Review Article

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Studying Kidney Diseases at the Single-Cell Level

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Keywords

Single-cell technology · Kidney disease · Immune cell · Kidney organoid · Allograft

Abstract

Background: The kidney is a highly complex organ that performs diverse functions that are essential for health. Kidney disease occurs when the kidneys are damaged and fail to function properly. Single-cell analysis is a powerful technology that provides unprecedented insights into normal and abnormal kidney cell types and will transform our understanding of the mechanism underlying common kidney diseases. Summary: Our understanding of kidney disease pathogenesis is limited by the incomplete molecular characterization of cell types responsible for kidney functions. Application of single-cell technologies for the study of the kidney has revealed cellular heterogeneity, gene expression signatures, and molecular dynamics during the onset and development of kidney diseases. Single-cell analyses of kidney organoids and allograft tissues offer new insights into kidney organogenesis, disease mechanisms, and therapeutic outcomes. Collectively, a better understanding of kidney cell heterogeneity and the molecular dynamics of kidney diseases will improve diagnostic accuracy and facilitate the identification of novel treatment strategies in nephrology.

Key Message: In this review article, we summarize recent single-cell studies on kidney diseases and discuss the impact of single-cell technology on both basic and clinical nephrology research.

Introduction

The kidneys are 2 bean-shaped organs that are responsible for filtering waste products, excess water, and other impurities from the blood and producing urine. The kidneys also regulate pH, salt, potassium levels, and blood pressure; control the production of red blood cells; and activate a form of vitamin D that helps the body absorb calcium \cite{1, 2}. To date, an estimated 850 million people worldwide have kidney diseases, including chronic kidney diseases (CKD), acute kidney injury, kidney failure, and many other diseases \cite{3}. Kidney disease occurs when the kidneys are damaged and cannot perform their function. Damage may be caused by diabetes, high blood pres-
Kidney diseases can lead to other health problems, including weak bones, nerve damage, malnutrition, and heart disease. The current therapeutic strategies for patients still involve kidney transplantation or dialysis, which are costly [2, 3]. A variety of cells in the kidney, including epithelial, mesangial, endothelial, and neuronal cells, as well as a network of immune cells, interact to maintain normal kidney function. Insights into the heterogeneity of healthy kidneys and the process underlying kidney diseases will refine kidney molecular and histopathological phenotype definitions and support the development of new disease classifications. Single-cell technology has potential advantages for the determination of cell subtypes, states, and frequency changes during kidney disease onset and progression [4]. With the rapid development of high-throughput single-cell RNA sequencing (scRNA-seq), comprehensive cellular landscapes of normal kidneys have recently been constructed for precision medicine in nephrology [5–9]. The Kidney Precision Medicine Project (KPMP) was developed worldwide, aiming to obtain human kidney biopsies, create a kidney tissue atlas, define disease subgroups, and eventually identify critical cells, pathways, and targets for novel therapies [10]. On the basis of the relevant single-cell transcription datasets for the kidney, researchers have also analyzed the gene expression signatures of ACE2, TMPRSS2, and SLC6A19 in kidney cell subtypes, which is critical for understanding the pathogenesis of severe acute respiratory syndrome coronavirus 2 [11]. In this review, we will focus on (1) the development and application of single-cell technologies, (2) studying the occurrence and development of kidney diseases using scRNA-seq, (3) a molecular atlas of immune cells in kidney diseases, (4) the application of scRNA-seq on kidney organoids, and (5) insight into kidney allografts by scRNA-seq (Fig. 1).

**Development and Application of Single-Cell Transcriptomic Technology**

Individual cells are fundamental units of life. Single-cell profiling technologies revolutionize our ability to identify cellular compositions, track molecular dynamics, and reveal pathological mechanisms. In the early
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stages, microarray and qPCR technologies were employed for global single-cell gene expression analysis. Ti
etjen et al. [12] monitored the molecular expression profiles of individual neurons and progenitor cells and defined signaling pathways at a distinct developmental stage using a single-cell microarray. In addition, single-cell expression analysis identified several subtypes of developing cells with novel pancreatic genes, providing new insights into pancreas development [13]. Our group developed single-cell qPCR technology and applied it to investigate key regulatory genes involved in mouse blastocyst development and hematopoietic lineage differentiation [14, 15].

With the advent of next-generation sequencing technology, scRNA-seq exhibited obvious advantages over microarrays in distinguishing different isoforms, allelic expression, and new transcripts in single cells at a low cost [16]. In 2009, Tang et al. [17] first reported mRNA whole-transcriptome sequencing of a single cell, demonstrating the complexity of transcript variants in individual mouse blastomeres or oocytes. Smart-seq, which detects full-length transcripts in single cells, was developed in 2012, and by applying it to rare cells, researchers identified candidate biomarkers for melanoma circulating tumor cells. One year later, Smart-seq2 with improved reverse transcription, read coverage, bias, and accuracy was introduced. The latest development of Smart-seq3 greatly increased its sensitivity in detecting thousands of single-cell transcripts at allele and isoform resolution [18–20]. Unlike PCR-based amplification methods, CEL-seq captures efficient single-cell transcriptomics via multiplexed linear amplification. A study of early C. elegans embryonic development by CEL-seq revealed its reproducible and sensitive results at single-cell resolution [21, 22]. The first automatic platform for scRNA-seq is Fluidigm C1, which traps single cells in 96 or 384 chambers using a microfluidic system, followed by cell lysis, reverse transcription, and PCR amplification [23].

Since 2015, with the continuous advent of Drop-seq, inDrop, 10× Genomics, Seq-well, Microwell-seq, SPLiT-seq, and other technologies, single-cell research has completely entered the era of high-throughput, low cost, and automation. Drop-seq and inDrop separate individual cells into nanoliter-sized aqueous droplets and mix them with a unique barcode to label each cell [24, 25]. On the other hand, Cyto-seq, Seq-well, and Microwell-seq achieve high-throughput single-cell mRNA sequencing by capturing one cell and one barcoded bead in microwells [26, 27]. Our group constructed the first mouse cell atlas and human cell landscape at the single-cell level using Microwell-seq [5, 8]. Most recently, more high-throughput and simpler methods, called SPLiT-seq and sci-RNA-seq for scRNA-seq, have been created [28, 29]. These methods use cells or nuclei themselves as the reaction chamber through several rounds of split-and-pool for barcoding to label cells. All cells are then pooled for cDNA PCR amplification and sequencing. Cao et al. [30, 31] profiled the transcriptomes of approximately 2 million mouse organogenesis cells and 4 million human fetal cells using sci-RNA-seq3, providing a global view of mammalian developmental processes. Other single cell technologies covering genome, proteome and epigenome analyses have also flourished; but they are beyond the scope of this review.

Occurrence and Progression of Kidney Disease

The kidney can be affected by a number of prevalent and serious conditions, including acute kidney injury, glomerulonephritis, ascending infection (pyelonephritis), and cancer. In general, our understanding of kidney disease pathogenesis is limited by an incomplete molecular characterization of the cell types responsible for the organ’s specific functions. To narrow this knowledge gap, our group and Park et al. [6] constructed transcriptional maps of healthy mouse kidneys using scRNA-seq. Major subtypes of nephron epithelial cells, including podocytes, proximal tubule epithelial cells, the loop of Henle, distal tubule, and collecting duct cells, were identified [5, 6]. Research characterized a novel transitional cell type in collecting duct cell populations, confirming the findings in a previous scRNA-seq study of interconversion between intercalated and principal cells [32]. Moreover, Ransick et al. [33] profiled male and female adult kidneys and generated an anatomical atlas of mouse nephrons at the single-cell level. Our group has also performed Microwell-seq analyses of human fetal and adult kidney tissues. In addition to epithelial, endothelial, stromal, and tissue-resident immune cells, we found previously undescribed types of S-shaped body cells in the fetal kidney and a new transitional cell type in the adult kidney [8]. A single-cell molecular landscape of the kidney vasculature has revealed specialized expression signatures for building nephrons and has uncovered the pathogenesis of kidney disorders [34]. Furthermore, all cell types of kidney glomeruli from healthy mice and different disease models were identified using single-cell transcriptomic cluster analysis, and novel disease-related genes and regulatory pathways in disease models, such as the activated Hippo
pathway in podocytes after nephrotoxic immune injury, have been detected [35].

Kidney fibrosis is the hallmark of CKD, which affects more than 10% of the world population. In parabiosis models of kidney fibrosis, Kramann et al. [36] confirmed that monocytes contributed a small population of myofibroblasts using scRNA-seq, but that most myofibroblasts were derived from mesenchymal cells. Subsequently, Kuppe et al. [37] profiled approximately 135,000 transcriptomes of cells from the proximal and non-proximal tubules and further verified distinct subtypes of mesenchymal cells as the main contributors to fibrosis in the human kidney. Moreover, in a comparative analysis of healthy and fibrotic kidney single cells, a myofibroblast-specific gene, naked cuticle homolog 2 (NKD2), was identified as a potential therapeutic target in human kidney fibrosis [37]. On the basis of 402 kidney biopsies, urinary fibrinogen has been predicted to be a noninvasive biomarker in patients with CKD [38].

Diabetic nephropathy (DN) is characterized by damage to both the glomerulus and tubulointerstitium. However, relatively little is known about how cell states and frequencies change with disease onset and progression with regard to gene expression. To elucidate glomerular cell gene expression changes in mouse DN, Fu et al. [39] performed scRNA-seq analysis and identified several new potential markers of glomerular cells, including Magi2, Robo2, Ramp3, and Fabp4. Regulation of angiogenesis and migration pathways were altered in endothelial cells, while pathways of regulation of translation and protein stabilization were highly enriched in mesangial cells in diabetic kidney diseases (DKDs). Overall, molecular dynamic changes in endothelial and mesangial cells will help in identifying important pathophysiologic factors contributing to the progression of DN. Chung et al. [35] profiled the single-cell transcriptome of the glomerulus in a mouse diabetes model. Gene expression changes of mesangial cells and podocytes in ob/ob mice compared to control mice were revealed. Proliferation pathways were induced in mesangial cells and cell death-associated pathways were induced in podocytes, coinciding with the cell number proportional changes [35]. Integrated analysis of published DKD scRNA-seq datasets identified 17 hub genes, enriching our understanding of the molecular mechanisms underlying the pathogenesis of DKDs [40].

Moreover, unbiased single-nucleus RNA sequencing (snRNA-seq) has been performed on cryopreserved human diabetic kidney samples, generating 23,980 transcriptomes from 3 control and 3 early DN samples [41]. The results demonstrated cell type-specific changes in gene expression and suggested increased potassium secretion in human DN. A previous study comparing scRNA-seq and snRNA-seq of adult mouse kidneys showed a more effective capture capability for the latter. For instance, glomerular podocytes, mesangial cells, and endothelial cells were captured by snRNA-seq but not by scRNA-seq [42]. The utility of snRNA-seq minimizes enzymatic dissociation artifacts and can be performed on frozen samples, which is expected to lead to broader applications to accelerate studies of pathological mechanisms in diverse kidney diseases.

Precise cellular transcriptomes may reveal a kidney tumor cell of origin as well as the transcriptional trajectories underpinning malignant transformation. Young et al. [43] defined normal and cancerous human kidney cell types from a catalog of 72,501 single-cell transcriptomes. By identifying specific normal cell correlates of renal cancer cells (RCCs), a study provided evidence for the hypothesis that Wilms’ tumor cells were aberrant fetal cells and RCCs might be derived from a little-known subtype of proximal convoluted tubule cells. Hence, scRNA-seq provides a scalable experimental strategy for determining the characteristics of human kidney cancer cells at a precise cellular, quantitative molecular resolution.

**Molecular Atlas of Immune Cells in Kidney Disease**

Immune cells are a foundational component of human tissues, playing a vital role in physiological and pathological metabolism. Indeed, the ability of the immune system to recognize and respond to pathogenic or danger signals, or to malignant cells, is critical. The application of single-cell technologies to the study of immune cells in the kidney has the potential to facilitate a better understanding of the role of the immune system in healthy kidneys and disease pathogenesis, as well as the identification of novel treatment strategies. These efforts are beginning to map the complex immune landscape within the kidney and reveal the relationship between tissue-resident immune cells and their inter-activated neighboring cells.

In 2018, the application of scRNA-seq to mouse kidney tissues provided a comprehensive immune cell landscape, including resident macrophages, neutrophils, B and T lymphocytes, and NK cells [6]. One year later, unprecedented single-cell insights into the spatiotemporal organization of human kidney immune cells were established. A network of tissue-resident myeloid and lymphoid immune cells in fetal and adult kidneys was re-
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Recent studies have shown the role of immune cells in kidney disease models at single-cell resolution. The unilateral ureteral obstruction model is widely used in renal interstitial fibrosis, resulting in marked renal hemodynamic and metabolic changes with infiltration of macrophages and inflammatory cells into the renal interstitium. scRNA-seq has been employed in a reversible unilateral ureteral obstruction model to dissect the myeloid cell landscape during the progression and regression of fibrosis. Conway et al. [48] revealed myeloid cell heterogeneity, and the relative proportions of the subsets changing dynamically, with monocytes being recruited in early injury, Ccr2+ macrophages accumulating in late injury, and a novel Mmp12+ macrophage cluster acting during repair. Ischemia-reperfusion injury (IRI) is linked to inflammation and leukocyte recruitment in kidney diseases and transplantation. An experimental model of IRI was used to assess the functional role of group 2 innate lymphoid cells in the kidney, and the data suggested a redundant function for these cells in renal injury [49]. By applying scRNA-seq to a kidney transplantation IRI model, Kreimann et al. [50] found increased levels of systemic CXCL13 with subsequent infiltration of Cxcr5+ leukocytes. Kirita et al. [51] also applied snRNA-seq to mouse models of IRI to describe detailed cellular responses after injury and identified a distinct pro-inflammatory and pro-fibrotic proximal tubule cell state. Uric acid is the final oxidation product of purine metabolism, with a strong association with renal inflammation across several disease models. For instance, NLRP3 (NOD-, LRR-, and pyrin domain-containing 3) senses the signal of excessive uric acid, and the inflammasome is activated in hyperuricemic nephropathy [52]. However, the specific immune mechanisms of hyperuricemia-induced renal injury remain unclear. Construction of the inflammasome landscape at the single-cell level in hyperuricemia-induced renal injury models is expected in the near future.

Immune infiltrates are present within kidney tumors and are altered by the tumor microenvironment. A previous study revealed a population of tumor-infiltrating macrophages expressing vascular endothelial growth factor A and participating in a complex VEGF signaling circuit in RCC tissues [43]. This study illustrated the ability of scRNA-seq to address questions of tumor ontogeny and to identify putative pathophysiological mechanisms and cell signaling networks that may serve as targets for pharmacological treatment. Collectively, research has highlighted the advancement of scRNA-seq to provide insights into the immune system and the cellular networks that operate in healthy and diseased kidneys.

Application of scRNA-seq to Kidney Organoids

In addition to studies of diseased kidney tissues, kidney organoids have emerged as critical tools to investigate organogenesis and disease mechanisms, and potentially as a replacement tissue source to expedite therapeutic development. In parallel, advances in scRNA-seq have led to more detailed analyses of a variety of cell subgroups and gene expression changes in kidney organoids [53].

In general, a better understanding of kidney embryonic development at the single-cell level is necessary to guide the maturation of kidney organoids. A single-cell transcriptomics study of the human fetal kidney identified 22 cell types and corresponding marker genes [7].
Comparison of different developmental stages revealed continuous molecular dynamics in podocytes. Another single-cell analysis was performed on human fetal kidneys and embryonic stem cell-derived kidney organoids [54]. Comparative analysis of kidney single-cell developmental trajectories showed similar gene expression signatures in vivo and in vitro, except for late mature podocytes, suggesting an incomplete maturation process of podocyte organoids. Transplantation experiments have further indicated that optimization of the mesangial and vascular compartments can be performed using organoid models. A recent study used metanephric mesenchyme and ureteric bud-like cells to generate kidney organoids, revealing improved proximal tubule maturation and reduced off-target cell populations by scRNA-seq [55]. Consistently, Subramanian et al. [56] and Wu et al. [57] profiled human pluripotent stem cell (PSC)-derived kidney organoids by scRNA-seq. Compared to human fetal and adult kidney single-cell datasets, different kidney organoids displayed largely reproducible cell types but variable cell proportions because of off-target cells. Moreover, transcription factor network analysis uncovered the kidney organoid differentiation pathway, highlighting the power of single-cell technologies in characterizing and directing organoid differentiation.

Together, human kidney organoids are useful resources for mining disease models, underlying regulatory mechanisms, high-throughput drug screening, and ultimately regenerative therapy [58, 59]. These studies emphasize the potential utility of scRNA-seq in kidney model systems to elucidate in vivo physiological and pathological processes and provide guidance for future studies on kidney disease diagnosis and treatment.

**Insight into Kidney Allografts by scRNA-seq**

Allografts are one of the most effective methods in the clinical therapy of end-stage kidney diseases. Single-cell technologies offer opportunities to describe kidney cell types and states with accuracy and precision using a human biopsy specimen after allograft transplantation. The first published report of kidney allograft biopsy specimens profiled 8,746 single-cell transcriptomes and defined a diverse inflammatory response. Monocytes formed a nonclassical CD16+ group and a classic CD16- group; endothelial cells presented a resting state and 2 antibody-mediated rejection response states. These findings help us better understand immune rejection in kidney transplantation [60]. In a comparative study of recipient and donor-origin kidney single cells, distinct transcriptional gene expression of kidney biopsy specimens was profiled. Inflammatory activated macrophages and cytotoxic-expressed T cells in recipients were observed [61]. Similarly, Liu et al. [62] analyzed cells from chronic kidney transplant rejection and matched healthy adult kidneys at the single-cell level. Unsupervised clustering analysis revealed that increased numbers of immune cells and myofibroblasts may lead to renal rejection and fibrosis in the chronic kidney transplant rejection group [62]. Remarkably, a recent study profiled the first single-cell atlas of adult human urine and identified a SOX9+ kidney stem/progenitor cell population in the urine. Progenitor cells successfully proliferated and differentiated in vivo and gained some properties of tubular cells, providing a potentially useful resource for future kidney transplantation therapy [63]. Collectively, scRNA-seq technologies provide novel and deep insights into human renal transplantation rejection, ultimately improving diagnostic accuracy and accelerating the adoption of molecular biopsy interpretation.

**Conclusion**

Over the past decade, scRNA-seq has become an indispensable tool for transcriptome-wide analysis of differential gene expression to investigate physiological biology and determine the molecular dysregulation in diseases [64]. Unlike traditional discrete phenotypes, molecular features at the single-cell level may offer a systematic standard for describing kidney disease phenotypes. The application of single-cell technologies to healthy kidney tissues or clinical renal biopsy samples has provided comprehensive kidney molecular atlases and broadens our understanding in the nephrology. Single-cell atlases of the kidney will be an integral part of the international effort on the Human Cell Atlas, which aims at producing a comprehensive and systematic reference map of the human body [65]. In future studies, single-cell ultrahigh-throughput and spatial transcriptomic analyses are expected to expand our understanding of kidney. The integration of multi-omics data will further improve personalized diagnoses and treatments for kidney diseases.

**Conflict of Interest Statement**

The authors declare that they have no competing interests.
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Author Contributions

Mengmeng Jiang drafted the literature. Haide Chen and Guoji Guo provided valuable input into the finalization of the article. All authors read and approved the final manuscript.

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