Cardio-Pulmonary-Renal Consequences of Severe COVID-19

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Abstract
Severe acute respiratory syndrome coronavirus 2 has rapidly spread worldwide and resulted in the coronavirus disease 2019 (COVID-19) pandemic. The disease raised an unprecedented demand for intensive care support due to severe pulmonary dysfunction and multiorgan failure. Although the pulmonary system is the potential target of the COVID-19, recent reports have demonstrated that COVID-19 profoundly influences the cardiovascular system and the kidneys. Research studies on cadavers have shown that direct heart and kidney injury can be frequently seen in patients deceased due to COVID-19 infection. On the other hand, functional or structural dysfunction of the heart may deteriorate the renal function and vice versa. This concept is already known as the cardiorenal syndrome and may play a role in COVID-19. Proactive monitoring of micro- and macrohemodynamics could allow prompt correction of circulatory dysfunction and can be of pivotal importance in the prevention of acute kidney injury. Moreover, type and amount of fluid therapy and vasoprotective drug support could help manage these patients either with or without mechanical ventilator support. This brief review outlines the current evidence regarding the COVID-19-related renal and cardiorenal complications and discusses potential hemodynamic management strategies.

Introduction
Coronaviruses (CoVs) are enveloped, positive single-stranded RNA viruses that lead to respiratory, cardiac, or intestinal infections in animals and humans. They consist of 4 subgroups: α-, β-, δ-, and γ-CoVs. Humans could be infected by various α-CoV and β-CoV, although δ-CoV and γ-CoV mainly infect birds [1]. The critical importance of these viruses is their potential to cause pandemics. In the last 2 decades, 3 overwhelming outbreaks of viral pneumonia caused by CoVs were reported to spread around the world. First in 2002, severe acute respiratory syndrome virus originated in China and caused around 800 deaths worldwide, with an 11% mortality rate [2]. A decade later in 2012, Middle East respiratory syndrome virus was first reported in Saudi Arabia and affected 2,494...
people with a fatality rate of 34% [3] [1–55]. And again, almost a decade later, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) appeared in late 2019 again in China and was called coronavirus disease 2019 (COVID-19). Since then, COVID-19 has rapidly spread worldwide, leading to classification as a pandemic by the World Health Organization on March 11, 2020 [4].

When COVID-19 was first presented itself, the SARS-CoV-2 was thought to affect the upper and lower respiratory tract only. However, recent evidence suggests that COVID-19 is a multisystem inflammatory vasculopathy and centered around endothelial dysfunction [5, 6]. Similar to severe acute respiratory syndrome virus and Middle East respiratory syndrome virus did, SARS-CoV-2 directly and indirectly causes cardiovascular and renal complications [7–9]. The concomitant development of acute myocardial or kidney injury is associated with significantly worse outcomes [9, 10]. Underlying comorbidities such as hypertension, diabetes, obesity (i.e., metabolic syndrome), smoking, and cardiovascular diseases have been associated with increased risks for the development of severe complications of COVID-19 [11]. This is probably one of the reasons why the mortality and morbidity of COVID-19 infection is more prominent in the elderly population who have higher rates of these comorbidities [12].

The SARS-CoV-2 tightly binds to the host ACE-2 receptor by its S protein (S1). After that, cellular transmembrane serine proteases (TMPRSS) cleave the S1 protein so that the virus releases peptides to accomplish fusion to the host membrane [13]. Binding the ACE-2 receptor and TMPRSS is crucial for the virus to enter the cells, and this process occurs at the tissues which coexpress the ACE-2 receptor and TMPRSS [13, 14]. The heart, lungs, kidney, and intestines are abundant in the ACE-2 receptor and TMPRSS. Depending on the type of the affected tissue and the severity of the inflammation, the clinical manifestation varies by dyspnea, cough, fever, myalgias, respiratory failure, diarrhea, proteinuria, myocarditis, arrhythmia, and neurologic symptoms [15]. In this report, we briefly review the pathogenesis of COVID-19 and its direct and indirect effects and mutual interactions of the heart, lungs, and kidneys.

Cardiopulmonary Interaction and COVID-19

In contrast to classic acute respiratory distress syndrome (ARDS), a hypothesis suggested that the COVID-19-related ARDS has a unique pattern associated with either relatively low (type L) or normal/high (type H) pulmonary elastance. L-type is characterized by high lung compliance and does not require high positive end-expiratory pressure (PEEP) and can tolerate high tidal volume without an increase in plateau and driving pressures. On the other hand, H-type has a low lung compliance and shows a similar pattern to classic ARDS [16].

Determining the phenotypes of lung injury is clinically necessary to design the ventilator strategy and direct the hemodynamic management. Mechanical ventilation and applied settings (PEEP and tidal volume) influence highly the intrathoracic pressure and therefore the preload and afterload of the right and left ventricles [17]. Especially the right ventricle, compared to the left ventricle, is highly sensitive to the increase in afterload [18]. Therefore, the amount of the applied PEEP level, the influence of alveolar pressure (Palv) transmission on pleural pressure (Ppl), and the lung’s compliance can have a significant impact, especially on the right heart and risks of right heart failure [18, 19]. This is one of the reasons that right ventricular (RV) dysfunction and failure should be actively monitored in patients with severe COVID-19 lung involvement. Even in nonprogressed COVID-19 stages, the RV function has shown to deteriorate depending on the pulmonary arterial pressure, direct or indirect lung injury, hypoxic vasoconstriction of the pulmonary microvasculature, micro- or macrovascular thrombotic processes, myocardial injury, and mechanical effects of the mechanical ventilator [20]. The presence of venous congestion and obstructed lymphatic drainage due to RV dysfunction can lead to decreased secondary organ perfusion pressure due to venous pressure-induced tamponade, especially in kidneys potentially provoking acute kidney injury (AKI) [21, 22].

Renal Injury and Cardiorenal Interactions in COVID-19

Initially, a low incidence of AKI has been reported in patients with confirmed COVID-19 [23]. Guan and colleagues [24] reported an AKI incidence of between 0.1 and 6% in around 1,100 patients with COVID-19 in China. A large portion of these patients had normal kidney functions, and the reported data seem similar to what would be expected from patients hospitalized with acute illnesses. However, recent reports show a higher incidence of AKI depending on the ethnicity, country, and the severity of the disease in COVID-19 [25]. Critically ill COVID-19 patients had increased risk for AKI 7–56%,
where the development of AKI was highly related to increased mortality, even up to 80% among ICU population [9, 26]. Given the impact of AKI in patients with COVID-19, experts on the kidney have designed a consensus report that can help in disease management [27].

Independently from its origin, ARDS patients have increased risk for the development of AKI [28]. In fact, AKI is the most prevalent extrapulmonary organ failure in ARDS [29]. The responsible mechanism includes an inflammatory and hemodynamic crosstalk between the lung, heart, and kidneys [29, 30]. The kidney and heart display a bidirectional interaction whereby an acute or chronic dysfunction in one affects the other. This concept is known as cardiorenal syndrome (CRS) and has been well defined in several preclinical and clinical studies [31]. However, the pathophysiologic basis of COVID-19-related organ dysfunction is complicated and characterized by unique changes that are specific for COVID-19. The responsible mechanisms that might be responsible for the development of COVID-19-related CRS are summarized in Figure 1.

AKI caused by cardiac dysfunction due to ARDS itself or direct cardiac injury can be classified as type 1 CRS. On the other hand, the occurrence of heart and kidney injury due to the systemic effects of inflammation constitutes type 5 CRS’s pathophysiology. From the point of the microvascular level, the cytokines released secondary to systemic inflammation, increased vascular permeability due to endothelial activation, renovascular microthrombus formation, intravascular fluid depletion or on the contrary venous congestion secondary to volume overload, and excess vasoactive or fluid therapy all could contribute to the deterioration in kidney function [32, 33]. Therefore, the kidneys that already have physiologically low tissue oxygen levels could become even more vulnerable to
widely used to assess cardiac function and predict fluid responsiveness in these patients [41]. Moreover, hemodynamic parameters such as blood pressure, cardiac output, pulse pressure variation, pleth variability index, and systemic vascular resistance can noninvasively be measured. Although noninvasive monitoring techniques have several limitations, they could be used, especially in limited access to resources and qualified healthcare workers.

During their stay in the ICU, almost >30% of patients require vasopressor therapy [26, 42, 43]. Due to its infective origin, COVID-19-related shock could be classified as a septic shock where the underlying reason for the hypotension is of a distributive shock [44]. Second, COVID-19 patients could be prone to relative hypovolemia due to restrictive fluid administration, excessive insensible fluid loss, or diuretic therapy aimed at keeping the patients dry [23]. Third, secondary to increased pulmonary arterial pressures (secondary to ARDS), need of mechanical ventilation with high pressures or frequently encountered pulmonary embolism may induce right heart failure and compromise secondary organ perfusion including the liver, intestines, and the kidneys [18, 20]. Last but not least, direct cardiac injury due to myocarditis or myocardial depression could decrease heart pump function and exacerbate tissue hypoperfusion [10, 45].

In clinical practice, recognition of the underlying pathophysiologic mechanisms causing the deterioration in hemodynamics could be achieved by measuring dynamic and static hemodynamic parameters. Dynamic hemodynamic indices have been shown to be superior to static hemodynamic indices [46]. Therefore, continuous hemodynamic monitoring and regular echocardiographic imaging is pivotal. Unfortunately, only continuous monitoring of arterial pressure is not enough to assess heart-lung interaction although it seems to be common practice in COVID-19 patients today. Continuous hemodynamic monitoring by inserting a pulmonary arterial catheter or less-invasive thermodilution techniques such as pulse contour cardiac output monitoring including the measurement of the extravascular lung water and pulmonary vascular permeability index might be helpful due to providing comprehensive hemodynamic data.

The techniques mentioned above are used to estimate “classic” macrohemodynamic parameters, which might not always show parallel changes with microcirculatory indices. This concept is known as “loss of hemodynamic coherence” and well described, especially in critically ill patients [47]. Since the main target for optimizing the macrocirculation is to provide sufficient oxygen and blood flow to the tissues, there will be need of novel tech-
Techniques to assess the microcirculation [48, 49]. Hand-held vital microscopes might be good candidates for the bedside evaluation of microcirculation in critically ill COVID-19 patients. Hand-held vital microscopes enable direct monitoring of the microvessels and might allow a point of care assessment of the applied therapy using newly introduced automatic analysis called MicroTools [48–50]. Besides, serum lactate level, capillary refill time, and mottling score could also indirectly monitor the microcirculation [51].

**Proposal for a Proactive Hemodynamic Management of COVID-19**

Prevention of hospitalization and subsequently severe clinical complications would be of the uttermost importance. Unfortunately, no single effective treatment or medication is yet available except prevention with novel vaccines. Prehospital initiation of the sequenced multidrug therapy including adjuvant nutraceuticals, the combination of anti-infective therapy, inhaled/oral corticosteroids, antiplatelets/anticoagulants, and supplemental oxygen along with active telemonitoring have been proposed to mitigate the intensity and duration of COVID-19-associated symptoms and complications. This could decrease the majority of hospitalizations and, by extension, development of cardio-pulmonary-renal complications and deaths [52, 53].

Once progressively ill, advanced hemodynamic management of COVID-19 patients should be applied in critical care settings with access to conventional and complex treatment. Surviving Sepsis Campaign guideline for COVID-19 published by the European Society of Intensive Care Medicine experts suggests a conservative fluid strategy in critically ill COVID-19 patients [44]. Albeit fluid therapy in critical illness aims at achieving adequate blood flow required for tissue oxygenation, both excess and inadequate fluid can cause tissue hypoxia. Therefore, fluid therapy of critical illness might be considered as a 2-edged sword, which makes monitoring the amount of fluid given and monitoring the fluid responsiveness critical at the bedside. Furthermore, due to the endothelial damage in COVID-19, excessive fluid therapy could be harmful leading to tissue edema and pulmonary capillary leakage [54]. Moreover, an increase in right heart pressure might facilitate the deterioration in ventilation/perfusion mismatch and worsen the arterial and tissue hypoxia [20, 21, 31]. On the other hand, inadequate fluid therapy leads to hypovolemia and tissue hypoxia [9]. Targeting central venous pressure <10 mm Hg could avoid hypervolemia [21].

If the patient is hypotensive despite adequate fluid therapy, a vasopressor should be added to therapy instead of giving more fluids [44]. Norepinephrine is preferred as the first-line vasoactive agent, and titrating the dose until the mean arterial pressure reaches 60–65 mm Hg seems better than targeting a higher mean arterial pressure level. Vasopressin or epinephrine could be the second-line therapy if norepinephrine is insufficient. Having low ScvO₂ despite adequate hemoglobin level should remind adding dobutamine to the therapy to compensate the failing heart [44]. For ongoing cardiopulmonary failure and shock, extracorporeal membrane oxygenation should be considered given the clinical successes with extracorporeal real membrane oxygenation in the treatment of critically ill H1N1 influenza epidemics [55].

**Conclusion**

In this brief report, we summarize the pathogenesis and current evidence of renal and cardiorenal complications in patients with COVID-19. Acute kidney injury and cardiac injury are common in critically ill COVID-19 patients and are associated with high mortality. Understanding the interaction between 2 organs in COVID-19 and recognizing the factors affecting organ functions can provide more accurate treatment and monitoring approaches. Proactive micro- and macrohemodynamic monitoring and timely intervention could prevent progressive multiorgan failure, including the AKI and its dire sequelae. Future studies focusing on the disease pathophysiology of secondary organ injury are needed to learn more about the renal and cardiorenal interactions in COVID-19.

**Statement of Ethics**

The study is exempt from ethics committee approval due to the nature of the study. This is a review article and does not include human or animal subjects' material or data.

**Conflict of Interest Statement**

The authors have no conflicts of interest to declare.
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Author Contributions

G.G. and K.C.: conception, design, and drafting of the work. G.G., C.I., A.T., and K.C.: revising critically for important intellectual content. G.G., C.I., A.T., and K.C.: final approval of the version to be published.

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