ACE2, Metformin, and COVID-19

Atul Malhotra, Mark Hepokoski, Karen C. McCowen, John Y-J Shyy

Department of Medicine, University of California, San Diego, La Jolla, CA, 92023

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To whom correspondence should be addressed to:

John Y.-J. Shyy, Ph.D., Division of Cardiology, Department of Medicine, University of California, San Diego, 9500 Gilman Drive, La Jolla, CA
jshyy@health.ucsd.edu

Atul Malhotra, MD, Division of Pulmonary, Critical Care and Sleep Medicine, Department of Medicine, University of California, San Diego, 9500 Gilman Drive, La Jolla, CA
amalhotra@health.ucsd.edu
**SUMMARY**

COVID-19 is becoming a leading cause of mortality throughout the world, and few effective therapies are currently available. Angiotensin converting enzyme 2 (ACE2) is essential to COVID-19 pathogenesis, as the binding of SARS-CoV-2 spike protein (S protein) is required for viral entry and development of COVID-19. ACE2 regulates the protective arm of the renin-angiotensin-aldosterone system (RAAS) that endows anti-hypertensive and anti-inflammatory effects in the cardiovascular and pulmonary systems. Preclinical data suggest ACE2 might be downregulated after SARS-CoV-2 binding, and treatments that increase ACE2 may prevent cardiopulmonary injury. Development, testing, and mass production of novel ACE2 therapies may take years, while more effective treatments for COVID-19 are needed urgently. Metformin is a widely available anti-diabetic agent that has an excellent safety profile, and clinical and preclinical data suggest metformin may offer cardiopulmonary protection in COVID-19 via enhanced ACE2 expression.
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With the COVID-19 pandemic a threat to human health worldwide, there is an urgent need to identify treatments that are safe and available immediately. The virus responsible for COVID-19, SARS-CoV-2, has a spike protein (S protein) binding domain that is nearly identical in structure to the virus responsible for prior outbreaks of severe acute respiratory syndrome (SARS), SARS-CoV-1.¹ However, the binding affinity of SARS-CoV-2 for the angiotensin converting enzyme 2 (ACE2) receptor appears to be 10 times stronger, which may partially explain its rapid spread throughout the world.² ACE2 binding in pneumocytes and enterocytes is required for viral entry, yet paradoxically, treatments that increase ACE2 may be beneficial in mitigating the complications of COVID-19. In the renin–angiotensin–aldosterone system (RAAS), Angiotensin converting enzyme 2 (ACE2)–Ang 1-7–Mas acts as the protective arm to counteract the deleterious arm of ACE1–Ang II to reduce systemic and pulmonary hypertension, and activate anti-inflammatory pathways after tissue injury.³ Interestingly, SARS-CoV-1 was found to decrease ACE2 expression after binding, and low levels of ACE2 have been implicated in various cardiovascular impairments and acute respiratory distress syndrome (ARDS).⁴ Recently, treatment with human recombinant ACE2 (hrACE2) showed positive results in a human organoid model of SARS-CoV-2⁵, but safety and efficacy of hrACE2 still need to be determined in human studies and timely production might not meet the need for patients. Here, we discuss the rationale for metformin as a safe and currently available therapy that is known to increase ACE2 and may offer cardiopulmonary benefit in patients suffering from COVID-19.

Data collected from cohorts in Wuhan, China indicate high comorbidity of hypertension, diabetes, and cardiovascular diseases among COVID-19 patients.⁶ The prevalence of hypertension in COVID-19 draws tremendous attention to using RAAS inhibitors, i.e., ACE
inhibitors (ACEI) and angiotensin receptor blockers (ARBs), in treating COVID-19 symptoms and the underlying diseases.\textsuperscript{7} Ample preclinical and clinical studies have been conducted to investigate the effects of ACEI and ARBs on the expression and activity on ACE2.\textsuperscript{8} However, these studies have generated conflicting results and the effects of these drugs on ACE2 in humans remain inconclusive.\textsuperscript{9} Nonetheless, consensus is emerging that potential benefits override the potential harms of RAAS inhibitors in COVID-19\textsuperscript{7}, possibly via ACE2 augmentation.

Despite the American Heart Association statement advocating “not to add or remove any RAAS-related treatments, beyond actions based on standard clinical practice”, we have little knowledge on the benefits or risks of other drugs known to modulate ACE2, such as anti-diabetic agents. Amid the comorbidity of diabetes and high body mass index (BMI) in COVID-19, this information is urgently needed. As first line anti-diabetes agents, the biguanide class of drugs, (metformin is the only available currently), inhibit gluconeogenesis. Although clinical evidence demonstrating the effect of metformin in RAAS homeostasis in humans is lacking, there are plenty of data from animal and cultured cell experiments supporting that metformin antagonizes the effects of the ACE1–Ang II arm of RAAS.\textsuperscript{10} Furthermore, large cohort studies have suggested that metformin offers pleiotropic cardiovascular protection. In the United Kingdom Prospective Diabetes Study (UKPDS), metformin therapy in obese patients with type II diabetes was associated with a lower risk for myocardial infarction compared to control.\textsuperscript{11} In an observational study of patients with diabetes and high cardiovascular risk, all-cause mortality was significantly reduced in metformin-users compared to non-users, although curiously there were no differences in hard cardiovascular end-points.\textsuperscript{12} Cardiovascular complications, such as
non-ischemic cardiomyopathy, have been described as a complication of COVID-19, but the mechanisms involved, including the potential role of ACE2 depletion, remain unclear.

Beyond the potential cardiovascular benefits, several studies have suggested metformin may provide pulmonary protection following SARS-CoV-2 infection. Lung injury, more commonly referred to as ARDS, is one of the most serious complications of COVID-19. Interestingly, metformin has been shown to attenuate lung injury in animal models of lipopolysaccharide\textsuperscript{13}, hyperoxia\textsuperscript{14}, and ventilator induced lung injury.\textsuperscript{15} Other potential mechanisms include metformin alteration of the microbiome or mitochondrial function.\textsuperscript{16,17} In a retrospective study of human subjects, Jo et al. showed an interesting trend in ARDS mortality comparing metformin users vs. non-users (42.4% vs. 55.3%, p=ns) even though metformin was discontinued on admission in the cohort examined.\textsuperscript{18} To our knowledge, metformin has not been studied prospectively in the treatment of ARDS, but these pre-clinical and retrospective data are intriguing.

The molecular basis for the beneficial effects of metformin on the cardiovascular and pulmonary systems likely involves metformin activation of AMP-activated protein kinase (AMPK).\textsuperscript{10} Functioning as an energy gauge, AMPK regulates metabolic homeostasis from cellular to whole body levels. AMPK not only phosphorylates key molecules regulating metabolism (e.g., acetyl-CoA carboxylase, sterol regulating element binding proteins), but also those involved in cardiovascular health (e.g., eNOS). Metformin and ARBs (e.g., telmisartan) can activate AMPK, which phosphorylates ACE2 on Ser-680.\textsuperscript{10,19} This post-translational modification of ACE2 decreases its ubiquitination and therefore extends the half-life of ACE2 which may offer lung protection. Indeed, transgenic mice overexpressing the phosphomimetic ACE2 S680D exhibit less damage in pulmonary vasculature under injurious conditions\textsuperscript{10}, and
ARBs were found to attenuate lung injury after exposure to the S protein of SARS-CoV-1.\textsuperscript{20} Due to similar interactions between ACE2 to SARS-CoV-1 and SARS-CoV-2, it is reasonable to suspect that metformin may offer similar lung protection in COVID-19. A second notion is that post-translational modifications may change the 3-D conformation of the extracellular domain of ACE2, thereby decreasing SARS-CoV-2 viral recognition.

Although diabetes is considered a risk factor for COVID-19, prior clinical studies have suggested diabetes patients may have less risk of lung injury than matched controls.\textsuperscript{21} The reason for this discrepancy is unclear, but at least in theory diabetes medications, such as metformin, could offer protection from ARDS. However, a recent meta-analysis demonstrated conflicting results\textsuperscript{22} and the possible protective effect of diabetes appears to be present in both type 1 and type 2 DM, suggesting that metformin use was not the underlying mechanism (since metformin is not generally used in type 1 DM). Nonetheless, we believe that the use of metformin to prevent ARDS in the COVID-19 context deserves further study. The use of metformin in critically ill patients is complex, since rare reports of metformin induced lactic acidosis exist. However, such anecdotes would not preclude a preventative strategy using metformin assuming risk/benefit data were supportive.

We hypothesize that patients with COVID-19 taking metformin will have higher circulating ACE2 levels, and lower morbidity and mortality compared to matched controls. We propose to test this hypothesis through a combination of retrospective cohort studies, and prospective translational studies evaluating the ACE2 axis in COVID-19 patients. First, the effect of metformin on cardiopulmonary outcomes needs to be evaluated in a cohort of patients with COVID-19. These data could potentially be evaluated retrospectively in databases that are currently available. Next, we would evaluate circulating ACE2 levels and both arms of the
RAAS axis prospectively in patients admitted with COVID-19. These data would allow an assessment of the effect of metformin on circulating ACE2 levels, as well as the ability of circulating ACE2 levels to predict morbidity and mortality due to COVID-19. Noticeably, metformin induction of ACE2 in the cardiovascular system is mainly validated in rodent experiments. The experimental doses used in rats or mice (i.e., 100-200 mg/kg/day) are likely correspondent to the therapeutic doses in human (800-1600 mg/day). Thus, these mechanisms described are expected to be observed at diabetic doses of metformin therapy. If our hypothesis is supported by these studies, it would provide a strong foundation for rational investigations, including clinical trials, focused on metformin as a safe and immediately available treatment for patients infected with SARS-CoV-2. While this work was submitted for review, we noted an observational study from Wuhan, China, which demonstrated lower hospital mortality in COVID-19 patients that were being treated for diabetes with metformin, compared with patients not receiving metformin.²³

Limitations of Study: We acknowledge the need for clinical corroboration of our discussion. Despite some potentially compelling rationale, we acknowledge a number of unanswered questions:

a. Does metformin attenuate the risk of poor cardiopulmonary outcomes associated with DM with and without obesity in COVID-19?

b. Does metformin increase circulating ACE2 levels, and do circulating ACE2 levels predict outcomes in patients with COVID-19?

c. Is metformin therapy a safe and effective treatment for patients infected with SARS-CoV-2 who were not previously taking metformin?
Does metformin treatment have clinical benefit in patients with COVID-19 who are not diabetic or obese?

In summary, ACE2 is an anti-inflammatory peptide in the protective arm of the RAAS system that offers beneficial cardiopulmonary effects. Although binding of ACE2 receptors and subsequent downregulation appears to be a key feature in the pathophysiology of SARS-CoV-1 and SARS-CoV-2 infections, therapies that enhance ACE2 have great potential to benefit patients suffering from COVID-19. Metformin is a commonly used antidiabetic agent that is known to enhance ACE2 expression via AMPK activation. Metformin has an excellent safety profile and wide availability throughout the world which allows for immediate, wide distribution if proven to be effective as a monotherapy, or in combination with antiviral agents, such as remdesivir.
REFERENCES


