COVID-19 and the kidney: what we think we know so far and what we don’t

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Abstract
The novel coronavirus disease infection (COVID-19) outbreak that was declared a global pandemic in March 2020 had led to an internationally variable but concerning incidence of COVID-associated acute kidney injury (AKI), with prevalence reported as high as 46% in large cohorts of hospitalized patients. Variability in AKI may be explained by differences in traditional risk factors for AKI, heterogeneity among patient cohorts, and differences in racial and ethnic groups. Further, AKI requiring kidney replacement therapies (KRT) has been associated with increased mortality. Proposed mechanisms of kidney injury include direct viral-induced tubular or glomerular injury, sepsis-associated AKI, and thrombotic disease. Kidney pathology include acute tubular injury, glomerular fibrin thrombi, pigmented tubular casts, and collapsing focal segmental glomerulosclerosis. “Viral-like” particles have been observed in renal samples at electron microscopy and viral RNA has been identified in both glomerular and tubular compartments of kidney specimens, but the link between viral presence and injury remain unclear. Though the link between AKI and poor outcomes is clear, prevalence and outcomes of COVID-19 in patients with chronic kidney disease and end stage kidney disease has not yet been reported. In patients on immunosuppression like those with kidney transplants or glomerular disease, COVID-19 has presented a management dilemma. Herein, we review the existing literature on kidney disease in COVID-19 and discuss what remains to be learned.

Keywords SARS-CoV2 · Kidney transplant · Acute kidney injury · COVID-19

Introduction
The novel coronavirus disease infection (COVID-19) outbreak, caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV2), began in Wuhan, Hubei Province, China in December 2019 and was declared a global pandemic on March 11, 2020 by the World Health Organization (WHO). Though the initial focus of research has been on pulmonary pathophysiology, observed acute kidney injury (AKI) prevalence an associated morality [1] has been high in some reports—particularly AKI requiring kidney replacement therapies (KRT). Herein, we discuss what has been reported about kidney disease in COVID-19, and what remains to be discovered and lies ahead.

The true incidence of acute kidney injury?
Though findings from several patient populations have been reported, the true incidence of AKI in COVID-19 remains unknown. Existing data, while heterogenous with respect to population size, location, severity of illness, and definitions of AKI, show a wide range of rates of AKI occurrence in patients with COVID-19 from 1 to 46% (Fig. 1 [1–18]). At our institution, the incidence of AKI in 3325 patients with COVID-19 was 46%—with 20% requiring KRT [19]. In the largest cohort of 5449 hospitalized patients with COVID-19 studied so far, the proportion of patients with AKI any stage was 1993 (31.1%) [14]. Few European data are currently available on this topic. In a cohort of 62 patients not on maintenance dialysis in Lombardy, Italy, 40% developed AKI and 10% required KRT [18]. In this cohort, a mortality rate of 88% was
reported in patients with advanced chronic kidney disease (CKD). In a large tertiary care hospital (Parma University Hospital, Parma, Italy) with 800 medicine beds and 60 COVID-19-dedicated ICU beds, 120 COVID-19 patients were admitted to the ICU between February 23rd to May 5th. AKI (as defined by the 2019 Kidney Diseases: Improving Global Outcomes Consensus Conference [20]) was observed in 26/120 (21.7%) before or at ICU admission, and in 29/126 (24.2%) during ICU stay; four patients (4/120, 3.3%) underwent KRT, performed as sustained low-efficiency dialysis (SLED) with regional citrate anticoagulation (personal communication).

Not surprisingly, there are noticeable differences in AKI rates between not only different countries, but also among those studies that included all hospitalized patients compared to those which analyzed only patients required intensive care. Similarly, the percentage of those requiring KRT widely varied from 0 to 23.2% [21]. Potential explanations for these differences include the prevalence of co-morbid conditions and heterogeneity along racial and ethnic lines, local institutional policies about KRT timing, the use of extracorporeal KRT beyond classical “nephrological” indications. One study reported that of the five New York City boroughs, the Bronx had not only higher rates of mortality, but also the highest proportion of racial and ethnic minoritized groups, most persons living in poverty, and lowest levels of educational attainment [22].

**Fig. 1** Incidence of acute kidney injury (AKI) reported in China (black bars), Ireland, United Kingdom, Wales, and United States. Three studies included only patients requiring intensive care [1, 12, 13] while others included hospitalized patients.

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**A proposed pathogenesis of kidney injury**

In a relatively short period of time, several case reports and two series of post-mortem tissue sample analysis from China have begun to shed some light on potential mechanisms of kidney injury associated with COVID-19, even though many open issues remain (Table 1 [23–26]).

It is still not clearly ascertained whether SARS-CoV-2 directly damages the kidney injury or if kidney injury results from complications arising in the course of the viral infection outside the kidney. Reports of urinary abnormalities in COVID-19 patients, positive staining of tubules with viral antigens and complement components in one autopic series, visualization of viral particles in tubular epithelial cells and podocytes on ultrastructural examination in other kidney biopsies and isolation of SARS-CoV-2 in urine, raised the possibility of a “SARS-CoV-2 nephropathy” (Table 1). In a kidney histological analysis of the largest autopsy series so far on 26 patients with COVID-19 from China [27], acute tubular injury ranging from mild to severe was almost always observed. Three biopsy specimens revealed segmental glomerular fibrin thrombi, 3 had pigmented tubular casts, and 7 were thought to have viral-like particles identified under electron microscopy (EM). What was perhaps most interesting of this study was that only 9/26 (34.6%) showed clinical
Table 1  Summary of evidence in favor of and against direct SARS-CoV2-mediated pathogenesis of acute kidney injury (AKI)

| Suggests direct viral involvement | Against direct viral involvement | Comments |
|---------------------------------|---------------------------------|----------|
| Urinalysis abnormalities         | Transient proteinuria and hematuria are common in critically ill patients | Pathophysiology of proteinuria during septic shock remains still largely unclear. Release of several proinflammatory cytokines into the systemic circulation may lead to: a. Generalized loss of endothelial cells barrier integrity and subsequent capillary leak; b. Altered tubular handling of filtered albumin, promoting albuminuria; c. Podocyte cytoskeleton disruption and apoptosis |
| Urinalysis abnormalities (proteinuria, hematuria) commonly occur in the patients with COVID-19 [17] | | |
| Glomerular and tubulointerstitial injury | Immunohistochemical analysis of postmortem kidney biopsies in AKI associated with sepsis shows neutrophils and macrophages in the glomeruli, and neutrophils in the tubulointerstitium. Cell proliferation and fibrin deposition are pronounced in the glomeruli and tubulointerstitium of sepsis-associated AKI [23, 24] | Considering the many complications that can induce acute tubulointerstitial injury in ICU patients with ARDS and AKI, the causal relationship between viral infection and tubular damage is questionable |
| Evidence of diffuse proximal tubule injury with loss of brush border, non-isometric vacuolar degeneration, and necrosis was reported in patients with COVID-19. In addition, erythrocyte aggregates obstructing the lumen of capillaries without platelet or fibrinoid material were present [22] | Absence of viral detection by in situ RNA analysis in the kidney biopsies from two COVID-19 positive cases with collapsing glomerulopathy [33, 34] | |
| Postmortem biopsies from patients with AKI showed severe ATN and lymphocyte/macrophage infiltration, along with positive viral antigen NP staining of tubules and complement components [5] | Absence of virus in serum, urine and kidney tissue by real-time RT-PCR, despite repeatedly positive nasopharyngeal swabs in a critically ill patient with ARDS and AKI [31] | |
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| Viral kidney tropism | Putative coronavirus particles were present, though all other tests for coronavirus in kidney were negative [27] | Many cellular components and structures can masquerade as viruses. In fact, transmission EM of tissue sections are not specific or sensitive for viral particle detection, as many intracellular structures observed in podocytes and endothelial cells resemble viruses (“viral-like particles”). For example, endothelial tubuloreticular inclusions also called myxovirus-like particles [22] |
| Clusters of coronavirus-like particles with distinctive spikes in the tubular epithelium of the kidney were observed in EM specimens from an autopsy series [5, 22] and a native kidney biopsy [27] | Multivesicular bodies (MVB) mimicking SARS-CoV-2 can be found in podocytes from COVID-19 negative patients, irrespective of the underlying kidney disease [22] | Purported virus particles in the cytoplasm of kidney tubular epithelium and podocytes could represent clathrin-coated vesicles, which are normal cell organelles involved in intracellular transport [23] |
| Viral nephrotoxic effects | In a COVID-19 patient with collapsing glomerulopathy, “viral particles” were described as tubuloreticular inclusions in the endothelial cells and podocytes (with the appearance of MVB), but no viral RNA signal was detected in biopsy tissue [27] | Purported virus particles in the cytoplasm of kidney tubular epithelium and podocytes could represent clathrin-coated vesicles, which are normal cell organelles involved in intracellular transport [23] |
| SARS-CoV-2 viral load has been quantified in autopsy kidneys, together with single cell RNA sequencing of viral receptors in kidney specimens and in situ hybridization. This analysis has revealed viral RNA in microdissected portions of the same specimens, thus suggesting both viral entry and replication into kidney cells [29] | Viral entry and replication do not necessarily translate into cytopathic damage, as best exemplified by other viral infections such as HIV [24] | There has not yet been reproducible proof of viral tropism with active infection, which would require the concurrent demonstration of kidney cell infection (i.e. viral entry) together with viral replication/cytopathy |
| Viral nephrotoxic effects | SARS-CoV, whose genome is largely similar to that of SARS-CoV-2, was shown to be capable of entering kidney cells and replicating—but did not cause apparent harm [25] | |

ICU intensive care unit, ARDS acute respiratory distress syndrome, ATN acute tubular necrosis, RNA ribonucleic acid, EM electron microscopy, COVID-19 novel coronavirus disease infection
signs of AKI and 5/26 (19.2%) required KRT. Acute tubular necrosis may be related to a similar presentation as seen with sepsis-induced AKI, mediated by the inflammatory milieu [28, 29]. Sixteen patients out of 26 (61.5%) of patients were hypotensive or required vasopressor support, thus making poor perfusion another likely mechanism of tubular injury. Could SARS-CoV2 itself be playing a role in tubular injury? Viral entry via angiotensin converting enzyme 2 (ACE2), expressed on proximal tubular cells, seems to support such a hypothesis [30, 31]. Additional support of this theory comes from the observation of spherical viral particles characteristic of viral inclusions on EM [27, 32, 33] as well as viral ribonucleic acid (RNA) detection [6] in the kidney. Of note, the presence of presumed viral particles and RNA have not been consistently reported throughout the literature (Table 1), and many cell structures could mimic the virus itself (Table 1). Whether or not the presence of virus is an incidental finding or indeed contributing to pathology has not been determined. One recent study detected SARS-CoV-2 in the kidneys of autopsy specimens using in situ hybridization and indirect immunofluorescence [34]. They noted preferential targeting of glomerular cells in the kidney, with viral detection also in liver, heart, brain, blood and most highly in the respiratory tract. It is likely, as suggested by the authors, that multiorgan tropism influences the course of COVID-19.

The presence of fibrin thrombi typically points towards a diagnosis of thrombotic microangiopathy (TMA), though the hypercoagulable state in patients with COVID-19 raises the likelihood of thrombi both in pulmonary blood vessels [35] and within microvasculature of the kidney. It is also possible that thrombotic occlusion of large kidney blood vessels (e.g. renal artery, renal vein) may contribute to kidney injury. A kidney biopsy from a critically ill patient with COVID-19 showed a classical pattern of acute tubular injury (ATI) with marked regenerative changes and focal acute tubular necrosis (ATN) [36].

Evidence of glomerular injury in patients with COVID-19 also exist, as documented by the presence of proteinuria or hematuria in up to 60% of the cases in some series [37]. Though not reported in the autopsy series, several reports of collapsing focal segmental glomerulosclerosis (FSGS) have been published [32, 38, 39]. Two out of 3 reported the presence of 2 high-risk APOL1 alleles in these patients. Similar to other viral-associated collapsing FSGS (e.g. human immunodeficiency virus (HIV), parvovirus 19) tubuloreticular inclusions suggesting systemic interferon activity [40] were reported.

Like the majority of kidney diseases, the mechanisms are most likely to be multifactorial and data thus far point to contributions from direct viral infection, inflammatory syndrome-mediated injury, hemodynamic instability, and perhaps the hypercoagulable state.

COVID-19 and special populations

Chronic kidney disease (CKD) and end-stage kidney disease (ESKD)

It is not yet known whether CKD or ESKD are a significant risk factors for COVID-19 infection and associated hospitalizations. While the development of AKI is associated with increased in-hospital mortality, hospitalized patients with a prior history of ESKD do not appear to require the same rate of intensive unit level care anecdotally. Though immune dysfunction (both immunosuppression and overactivation) have been described in this population [41], it is possible that ESKD patients do not mount the same cytokine storm response implicated in multiorgan dysfunction noted in critically ill patients. Mortality data on this patient populations remains limited, though a 30% mortality rate was reported in a cohort of 20 hemodialysis patients with COVID-19 [18]. In a large cohort of patients in New York City, patients with CKD and either atrial fibrillation, heart failure, or ischemic heart disease had a higher mortality rate. [42]. In an Italian cohort, 94 patients on hemodialysis with COVID-19 who were managed as either inpatient or outpatient had mortality rates of 42% and 8%, respectively [43]. At our outpatient hemodialysis center, we found an incidence and mortality rate of 17.5% and 3.6%, respectively.

Transplant

Kidney transplant recipients appear to be at higher risk for hospitalization and death from COVID-19. Similar to incidence rates of AKI, reported mortality rates in this population have varied and as high as 28% [22, 44]. There is an absence of consensus about how to adjust immunosuppression in Covid-19 kidney transplant recipients. While immunosuppression may prevent an effective anti-SARS-CoV-2 T cell response, it may also allow controlling inflammatory response that is, at least in part, responsible for COVID-19 related mortality. Calcineurin inhibitors, the backbone of the majority of immunosuppression regimens, have shown in vitro activity against coronaviruses—though clinical data are lacking [45]. At our center, our approach has been to maintain both corticosteroids and anti-metabolite immunosuppressive therapy while lowering calcineurin trough targets. While most centers reduce immunosuppression, approaches vary significantly from institution to institution [46].
Glomerular disease

The impact of COVID-19 on glomerular disease patients remains unclear. An individualized risk–benefit analysis to reduce, defer, or maintain immunosuppression seems appropriate. Some centers, including our own, have noted a marked reduction in rapidly progressive glomerulonephritis cases and urgent kidney biopsies during widespread home quarantining—supporting an environmental component in exacerbating glomerular disease [47].

A look to the future

Several months into the COVID-19 pandemic, there is much that we do not understand. The relationship between acute kidney injury and multiorgan failure is unclear. Details on the contribution of hypercoagulability, complement activation and endothelial injury to organ dysfunction are lacking. Presentation of a catabolic state in patients with COVID-19 [48] raises the possibly of acquired metabolic or mitochondrial disorders, in addition to the impact of cytokines and the inflammatory state on these pathways. Though this catabolic state seems to mimic rhabdomyolysis, creatine phosphokinase (CPK) levels have not been found to be dramatically elevated. While ACE2 plays an important role in viral cellular entry, proposed angiotensin dysregulation, innate and adaptive immune pathway activation, and hypercoagulation have emerged as proposed mechanisms of kidney and other organ injury [49]. Most concerning is the high level of residual kidney dysfunction in patients who have recovered from COVID-19. We are yet to fully appreciate their long-term risk for progression to end-stage kidney disease or whether the virus like HIV, could use kidney epithelial cells as a reservoir. A better understanding of kidney disease pathophysiology may aid in the development of effective therapies. As we move forward, it is plausible that computational biology will be important in understanding COVID-19 pathophysiology in identifying both therapies and innovative strategies to design a vaccine [50].

Compliance with ethical standards

Conflict of interest This manuscript has not been published previously, in whole or part. The authors have no conflicts of interest to disclose.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

Informed consent For this type of study formal consent is not required.

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