Immune-mediated tubule atrophy promotes acute kidney injury to chronic kidney disease transition

Incomplete repair after acute kidney injury can lead to development of chronic kidney disease. To define the mechanism of this response, we compared mice subjected to identical unilateral ischemia-reperfusion kidney injury with either contralateral nephrectomy (where tubule repair predominates) or contralateral kidney intact (where tubule atrophy predominates). By day 14, the kidneys undergoing atrophy had more macrophages with higher expression of chemokines, correlating with a second wave of proinflammatory neutrophil and T cell recruitment accompanied by increased expression of tubular injury genes and a decreased proportion of differentiated tubules. Depletion of neutrophils and T cells after day 5 reduced tubular cell loss and associated kidney atrophy. In kidney biopsies from patients with acute kidney injury, T cell and neutrophil numbers negatively correlated with recovery of estimated glomerular filtration rate. Together, our findings demonstrate that macrophage persistence after injury promotes a T cell- and neutrophil-mediated proinflammatory milieu and progressive tubule damage.

Acute kidney injury (AKI) is the syndromic term used to describe the abrupt reduction in glomerular filtration rate (GFR) caused by an insult such as ischemia, sepsis, nephrotoxin exposure or urinary obstruction. Based on animal models, these insults often precipitate a rapid innate immune response dominated by neutrophil and proinflammatory macrophage recruitment with significant tubular and endothelial injury, leading to GFR reduction even after correction of the inciting event\(^1\). Under ideal conditions (e.g., transient insults in young healthy subjects), repair pathways that include neutrophil egress and a switch to reparative macrophage activation then promote clearance of cell debris and proliferation of surviving tubular cells, leading to restoration of nephron structure and GFR\(^7\)–\(^10\). However, in many cases this repair phase is incomplete and includes maladaptive processes such as interstitial fibrosis and tubule atrophy\(^1\). As a result, people who survive AKI are at an 8.8-fold increase in risk for chronic kidney disease (CKD) and a 3.3-fold increase in risk for end stage renal disease (ESRD)\(^12\)–\(^14\). To date, no therapy is available to improve the post-injury repair process or prevent progression from AKI to CKD. Defining the signals that promote maladaptive kidney repair following AKI may therefore help us identify therapeutic targets to prevent or even reverse tubule atrophy and the progression to CKD.

Investigations of animal models in which the initial injury stimulus is sustained, such as unilateral ureteral obstruction (UOO), have shown that macrophages progressively accumulate in the kidney interstitium adjacent to unrepaired tubules and promote kidney fibrosis\(^15\)–\(^17\). However, the underpinnings of kidney fibrosis and atrophy following transient insults, as is frequently seen in hospitalized patients, are currently unknown. In models of transient injury such as bilateral ischemia-reperfusion injury (IRI), or unilateral IRI with the contralateral nephrectomy (IRI/CL-NX), kidneys undergo a biphasic response of initial tubule cell death driven in part by an innate proinflammatory activation, followed by tubule repair and restoration of normal or near-normal function\(^7\)–\(^10\). However, using the model of unilateral IRI with the contralateral kidney left intact (U-IRI), we and others have shown that the injured kidney undergoes progressive atrophy and fibrosis rather than successful repair\(^18\)–\(^21\). These transiently injured kidneys continue to exhibit large numbers of macrophages long after the tubule repair phase is complete (days 7–10). Knock-out of either the macrophage...
survival factor chitinase 3-like 1 (Chil3I or Bpr-39) or Ccc2, the chemokine receptor for monocyte chemoattractant protein-1 (Mcp1 or Cc2d) reduces macrophage numbers and the degree of fibrosis, but does not significantly impact the degree of tubule or kidney atrophy. This disconnection between fibrosis and atrophy suggests that the progressive loss of kidney tubules during AKI-to-CKD transition involves mechanisms in addition to accumulation of extracellular matrix.

To better understand the pathogenesis of AKI-to-CKD transition and specifically the mechanism of kidney tubule atrophy, we compared the kidney response to identical times of ischemic injury between mice subjected to U-IRI (to induce atrophy) or IRI/CL-NX (to induce adaptive repair). We performed single-cell-RNA-sequencing (scRNA-seq) analysis on day 7, 14, and 30 after injury to identify major cell types in the kidney and the differential transcriptional response between the models in each cell type. We confirmed our previous finding that U-IRI leads to macrophage persistence beyond the period of normal repair, and now show that these macrophages express high levels of T cell and neutrophil activating chemokines including Cxcl16 and Mcp2 (Ccl2), corresponding with a second wave of infiltrating Ccr6 + T cells and Ccl2 + neutrophils in the interstitium. This late recruitment of T cells and neutrophils closely associated with a proinflammatory milieu (including Tnf and Il1b) Concomitantly, the tubular cells from U-IRI kidneys expressed a different profile of injury markers including vascular cell adhesion molecule 1 (Vcam1) and class I & II major histocompatibility (MHC) genes (rather than Kim1), and exhibited a dedifferentiated expression profile, correlating with late kidney atrophy. Depletion of T cells and neutrophils together, but not individually, was found to attenuate the second wave of injury and partially restore tubule mass in the U-IRI model. Consistent with the mouse models, we found that increasing numbers of T cells and neutrophils in the renal interstitium at the time of renal biopsy for AKI negatively correlated with 6-month recovery of GFR. Together, these findings suggest that failed tubule repair leads to macrophage persistence with a second wave of T-cell and neutrophil-dependent proinflammatory immune activation that induces secondary tubule injury and promotes kidney atrophy during AKI-to-CKD transition.

Results

U-IRI leads to tubule atrophy and tubule cell dedifferentiation

Phenotypically, the injured kidneys from the IRI/CL-NX mice had hypertrophied by day 7 after injury; whereas the injured kidneys from the U-IRI mice were equal in size to controls on day 7, but progressively atrophied afterwards (Fig. 1a and Supplementary Fig. 1a–c). By day 30, IRI/CL-NX kidneys weighed 38% more than age-matched control kidneys and the same as kidneys 30 days after contralateral nephrectomy alone (Supplementary Fig. 1d), while U-IRI kidneys weighed 38% less than control kidneys and 69% less than IRI/CL-NX kidneys (Fig. 1b). This resulted in a 38% and 46% decrease in cross-sectional area of the U-IRI kidneys relative to control and IRI/CL-NX kidneys, respectively (Fig. 1c). Staining for the general tubular epithelial cell marker KSP-cadherin revealed a 56% reduction in absolute tubular area 30 days after U-IRI as compared to IRI/CL-NX (Fig. 1d, e), however, the proportion of renal parenchymal area comprised of tubules was only 12% lower in the atrophied U-IRI kidneys than in the hypertrophied IRI/CL-NX kidneys (Fig. 1e). This suggests that kidney atrophy after AKI is predominantly due to reduced tubular mass rather than replacement of tubule epithelia by fibrosis. To assess the differentiation state of the remaining tubules, LTL and megalin staining were performed to quantify the area of preserved proximal tubule brush border. This showed that PT brush border area comprised 29% less of the remaining renal parenchymal area 30 days after U-IRI compared to IRI/CL-NX (Fig. 1f). Quantitative PCR analysis for the mature proximal tubular brush border constituents megalin (Lp2), NaPi-IIa (Slc34a1) and NaDC3 (Slc34a3) along with IF staining for megalin protein confirmed that proximal tubule redifferentiation was significantly reduced at 14 and 30 days in the U-IRI kidneys as compared to IRI/CL-NX kidneys (Fig. 2a–c). Of note, mice subjected to IRI/CL-NX and mice subjected to U-IRI had equivalent creatinine levels on days 14 and 30, with BUN values that approached equivalence, suggesting nearly complete recovery of filtration function by the previously injured IRI/CL-NX kidneys as compared to the uninjured kidneys in the U-IRI mice (Supplementary Fig. 1e, f).

Both mRNA and protein expression levels of the proximal tubule acute injury marker Kim1 (Haucre1) were equally upregulated on day 1 after IRI in both models, confirming that the initial ischemic injury was equivalent following U-IRI and IRI/CL-NX (Fig. 2a–c and Supplementary Fig. 2). Of note, serum levels of both Kim1 and Ngal were significantly higher on day 1 after IRI/CL-NX than after U-IRI, likely due to clearance by the intact contralateral kidney in mice subjected to U-IRI (Supplementary Fig. 3). Serum levels of both Kim1 and Ngal on day 1 were equivalent between survivors and non-survivors in the IRI/CL-NX mice, indicating equivalent initial injury between the surviving IRI/CL-NX and U-IRI comparison groups on days 7, 14, and 30 (Supplementary Fig. 3b, d). However, Kim1 remained substantially higher in the kidneys subjected to U-IRI on days 7 and 14 at the mRNA level and day 14 at the protein level (Fig. 2a–c). As compared to Kim1, Vcam1 expression was not upregulated on day 1 after IRI in either model but markedly increased afterwards, and was significantly higher in U-IRI kidneys than IRI/CL-NX kidneys from day 7 to day 30 (Fig. 2a), correlating with the period of progressive atrophy seen in this model (Fig. 1b). Together, these data suggest that tubular cells are subjected to equivalent initial ischemic injury in both models correlating with increased Kim1 expression, followed by delayed injury and failure to redifferentiate that predominates in the U-IRI model and correlates with increased Vcam1 expression.

Kidney atrophy is preceded by a second wave of immune activation and proximal tubule cell loss

As the difference in kidney injury marker expression between the two models became apparent on day 7 (Fig. 2a), we performed single-cell RNA sequencing (scRNA-seq) of the injured kidneys on days 7, 14, and 30 after IRI to identify the transcriptional and cellular differences underlying tubule repair and tubule atrophy. Unsupervised clustering generated twenty seven distinct cell types with gene expression profiles identifying them as proximal tubule (PT-S1, S2, and S3), injured PT, thick ascending limb (TAL), DCT/CNT (distal convoluted tubule/connecting tubule), collecting duct-principal cells (CD-PC), collecting duct-intercalated cells (CD-IC), endothelium, myofibroblasts, six clusters of macrophages (monocyte, infiltrating, M1, M2, proliferating, and resident macrophages), four clusters of dendritic cells (CD-DC), four clusters of myeloid cells (plasmacytoid DC (pDC), conventional DCs (cDC1 and CDC2), and proliferating cDC1), two clusters of neutrophils (Ngal high and low), four clusters of T cells [naive, Cd4 + T helper/regulatory T (Th/Treg), Cds8a+ cytotoxic T/natural killer T (Tc/NKT), and proliferating T cells], and B cells (Fig. 3a, b).

The kidney atrophy seen on day 14 and 30 after U-IRI was found to correspond to a decrease in tubular cell populations relative to the total cells present, with the greatest loss occurring in the PT cell compartment beginning by day 7 (Fig. 3c, Supplementary Fig. 4, and Supplementary Table 2). Along with tubular cell loss, U-IRI kidneys exhibited an increased percentage of immune cells including macrophages, cDCs, neutrophils, and T cells as compared to the IRI/CL-NX kidneys. Of the immune cells identified, macrophages and DCs were the predominant cell types in the kidney on day 7 after injury in both models. However, both neutrophils and T-cell populations markedly increased by day 14 after U-IRI as compared to those cells in IRI/CL-NX kidneys.

Consistent with the scRNA-seq analysis, quantitative PCR of whole-kidney RNA revealed that markers for macrophages (Adgre1, ...
Fig. 1 | U-IRI leads to tubule atrophy. Wild-type mice were subjected to 27 min of ischemia/reperfusion injury (IRI) with contralateral nephrectomy (IRI/CL-NX) or unilateral IRI (U-IRI) and sacrificed on day 1, 7, 14, and 30 after injury. a Kidney-to-body weight ratios were determined on day 1, 7, 14, and 30 after injury. Data are presented as mean ± SD. n = 10 kidneys/time point. p < 0.0001 between models and in time series (by two-way ANOVA); ****p < 0.0001 in the indicated subgroup analyses (by Bonferroni multiple comparison). b Kidney weights were determined on day 30 after injury. CTRL, age-matched control. Data are presented as mean ± SD. n = 9 control kidneys. n = 10 kidneys/model. p < 0.0001 by one-way ANOVA; ****p < 0.0001 in the indicated subgroup analyses (by Tukey multiple comparison). c Midline kidney cross-section area was determined on day 30 after injury. CTRL, age-matched control. Data are presented as mean ± SD. n = 9 control kidneys. n = 10 injured kidneys/model. p < 0.0001 by one-way ANOVA; **p < 0.01 (p = 0.0075), ****p < 0.0001 in the indicated subgroup analyses (by Tukey multiple comparison). d Midline kidney sections on day 30 after IRI were co-stained with kidney-specific (KSP)-cadherin (red), uromodulin (UMOD, green), and DAPI (blue). Scale bars, 1 mm. e KSP-cadherin-positive area as in d was quantified for the entire kidney section (left panel) and as a percentage of the section area (right panel). Data are presented as mean ± SD. n = 10 kidneys/model. *p < 0.05 (p = 0.0424), ****p < 0.0001 by unpaired two-tailed t-test. f Midline kidney sections underwent IHC staining for lotus tetragonolobus lectin (LTL, dark gray) on day 30 after IRI. Scale bars, 1 mm. g LTL-positive area as in f was quantified for the entire kidney section (left panel) and as a percentage of the section area (right panel). Data are presented as mean ± SD. n = 10 kidneys/model. ****p < 0.0001 by unpaired two-tailed t-test. h Midline kidney sections underwent IHC staining for megalin (dark gray) on day 30 after IRI. Scale bars, 1 mm. i Megalin-positive area as in h was quantified for the entire kidney section (left panel) and as a percentage of the section area (right panel). Data are presented as mean ± SD. n = 10 kidneys/model. ****p < 0.0001 by unpaired two-tailed t-test.
(F4/80) and Cd68, dendritic cells (Itgax), neutrophils (Ly6g) and T cells (Cd3ε, Cd4 and Cd8α) significantly increased in the U-IRI model after day 7, with T cell, macrophage and dendritic cell markers remining high through day 30 (Fig. 4a and Supplementary Fig. 5a). IHC staining for F4/80, CD11c, Ly6G, CD3ε, CD4, and CD8α on day 14 after injury confirmed that the numbers of interstitial macrophages, DCs, neutrophils, CD4 + T helper cells, and CD8+ cytotoxic T cells were significantly higher in the cortex of U-IRI kidneys as compared to contralateral and IRI/CL-NX kidneys, and all but Ly6G were higher in the outer medulla in the U-IRI model (Fig. 4b, c and Supplementary Figs. 6–11). Together, these data demonstrate that macrophages and dendritic cells are the major cell responders in the first 7 days after IRI in both injury models, with the number of macrophages, T cells and neutrophils plateauing between 7 and 14 days in the IRI/CL-NX model whereas macrophage numbers continue increasing between days 7 and 14 in the U-IRI model accompanied by a second wave of immune activation involving recruitment of neutrophils and T cells. The macrophages present in injured kidneys after day 7 express decreasing levels of proreparative arginase 1 (Arg1)28 with increased profibrotic gene expression predominantly in the U-IRI kidneys (Supplementary Fig. 5b–f).
Chemokine-receptor interactions define the immune signature of the second wave of inflammation

To identify pathways mediating the second wave of immune cell recruitment following IRI, we analyzed the gene expression of homing receptors and their known chemokine ligands in the integrated single-cell dataset from both injury models. Macrophages predominantly expressed Ccr2, Ccr5 and Cx3cr1; DCs expressed Ccr2 and Xcr1; neutrophils expressed Ccr1, Cxcr2, and Cxcr4; and T cells expressed Cxcr3 and Cxcr6 (Supplementary Fig. 12). Corresponding homing chemokines for macrophages included the Ccr2 ligands Ccl2, Ccl7, and Ccl12 (expressed by macrophages themselves); the Ccr5 ligands Ccl3, Ccl4 and Ccl8 (made by macrophages and/or neutrophils); and the Cxcr3 ligand Cxcl11 (expressed at low levels by injured PT and myofibroblasts) (Supplementary Figs. 13 and 14). Chemokines for neutrophil recruitment and activation included the Ccr1 ligands Ccl3, Ccl5, Ccl8 and Ccl9 (expressed by neutrophils, CD8a+ T cells, macrophages and dendritic cells, respectively); the Cxcr2 ligand Cxcl2 (expressed by neutrophils); and the Cxcr4 ligand Cxcl12 (expressed by myofibroblasts at a low level). Finally, T-cell recruitment and activation appears to predominantly depend on the Cxcr6 ligand Cxcl16 (expressed by macrophages) since the other potential T-cell recruitment chemokines were expressed either at low levels (Ccl9 and Ccl10) or were not
detected (Cxcl11) (Supplementary Figs. 13 and 14). Together, the data suggests that infiltrating, M1, and M2 macrophages are the predominant cells to express immune-recruiting chemokines in the injured kidneys.

To better understand how macrophages signal in response to unresolved injury, we identified the differentially expressed genes (DEG) of infiltrating, M1, and M2 macrophages between the U-IRI and IRI/CL-NX kidneys. On day 14, we found a set of chemokines including Ccl2, Ccl7, Ccl8, and Ccl12, that were significantly upregulated in all three macrophage populations from the U-IRI kidneys as compared to the IRI/CL-NX kidneys (Fig. 5a–d). The upregulation of these chemokines by day 14 in U-IRI kidneys correlates with the second wave of immune cells observed on days 14 and 30 in this model (Fig. 4a).

Quantitative PCR for the macrophage receptor Ccr2 and its activating chemokines Ccl12, Ccl7 and Ccl2; the neutrophil receptor Ccr1 and its activating chemokine Ccl8; and the T-cell receptor Cxcr6 and its activating chemokine Cxcl16 from whole-kidney mRNA confirmed the significant increase of these ligand-receptor pairs between days 14 and 30 in the U-IRI injury model (Fig. 5e and Supplementary Fig. 15).

To identify which of these ligand-receptor interactions resulted in functional immune cell responses, we utilized the NicheNet ligand-receptor-target algorithm developed by Saeys and colleagues30,31. Since the chemokines were predominantly expressed by macrophages at 14 days (Supplemental Fig. 13), we focused on immune cell-secreted ligands. This analysis revealed extensive chemokine-receptor interactions between infiltrating, M1, and M2 macrophage-
secreted ligands and their receptors on neutrophils (Ccr1, Cxcr2, and Cxcr4), Cd4+ Th/Treg, and Cd8a+ Tc/NKT (Ccr2, Ccr5, Cxcr3, and Cxcr6) (Fig. 6a, b and Supplementary Figs. 16 and 17). For example, Ccl3, Ccl4, and Ccl8 that were significantly upregulated in M1 and M2 macrophages on day 14 after U-IRI are predicted to interact with Ccr1-expressing cells such as PMN cluster #1. As a result, the pathways involved in leukocyte (neutrophils, monocytes, and lymphocytes) chemotaxis were significantly enriched in the infiltrating and M1 macrophages (Fig. 6c). The functional role of these macrophage-secreted chemokines in recruiting the second wave of T cells and PMNs was supported by numerous examples of both CD3ε+ T cells and Ly6G+ PMNs found adjacent to F4/80+ macrophages in kidneys 14 days after U-IRI (Fig. 6d). Taken together, these data show that macrophages that persist beyond day 7 in U-IRI kidneys are selectively activated to express high levels of chemokines that are predicted to promote a second wave of inflammatory neutrophil and
T-cell recruitment, resulting in a local nidus for extensive inter- and intra-immune cell cross-talk.\textsuperscript{32,33}

Late immune activation promotes tubule oxidative stress and secondary injury

To understand the impact of this second wave of immune activation on tubule injury and atrophy, we determined the DEG in PMNs, Tc/NKT, and Th/Tregs between the U-IRI and IRI/CL-NX kidneys. On day 14, we found that PMNs in the U-IRI kidney significantly upregulated inflammatory gene expression including \textit{Il1b}, \textit{Il1f9}, \textit{Tnf}, \textit{Tnfaip3}, \textit{Ifitm1}, and \textit{Lcn2} (Fig. 7a, b). Cd8+ Tc/NKT cells in the U-IRI kidney expressed significantly higher levels of T-cell activation genes such as \textit{Tyrobp}, \textit{Klre1}, \textit{Serpinb9}, \textit{Ripor2}, \textit{Fastl}, \textit{Xbp1}, and \textit{Zfp36l2} (Fig. 7c). Cd4+ Th/Treg cells in the U-IRI kidneys also showed increased expression of T-cell activation genes such as \textit{Fyn}, \textit{Gpr183}, \textit{Laptm5}, \textit{Lax1}, \textit{Lfng}, \textit{Itk}, \textit{Trp53}, and \textit{Zfp36l1} (Supplemental Fig. 18a). The upregulation of these genes is predicted to activate pathways involved in production of interleukin 1, 6, and 12, interferon \(\gamma\), nitric oxide, and superoxide anion in PMNs (Supplemental Fig. 18a, c) on day 14 after U-IRI. Quantitative PCR analysis confirmed that the mRNA expression levels of \textit{Il1b}, \textit{Tnf}, \textit{Fastl},...
**Fig. 7 | U-IRI promotes late PMN- and T- cell-mediated inflammation.** Wild-type mice were subjected to 27 min of ischemia/reperfusion injury (IRI) with contralateral nephrectomy (IRI/CL-NX) or unilateral IRI (U-IRI) and sacrificed on day 7, 14, and 30 after injury. Injured kidneys and normal control kidneys were harvested for single-cell-RNA-sequencing analysis as shown in Fig. 3. 

a–c Volcano plots demonstrating differential gene expression in U-IRI compared to IRI/CL-NX derived polymorphonuclear neutrophil cluster #1 (PMN #1) (a), PMN cluster #2 (b), and Cd8a+ cytotoxic T/natural killer T (Tc/NKT) cells (c) on day 14 after injury. 

d–f Based on differentially expressed genes (DEG) between U-IRI and IRI/CL-NX kidneys on day 14 after injury, the top relevant enriched gene ontology (GO) terms for PMN cluster #1 (d), PMN cluster #2 (e), and Cd8a+ Tc/NKT cells (f) in the U-IRI kidneys are visualized in the dot plots. 

Based on DEG between U-IRI and IRI/CL-NX kidneys on day 14 after injury, the potential ligands expressed by the PMNs, Cd8a+ Tc/NKT, and Cd4+ T helper/regulatory T (Th/Treg) cells were linked to their corresponding potential target genes for the injured proximal tubule (PT) cells and visualized by a chord diagram.
and class I genes (H2-K1, H2-Ab1, H2-Eb1, and Cd74) and class II genes (H2-DR, H2-KI) were upregulated in U-IRI tubular cells, while anti-oxidative stress genes (Gsta2, Gstm1, Gsp1, and Cd74) and class I genes (H2-DR1 and H2-KL) were upregulated in U-IRI tubular cells, and class II gene expression in the injured PT and TAL, DCT/CNT, and CD; Fig. 8c). The increase persisted through day 30 and was recapitulated to a lesser degree in most non-canonical injury markers after U-IRI was observed from day 7 (Supplementary Tables 4 and 5). Ligand-receptor-target analyses revealed that PMNs, Tc/NKT, and Th1/Th2 are projected to interact with MHC class I expression is associated with the increase of nephron segments (PT, TAL, DCT/CNT, and CD; Fig. 8c). Consistent with the single-cell data, quantitative PCR analysis confirmed that performing U-IRI with the contralateral kidney intact leads to a different outcome in which the absolute relative in eGFR at 6 months after AKI revealed a strong negative correlation between inflammatory cell number and eGFR recovery following AKI (Fig. 10c, d).

**Discussion**

Kidney injury has been extensively investigated using mouse models of either bilateral IRI or IRI/CL-NX to develop our current mechanistic understanding of tubule injury and repair including mechanisms of cell death, the innate immune response to injury, proliferation of surviving tubular cells to replace those that are lost, clearance of casts and restoration of GFR23–25. Recently, our group and others have shown that performing U-IRI with the contralateral kidney intact leads to a different outcome in which fibrosis and kidney atrophy predominate rather than tubule repair and restoration of function9,36.

In this study, we show that the initial macrophage, DC, neutrophil and T-cell response to ischemia/reperfusion injury is similar during the first 7 days after injury in the two models. However, after 7 days the responses markedly diverge with a surge in T cells and neutrophils observed in the U-IRI kidneys as compared to IRI/CL-NX kidneys. The divergence becomes detectable on day 7 with continued Kim1 expression by the proximal tubule and de novo upregulation of Vcam1, and ends with extensive kidney atrophy by day 30. As previously reported by our group and others, the decreased kidney size following U-IRI is accompanied by excessive macrophage persistence and a transition to a Pdgfb+ and Tgfβ1+ profibrotic expression profile26–28. However, quantitative analysis of the remaining tubule area relative to total kidney area suggests that the major phenotypic change underling kidney atrophy after U-IRI is tubule loss rather than increased fibrosis (Fig. 1). In fact, the transition of macrophages to a profibrotic phenotype was seen in both models of IRI (Supplementary Fig. 5b, f), consistent with evidence that even "reparative" models of kidney injury such as IRI/CL-NX or bilateral IRI lead to increased kidney fibrosis23–25.

Similar to the results of Liu and colleagues who studied the slowly progressive kidney atrophy and fibrosis seen following bilateral IRI23, we find that the accelerated renal atrophy following U-IRI correlates with sustained tubular injury and inflammatory cell activation. By performing single-cell sequencing and cell-cell ligand-receptor-target interaction analyses, we now demonstrate that this delayed increase in inflammation in the U-IRI kidneys correlates with the unique chemo- kine (Ccl2, Ccl7, Ccl8, Ccl12, and Cxcl16) expression by macrophages, that in turn can recruit more macrophages23, as well as a second wave of infiltrating proinflammatory neutrophils and T cells between 7 and 30 days after injury.

Initial neutrophil recruitment and transmigration peaks at ~24 h after IRI and promotes reactive oxygen species (ROS) production, activation of resident mononuclear phagocytes, and differentiation of patients with AKI. To assess this, ten kidney biopsies from patients with acute kidney injury (AKI), an estimated GFR (eGFR) >60 mL/min/1.73m2 prior to AKI episode, and no history or pathologic indication of glomerulonephritis (GN), diabetic kidney disease (DDK), or acute interstitial nephritis (AIN) were selected from the Yale kidney biopsy biorepository (Supplementary Tables 7 and 8)24–25. For these patients, eGFR declined by 161 (36, 71) mL/min from the time of AKI from their pre-AKI level with a spectrum of renal fibrosis from partial to full eGFR recovery 6 months after AKI (Fig. 10a and Supplementary Tables 7 and 8). We quantified the number of T cells and neutrophils in sections from each of these kidney biopsies, and compared to those seen in kidney biopsies from healthy living kidney donors26. As expected, very few T cells or neutrophils were found in kidneys from living donors, whereas T cells and neutrophils were easily detected in the kidney interstitium of some patients at the time of biopsy for AKI (Fig. 10b). Nonparametric Spearman correlation analysis of the number of T cells or neutrophils (quantiﬁed as a percentage of total cells identiﬁed in the cortex) with either the absolute or relative increase in eGFR at 6 months after AKI revealed a strong negative correlation between inflammatory cell number and eGFR recovery following AKI (Fig. 10c, d).

**Accumulation of T cells and neutrophils is negatively associated with GFR recovery in patients with AKI**

Our animal model findings prompted us to investigate the relevance of T-cell and neutrophil accumulation for recovery of kidney function in
Fig. 8 | Identifying the injury signature of inflammation-induced tubular stress.

Wild-type mice were subjected to 27 min of ischemia/reperfusion injury (IRI) with contralateral nephrectomy (IRI/CL-NX) or unilateral IRI (U-IRI) and sacrificed on day 7, 14, and 30 after injury. Injured kidneys and normal control kidneys were harvested for single-cell-RNA-sequencing analysis as shown in Fig. 3. a, b Volcano plots demonstrating the differential gene expression in cells from segment 1 of the proximal tubule (PT-S1, a) and injured PT (b) 14 days after U-IRI compared to IRI/CL-NX. c The distribution and relative expression of the injury markers (Havcr1, Vcam1, and Lcn2), anti-oxidative stress and detoxification genes (Gatm, Gsta2, Miox, Gpx1, Gpx3, and Gpx4), and the major histocompatibility complex class II (H2-Aa, H2-Ab1, H2-Eb1, and Cd74) and class I (H2-D1 and H2-K1) are visualized in a dot plot. d Quantitative RT-PCR analysis for indicated genes was performed on whole-kidney RNA harvested on day 0, 1, 7, 14, and 30 after injury. Data are presented as mean ± SD. n = 10 kidneys/time point/model. Two-way ANOVA is summarized in Supplementary Table 1. ****p < 0.0001 at the indicated time points (by Bonferroni multiple comparison). e Based on differentially expressed genes (DEG) between U-IRI and IRI/CL-NX kidneys on day 14 after injury, the top relevant enriched gene ontology (GO) terms for injured proximal tubule (PT) in the U-IRI kidneys are visualized in the dot plots.
recruited monocytes into proinflammatory macrophages, all of which contribute to the early components of reperfusion injury. In models of injury where repair pathways predominate, CD169+ macrophages limit subsequent neutrophil infiltration in the kidney by downregulating intercellular adhesion molecule-1 (ICAM-1) expression on vascular endothelial cells, and neutrophil numbers in the interstitium dramatically decrease after day 3, corresponding to the period when reparative macrophages become prominent. In contrast, intrarenal T cells are not typically seen in large numbers in the first week after IRI, but have been identified in the late stages of IRI where they can promote upregulation of proinflammatory cytokine expression including IL1b, IL6, Tnf, and Ifng. We now show that in a model where atrophy predominates, this late neutrophil and T-cell infiltrate correlates with sustained tubular cell dedifferentiation and the increased expression of multiple homing chemokines by macrophages as well as the infiltrating T cells and neutrophils themselves.

Infiltrating T cells can be activated through antigen-independent mechanisms by inflammatory cytokines and reactive oxygen intermediates as well as through interaction with antigen-presenting cells (APCs). Our results illustrate that in response to the late cell stress induced by inflammatory cytokines, proximal tubular epithelial cells in U-IRI kidneys upregulated MHC class II (H2-Aa, H2-Ab1, H2-Eb1, and...

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**Fig. 9 | Dual depletion of T cells and neutrophils attenuates kidney tubule atrophy.** WT mice were treated as described in Methods with either PBS or a combination of antibodies (Ab)-against Thy1.2 and Ly6G beginning 5 days after unilateral ischemia/reperfusion injury (U-IRI) and sacrificed on day 30. 

- **a** The IRI kidney sections were immunostained with CD3ε and Ly6G. Nine kidneys/group were sectioned and stained, and representative images are shown. Scale bars, 25 μm.
- **b** CD3ε-, CD8α-, and Ly6G-positive areas were quantified using Image J. Data are presented as mean ± SD. n = 9 kidney sections/group. ****p < 0.0001 by unpaired two-tailed t-test.
- **c** Kidney sections from day 30 after U-IRI ± dual T-cell/neutrophil depletion were immunostained with lotus tetragonolobus lectin (LTL, dark gray). Nine kidneys were sectioned and stained and representative images are shown. Scale bars, 1 mm.
- **d** LTL-positive area was quantified from the entire section (left panel) and as a percentage of the section (right panel). Data are presented as mean ± SD. n = 9 kidneys/group. ***p < 0.001, ****p < 0.0001 by unpaired two-tailed t-test.
- **e** Kidney-to-body weight ratios on day 30 following U-IRI ± dual T-cell/neutrophil depletion. Data are presented as mean ± SD. n = 9 kidneys/group. **p < 0.01 by unpaired two-tailed t-test.
- **f** Quantitative RT-PCR analysis for Lrp2 and Slc34a1 was performed on whole-kidney RNA from U-IRI mice ± dual T-cell/neutrophil depletion. Data are presented as mean ± SD. n = 9 kidneys/group. *p < 0.05 by unpaired two-tailed t-test.
 indirectly in vivo47, suggesting that the simultaneous recruitment of neutrophils that increase to express cytotoxins such as perforin, granzymes, and FAS ligand are involved in positive regulation of lymphocyte activation4.

These FR-PTCs were shown to increase late tubule dedifferentiation, sustained injury and atrophy seen following U-IRI. Interestingly, neutrophils that increase expression and collecting duct-vcam1 expression gradually decreased despite the tubular atrophy, whereas renal function after U-IRI49. In addition, the store-operated calcium entry (SOCE) channel, Orai1, participates in the activation of Th17 cells and DCs.

Alternatively, the partial response to T cell and PMN depletion may reflect the dichotomy of T-cell responses to kidney injury48. It has been shown that CCR5 was upregulated in the CD3+ infiltrating T cells, and blockade of CCR5 using a neutralizing antibody in mice protected from injury. However, deletion of T cells and neutrophils did not fully prevent kidney atrophy when compared to the contralateral kidneys and control kidneys. This partial effect may be due to our inability to fully eliminate T cell and neutrophil infiltration (~80% reduction based on immunostaining), or because administration of the anti-Thy1.2 and anti-Ly6G antibodies does not prevent accumulation of macrophages and DCs.

The question that our single-cell sequencing data raised is whether or not the late influx of T cells and neutrophils, occurring after preoperative macrophage activation has waned, is responsible for the late tubule dedifferentiation, sustained injury and atrophy seen following U-IRI. Experiments in which we selectively depleted either T cells alone or neutrophils alone failed to reduce the progressive kidney and tubule atrophy (Supplemental Figs. 25 and 26), but when we depleted both T cells and neutrophils simultaneously there was a significant reduction in the degree of tubule atrophy following U-IRI with preservation of proximal tubule brush border and differentiation (Fig. 9). However, depletion of T cells and neutrophils did not fully prevent kidney atrophy when compared to the contralateral kidneys and control kidneys. This partial effect may be due to our inability to fully eliminate T cell and neutrophil infiltration (~80% reduction based on immunostaining), or because administration of the anti-Thy1.2 and anti-Ly6G antibodies does not prevent accumulation of macrophages and DCs.

Finally, we show that patients who underwent a renal biopsy at the time of AKI exhibit a strong negative correlation between the number of infiltrating neutrophils and T cells present at the time of biopsy and recovery of GFR, green square in a], and a healthy living donor are shown. Scale bars, 50 μm. c Correlation between absolute eGFR increase (ΔeGFR) within 6 months after biopsy and T-cell infiltrate (%) or neutrophil infiltrate (%) at the time of AKI biopsy was determined by nonparametric Spearman correlation coefficient r. d Correlation between relative eGFR increase (fold change) within 6 months (6 m) after biopsy and T-cell infiltrate (%) or neutrophil infiltrate (%) at the time of biopsy was determined by nonparametric Spearman correlation coefficient r.

Fig. 10 | Accumulation of T cells and neutrophils negatively associates with GFR recovery in patients with AKI. a The estimated GFR (eGFR) was determined at reference, biopsy (AKI), and 6-month follow-up (6 m F/U). b Biopsy sections from each patient were immunofluorescence-stained with anti-CD3ε (T-cell marker) and anti-megalin (PT marker, left panel) or anti-CD66b (PMN marker, right panel) and the percentage of nuclei positive for CD3ε and CD66b calculated for all sections. Representative images of biopsies from a patient with full recovery [Case #0252, 150% recovery of GFR, magenta triangle in a], low-recovery [Case #0284, 52%
the subsequent recovery of estimated GFR (Fig. 10). While acknowledge
ing the limitations of eGFR in patients with AKI, our findings are con-
istent with studies showing large numbers of terminally differ-
entiated CD4+ T helper and CD8+ cytotoxic T cells in patients with 
ESRD5, and raise the ultimate question of whether or not these data 
may provide new targets for slowing or even preventing the progres-
sion from AKI to CKD in humans. By analyzing the chemokine-receptor 
interactions among cell types in the injured mouse kidneys, we iden-
tified several potentially important signaling interactions. For instance, 
interactions were identified between infiltrating macrophages and 
dendritic cells through CCL2-CCR2 signaling, and between macro-
phages, DC and T cells via CXCL16-CXCR6 signaling. We recently 
reported that blocking CCL2-CCR2 signaling by knock-out of 
Ccr2 in vivo depletion of T cells and neutrophils 
To deplete T cells and neutrophils, wild-type mice were i.p. injected 
with 200 µg anti-mouse Thy1.2 (CD90.2) antibody (clone 30H12, 
BioXcell) once in 3 days and 200 µg anti-mouse Ly6G (clone IAB, 
BioXcell) (n = 9 mice) every other day or PBS control (n = 9 mice) 
starting on day 5 after U-IRI. The depletion efficacy was confirmed 
at the kidney level using immunohistochemistry against CD3ε, CD4, 
CD8a, and Ly6G, respectively, and qPCR analysis for Cd3e, Cd4, Cd8a, 
and Ly6g, respectively, on the whole-kidney RNA at the end point (day 
30 after U-IRI).

ELISA of serum KIM-1 and NGAL levels 
Serum KIM1 and NGAL concentrations were measured using mouse 
TIM-1/KIM-1/HAVCR and Lipocalin-2/NGAL quantikine ELISA kits (R&D 
Systems) according to the manufacturer’s instructions.

Preparation of single-cell suspension
Euthanized mice were perfused with chilled 3× PBS (10 mL) via the left 
ventricle. Kidneys were harvested, minced into -1 mm³ cubes, and 
digested using Liberase™ (100 µg/mL) and DNase I (10 µg/mL) (Roche 
Diagnostics) for 25 min at 37 °C. Reaction was deactivated by adding 
chilled DMEM with 10% FBS. The solution was then passed through a 
40-µm cell strainer. After centrifugation at 300 × g for 10 min at 4 °C, 
the cell pellet was resuspended in chilled DMEM with 10% FBS and 
passed through another 40-µm cell strainer. The dead cells were 
removed using Dead Cell Removal Kit (Miltenyi Biotec). Cell number 
and viability were analyzed using trypan blue staining (Invitrogen). 
This method generated single-cell suspensions with greater than 80% 
viability.

Single-cell-RNA-sequencing (scRNA-seq) library generation and 
sequencing
scRNA-seq library and sequencing were performed at the Yale Center 
for Genome Analysis (YCGA). Briefly, single cells, reagents and a single 
GeL Bed containing barcoded oligonucleotides were encapsulated 
into nanoliter-sized GeL Bead in Emulsion (GEM) using the GemCode™ 
Technology 10× Genomics. Lysis and barcoded reverse transcription 
of polyadenylated mRNA from single cells were performed inside each 
GEM. The scRNA-seq libraries were finished in single bulk reaction. 
The CDNA libraries were constructed using the 10x Chromium™ Single-cell 
3’ Library Kit. Qualitative analysis was performed using the Agilent 
Bioanalyzer High Sensitivity DNA assay as shown in Supplementary 
Fig. 24. The final libraries from IRI/CL-NX and U-IRI kidneys were 
sequenced on an Illumina HiSeq 4000 sequencer. Cell Ranger version 
5.0.1 was used to process Chromium single-cell 3’ RNA-seq output and 
align the Read to the mouse reference transcriptome (mm10-2020-A), 
al of which were provided by the YCGA.
scRNA-seq data analysis

Downstream data analysis was performed using the Seurat v4.0 R package. The Seurat integration strategy was performed to identify common cell types and enable comparative analyses between IRI/CL-NX and U-IRI kidneys at each time point.14,15 Briefly, all the datasets were first merged for the quality control (QC) analysis. Poor quality cells were defined as unique genes and <500 unique molecular identifier (UMI) counts (likely cell fragment) and >100,000 UMI (potentially cell duplet) were excluded. Cells were excluded if their mitochondrial gene percentages were over 50%. Low-complexity cells like red blood cells with <0.8 log10 genes per UMI counts were also excluded. Only genes expressed in 5 or more cells were used for further analysis. The QC filters resulted in a total of 95,343 cells with a median of 2743 UMI counts per cell at a sequencing depth of 44,719 genes across 95,343 cells. The merged dataset was split, normalized, cell cycle scored, SCTransformed, and integrated using integrated anchors.15 Confounding sources of variation including mitochondrial gene content were removed for downstream clustering analysis.

Principal component analysis (PCA) was performed on the scaled data. The top 20 principal components were chosen for cell clustering and neighbors finding with k-param = 20, perplexity of 30, and resolution of 0.8. The Uniform Manifold Approximation and Projection (UMAP) was used to visualize the single cells in two-dimensional space. Each cluster was screened for marker genes by differential expression analysis based on the non-parametric Wilcoxon rank sum test for all clusters with genes expressed in at least 25% of cells either inside or outside of a cluster. Based on the kidney cell and immune cell lineage-specific marker expression, eighteen cell clusters were identified. Both IRI/CL-NX and U-IRI datasets at each time point could be split from the integrated dataset for differential analyses. The average expression of both IRI/CL-NX and U-IRI cells was plotted using volcano plots. The outliers were used to identify the genes that were differentially expressed between the models in each cluster. All the changes in gene expression were visualized in dot plots or volcano plots. Gene set enrichment analyses were performed using the ClusterProfiler and Gene Ontology (GO) Resource. The potential ligand-receptor interaction analyses were performed using the NicheNet R package3 by linking potential ligands expressed by sender cells to their target genes that were differentially expressed by receiver cells and their corresponding receptors. The ligand-receptor pairings for each cell type were visualized by a chord diagram using the R package circlize.59

Quantitative PCR analysis

Whole-kidney RNA was extracted with an RNeasy Mini kit (Qiagen) and reverse transcribed using the iScript cDNA Synthesis Kit (Bio-Rad). Gene expression analysis was determined by quantitative real-time PCR using an iCycler IQ (Bio-Rad) and normalized to hypoxanthine-guanine phosphoribosyltransferase (Hprt). The primers included previously published sequences,16 and those provided in Supplementary Table 9. The data were expressed using the comparative threshold cycle (∆CT) method, and the mRNA ratios were given by 2−∆CT.

Immunohistochemistry (IHC) and immunofluorescence (IF)

Kidneys were fixed in 10% neutral-buffered formalin and embedded in paraffin. Deparaffinized kidney sections were rehydrated in graded alcohols (100%, 95%, 90%, 80%, and 70%) and microwaved in citrate buffer antigen retrieval for 20 min. Endogenous peroxidase activity quenched by incubation in 3% (vol/vol) H2O2 for 15 min. The sections were incubated overnight at 4°C with the primary antibody. After washing with TBST, the sections were incubated with biotinylated secondary antibody (Vector Laboratories) followed by VECTASTAIN Elite ABC system (Vector Laboratories). DAB (Vector Laboratories) and hematoxylin (Vector Laboratories) were used as the chromogen and the nuclear counterstain, respectively. The primary antibodies were omitted as negative controls. Primary antibodies were as follows: F4/80, CD11c, Ly6G, CD3ε, CD4, CD8α (#70076, #97585, #87048 S, #99940, #25229, and #98941; Cell Signaling Technology, respectively, 1:100 dilution) and primary rabbit polyclonal antibody against megalin (anti-MC22060 original obtained from Dr. Daniel Biemesderfer, Yale Section of Nephrology). Lotus tetragonolobus lectin (LTL) was detected by IHC using primary antibodies against biotinylated LTL (#81325, Vector Laboratories, 1:200 dilution). All the IHC staining slides were scanned using Aperio LV1 Real-time slide scanner and processed using ImageScope software. Six independent fields in cortex and four independent fields in outer medulla were analyzed per kidney section. The percent area of F4/80-, CD11c-, Ly6G, CD3ε, CD4+, and CD8α-positive staining was quantified using IHC Profiler in ImageJ (NIH).

Kidney-specific cadherin (KSP-Cadherin) and UMOD were detected by IF using primary antibodies against UMOD (#sc-20631, Santa Cruz Biotechnology, 1:100 dilution) and primary mouse monoclonal antibody against KSP-Cadherin (clone:4H16/1.66, kindly provided by Dr. Robert Brent Thomson, Yale Section of Nephrology, 1:100 dilution). All the IF staining slides were scanned by Pannoramic 250 FLASH III (HiBio Inc, MA), and the images were processed using CaseViewer software. Megalin and KIM-1 were detected by IF using primary antibodies against megalin (anti-MC22060, 1:100 dilution) and TIM-1/KIM-1/HAVCR (#AF1817, Novus Biologicals, 1:100 dilution). Six to ten random fields in cortex were imaged per kidney section. The percent area of megalin and KIM-1-positive staining was quantified using ImageJ.

Confocal microscopy

Three mice were sacrificed 14 days after U-IRI. The kidneys were perfused with 4% paraformaldehyde (PFA) and embedded in optimum cutting temperature (OCT) compound (Tissue Tek). Kidneys were cryosectioned at 5 μm thickness and mounted on Superfrost slides. Sections were washed with PBS, blocked with 10% normal donkey serum, and then stained with primary antibodies against F4/80 (#MCA497GA, Clone:A3-1, Bio-Rad, 1:100 dilution) and Ly6G or CD3ε (#87048 S and #99940, Cell Signaling Technology, 1:100 dilution). The sections were mounted with VECTASHIELD® HardSet® Antifade Mounting Medium with DAPI (4,6-diamidino-2-phenylindole). The fluorescence images were obtained by confocal microscopy (Zeiss LSM 880).

Western blot analysis

Kidney lysates were fractioned using a RIPA lysis and extraction buffer (Thermo Fisher Scientific) fixed with cOmplete, EDTA-free, protease inhibitor (Roche). Protein concentration was measured using Bio-Rad Protein Assay. Fifty micrograms of protein was separated by SDS electrophoresis using 7.5% polyacrylamide gel and transferred to an Immobilon PVDF membrane (Millipore). Membrane was blocked with 5% non-fat milk in TBST for 2 hrs and probed overnight at 4°C with the primary antibody against TIM-1/KIM-1/HAVCR (#AF1817, Novus Biologicals). After washing the membrane with TBST, it was incubated for 1 h at room temperature with HRP-conjugated secondary antibody (Thermo Fisher Scientific). The membrane was developed using the ECL Detection System (Thermo Fisher Scientific) and imaged using the Odyssey FC Imaging System (LI-COR Biosciences). Next, the membrane was washed with TBS and then immersed and incubated in Restore™ Plus Western Blot Stripping Buffer (Thermo Fisher Scientific) for 15 min. After TBST wash, the same membrane was blocked with 5% non-fat milk in TBST for 2 hrs and probed overnight at 4°C with the primary antibody against GAPDH (HRP Conjugate) (#51332; Cell Signaling Technology). The membrane was developed using the ECL Detection System and exposed to autoradiographic films. The films were scanned on an Epson Perfection V500 Photo Scanner. Densitometry was analyzed using ImageJ. The uncropped scans of all blots were supplied in the Source Data files.
Human kidney biopsy samples were obtained in accordance with the policies of Yale University’s Human Investigations Committee (HIC approval number 11110009286). Written informed consent for research use of patient samples was obtained from each patient by study team. Living donor kidney biopsies were collected under a protocol approved by the Yale University institutional review board (Protocol 2000022915). Informed consent was obtained for all patients before enrollment. Following organ harvest, an 18-gauge core or wedge biopsy was collected intraoperatively prior to organ transplantation. No compensation was provided for the study participation. The biopsied AKI samples obtained between 2015 and 2018 were acquired retrospectively after chart review. All data were deidentified and maintained on a secure database. We estimated GFR (eGFR) using the CKD-epidemiology equation. Ten cases were identified using the following criteria: baseline eGFR ≥60 mL/min/1.73 m², presence of AKI, exclusion of those with GN, DKD, or AIN, availability of biopsy sample, and availability of 6-month follow-up eGFR (8/10 with pathologic diagnosis of ATI, acute tubular injury). CD3e and CD68 cells were detected by IF using primary monoclonal antibodies against CD3e (#NBP2-33357, Novus Biologicals, 1:100 dilution) and CD68 (#305102, BioLegend, 1:100 dilution), respectively, and proximal DAPI + nuclei in the same cortical region. and neutrophils in the entire sample cortex were counted in a blinded Yale Center for Advanced Light Microscopy Facility, and the images cortical regions on the biopsy specimen. All the IF staining sections pathologic diagnosis of ATI, acute tubular injury). CD3 presence of AKI, exclusion of those with GN, DKD, or AIN, availability of biopsy sample, and availability of 6-month follow-up eGFR (8/10 with pathologic diagnosis of ATI, acute tubular injury). CD3e and CD68 cells were detected by IF using primary monoclonal antibodies against CD3e (#NBP2-33357, Novus Biologicals, 1:100 dilution) and CD68 (#305102, BioLegend, 1:100 dilution), respectively, and proximal tubules were visualized by IF using polyclonal antibody against LRP2 (#19700-1-AP, Thermo Fisher Scientific, 1:100 dilution) to identify renal cortical regions on the biopsy specimen. All the IF staining sections were tile-scanned by Zeiss LSM 880 with Airyscan Microscope at the Yale Center for Advanced Light Microscopy Facility, and the images were processed using Zeiss Zen lite software. The number of T cells and neutrophils in the entire sample cortex were counted in a blinded manner and normalized to the total number of cells (identified by DAPI + nuclei) in the same cortical region.

Statistical analysis
The data were expressed as means ± standard deviation (SD). Two-group comparison was performed by Student t-test. Multigroup comparison was performed by one-way analysis of variance (ANOVA) for group mean comparison followed by Tukey’s multiple comparison test for subgroup comparison. Two-group time-course comparison was performed by two-way ANOVA for model comparison to test whether there is a difference between the models and in the time-course, followed by Bonferroni post-tests or Tukey’s post-tests for subgroup comparison (Supplementary Tables 1 and 3). Correlation of gene expression was performed by Pearson correlation coefficient r with two-tailed p-value (Supplementary Tables 4–6). Correlation of T cells/neutrophils and eGFR was performed by nonparametric Spearman correlation coefficient r with two-tailed p-value. All the statistical analysis was performed using Prism 8 (GraphPad Software). A value of p < 0.05 was considered statistically significant.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The single-cell RNA-sequencing data generated in this study have been deposited and publicly available in the GEO database under accession code GSE197626. Source data are provided with this paper as a Source Data file. The remaining data are available within the Article or from the authors upon request. Source data are provided with this paper.

Code availability
The analysis code has been deposited and publicly available in Zenodo at https://zenodo.org/record/6794806#.Yli9Aa2MwJ or from the authors upon request.

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Author contributions
L.X. designed and performed the primary experiments, analyzed the data, and wrote the manuscript. J.G. in part performed single-cell preparation, data interpretation, and edited manuscript. D.G.M. obtained human samples, performed patient chart review, and edited the manuscript. L.G.C. oversaw the project and edited the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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