PERSPECTIVE

Does COVID19 Infect the Brain? If So, Smokers Might Be at a Higher Risk

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ABSTRACT

COVID19 is a devastating global pandemic with epicenters in China, Italy, Spain, and now the United States. While the majority of infected cases appear mild, in some cases, individuals present serious cardiorespiratory complications with possible long-term lung damage. Infected individuals report a range of symptoms from headaches to shortness of breath to taste and smell loss. To that end, less is known about how the virus may impact different organ systems. The SARS-CoV2 virus, which is responsible for COVID19, is highly similar to SARS-CoV. Both viruses have evolved an ability to enter host cells through direct interaction with the angiotensin converting enzyme (ACE) 2 protein on the surface of many cells. Published findings indicate that SARS-CoV can enter the human nervous system with evidence from both postmortem brains and detection in cerebrospinal fluid of infected individuals. Here, we consider the ability of SARS-CoV2 to enter and infect the human nervous system based on the strong expression of the ACE2 target throughout the brain. Moreover, we predict that nicotine exposure through various kinds of smoking (cigarettes, electronic cigarettes, or vape) can increase the risk for COVID19 neuroinfection based on known functional interactions between the nicotinic receptor and ACE2. We advocate for higher surveillance and analysis of neurocomplications in infected cases.

SIGNIFICANCE STATEMENT

The COVID19 epidemic has spurred a global public health crisis. While many of the cases requiring hospitalization and intensive medical care center on cardiorespiratory treatment, a growing number of cases present neurological symptoms. Viral entry into the brain now appears a strong possibility with deleterious consequences and an urgent need for addressing. COVID19 belongs to a family of coronaviruses (CoV) that have evolved in various species (Lam et al., 2016). Human-infecting CoV such as SARS and COVID19 have acquired ability to bind the angiotensin converting enzyme (ACE) 2 on epithelial cells as a primary mechanism of entry into the host (Qi et al., 2020). Critical cases of COVID19 infection commonly manifest as cardiopulmonary symptoms and in severe cases advance into organ failure and sepsis as a result of a “cytokine storm” over activation of the immune system (Guan et al., 2020). The case fatality rate is still unclear but is likely anywhere from 0.4% to 4% and depends critically on the ability of public health systems to provide intensive supportive care. Recently, we have raised the question of whether nicotine exposure through cigarette smoke (and other formulas) is a comorbidity factor in COVID19 infection and clinical outcome (Olds and Kabbani, 2020). Functional interactions between nicotine exposure and ACE2 expression in lungs and other organ systems such as heart and kidneys as well as nicotine and other components of the renin angiotensin system (RAS) suggest that smoking can promote COVID19 cellular entry through nicotinic acetylcholine receptor (nAChR) signaling. Notably, nAChRs are known to be on many of the same cells that express ACE2 in the lungs, kidneys, circulation, and brain (Changeux, 2010; Tolu et al., 2013; Nordman et al., 2014). Thus, smoking can impact COVID19 pathophysiology and clinical outcome in several organ systems.

Differential host factors such as age, health, simultaneous infection, and genetics are known determinants of susceptibility to a viral infection. Smoking is a strong factor in predicting an individual’s likelihood of developing and managing a viral infection and especially a respiratory infection (Razani-Boroujerdi et al., 2004; Eddleston et al., 2011). Here, we raise the question of nicotine-associated comorbidity to COVID19 in the context of the brain based on published evidence that the viral target receptor ACE2 is expressed in the brain and functionally interacts with nAChRs (Ferrari et al., 2007; Oakes et al., 2018). We consider if neural cells, such as epithelial cells, are more vulnerable to infection in

ABBREVIATIONS: ACE, angiotensin converting enzyme; CoV, coronavirus; COVID19, coronavirus disease 19; nAChR, nicotinic acetylcholine receptor; RAS, renin angiotensin system; SARS, severe acute respiratory syndrome.
smokers because nicotine stimulation of the nAChR can increase ACE2 expression within them (Olds and Kabbani, 2020). This issue is critical because evidence shows that mRNA from the closely related SARS virus, which also binds ACE2 as a mechanism of cell entry, was detected in brain and cerebrospinal fluid of infected individuals (Zhang et al., 2003; Chong et al., 2004; Inoue et al., 2007). Moreover, SARS ability to enter neurons is established in experimental systems using recombinant human ACE2 as the point of entry (Netland et al., 2008; Kaparianos and Argyropoulou, 2011).

Now is the time to ask if infection with COVID19 can result in long-term neural damage in both symptomatic and asymptomatic individuals and if chronic nicotine exposure through smoking habits and addiction increases risk of developing COVID19-associated neuropathology through interactions between nAChRs and ACE2 in neurons and glia.

Functional interaction between nicotine and components of the RAS (such as ACE2) are well established in several organ systems including the lungs where smoking is found to impact cardiopulmonary health (Ferrari et al., 2008; Virdis et al., 2010; Fuchs et al., 2018). Similar RAS components also exist in the human brain, and nicotine exposure is documented to modulate RAS activity in areas such as the hypothalamus and brain stem, leading to changes in endocrine release as well as hypertension, respectively (Ferrari et al., 2007; Oakes et al., 2018). In the brain, as elsewhere, ACE2 metabolizes angiotensin II to produce angiotensin 1–7, and this process occurs in neurons as well as astrocytes (Hung et al., 2016). ACE2 signaling is widely thought to oppose oxidative stress and neuroinflammation and disruption in ACE function, and balance can drive neurodegeneration of dopaminergic neurons (Labandeira-García et al., 2014). ACE activity may also contribute to cortical cholinergic pathways and participate in the progression of Alzheimer disease (Keoh et al., 2009). Figure 1 depicts areas of notable ACE2 mRNA expression such as the cortex, striatum, hypothalamus, and brainstem within the adult human brain as based on microarray data from the Allen Brain Atlas (Jones et al., 2009). These regions, which are known to also express various types of nAChRs (Dani and Bertrand, 2007), are putative sites for primary infection with COVID19 in the human brain. Interactions between nAChRs and ACE2 have been studied in several of these regions including the ventrolateral medulla (Deng et al., 2019), and smoking may lead to enhanced viral infection through the ability of nicotine to upregulate nAChRs in regions such as the lungs (Plummer et al., 2005; Thorgeirsson et al., 2008; Changeux, 2010). In this case, upregulation of nAChRs in either/both neurons and astrocytes could promote greater viral entry and replication through augmented ACE2 expression in the cell (Fig. 2A).

In conclusion, we suggest that were COVID19 to breach the blood brain barrier (as evidence suggests for SARS) during the course of ongoing infection, interactions with ACE2 in multiple brain regions would present the virus the opportunity to infect the brain of COVID19 patients. Disruptions to the blood brain barrier may especially be more probable in severe cases of infection and associated with strong immunologic responses such as the cytokine storm pathologies as well as adenovirus coinfection (Channappanavar and Perlman, 2017). In one scenario, neural cells could serve as latent reservoirs for the virus but also produce debilitating neurologic symptoms with adverse affects on vascular homeostasis, adaptive immunologic responses, and cognitive function (Fig. 2). Supporting our concern, a recent report suggests that over 30% of 814 retrospectively studied COVID19 patients showed neurologic symptoms (Mao et al., 2020). The above supports the notion that COVID19 may pose a risk for brain infections and suggests that smokers as well as immune compromised individuals are at higher risk for neurologic complications. The evidence calls for careful studies on postmortem COVID19 patients’ brains and for continued surveillance for signatures of the virus beyond lung, cardiac, and renal tissue.

![Figure 1](https://example.com/figure1.jpg)
Future studies should also include a postmortem analysis of brains from smokers versus nonsmokers.

Authorship Contributions

Wrote or contributed to the writing of the manuscript: Kabbani, Olds.

References


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Fig. 2. Mechanisms of ACE2 entry and COVID19 infection in neural cells. (A) A role for nicotine-associated upregulation of nAChRs in ACE2 expressing astrocytes and neurons. Based on published findings, upregulation of nAChRs can also increase ACE2 cell surface expression, thereby enhancing viral entry and infection. (B) A putative clinical timeline for progression of COVID19 into the brain.