The Nucleotide Oligomerization Domain-Like Receptors in Kidney Injury

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Key Words
Inflammation · Kidney disease · NLRP3 · NOD2 · Pattern recognition receptors

Abstract
Background: Inflammation is a hallmark of almost all forms of renal injury and the activation of the innate immune system is of importance in the development of many kidney diseases. Pattern recognition receptors (PRRs) act as sensors of the innate immune system to detect pathogen- or damage-associated molecular patterns, which initiate immune responses to resolve infections and repair damaged tissues. Abnormalities in PRR activation will lead to excessive inflammation. Summary: Nucleotide oligomerization domain (NOD)-like receptors (NLRs) are recently identified intracellular PRRs that are essential to innate immune responses and tissue homeostasis. A better understanding of the function of NLRs will provide unexpected opportunities to develop new therapies for kidney diseases by modulation of the innate immune system. Key Messages: NLRs are constitutively expressed in the kidney and emerging evidence has shown that activation of NLRs plays an important role in the pathogenesis of renal injury. Among NLRs, NOD2 and NLRP3 inflammasome are the best characterized members in the kidney. In this review, we summarize current knowledge about the pathological mechanisms that are related to NOD2 and NLRP3 inflammasome in various kidney diseases by their canonical and non-canonical effects and discuss the opportunities of pharmacological targeting of NLR-mediated signaling pathways at multiple levels for the treatment of renal disease.

Introduction

Emerging evidence has implicated that the activation of innate immune signaling and inflammatory responses are of importance in the pathogenesis of renal disease. In contrast to the adaptive immune system, the innate immune system plays a key role as a first response to pathogens or tissue injury [1–3], which is initiated by pattern recognition receptors (PRRs) in the recognition of pathogen-associated molecular patterns (PAMPs) and damage-associated molecular patterns (DAMPs) that are independent of the clonal expansion of antigen-specific lymphocytes [4]. Although the activation of PRRs can induce the recruitment of immune cells, initiate tissue repair and activate adaptive immune responses, inappropriate or chronic PRR activation results in excessive inflammation and exacerbates tissue damage [5–7]. PRRs are classified into several subfamilies, which are composed of C-type lectin-like receptors, Toll-like receptors,
Nucleotide oligomerization domain (NOD)-like receptors (NLRs), retinoic acid inducible gene I-like receptors and absent in melanoma 2-like receptors [8]. Among them, NLRs are recently identified intracellular PRRs [9, 10] and 22 members have been identified in humans and 34 in mice which contain a highly conserved nucleotide binding and oligomerization (NACHT) domain that is required for self-oligomerization of the proteins upon ligand recognition, a C-terminal leucine-rich repeat that acts as a sensing domain, and an N-terminal effector domain that mediates downstream signaling such as caspase activation and recruitment domain (CARD) or a pyrin domain (PYD) [11] as shown in figure 1. Among them, NOD1/2 can activate NF-κB and MAPK signaling pathways in response to peptidoglycan-related molecules [12], resulting in the transcription of a large number of proinflammatory factors and mediating inflammatory processes to establish an appropriate immune response [13]. Another important class of the NLR family, such as NLRP1, NLRP3, NLRP6 or NLRP12, can oligomerize to form an inflammasome with the PYD-CARD adaptor protein ASC (apoptosis-associated speck-like protein containing a CARD) and caspase-1, which are associated with a wide range of physiological and pathological processes by activation of caspase-1 [14]. Activation of caspase-1 promotes the processing and secretion of IL-1β and IL-18 contributing to the inflammatory response [15]. NLRs are not only expressed in immunocytes, but also widely distributed in various renal parenchymal cells, which induce innate immune responses by the recognition of exogenous PAMPs or endogenous DAMPs. A growing number of studies have demonstrated that NLRs are involved in the pathogenesis of various kidney diseases [16–20]. Among the NLRs, NOD2 and NLRP3 inflammasome are the best characterized members in the kidney. In this review, we focus on NOD2 (fig. 2) and NLRP3 (fig. 3) and their canonical and non-canonical effects in acute kidney injury (AKI) and chronic kidney disease (CKD) and discuss the opportunities of targeting of NLR-mediated signaling pathways at multiple levels for the treatment of renal disease.

**NLRs and AKI**

AKI is a common complication associated with high morbidity and mortality, which is often induced by ischemia/reperfusion injury (IRI), sepsis or toxins. AKI involves renal inflammation related to the activation of innate immunity. NLRs are constitutively expressed in the kidney and emerging evidence has shown that the activation of NLRs plays an important role in the pathogenesis of AKI [21].

**Renal IRI**

One of the most common cause of AKI is renal IRI, which mainly attributes the organ damage to the inflammatory and oxidative stress responses induced by renal ischemia and subsequent reperfusion. Numerous studies have indicated that inflammatory processes engaging both innate and adaptive immune responses are responsible for the initial renal injury such as tubular apoptosis or necrosis and mediate long-term structural changes including interstitial fibrosis or repair [22, 23]. Shigeoka et al. [24] for the first time reported that NOD1 and NOD2 were present in renal tubular epithelial cells, and that gene deficiency of NOD1 or NOD2 protected against renal IRI and the simultaneous deficiency of NOD1 and NOD2

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**Fig. 1.** Classification and molecular structures of NLRs. AD = Activator domain; BIR = baculovirus inhibitor of an apoptosis protein repeat; FIIND = function to find domain; LRR = leucine-rich repeat; NACHT = NAIP (neuronal apoptosis inhibitor protein), C2TA (MHC class 2 transcription activator), HET-E (incompatibility locus protein from Podospora anserina) and TP1 (telomerase-associated protein); NAD = NACHT-associated domain.
provided even greater protection, which was associated with reduced tubular epithelial apoptosis and inflammation. A recent study from our group identified that deficiency of progranulin (PGRN), an autocrine growth factor, exacerbated renal injury after ischemia/reperfusion and treatment of mice with recombinant PGRN resulted in a dramatic reduction in renal dysfunction. Interestingly, we observed an increase in NOD2 expression after renal IRI and a negative correlation between PGRN and NOD2 expression in the kidney. Considering that NOD2

Fig. 2. Summarized recent findings on the role of NOD2 and its canonical and non-canonical effects in kidney injury.

Fig. 3. Summarized recent findings on the function, activation and canonical and non-canonical effects of NLRP3 in kidney injury. EMT = Epithelial-mesenchymal transition.
initiates inflammatory responses largely dependent on NF-kB activation [25], we further detected the effect of PGRN on NF-kB and found that NOD2-dependent activation of NF-kB was also negatively regulated by PGRN. Together, these results suggest that PGRN may be an innovative therapeutic strategy for treating patients with AKI by targeting NOD2-mediated inflammatory responses [23].

NLRP3 has been demonstrated to contribute to renal IRI [26, 27]. It was found that NLRP3 and adaptor protein ASC were highly expressed in renal tubular epithelium, and the absence of NLRP3, but not ASC or the downstream inflammasome targets, protected from renal IRI, indicating that NLRP3 contributes to renal IRI by a direct effect on renal tubular epithelium, which is independent of inflammasome-induced proinflammatory cytokine production [26]. In addition, a very recent study by Baker et al. [28] explored the role of NLRP3 in the repair phase of renal ischemia/reperfusion and investigated the contribution of leukocyte- versus renal-associated NLRP3 by studying bone marrow chimeric mice. It was found that although both renal- and leukocyte-associated NLRP3 were detrimental to renal function after renal IRI, this was through different mechanisms. Leukocyte-associated NLRP3 was associated with increased tubular epithelial apoptosis, whereas renal-associated NLRP3 impaired the tubular epithelial repair response, suggesting NLRP3 as a negative regulator of resident tubular cell proliferation in addition to its detrimental role in renal fibrosis and inflammation.

**Toxin-Induced AKI**

The kidney is susceptible to toxin injury. Cisplatin is a major course of toxin-induced AKI, which triggers the apoptosis of renal tubular epithelial cells followed by inflammation and fibrosis. In our previous studies, we found that PGRN protected against cisplatin-induced AKI, which was also associated with NOD2-mediated signaling (unpublished data). Interestingly, Kim et al. [27] provided evidence showing that NLRP3-deficient mice were protected against ischemic but not cisplatin-induced AKI. They found that although in the kidneys from mice with cisplatin-induced AKI, inflammasome components ASC and caspase-1 was upregulated, the increase in caspase-1 in kidney and proximal tubules was not associated with an increase in NLRP3 protein. Furthermore, they noted that NLRP3-deficient mice were not protected against cisplatin-induced AKI, despite previous studies having demonstrated that caspase-1 is a mediator of cisplatin-induced AKI.

**Sepsis-Induced AKI**

AKI is one of the most important factors determining morbidity and mortality in the prognosis of sepsis. Unlike the pathogenic mechanisms of other causes of AKI, sepsis is associated with an entire orchestra of cellular mechanisms, which potentiate each other and ultimately induce clinical AKI. The microcirculation is the important physiological compartment where these mechanisms exert their integrated and deleterious action [29]. Although the role of NLRs in renal injury in response to sepsis is not well known, emerging studies have reported the participation of NLRs in sepsis-induced AKI. A study from Cao et al. [30] investigated the role of NLRP3 inflammasomes in renal injury in a cecal ligation and puncture model of sepsis-induced AKI. It was found that cecal ligation and puncture upregulated NLRP3, ASC and caspase-1 expression and caspase-1 activation in the kidney and that genetic deletion of NLRP3 reversed the cecal ligation and puncture-induced increases in creatinine and neutrophil infiltration. Moreover, very recent studies have also reported that mangiferin [31] and exogenous carbon monoxide [32] attenuated sepsis-induced AKI via inhibition of NLRP3 inflammasome activation.

**NLRs and CKD**

Inflammation is a hallmark of almost all forms of CKD and the innate immune system participates in many inflammatory processes during the development of CKD. Several PRRs have been identified as contributing to the development of a CKD-specific proinflammatory microenvironment. In tissue from human renal biopsies, a wide variety of CKD exhibited increased expression of NOD2 and NLRP3 mRNA, including IgA nephropathy, focal segmental glomerulosclerosis, lupus nephritis, minimal change disease and diabetic nephropathy, which correlated with renal function [21, 33], strongly supporting the role for NLRs in CKD. In this section, we review the current knowledge regarding NOD2 and NLRP3 inflammasome signaling and outline existing evidence on their functional roles in CKD.

**Renal Interstitial Fibrosis**

Progression of renal injury ultimately leads to renal interstitial fibrosis, which is characterized by activation and proliferation of renal interstitial fibroblasts as well as accumulation of extracellular matrix components. During the development of renal fibrosis, inflammatory responses are induced and involved in this process [34].

NOD-Like Receptors and Renal Function

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The unilateral ureteral obstruction model is usually employed to explore the mechanisms of renal fibrosis. Compared with wild-type mice, NLRP3-deficient mice had less tubular injury, inflammation and fibrosis after unilateral ureteral obstruction, accompanied by a reduction in caspase-1 activation and secretion of IL-1β and IL-18. Furthermore, bone marrow chimeras revealed that NLRP3 mediated the injurious and inflammatory processes in both immune and non-immune cellular compartments [33]. Additionally, Wang et al. [20] also reported that NLRP3 promoted TGF-β signaling and Smad activation in epithelial cells and that the effect of NLRP3 on TGF-β1 signaling was independent of the inflammasome components. They found that NLRP3 expression was increased in response to TGF-β1 and associated with epithelial-mesenchymal transition. Furthermore they demonstrated that TGF-β stimulation not only increased NLRP3 expression in a Smad3-dependent manner, but that NLRP3 and ASC were required for TGF-β-mediated Smad2 and Smad3 phosphorylation, supporting a novel role for NLRP3 in promoting TGF-β signaling and Smad activation in epithelial cells independent of the inflammasome [20]. On the contrary, Pulskens et al. [35] reported the non-canonical effects of NLRP3 following progressive renal injury induced by unilateral ureteral obstruction. They found that the deficiency of NLRP3 led to early tubular damage and interstitial edema, suggesting a protective role of NLRP3 against progressive renal injury by preserving the vascular barrier and epithelial integrity in the kidneys.

In fact, proteinuria is not only a common feature of CKD, but also an independent causative factor for the development of tubulointerstitial inflammation and fibrosis [36, 37]. Recent studies have indicated that the NLRP3 inflammasome contributes to albumin-induced lesions in tubular cells [38, 39]. Liu et al. [40] demonstrated that endocytic receptor-lysosome-mediated albumin reabsorption was involved in the tubular cell activation of NLRP3 inflammasome and tubulointerstitial inflammation. Moreover, studies from Zhuang et al. reported that NLRP3 inflammasome mediated albumin-induced renal tubular injury through impaired mitochondrial function [41] and further provided direct evidence indicating the important role of the albumin-NLRP3 inflammasome axis in mediating the impairment of renal tubular tight junctions and integrity [42]. Additionally, Fang et al. [43] also provided evidence showing the involvement of endoplasmic reticulum stress in albuminuria-induced inflammasome activation in renal proximal tubular cells.

Diabetic Nephropathy

Diabetic nephropathy is one of the major microvascular complications of diabetes mellitus and the most common cause of end-stage renal disease. Dysregulation of the innate immune response via NLRs has been implicated in the development and progression of diabetic nephropathy [4]. We have identified the upregulation of NOD2 in both renal parenchymal cells and infiltrating immune cells from renal biopsies of diabetic patients and high-fat diet/streptozotocin-induced diabetic mice. We further found that NOD2 deficiency ameliorated diabetic renal injury. Evidence for a functional contribution of NOD2 expression to renal function came from an in vitro study showing that NOD2 activation induced ERK1 and ERK2, JNK and p38 MAPK signaling as well as NF-κB, which was associated with high glucose-induced podocyte inflammatory responses and apoptosis. Interestingly, we also found that NOD2 activation induced podocyte insulin resistance, impaired insulin-induced glucose uptake and reduced nephrin expression in podocytes [21]. These results provided direct evidence on the contribution of NOD2 canonical and non-canonical effects in diabetic nephropathy [44]. Furthermore, we found that human antigen R (HuR), a RNA-binding protein which regulates mRNA cargos that contain adenylate-uridylate-rich elements (AREs) in the 3′-untranslated region [45], contributed to NOD2 mRNA stability in diabetic nephropathy [46]. Among four AUUUA regions (ARE1–4) of the 3′-untranslated region of NOD2, HuR can directly bind to ARE4 of NOD2 to enhance NOD2 mRNA stability and expression. Regarding the regulation of HuR activity in the kidney, NADPH oxidase-mediated oxidative stress has been reported to be associated with HuR translocation and activity [47, 48]. NOX4 is an important subunit of NADPH oxidase to produce renal reactive oxygen species (ROS) and deficiency of NOX4 results in renal protection from glomerular injury in diabetic mice [49, 50]. We further found that NOX4-mediated redox signaling contributed to the expression and translocation of HuR and NOD2 mRNA stability [46].

In addition to NOD2, NLRP3 inflammasome-mediated inflammatory responses have also been implicated in diabetic nephropathy. In patients with type 2 diabetes-associated nephropathy, NLRP3 expression was significantly increased in renal tubular epithelial cells, which positively correlated with urinary IL-1β and IL-18 levels [51]. Wang et al. [52] reported that the NLRP3 inflammasome was activated in the kidney of streptozotocin-induced diabetic rats and that suppression of
NLRP3 inflammasome activation significantly reduced renal inflammation and improved renal function. Baker et al. [53] further demonstrated that NLRP3 plays a key role in diet-induced nephropathy and renal cholesterol accumulation. It was found that NLRP3 deficiency ameliorated fructose-induced renal injury by reduced renal inflammation, fibrosis, albuminuria and hyperuricemia. Mechanically, under diabetic conditions, NLRP3 was activated by ROS or extracellular ATP [54]. Hyperglycemia induced the expression of the ATP receptor P2X4 in renal tubular epithelial cells of patients with type 2 diabetic nephropathy, which correlated with IL-1 cytokine release. ATP-P2X4 signaling was further found to mediate high glucose-induced activation of the NLRP3 inflammasome and to regulate IL-1 family cytokine secretion, resulting in interstitial inflammation [51]. Regarding the regulation of NLRP3 inflammasome by ROS, Gao et al. [55, 56] recently found that hyperglycemia-induced NADPH oxidase activation was driven by thioredoxin-interacting protein (TXNIP) which subsequently triggered NLRP3 inflammasome activation in podocytes and ultimately led to podocyte injury. A recent study also indicated that thrombomodulin domain 1 ameliorated diabetic nephropathy in mice via anti-NF-κB/NLRP3 inflammasome-mediated inflammatory responses, enhancement of NRF2 antioxidant activity and inhibition of apoptosis [57].

More interestingly, although it is generally accepted that hyperuricemia-induced NLRP3 activation of macrophages contributes to the progression of diabetic nephropathy [58], recent studies have unraveled the role of the activated NLRP3 inflammasome in glomerular resident cells and addressed the importance of NLRP3 inflammasome activation in non-myeloid-derived cells in diabetic nephropathy [59, 60]. It was found that abolishing NLRP3 or caspase-1 expression in bone marrow-derived cells failed to protect mice against diabetic nephropathy, as evidenced by the fact that albuminuria and mesangial expansion in db/db mice transplanted with NLRP3−/− or caspase-1−/− bone marrow increased to the same extent as those in control db/db mice. Conversely, NLRP3−/− mice were protected against diabetic nephropathy despite transplantation of wild-type bone marrow, indicating that the NLRP3 in renal resident cells significantly contributed to the pathogenesis of diabetic nephropathy. Further studies are required to use cell-specific deletion of the NLRP3 inflammasome to explore the role of inflammasome activation in glomerular and tubular epithelial cells individually.

Hyperhomocysteinemia-Induced Renal Injury

Hyperhomocysteinemia is one major metabolic disorder of amino acids, which is a medical condition characterized by an abnormally high level of homocysteine in the blood, conventionally described as >10 μmol/l. Hyperhomocysteinemia is considered an important independent risk factor in the progression of end-stage renal disease and in the development of cardiovascular complications related to end-stage renal disease [61, 62]. In our study, we found that NOD2 deficiency ameliorated renal injury in mice with hyperhomocysteinemia. We further discovered the non-canonical effects of NOD2 in mediating Ca2+ signaling showing that NOD2 regulated transient receptor potential cation channel 6 expression and activity via nphrin, resulting in intracellular Ca2+ release and podocyte cytoskeleton rearrangement and apoptosis. These results indicate that NOD2-mediated Ca2+ signaling is one of the critical signal transduction pathways that links innate immunity mediator NOD2 to homocysteine-induced podocyte injury. In experimental mice with hyperhomocysteinemia, Zhang et al. [63] found that NLRP3 inflammasome formation and activation in glomerular podocytes were detected at an early stage and that homocysteine-associated albuminuria, foot process effacement of podocytes, loss of podocyte slit diaphragm molecules and glomerulosclerosis at the late stage were significantly improved by inhibition of local ASC or caspase-1, indicating that NLRP3 inflammasome activation is an important molecular mechanism triggering podocyte injury and ultimately resulting in glomerulosclerosis. Further studies demonstrated that NADPH oxidase-mediated oxidative stress was importantly involved in the switching on NLRP3 inflammasomes in podocytes, which led to the downstream recruitment of immune cells, ultimately resulting in glomerular injury [64]. An important factor links changes in oxidative stress to NLRP3 activation is TXNIP, a negative regulator of the antioxidant thioredoxin. When ROS accumulates, TXNIP can sense ROS and time-dependently dissociate from thioredoxin to bind with NLRP3, leading to inflammasome formation and activation. Abais et al. [65] reported that inhibition of TXNIP prevented homocysteine-induced TXNIP protein recruitment to form NLRP3 inflammasomes and reduced caspase-1 activity in glomeruli of mice with hyperhomocysteinemia, indicating that TXNIP binding to NLRP3 is a key signaling mechanism for homocysteine-induced NLRP3 inflammasome activation.
Renal Injury Associated with Hypertension

Hypertension is now considered a chronic, low-grade inflammatory disease, with the kidney representing a major site of this inflammation. Although studies on NLRs in renal injury associated with hypertension are very limited, a very recent study by Krishnan et al. [66] highlights the crucial role of inflammasome activity in one kidney/deoxycorticosterone acetate/salt model of hypertension in mice. They found that deoxycorticosterone acetate/salt-induced hypertension in mice was associated with increased renal expression of NLRP3, ASC, caspase-1 and IL-1β and further demonstrated that renal inflammation and fibrosis in this model were dependent on inflammasome activity.

Crystalline Nephropathies

Crystals are particles of endogenous inorganic or organic composition that can trigger renal injury when deposited or formed inside the kidney [67]. NLRP3 inflammasome-mediated IL-1β secretion is recognized to be the essential pathophysiological element of crystal- and particle-induced inflammation and has been demonstrated to contribute to crystalline nephropathies. Mulay et al. [68] observed that calcium oxalate crystals induced renal inflammation by NLRP3. They found that calcium oxalate crystals triggered IL-1β-dependent innate immunity via the NLRP3/ASC/caspase-1 axis in intrarenal mononuclear phagocytes and directly damaged tubular cells, leading to the release of the NLRP3 agonist ATP. Consistently, in an animal model of progressive oxalate nephropathy by feeding mice a diet high in soluble oxalate, NLRP3 expression was significantly upregulated in the kidney and NLRP3-deficient mice were completely protected from the progressive renal failure and death that occurred in wild-type mice fed the diet high in soluble oxalate, which was associated with NLRP3-mediated inflammation rather than oxalate homeostasis by intestinal handling [69]. Further studies are needed to clarify whether all crystalline nephropathies such as adenine or urate nephropathy share the same mechanisms.

Summary and Future Perspectives

The NLR family is becoming increasingly recognized as integral to the pathogenesis of many renal diseases and their complications. This review highlights the current findings on NOD2- and NLRP3-mediated canonical and non-canonical signaling pathways in AKI and CKD (fig. 2, 3). Despite remarkable progress in NLRs, there are still numerous aspects that need to be understood in the kidney. First of all, in the kidney, the molecular mechanisms for individual PRRs to induce pleiotropic outcomes remain largely unknown. In addition, controversies exist on the certainty of detrimental or beneficial effects of some NLRs in different disease states or different experimental animal models. For example, a recent study indicates a protective role of NLRP3 against progressive renal injury by preventing early renal interstitial edema and vascular permeability in unilateral ureteral obstruction [35]. Second, much of our knowledge of NLRs is limited to experimental models, and it is thus necessary to elucidate their roles in patients under clinical investigation. Third, novel and selective NLR inhibitors are urgently needed. Although scientific progress in the field of NOD inhibitors, including the recently reported selective inhibitors of NOD1 and NOD2, has been made [70], the therapeutic potential of pharmacological modulation of NOD1 and NOD2 signaling needs to be further investigated. In addition, strategies for targeting the NLRP3 inflammasome pathway have been limited to inhibitors of IL-1β, specific inflammasome inhibitors are not yet available. Collectively, a better understanding of the function of NLRs and the development of NLR inhibitors will provide unexpected opportunities to develop new therapies for kidney disease by modulation of the innate immune system.

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Conflict of Interest Statement

All authors declare that they have no competing interests.

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