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Single-Nucleus RNA Sequencing Identifies New Classes of Proximal Tubular Epithelial Cells in Kidney Fibrosis

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Significance Statement (120 words)

Proximal tubular cells (PTCs) are central to normal kidney function, and to kidney regeneration versus organ fibrosis following injury. Single-nucleus RNA sequencing of fibrotic kidneys from mice exposed to aristolochic acid and naïve controls segregated PTCs to five abundant clusters, mapping to S1, S1-2, S2, S2-cortical S3, and medullary S3 segments. Four additional new PTC clusters were present, that showed increased abundance in fibrotic kidneys. The new PTC clusters exhibited gene expression signatures suggesting roles in renal injury responses and fibrosis progression. Pathway analysis revealed metabolic reprogramming, enrichment of cellular communication and cell motility, and immune activation. New PTC clusters also associated with fibrotic signalling to fibroblasts and inflammatory signaling to macrophages.
Abstract

Background: Proximal tubular cells (PTCs) are the most abundant cell type in the kidney. PTCs are central to normal kidney function, and to kidney regeneration versus organ fibrosis following kidney injury. This study determined PTC phenotype in renal fibrosis by single-nucleus RNA sequencing (snRNA-seq).

Method: Kidneys were harvested from naïve mice and mice with renal fibrosis induced by chronic aristolochic acid administration. Nuclei were isolated using Nuclei EZ Lysis buffer. Libraries were prepared on the 10X platform and snRNA-seq completed using the Illumina NextSeq 550 System. Genome mapping was carried out with high-performance computing.

Results: A total of 23,885 nuclei were analysed. PTC’s were found in five abundant clusters, mapping to S1, S1-2, S2, S2-cortical S3, and medullary S3 segments. Additional cell clusters were present (“new PTC clusters”) at low abundance in normal kidney and in increased number in kidneys undergoing regeneration/fibrosis following injury. These clusters exhibited clear molecular phenotypes, permitting labelling as, proliferating, New-PT1, New-PT2 and (present only following injury) New-PT3. Each of these clusters exhibited a unique gene expression signature, including multiple genes previously associated with renal injury response and
fibrosis progression. Comprehensive pathway analyses revealed metabolic reprogramming, enrichment of cellular communication and cell motility, and various immune activations in new PTC clusters. In ligand-receptor analysis, new PTC clusters promoted fibrotic signaling to fibroblasts and inflammatory activation to macrophages.

Conclusion: These data identify unappreciated PTC phenotype heterogeneity, and reveal novel PTCs associated with kidney fibrosis.
**Introduction**

The kidneys excrete waste and maintain homeostasis by blood filtration and subsequent selective reabsorption. The core functional unit of the kidney is the nephron, of which about one million are present in each human kidney. The nephron extends from the glomerulus, which is the site of blood filtration, to the collecting duct system through which nephrons converge to drain urine to the lower urinary tract. Ultrafiltrate is processed to urine via active transport, which underpins reabsorption of water, electrolytes, glucose, amino acids, and other constituents, and is balanced by secretion to achieve homeostasis. Highly specialized, sequential processing of urine occurs in discrete nephron segments, reflecting underlying cellular specialization(1).

The proximal tubules are the site for the majority of the resorptive activity, comprising active transport of an estimated 150 liters per day of solute-rich fluid in a healthy human(2). The scale of proximal tubular resorptive activity makes the kidney a highly metabolically active organ, second only in oxygen consumption to the brain(3). Proximal tubular cells (PTCs) are highly susceptible to ischemic and toxic injury, and their responses are central to progression and recovery in acute and chronic kidney injury of diverse etiology(4). PTCs play
a core role in dictating renal recovery and fibrosis responses following injury, and in chronic kidney disease (CKD)(5).

PTCs are a pre-eminent contributor to the overall cellular composition of the kidney(6). Single-cell RNA sequencing (scRNAseq) is a powerful technique in discriminating cellular characteristics in the kidney and elsewhere. Surprisingly little advance in knowledge about PTC phenotypes has so far come from scRNAseq, where these cells have typically been reported as a single cluster, often numerically much larger than other clusters analyzed in detail(6, 7).

We hypothesized that there may be underappreciated heterogeneity in PTCs, and used a clinically relevant mouse model of chronic aristolochic acid nephropathy (AAN) to investigate the responses of PTCs during kidney fibrosis. We have delineated PTC subtypes found in normal mouse kidney, and further describe novel PTC phenotypes associated with kidney fibrosis.
Methods

Mouse model of chronic aristolochic acid nephropathy (AAN)

Eight 8-9 weeks old C57BL/6 male mice bred by Charles River Laboratories were involved in the experiment. Animals were randomly allocated to control vs. AA injection. Chronic AAN was induced in four mice by intraperitoneal injection of aristolochic acid (2.5 mg per kg body weight, Sigma A5512) twice weekly for two weeks (Fig. 1A). Inflammation and tissue fibrosis developed after repetitive injuries, followed by tissue remodeling and fibrosis in the subsequent two weeks. Another four mice that did not receive intervention were used as naïve controls. Mice were housed with free access to chow and tap water on a 12-hour day/night cycle in a specific pathogen free environment. All mice were euthanized 4 weeks after the first inject of the AAN group. We perfused chilled PBS (1x) via left ventricle before kidney harvest. Both kidneys were harvested and processed, one for single-nucleus RNA sequencing (snRNA-seq) and one for histopathology. Serum creatinine measurement was performed using enzymatic mouse creatinine assay kit (Crystal Chem). Masson's trichrome stain for the formalin-fixed paraffin-embedded sections was performed for each mouse to confirm healthy or fibrotic status. Experiments were performed in line with institutional and
UK Home Office guidelines under the authority of an appropriate project licence.

**Isolation of nuclei**

One quarter of a harvested kidney from each mouse was minced into <2 mm pieces and transferred to a Dounce tissue grinder containing 2 ml of lysis buffer (Nuclei EZ Lysis buffer, Sigma NUC101) supplemented with protease inhibitor (Sigma 5892970001) and RNase inhibitors (Promega N2615 and Life Technologies AM2696)). Kidneys were homogenized and transferred into 50 ml tubes containing 2 ml of lysis buffer, incubated for 7 min on ice, filtered through a 40 am cell strainer, and centrifuged at 500g for 5 min at 4°C. The pellets were resuspended in 4 ml lysis buffer, incubated for another 7 min on ice, then centrifuged again at 500g for 5 min at 4°C. The pellets were then resuspended in 4 ml wash & resuspension buffer (1xPBS, 1.0% BSA, and RNase Inhibitor (Sigma 3335399001)) and filtered through a 20 am cell strainer. Samples were then processed immediately using the 10x Genomics single-cell library preparation protocol.

**Library preparation and RNA sequencing**

Library preparation was performed using Chromium Single Cell 3’ Reagent Kits v3 (10x Genomics) and cDNA quality was evaluated by fragment analysis (5200 Fragment Analyzer)
System, Agilent). RNA sequencing was carried out using the Illumina NextSeq 550 System.

**Bioinformatics analysis**

The sequencing data were processed using the zUMIs pipeline (version 2.3.0)(8). The pipeline was used to first discard reads with low-quality barcodes and UMIs, and then to map reads to the mouse reference assembly (Mus_musculus.GRCm38.95). The barcode-gene matrix generated by zUMIs was analyzed using the R package, Seurat (version 3.1.3)(9, 10).

In Seurat, cells for individual samples were retained if they contained $\geq 400$ genes and genes identified in $\geq 3$ nuclei. After merging the 4 naïve kidneys, and 4 AAN kidneys, cells were filtered again to remove nuclei expressing $\leq 400$ genes or $\geq 7500$ genes, or with mitochondrial gene expression $\geq 10\%$. The feature counts were normalized with scale.factor = 10000. The top 2000 variable genes were identified and scaled. The principal component analysis (PCA) result of the scaled data was obtained. The number of principal components (PCs) included in the downstream analysis was determined by identifying the knee point of the elbowplot generated after running JackStraw procedure. FindNeighbors and FindClusters (resolution 0.8 as default) functions were applied based on previously identified PCs to identify clusters of nuclei. T-Distributed Stochastic Neighbor Embedding (t-SNE) plot was used to
visualize clustering results of the naïve and the AAN dataset. DoubletFinder (version 2.0.2) was then used to exclude potential doublets (11). After doublet removal, the naïve and AAN datasets were integrated (dims = 1:50). The integrated dataset was then scaled and processed with PCA, *FindNeighbors* and *FindClusters* (resolution = 3.0). Final clustering results were visualized using Uniform Manifold Approximation and Projection (UMAP). For differential gene expression (DGE) analysis, we used *FindMarker* command (wilcox method as the default). Significance was defined as a gene with an adjusted p value < 0.05, a ≥ 0.25 average log fold difference between the two groups of cells, and with presence detected in at least 10% of cells in either of the two populations. P-value adjustment was performed using bonferroni correction based on the total number of genes in the dataset. Harmony (version 1.0), which is another batch integration method, was used to validate the pipeline of cell clustering and showed consistent results with Seurat (not shown). (12, 13)

The cell cycle analysis used *CellCycleScoring* to identify cells in the G2/M and S status. Cells with G2M.Score > 0.15 and G2M.Score > S.Score were assigned as G2M status. Cells with S.Score > 0.15 and S.Score > G2M.Score were assigned as G2M status. Cell with G2M Score <0.15 and S.Score < 0.15 were assigned as G1/G0 phase.
For trajectory analysis, we used the R package, Monocle3 (version 0.2.2.0)(14). The metadata from the integrated Seurat object and the top 2000 variable genes from the integrated assay were loaded to monocle. The analysis involved all PTC clusters and the proliferative PTCs identified in proliferative cells using default parameters and removing clusters where low gene numbers were detected (clusters 0 and 1, Supplementary Fig.1). The PTCs were re-clustered by monocle and then the trajectory analysis was performed using learn_graph function. The RNA velocity of each cell was calculated by velocyto.py and velocyto.R (version 0.6) using the spliced and unspliced RNA counts provided by 10X cellranger package (15). Pseudotime analysis was performed on the basis of the RNA velocity result.

We performed combined analysis of single-nucleus RNA sequencing results using this AAN induced CKD dataset and the murine ischemic reperfusion injury induced AKI dataset published by Kirita et al (16). The two datasets were merged and the top 2000 variable genes from the combined dataset were obtained for PCA. We used harmony as the batch integration method. The clustering result of all cells from both datasets was shown by UMAP. For PTC analysis, we analyzed PTCs from both datasets using Seurat integrate data function.

Pathway analysis
We conducted gene set enrichment analysis (GESA) to understand pathways of the new-PT clusters by using an R package, WebGestaltR (version 0.4.3)(17). We evaluated the pathway enrichment in four major functional databases: KEGG, Panther, Reactome and WikiPathways. The recommended False Discovery Rate (FDR) cutoff of 0.25 was used (https://www.gsea-msigdb.org/gsea/index.jsp).

**Ligand-Receptor analysis**

The analysis was based on a well-reviewed dataset with 2557 ligand-receptor pairs from a published study(18). PT S1-S3 were combined to normal PT and fibroblast-1, 2 were combined to fibroblast in ligand-receptor analysis. Ligands and receptors with an average fold-change ≥ 0.25 in the DGE analysis were paired. Pairing results of ligand genes of New-PT1, New-PT2, New-PT3 and receptor genes of fibroblast, immune cells and normal PT were shown using the cross products of ligand/receptor gene expression on heatmap. We also show individual ligand-receptor pairs with the ligands from the New-PT clusters and receptors from fibroblast, immune cell and normal PTCs using a circular visualization tool, circlize (version 0.4.8), in R(19).

**Immunohistochemistry (IHC) and immunofluorescence staining (IF)**
Formalin-fixed paraffin-embedded kidney sections from the AAN and the control group were used for IHC/IF stain. Antigen retrieval at 120°C for 20 minutes was carried out for rehydrated kidney reactions. For IHC stains, UltraVision LP HRP Polymer enhancer system was applied (Thermo Scientific, 12624007) for rabbit/mouse primary antibodies. Sections were blocked with 3% hydrogen peroxide then mouse on mouse block (Vector, MKB-2213-1) where needed, and Ultra V block. The sections were then stained with primary antibodies for Ki67 (Abcam, ab15580) and alpha-smooth muscle actin (Invitrogen, MA5-11547) followed by primary antibody enhancer, HRP Polymer, DAB and hematoxylin counter stain. For Havcr1 IHC stain, sections were blocked with 3% hydrogen peroxide, avidin/biotin block (Vector, SP-2001) and 10% donkey serum, and stained with goat anti-HAVCR1 primary antibody (R&D, AF3689). The sections were then stained with biotinylated donkey anti-goat secondary antibody (Abcam, ab6884), VECTASTAIN ABC-HRP enhancer kit (Vector, PK-4000), DAB and hematoxylin counter stain. For IF stains, sections were incubated with mouse on mouse block where needed and with 10% goat serum. Primary antibodies included anti-SLC4A4 (Invitrogen, PA5-57344), anti-VCAM1 (Invitrogen, MA5-11447), anti FODX1 (LifeSpan BioSciences, LS-B9155-LSP), anti-Akap12 (Abcam, ab49849), anti-WT1 (Sigma, MAB4234),
anti-NCAM1 (Abcam, ab220360), anti-Tenascin C (Abcam, ab108930) anti-P21 (novusbio, NBP2-29463) and anti-HAVCR1. For secondary antibodies we used goat anti-mouse or goat anti-rabbit Alexa Fluor 488 or 594 conjugated antibodies (Invitrogen). After primary and secondary antibody staining, the sections were stained with Hoechst 33342. TrueVIEW autofluorescence quenching kit (Vector, SP-8400-15) was used to diminish autofluorescence. Immunostained tissue slides were visualized and digitized using a confocal laser scanning microscope (LSM800, Carl Zeiss).

**Quantitative image analysis**

Immunostained tissue slides were visualized and digitized using Olympus DP27 5MP color camera attached to a Leica DMLA microscope or by a confocal laser scanning microscope (LSM800, Carl Zeiss). Images were analyzed with the ZEN2012 software (Zeiss) and quantification was performed with Qupath software(20). Pixel classifier was used to detect the collagen (cyan) stain and thresholds were determined to express the positive stain as a percentage of total area (in >10 fields of view per animal, n=3 per group). Quantification of the alpha-smooth muscle actin DAB signal was used to confirm fibrosis in the AAN model.
Nuclear Ki67 DAB signal was used to quantify proliferating cells as a percentage of all hematoxylin stained cells using QuPath’s cell detection algorithm.
Results

Sample processing and quality control

We isolated kidney nuclei immediately following terminal anesthesia and tissue harvesting. Rapid processing of tissue to nuclei was employed, to minimize artefacts arising during processing that may have limited discrimination of PTC sub clusters in previous reports of single cell sequencing from kidney (6, 21). The AAN group had significantly higher serum creatinine (Fig 1B). Histological evidence confirmed the presence of renal fibrosis in mice exposed to aristolochic acid (AA) and its absence in control animals (Fig. 1C-F). Compared to healthy kidneys, AAN kidneys demonstrated a significant increase in collagen deposition (cyan signal in Masson’s Trichrome stain) and alpha-smooth muscle actin DAB signal, indicating renal fibrosis (Fig. 1G, 1H). The AAN kidneys also had a higher percentage of Ki-67 positive cells (Fig 1I). Nuclei of kidneys acquired from mice exposed to AA to induce kidney fibrosis (AAN mice)) and ndeaïve controls were processed using the 10x platform. R packages, Seurat and DoubletFinder were used for data integration, quality control and doublet removal (Supplementary Data 1)(9-11). Downstream analysis included a total of 23,885 nuclei.

Nuclei clustering and cell-type identification
The clustering result was evaluated using UMAP (Fig. 2A, Supplementary Fig. 1). The number of nuclei comprising each cluster and the distribution on the UMAP of nuclei from each individual mouse are shown in Supplementary Data 2 and Supplementary Fig. 2. Nuclei from all mice contributed to every cluster, except for that subsequently labelled “New-PT3”, which was seen in four of four AAN-treated mice and none of four naïve controls.

We next used canonical markers of kidney cell populations to identify major cell types in the kidney: podocyte \((Nphs1)\), endothelial cells \((Flt1)\), mesangial cells \((Igfbp5)\), juxtaglomerular cells (JG cells) \((Ren1)\), PTCs \((Slc34a1)\), proliferative cells \((Top2a\) and \(Mki67)\), descending thin limb \((Aqp1)\), ascending thing limb \((Clcnka)\), thick ascending limb (TAL) \((Slc12a1\) and \(Umod)\), distal convoluted tubule (DCT) 1 \((Slc12a3)\) and 2 \((Slc12a3\) and \(Slc8a1)\), connecting tubule \((Slc8a1)\), principal cell (PC)-outer medullary collecting duct (OMCD) and inner medullary collecting duct (IMCD) \((Aqp2)\), intercalated cells type A \((Atp6v1b1\) and \(Slc4a1)\) and type B \((Atp6v1b1\) and \(Slc26a4)\), transitional epithelium \((Upk1b)\), immune cells \((Ptpre)\) and fibroblasts \((Pdgfrb\) and \(Cfh)\). Fig. 2B shows the expression level and the percentage of expression of the canonical genes per cluster. Regional-specific genes of mouse kidney identified by Ransick \textit{et al.} were used to localize clusters in cortical-medullary and outer-inner
medullary regions.(22) Fig 2C showed regional expression of Cyp2e1 in cortical PTCs and Cyp7b1 in medullary PTCs. The expression of canonical genes for PC-OMCD and PC-IMCD shows the PCs from the two different regions were well-clustered (Fig 2D).

*Clarifying the Pdgfrb+ clusters*

Some functionally distinct renal cell types share lineage and have many common features in their gene expression profiles, which has complicated their recovery in discrete clusters in recent landmark studies(21, 23). These include mesangial cells, JG cells and fibroblasts, which differentiate from Foxd1+ cortical stromal cells and acquire similar genetic features after profibrotic stimulation(24-26). Four clusters closely proximal to one another on UMAP showed expression of the shared markers for mesangial cells, fibroblasts and JG cells, Cfh, Fhl2 and Pdgfrb (Supplementary Fig. 3A, 3B). Mesangial cells were identified from expression of the mesangial-restricted Igfbp5 and Itga8 (http://www.proteinatlas.org)(27, 28) while JG cells were the only cluster expressing Ren1. Two renal fibroblast clusters were identified, of which the Fibroblast-1 cell number increased >200% in fibrotic kidney. We provisionally identified Fibroblast-1 as myofibroblast-containing, on the basis of Meis1 expression(29). Our dataset is limited in this regard by low Acta2 detection (Supplementary Fig. 3B) and further
characterization of renal fibroblast populations may benefit from supplementary technical approaches.

*Analysis of proliferative cells and cell cycle*

One cluster expressed markers of proliferative cells not seen in other clusters, notably including *Top2a* and *Mki67*. Uniquely, this cluster comprised four distinct and separated sub clusters of nuclei. These four sub clusters were located in close proximity to the endothelial, PTC, TAL and fibroblast clusters respectively (Supplementary Fig. 4A). Consistent with this cluster comprising proliferating cells from the adjacent clusters, the four sub clusters also expressed canonical markers for the cell type (e.g. *Flt1*-endothelial, *Slc34a1*-PTC, *Slc12a1*-TAL, *Pdgfrb*-fibroblast, Supplementary Fig. 4B). Cell cycle analysis identified cells in G2/M and S phase, localized mainly to the proliferative and immune cell clusters (Supplementary Fig. 4C). G2/M arrest of PTCs contributes to fibrogenesis after kidney injury(30), and cell number of G2M- and S-Phase PTCs increased in the AAN kidneys.

*Analysis of PTC sub clusters*

The proximal tubule is divided into segments S1, S2 and S3 on the basis of microscopic characteristics. The S1 segment is the longest, and comprises PTCs with extensive apical
microvilli, basolateral infoldings, cytoplasmic complexities, numerous long mitochondria, and a prominent endocytic region(31). These features are present but less evident in the S2 segment, which demonstrates a gradual transition from the S1 segments and additionally displays more numerous peroxisomes and larger secondary lysozymes. The S3 segment is more distinct, comprising simple cuboidal cells without the above features.

Five clusters were identified corresponding to proximal tubular segments S1-S3: “S1”, “S1-2”, “S2”, “S2-cortical S3” and “medullary S3”. These PTC clusters were labelled according to their expression of genes localized to PTC segments in previous studies. \textit{Slc34a1} encodes Na$^+$-Pi cotransporter 2a (NaPi-2a) which is localized in fully differentiated PTCs along all segments except for medullary S3(22, 32, 33). \textit{Slc5a2} encodes a low affinity/high capacity Na$^+$/glucose cotransporter, sodium/glucose cotransporter 2 (Sglt2), which is responsible for 80-90\% of the glucose reabsorption in S1(34). \textit{Slc22a6} encodes organic anion transporter 1 (Oat1) in S2, which plays a critical role in drug and xenobiotic elimination, and has been linked to AAN(33, 35-38). \textit{Slc13a3} encodes sodium-dependent dicarboxylate transporter (NaDC3) in S2, which is responsible for transport of succinate and other Krebs cycle intermediates(39). \textit{Slc7a13}, also known as Agt1, is an amino acid transporter localized to the apical membrane of
the S3 segment and is considered an S3-canonical gene(33, 40). The expression levels of segment-specific solute transporter related genes were reviewed to validate established PTC categories. Relevant genes were identified in the first instance from the study from the Knepper lab, in which bulk deep sequencing was applied to microdissected renal tubules, to identify nephron segment-specific transcriptomes(1). Additional studies are indicated for validation of key anchor genes below. These included markers for S1 (Slc5a2, Slc5a12), S2 (Slc22a6, Ca4, Slc13a3), S3 (Slc5a10, Slc7a13, Atp11a), medullary S3 (Cyp7b1, Slc6a13, Slc34a1(-)), and pan-PTC markers (Slc34a1, Lrp2, Slc4a4) (Fig. 3A, Supplementary Fig. 5A)(1, 22, 41-43). The differences in genes enriched in each PT cluster supported their attribution to specific PT segments (Supplementary Fig 6).

Identification of new PTC clusters

Four additional clusters were located in close proximity to clusters S1, S1-2, S2, S2-cortical S3, medullary S3 on UMAP. First, the largest component of the distributed proliferative cluster. This part of the proliferative cluster, labelled proliferative PTCs, expressed proximal tubular markers and proliferative marker genes, including Ki67 and Cdca3, that Park et al. highlighted as identifying a novel cell type in normal mouse kidney, and that Wu et al.
identified as proliferative-PT in the unilateral ureter obstruction model of renal fibrosis(6, 21).

The other three clusters showed strong signals for genes upregulated in tubules following kidney injury in previous bulk sequencing and other historical approaches (Vcam1, Haver1 and Akap12), demonstrated expression of canonical PTC genes, and were labelled as New-PT clusters. Clusters New-PT1 and 2 were increased in abundance in kidneys undergoing fibrosis following injury, in which circumstances the “New-PT3” were also found. The discrete gene expression signatures of New-PT1-3 clusters included genes associated with renal injury response and fibrosis progression (Fig. 3A, Supplementary Fig. 5A). DGE analysis comparing each individual cluster to all other clusters of the whole dataset detected 786 differentially expressed genes in New-PT1, 637 in New-PT2 and 1318 in New-PT3 (Supplementary Data 3). Analysis of adjusted \( p \)-value and average log-fold change of differentially expressed genes in New-PT 1-3 clusters demonstrated that each had a discrete identifying signature (Fig. 3B-D).

**Characteristics of the new classes of PTCs**

Trajectory analysis was performed to infer paths of cell state transitions within the PTC clusters. Trajectory analysis showed a continuous change in RNA profile in normal PTC’s, ordered along the anatomical axis from S1 to S3 tubular segments (Fig. 4A). RNA velocity
analysis was then employed, in which the RNA processing activity evident in the transcriptome of each cell is used to evaluate transcriptional reprogramming and to predict the future state of cells. Represented graphically by an arrow, the velocity of each cell indicates its rate and direction of transcriptional change. RNA velocity analysis showed that the New PT 1-3 and the proliferative PTCs exhibited strong directional change towards other states, while the normal PT clusters exhibited stable transcriptional profiles, concordant with mature cell phenotypes (Fig. 4B). The data further indicated that NewPT1 is an intermediate cell type, from which cells may differentiate in two directions, “NewPT1 - NewPT2 - proliferative PT - normal PTCs”, and, “NewPT1 - NewPT3” (Fig. 4B). Pseudotemporal ordering was also performed (Fig. 4C.) with time zero set at NewPT1, based on the results of the RNA velocity analysis. An ordered progression of cell states in pseudotime was seen, from New-PT states through anatomically distinct tubular segments (Fig. 4C.).

Genetic profiles suggested that the cells comprising these new PTC classes were dedifferentiated. Expression of Kim-1, Vcam1, CD44, Anax3, Akap12 and Ncam1 has been reported in dedifferentiated PTCs in the literature(21, 44, 45). Current evidence suggests that, following kidney injury, surviving differentiated PTCs can transform to dedifferentiated PTCs
then undergo proliferation and re-differentiation to restore normal proximal tubular morphology and function(32). We identified clusters labelled New-PT1 and New-PT2 as well as proliferative PTCs in normal kidney, suggesting that this dedifferentiation-proliferation-differentiation process occurs in normal circumstances.

Trajectory and RNA velocity analysis suggested that the New-PT1 represented an intermediate cell type bridging normal PTCs, New-PT2, New-PT3 and proliferative-PT. New-PT1 expressed injury markers, including Vcam1, Pdgfb, Pdgfd, Bmp6, Il34, Itgb6, and Itgav. New-PT1 also partially preserved typical PTC markers, Slc4a4 and Slc5a10. Some cells in New-PT1 also expressed Haverl, this subset of Haverl-positive New-PT1 cells were only identified in AAN mice (Supplementary Fig. 5B). The New-PT1 cluster was labelled as “dedifferentiated-intermediate PTCs”. New-PT2 cells expressed genes characteristic of the developing kidney and reactivated during tubular regeneration including Ncam1, Tnc, Tgfr3, Foxd1 and Wt1, and was labelled as “dedifferentiated-regenerating PTC”. New-PT3 cells were detected only in AAN mice, indicating a PTC phenotype specific to progressive kidney fibrosis. Haverl, also known as Kim1, was a prominent marker of New-PT3, which also had prominent expression of Chd2. Apart from genes related to kidney injury and fibrosis, this cluster
expressed Ckd6 and Cdkn1a (also known as p21), associated with cell cycle and also with cell death. The senescence-associated secretory phenotype (SASP) pertains to senescent cells with cell-cycle arrest that remain metabolically active and release senescence-associated proteins(46, 47). SASP-related genes, including Cdkn1a (P21), Cdkn2b (P15), Tp53, Tgfb1, Serpine1 (Pai1), Ccl2 (Mcp1), Cxc1, and Ccn2, were enriched in New-PT3 (Supplementary Fig. 7). This cluster was identified as “dedifferentiated-senescent PTC”.

Microscopic validation for New-PT markers was performed (Fig. 5). Occasional PTC’s exhibiting VCAM-1 and SLC4A4 positive staining were identified as New-PT1 and were present in normal and AAN kidney (Figure 5A, B). HAVCR1 and P21 were selected as NewPT3 markers. Consistent with the snRNAseq dataset, these were not detected in normal kidney, but were seen in a proportion of PTCs in AAN kidneys (Fig 5 C, D), in keeping with the presence of the New-PT3 cluster only following injury. Microscopic evaluation of genes selected as New-PT2 markers identified not only PTC but also glomerular parietal epithelial cell staining. Therefore an extended set of markers, comprising FOXD1, AKAP12, WT1, TNC and NCAM1, were used to validate the New PT-2 cluster. Tubular staining for each marker was evident, with co-expression in rare tubular cells discernible through merged signal in normal
mouse kidney, and in increased numbers of cells in kidneys from the AAN group (Fig 5 E, G, I, K). Co-expression of markers was also evident through merged signal in parietal epithelial cells, consistent with the presence also of parietal epithelial cells in the New-PT2 cluster (Fig 5 F, H, J, L). As expected from its known expression pattern, and consistent with the snRNAseq data, WT1 was also detected in podocytes (Fig 5 J, L). Similarly, TNC was seen in mesangial cells, as well as demonstrating strong interstitial staining in the fibrotic kidney (Supplementary Figure 8).

For further validation, we performed combined analysis of single-nucleus RNA sequencing results using this AAN-induced CKD dataset and the murine ischemia reperfusion injury-induced AKI dataset published by Kirita et al (16). Results of cell clustering and cell-type identification were comparable (Supplementary Fig. 9A) with significantly less injured and severely injured PTCs in CKD (Supplementary Fig. 9B). UMAP of PTCs also addresses the relationship of new PT clusters identified in the two studies. The New PT clusters identified in the current paper cluster in the combined analysis with the failed repair PT’s identified in the Kirita dataset, while the proliferative PT’s cluster with the repairing PT’s. The severely
injured PT cluster identified by Kirita et al in their ischemic AKI model was not found in our dataset.

**Pathway analysis of the new classes of PTCs**

Pathway enrichment analysis was carried out for a further understanding of the molecular interaction network of New-PT1-3. Functional and signaling pathway analysis in KEGG showed a general reduction of metabolic pathways among the three New-PT clusters (Fig. 6) (https://www.kegg.jp/kegg/). Environmental information processing was enriched, including signal transduction (KEGG category 3.2) and signaling molecules and interaction (KEGG category 3.3) pathways, except for the AMPK signal pathway, which acts as a sensor of cellular energy status. Consistent with categorization as "dedifferentiated-senescent PTCs", New-PT3 showed enrichment of cell growth and death (KEGG category 4.2), including cell cycle, apoptosis, p53 signaling pathway and cellular senescence. The cellular community (KEGG category 4.3) and cell motility (KEGG category 4.3) pathways were highly evident in the three new classes of PTCs, corresponding to their dedifferentiated phenotype. Immune system-activation pathways were also enriched in each new-PT cluster. A document summarizing the
pathway analysis results based on KEGG, Panther, Reactome, and WikiPathways databases is provided in Supplementary Data 4.

*Intercellular cross-talk between the new classes of PTCs and other cell types*

Ligands and receptors detected in clusters under analysis with a positive average fold-enrichment were paired, to elucidate intercellular signal transduction networks. For the purposes of this analysis, PT S1-3 were combined to “normal PT”, and fibroblast-1, -2 were combined to “fibroblast” (Fig 7A). We summarized ligand-receptor pairs for ligands from the New-PT clusters and receptors from fibroblast, immune cell and normal PT clusters (Fig. 7B-D, Supplementary Fig. 10, Supplementary Data 5). The New-PT clusters expressed multiple profibrotic signals to fibroblasts, and analysis of the interaction of the New-PTs with immune cells showed several ligand-receptor pairs associated with macrophage activation. New-PTs also expressed signals to EGF receptors, expressed abundantly by normal PTCs. In comparison, normal PTs demonstrated low expression of ligands, when compared to New-PTs. The strongest ligand signal from normal PTs was for netrin, a secreted laminin-related protein that is linked to suppression of inflammatory and injury signals in the kidney. The expression level of ligands and receptors in each cluster were shown in Supplementary Fig.11.
Discussion

Here, we have characterized the cellular composition of adult mouse kidney, comparing healthy animals to those recovering from toxic proximal tubule injury caused by aristolochic acid. Our experimental approach benefitted from refinements introduced by other investigators, demonstrating the benefits of rapidly processing unsorted whole kidney nuclear preparations prior to transcriptomic profiling. We were thus able to delineate major populations of cells that have proved challenging to resolve in previous studies, including mesangial cells, fibroblasts and juxtaglomerular cells. Furthermore, we identified proliferating cells of distinct lineages.

PTCs make a predominant contribution to the wet weight of normal kidney, are highly metabolically active, and play central roles in kidney recovery versus fibrosis following injury. Existing single cell analyses from kidney tissue have grouped PTCs in a single large cluster, often numerically dwarfing all other clusters presented. Here, we have identified five clusters of PTCs abundant in normal kidney tissue, each mapping to proximal tubule segments on the basis of canonical marker expression. These comprised clusters with expression profiles consistent only with a single tubular segment (S1, S2, medullary S3), as well as those enriched in genes found across neighboring segments (S1-2 and S2-cortical S3). These data uncover
significant complexity in PTC phenotype, and provide an expression map of abundantly expressed genes within the major cell phenotypes at single cell resolution.

Kidneys from animals treated with recurrent doses of AA to induce renal injury and subsequent fibrosis contained an increased representation of proliferating cells, immune cells and fibroblasts(48). We further identified three new PTC clusters more prominent in kidney undergoing fibrosis. The first of these, New-PT1, displayed an expression profile intermediate between canonical PTC clusters and clusters New-PT2 and New-PT3, and trajectory analysis further suggested that this cluster may represent PTCs in transition between canonical and these rarer phenotypes. Intriguingly, clusters New-PT2 and New-PT3 demonstrated enriched expression of a panel of genes expressed in proximal tubules following injury. Rather than the diffuse expression in PTC suggested by prior bulk analyses, however, our data reveal restricted expression of specific markers by cluster. Cluster New-PT2 expressed multiple genes associated with tubular regeneration following injury and was labelled dedifferentiating-regenerating PTCs on this basis. Within these regeneration-associated genes, Ncam1 is an early nephron progenitor marker that is also seen in proximal tubules after injury, and may contribute to recovery of PTC function(45, 49). Tnc protects against kidney injury and promotes tubular
regeneration(50). *Tgfbr3* attenuates TGF-beta signaling through processes including glycosaminoglycan modifications of the type I and type II TGF-beta receptors(51). New-PT2 also demonstrated enriched expression of *Foxd1* and *Wt1*, genes reactivated during tubular regeneration processes. Cluster New-PT3 exhibited unique enrichment for *Haverl*, a transcript that is nearly undetectable in normal kidney but occurs promptly after acute kidney injury. *Haverl* expression may be upregulated chronically after kidney injury, and its persistent expression leads to renal fibrosis(52, 53). New-PT3 cells also expressed other genes linked to fibrotic responses in tubular cells, namely *Cdh2* (*N-Cadherin*), which is associated with fibroblast growth factor signaling and cell invasiveness, and several genes linked to the SASP, including *Cdkn1a* (*P21*), *Cdkn2b* (*P15*), *Tp53*, *Tgfβ1*, *Serpine1* (*Pai1*), *Ccl2* (*Mcp1*), *Cxc1*, and *Ccn2*. New-PT3 were accordingly labelled dedifferentiated-senescent PTC’s.

At microscopy, combined expression of selected genes was confirmed in rare PTCs in normal kidney for NewPT1 and New PT2 markers, and at increased number for NewPT1-3, providing validation of identified expression phenotypes. NewPT2 markers were also identified in glomerular parietal epithelial cells, consistent with the presence also of parietal epithelial cells in the NewPT2 cluster. Shared expression of identifying markers has previously
been noted for scattered tubular cells and for a proportion of parietal epithelial cells (44) and as a consequence, the potential for a proportion of parietal epithelial cells to represent tubule-committed progenitor cells has been suggested. (54) Combined analysis with the data of Kirita et al provided further support for discrete, minority PT phenotypes, most notably post-injury. Alignment of the New-PT3 subset of cells identified in the current work with the failed-repair PT’s identified in the work of Kirita et al supports their identification of this as a distinct and important phenotype in fibrosis following renal injury. Definitive establishment of the contribution of this and the other identified PTC phenotypes in renal recovery versus fibrosis after injury will be an important future direction for research.

In summary, these data identify principle cellular phenotypes existing in the proximal tubule of the kidney. They further uncover PTC clusters with discrete, fibrosis-associated phenotypes delineated by unique expression profiles of disease-associated markers.
Data availability

RNA sequencing data and the annotated barcode-gene matrices are available from the ArrayExpress database, accession code E-MTAB-9390.

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Author disclosures

The authors have nothing to disclose.
Supplementary Data Table of Contents

Supplementary data 1. Summary of genome mapping results. Table shows details of nuclei and UMIs detection of each sample before data integration.

Supplementary data 2. Nuclei clustering results. Table shows the number of nuclei comprising each cluster from each individual mouse and the nuclei number left in each filtering step.

Supplementary data 3. Results of differentially expressed gene analysis (DGE). Results of DGE of each cell type by comparing the RNA expression profiles of one type of cell to all other cells of the dataset.


Supplementary data 5. Ligand-receptor interaction. Result of ligand-receptor analysis show pairs of the new-PTs ligands and fibroblast, immune cell and normal PTC (all normal PTC clusters) receptors.
Reference


43. Thiagarajan RD, Georgas KM, Rumballe BA, Lesieur E, Chiu HS, Taylor D, et al.:


