Epigenetic Changes in the Acute Kidney Injury-to-Chronic Kidney Disease Transition

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Abstract
Previously acute kidney injury (AKI) had been believed to be a transient event, and recovery from AKI had been thought to lead to no consequences. However, recent epidemiological studies have shown that even if there is complete recovery of the kidney function, AKI can eventually result in chronic kidney disease (CKD) and eventually in end-stage kidney disease in the long term. Transition of AKI to CKD is mediated by multiple mechanisms, including aberrant cell cycle arrest and hypoxia. Hypoxia of the kidney is induced by rarefaction of the peritubular capillaries after AKI episodes, and induces inflammation and fibrosis. It should also be noted that epigenetic changes are closely related to hypoxia, and epigenetic changes induced by hypoxia, called “hypoxic memory” can explain the AKI-to-CKD transition in the long term after complete recovery from the initial AKI episode. Targeting hypoxia and subsequent epigenetic changes are promising strategies to block the transition from AKI to CKD.

Introduction
Previously acute kidney injury (AKI) had been believed to be a transient event, and patients who survive an episode of AKI had been thought to regain complete kidney function with excellent long-term prognoses. However, recent epidemiological studies and meta-analyses have shown that even if the kidney function is completely restored, AKI can lead to chronic kidney disease (CKD) and eventually result in end-stage kidney disease in the long term. The AKI-to-CKD transition was also supported by a number of animal studies, which showed development of decreased function and fibrosis of the kidney due to maladaptive repair after recovery from AKI [1].

Mechanisms of the AKI-to-CKD Transition
Mechanisms underlying the AKI-to-CKD transition are a focus of intensive researches. Several mechanisms of the AKI-to-CKD transition have been proposed, such as aberrant cell cycle arrest and hypoxia [2]. Hypoxia is important in the pathogenesis of kidney disease because kidney is physiologically hypoxic primarily due to an arterial-venous diffusional oxygen shunt that permits the kidney to extract no more than 10% of the oxygen delivered by the renal artery [3, 4]. Hypoxia of the kidney is induced by rarefaction of the peritubular capillaries, and some capillaries lack perfusion and are nonfunctional after AKI.

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Hypoxia induces sterile inflammation and fibrosis, and fibrosis, in turn, aggravates hypoxia because fibrosis leads to loss of capillaries and increases the distance between capillaries and resident tubular cells, leading to reduced oxygen diffusion efficiency.

Hypoxia-inducible factor (HIF) serves as a master regulator of adaptive responses against hypoxia. HIF is a dimeric protein complex consisting of alpha (HIF-α) and beta (HIF-β, also referred to as aryl hydrocarbon receptor nuclear translocator) subunits, and enhances the transcription of >100 target genes controlling hematopoiesis (e.g., erythropoietin), angiogenesis (e.g., vascular endothelial growth factor [VEGF]), and anaerobic metabolism (e.g., glucose transporters and glycolytic enzymes).

Levels of HIF are regulated by the oxygen-sensing HIF prolyl hydroxylases (PHD), which belong to the 2-oxoglutarate-dependent protein family. Under normoxic conditions, two conserved proline residues in HIF-α are hydroxylated by PHD. Hydroxylated HIF-α is recognized by von Hippel-Lindau tumor suppressor protein, resulting in immediate proteosomal degradation. In contrast, under hypoxic conditions, PHD cannot hydroxylate HIF-α, thus leaving HIF-α intact. HIF-α is then able to translocate to the nucleus where it binds and forms a heterodimer with HIF-β, which is constitutively expressed irrespective of oxygen tension.

To add more clarity to the contribution of inflammation to kidney injury, we studied the role of vascular adhesion protein-1 (VAP-1) in an animal model of AKI. VAP-1 acts as an adhesion molecule as well as an ectoenzyme that catalyzes oxidative deamination of primary amines, generating hydrogen peroxide (H₂O₂) in the extracellular space. VAP-1 is expressed predominantly in pericytes, which release enzymatically active VAP-1. Inhibition of VAP-1 decreased neutrophil infiltration and ameliorated renal ischemia-reperfusion injury in rats [6]. We also found that CCAAT/enhancer-binding protein δ (CEBPδ), a transcription factor and inflammatory response gene, is a novel regulator of HIF-1, a master regulator of defensive mechanisms against hypoxia. Mechanistically, CEBPδ directly binds to the HIF-1α promoter and enhances its transcription. Notably, CEBPδ is rapidly inducible by inflammatory cytokines, such as IL-1β in an NF-κB dependent manner, which not only increases HIF-1α expression during hypoxia, but is also indispensable for the non-hypoxic induction of HIF-1α [7].

**Fig. 1.** Schematic view of the AKI-to-CKD transition mediated by hypoxic memory. Hypoxia is recorded as epigenetic changes in the cell and has a long-term effect.

**Hypoxic Memory**

Technological advances using high-throughput sequencing have allowed us to determine the expression profile of genes, the binding nature of transcription factors, and histone modifications in a genome-wide manner, and demonstrated that gene expression is intricately regulated by DNA methylation, histone modification, changes in chromosome conformation, long non-coding RNAs, and microRNAs [8]. These epigenetic changes induced by hypoxia are stored in cells as “hypoxic memory” [9], and can induce the AKI-to-CKD transition in the long term after complete recovery from the initial AKI episode (Fig. 1). Studies utilizing animal models of AKI showed that hypoxia-induced epigenetic changes promote proinflammatory and profibrotic gene expression, such as monocyte chemoattractant protein-1, TGF-β1, and collagen. In the same context, diabetic milieu induces long-term consequences potentially via epigenetic changes as “metabolic memory” [10].

JmjC-domain-containing histone lysine demethylases (JmjC-KDMs) also belong to the 2-oxoglutarate oxygenase family like PHD, and they play a key role in epigenetic changes via modulating the methylation levels of histone tails. We previously performed genome-wide analysis of HIF1 binding sites (chromatin immunoprecipitation with deep sequencing) of endothelial cells exposed to hypoxia and observed that lysine (K)-specific demethylase 3A is recruited to the GLUT3 (SLC2A3) locus in an HIF1-dependent manner, demethylates H3K9me2, and induces chromosomal conformational changes so as to upregulate its expression [11].

We also exposed tubular cells to hypoxia and identified novel HIF-1 downstream epigenetic factors that may
play important roles in the kidney. Our RNA-seq identified lncRNAs that are upregulated under hypoxic condition, and chromatin immunoprecipitation-seq analysis demonstrated that HIF-1 also binds to the lncRNAs under hypoxia. We identified a novel lncRNA, DARS-AS1 (aspartyl-tRNA synthetase anti-sense 1), which is upregulated only under hypoxia in an HIF-1-dependent manner and inhibits apoptotic cell death in renal tubular cells [12].

Treatment Targeting the AKI-to-CKD Transition

Treatments that successfully decrease the severity of AKI should block the transition from AKI to CKD because AKI severity is associated with subsequent development of CKD. Studies utilizing HIF knockdown mice showed aggravation of AKI, suggesting that HIF activation can serve as a therapeutic target for AKI [13]. To support this, various pharmacological approaches to PHD inhibition improved AKI models, at least as preventive strategies. Animal studies showed that HIF activation prevented the AKI-to-CKD transition. PHD inhibition not only reduced the severity of I/R-induced AKI, which should effectively suppress the AKI-to-CKD transition, but also blocked the mechanisms of the AKI-to-CKD transition itself, probably by promoting repair of renal I/R injury. Various PHD inhibitors are now in clinical trials of anemia in CKD and are expected to be available at the bedside soon [14]. It remains unknown about which HIF target genes are responsible for providing protection in AKI; however, and we need more in-depth knowledge to apply HIF augmentation strategy in AKI patients.

Treatment to suppress the AKI-to-CKD transition can target the amelioration of capillary rarefaction. Treatment with VEGF-121 attenuated the loss of peritubular capillaries with subsequent suppression of the AKI-to-CKD transition and fibrotic changes in a rat I/R injury model, although VEGF-121 did not affect acute kidney damage. Another angiogenic factor, angiopoietin-1, also preserved peritubular capillaries and blocked the AKI-to-CKD transition. Thus, the preservation of peritubular capillaries early in the AKI process leads to the maintenance of oxygen levels in the kidney, protecting the kidney from the AKI-to-CKD transition.

Another fascinating approach against the AKI-to-CKD transition is to target epigenetic changes directly [15]. Recent studies targeting a histone methyltransferase that induces histone H3 lysine 27 trimethylation showed that pharmacologic inhibition of the histone methyltransferase improved kidney fibrosis in a model of unilateral ureteral obstruction [16]. We also found that pharmacologic intervention targeting histone modification can improve kidney fibrosis in a model of the AKI-to-CKD transition (Mimura, Hirakawa, Tanaka, Nangaku: manuscript in preparation). However, specificity of genomic sites targeted by epigenetic intervention can be an issue, and epigenetic therapies in kidney disease need more investigations in the future.

Summary

Hypoxia serves as a key player in AKI pathophysiology and is the final common pathway from CKD to ESKD. Hypoxia also plays an important role in the AKI-to-CKD transition. Epigenetic changes induced by hypoxia in the kidney are likely to be crucial in the pathophysiology of the AKI-to-CKD transition as “hypoxic memory.” Therapeutic strategies targeting hypoxia, such as HIF activation, and those targeting epigenetic changes will be effective in blocking the AKI-to-CKD transition.

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Disclosure Statement

The authors have no conflicts of interest to declare.

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