from the corresponding authors upon request.

DATA AVAILABILITY

The data reported in this publication, including the clinical and serological data of the study participants, are deposited in the ImmPort repository (accession code SDY997). The raw single-cell RNA-seq data are also deposited in dbGAP (accession code phs001457.v1.p1). The processed data can be viewed using an interactive browser at https://immunogenomics.io/ampsle, https://immunogenomics.io/cellbrowser/, and https://portals.broadinstitute.org/single_cell/study/amp-phase-1.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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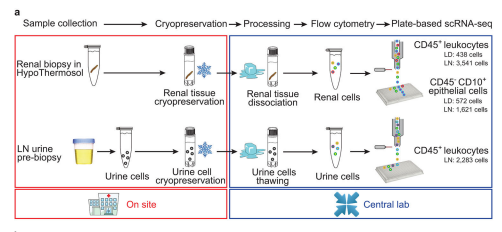
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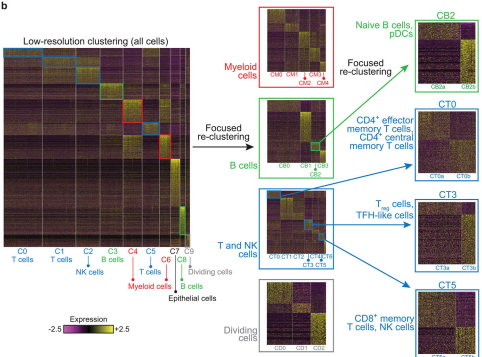


Figure 1.

An overview of the approach used for analyzing the cellular contents and molecular states of kidney and urine samples. **a,** Pipeline for collecting and processing kidney and urine samples. Both types of samples were frozen on collection, then shipped to a central processing site to minimize batch effects. **b,** Stepwise clustering of kidney cells. Initially, all cells were analyzed together (left heatmap), and the identified clusters were labeled as containing either myeloid cells (red), B cells (green), T cells or NK cells (blue), dividing cells (gray) or epithelial cells. Each lineage, with the exception of epithelial cells, was then analyzed separately (middle heatmaps), to identify finer clusters. One B-cell cluster and three T-cell clusters were further re-clustered separately, to generate an even finer description of cell subsets (right). LD, living donor.

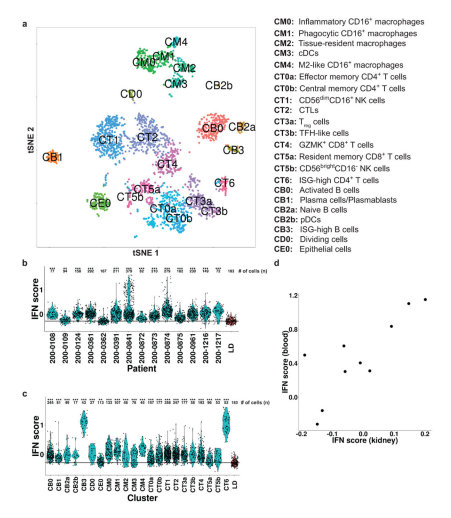


Figure 2.

A summary of the stepwise clustering of kidney cells. **a,** Twenty-two clusters were identified; their putative identities are specified on the right. **b,** The distribution of the interferon response score in all patients with LN (blue), compared with the cells of the LD controls (red). **c,** The distribution of the interferon response score in all cells of patients with LN, separated into clusters (blue), compared with cells of the LD controls (red). In both (b) and (c), *** - FDR-corrected p-value < 0.001; ** - FDR-corrected p-value < 0.01 (two-tailed Mann-Whitney U-test). The number of cells (n) used in each comparison is specified above the plot. The horizontal line designates the median interferon response score over the cells of the LD controls. **d,** A comparison of the interferon response score in kidney and in blood in 10 patients with LN for whom corresponding blood and kidney samples were available. The kidney score was calculated as the average over all kidney cells per compared patient; the blood score was calculated based on bulk RNA-seq data of total PBMCs. IFN, interferon.

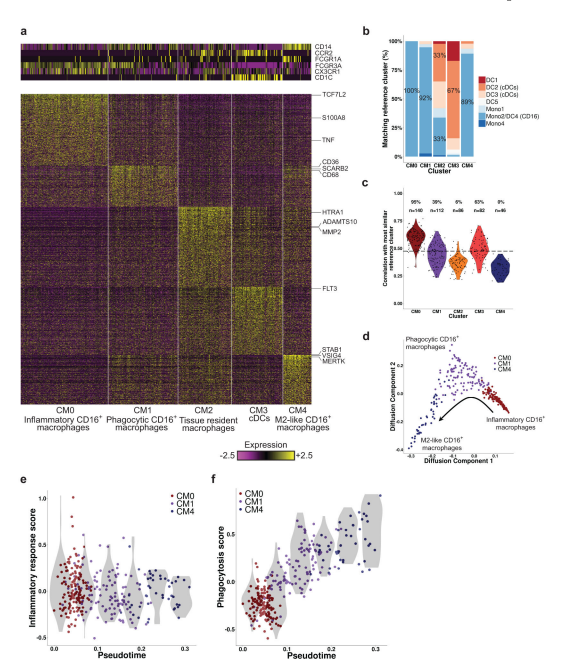


Figure 3.
Focused analysis of kidney myeloid cells. **a**, Five clusters of myeloid cells were identified. The heatmaps show the expression of either canonical lineage markers (top) or genes differentially upregulated in each cluster (bottom). **b**, The results of classifying the kidney myeloid cells by correlating their gene expression to a set of ten reference clusters (Mono1-Mono4, DC1-DC6), taken from Villani *et al.*¹² For each of the Five clusters identified in our data, the bars denote the percentage of cells most similar to each of the reference clusters. The percentage of cells mapped to the most frequent reference cluster in each case is specified on the corresponding bar. **c**, The distribution of highest Pearson correlation values per kidney myeloid cell, when compared with the reference clusters in Villani *et al.* The

percentage of cells in each cluster for which the correlation score was above the assignability threshold is specified above the plot, followed by the number of cells in the cluster (n); the assignability threshold itself is denoted by the horizontal dashed line. **d**, The cells of clusters CM0 (red), CM1 (purple) and CM4 (blue), presented in two dimensions using diffusion maps. The arrow represents the direction of the putative transition between these three clusters, as explained in the text. **e**, The change in the inflammatory response score, calculated as the average scaled expression of several pro-inflammatory genes, along the trajectory shown in d; "pseudotime" represents the ordering of the cells along this trajectory. The violin plots (shades) show the distribution of expression levels in equally-spaced intervals along the pseudotime axis (and do not directly correspond to cell clusters). **f**, Same as e, but with regard to a set of genes associated with phagocytosis.

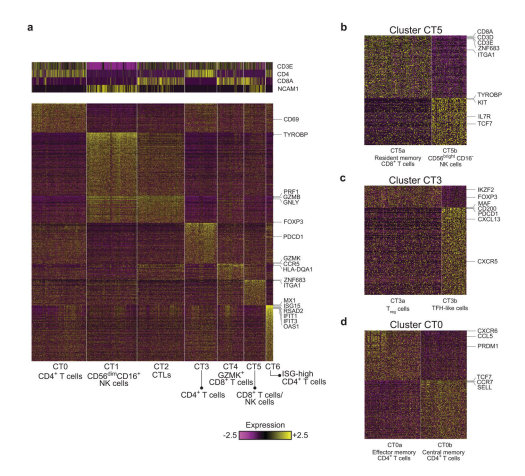


Figure 4.Focused analysis of kidney T cells and NK cells. **a,** Preliminary analysis identified seven clusters. The heatmaps show the expression of either canonical markers defining T-cell and NK cell subsets (top) or genes differentially upregulated in each cluster specifically (bottom). **b,** The splitting of cluster CT5 into two subclusters, representing resident memory CD8⁺ T cells (CT5a) and CD56^{bright}CD16⁻ NK cells (CT5b). **c,** Cluster CT3 can be split into two subclusters, putatively corresponding to CD4⁺ T_{reg} cells (CT3a) and TFH-like cells (CT3b). **d,** Analyzing the cells of cluster CT0 reveals two populations of cells, one putatively identified as early effector memory CD4⁺ T cells (CT0a), the other late central memory CD4⁺ T cells (CT0b).

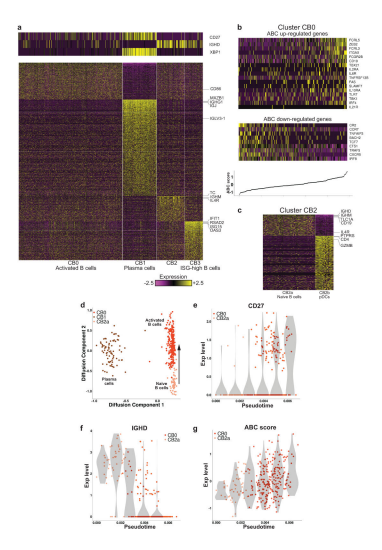


Figure 5.

Focused analysis of kidney B cells. a, Preliminary analysis identified four clusters. The heatmaps show the expression of either canonical markers defining B-cell subsets (top) or genes differentially upregulated in each cluster specifically (bottom). b, The expression of genes previously found to be differentially expressed in ABCs. The top heatmap pertains to genes known to be upregulated in ABCs, the bottom heatmap to genes downregulated in this subset. Columns are sorted by the ABC score, defined as the difference between the average expression of these two sets of genes. The bottom panel shows the ABC score per cell, such that each point on the line corresponds to the heatmap column directly above it. c, Cluster CB2 split into two subclusters, one corresponding to naive B cells (CB2a), the other to pDCs (CB2b). d, Projection of the cells in clusters CB0, CB1 and CB2a onto a two-dimensional plane, using diffusion maps. The arrow represents the hypothesized direction of transition along the trajectory from naive to activated B cells. e-f, The changes in the expression of CD27 and IGHD along the trajectory shown in d. g, The change in the ABC score along this trajectory. In (e)-(g), the violin plots (shades) show the distribution of expression levels in equally spaced intervals along the pseudotime axis (and do not directly correspond to cell clusters).

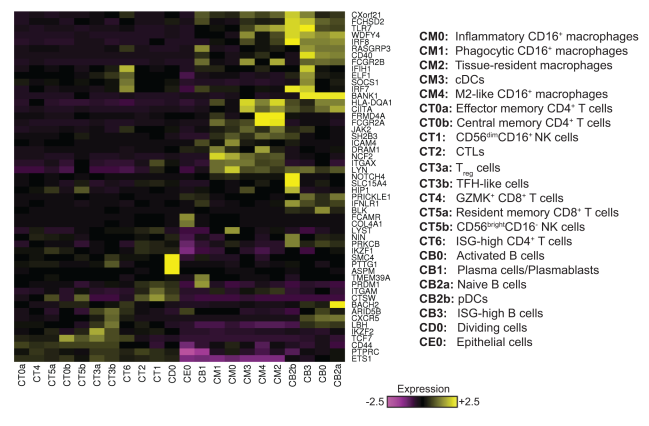


Figure 6.

The expression of GWAS genes in LN kidneys. The heatmap shows, for each gene, the scaled average expression over all cells in each cluster. Included are genes previously indicated in lupus by GWAS, considering only genes that demonstrated high variability across clusters in our data. Both rows and columns are clustered based on Euclidean distance.

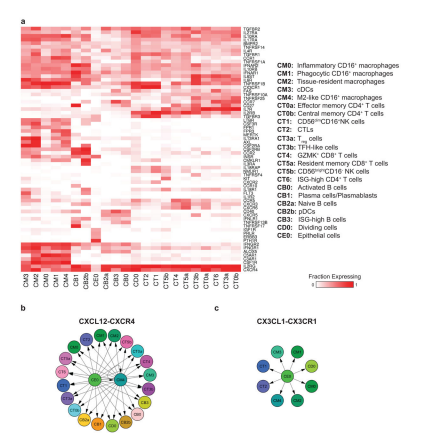
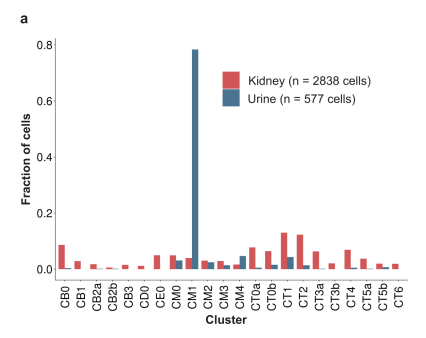


Figure 7. Chemokine- and cytokine-mediated cellular networks. **a,** The pattern of chemokine receptor expression over the cell clusters. The color codes for fraction of cells expressing each receptor. Shown are receptors that are expressed in at least 30% of the cells of at least one cluster. Both rows and columns are clustered based on Euclidean distance. **b,** The producers-consumers cellular network corresponding to the chemokine *CXCL12* and its receptor *CXCR4*. **c,** The producers-consumers cellular network of the chemokine *CX3CL1* and its receptor *CX3CR1*.



CM0: Inflammatory CD16* macrophages CM1: Phagocytic CD16* macrophages CM2: Tissue-resident macrophages

CM3: cDCs

CM4: M2-like CD16⁺ macrophages CT0a: Effector memory CD4⁺ T cells CT0b: Central memory CD4⁺ T cells CT1: CD56^{dim}CD16⁺ NK cells

CT2: CTLs
CT3a: T_{reg} cells
CT3b: TFH-like cells
CT4: GZMK+ CD8+ T cells

CT5a: Resident memory CD8+ T cells

CT5b: CD56brightCD16- NK cells CT6: ISG-high CD4+ T cells CB0: Activated B cells

CB1: Plasma cells/Plasmablasts

CB2a: Naive B cells CB2b: pDCs CB3: ISG-high B cells

CD0: Dividing cellsCE0: Epithelial cells

b		Kidney							
		CM0	CM1	CM2	CM3	CM4	CT0b	CT1	CT2
	CM0 (n = 18)	0.92	0.90	0.83	0.85	0.80	0.73	0.74	0.75
	CM1 (n= 452)	0.90	0.95	0.90	0.90	0.88	0.75	0.75	0.76
	CM2 (n= 14)	0.82	0.87	0.89	0.85	0.86	0.72	0.70	0.71
	CM3 (n= 8)	0.80	0.83	0.86	0.90	0.79	0.76	0.72	0.74
	CM4 (n= 27)	0.80	0.85	0.84	0.81	0.87	0.69	0.68	0.68
	CT0b (n= 9)	0.72	0.71	0.72	0.77	0.65	0.92	0.82	0.87
	CT1 (n= 25)	0.77	0.76	0.75	0.79	0.71	0.86	0.95	0.93
	CT2 (n= 8)	0.74	0.73	0.73	0.77	0.68	0.87	0.89	0.91

Figure 8.

Comparison of immune cells extracted from urine samples and from kidney samples. **a,** The relative frequency of each cluster in urine and in kidney. **b,** Pearson correlation values between gene expression data of urine and kidney clusters, computed using the average gene expression taken over the cells in each cluster, and considering only clusters that had at least five urine cells.