The neuroinvasive potential of SARS-CoV2 may play a role in the respiratory failure of COVID-19 patients

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Abstract
Following the severe acute respiratory syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV), another highly pathogenic coronavirus named SARS-CoV-2 (previously known as 2019-nCoV) emerged in December 2019 in Wuhan, China, and rapidly spreads around the world. This virus shares highly homologous sequence with SARS-CoV, and causes acute, highly lethal pneumonia coronavirus disease 2019 (COVID-19) with clinical symptoms similar to those reported for SARS-CoV and MERS-CoV. The most characteristic symptom of patients with COVID-19 is respiratory distress, and most of the patients admitted to the intensive care could not breathe spontaneously. Additionally, some patients with COVID-19 also showed neurologic signs, such as headache, nausea, and vomiting. Increasing evidence shows that coronaviruses are not always confined to the respiratory tract and that they may also invade the central nervous system inducing neurological diseases. The infection of SARS-CoV has been reported in the brains from both patients and experimental animals, where the brainstem was heavily infected. Furthermore, some coronaviruses have been demonstrated able to spread via a synapse-connected route to the medullary cardiorespiratory center from the mechanoreceptors and chemoreceptors in the lung and lower respiratory airways. Considering the high similarity between SARS-CoV and SARS-CoV2, it remains to make clear whether the potential invasion of SARS-CoV2 is partially responsible for the acute respiratory failure of patients with COVID-19. Awareness of this may have a guiding significance for the prevention and treatment of the SARS-CoV-2-induced respiratory failure.

KEYWORDS
cell susceptibility, coronavirus, dissemination, nervous system

1 | INTRODUCTION

Coronaviruses (CoVs), which are large enveloped non-segmented positive-sense RNA viruses, generally cause enteric and respiratory diseases in animals and humans.1 Most human CoVs, such as hCoV-229E, OC43, NL63, and HKU1 cause mild respiratory diseases, but the worldwide spread of two previously unrecognized CoVs, the severe acute respiratory syndrome CoV (SARS-CoV) and Middle East respiratory syndrome CoV (MERS-CoV) have called global attention to the lethal potential of human CoVs.2 While MERS-CoV is still not eliminated from the world, another highly pathogenic CoV, currently named SARS-CoV-2 (previously known as 2019-nCoV), emerged in December 2019 in Wuhan, China. This novel CoV has caused a national outbreak of severe pneumonia (coronavirus disease 2019 [COVID-19]) in China, and rapidly spreads around the world.
Genomic analysis shows that SARS-CoV-2 is in the same beta-coronavirus (βCoV) clade as MERS-CoV and SARS-CoV, and shares highly homologous sequence with SARS-CoV. The public evidence shows that COVID-19 shares similar pathogenesis with the pneumonia induced by SARS-CoV or MERS-CoV. Moreover, the entry of SARS-CoV-2 into human host cells has been identified to use the same receptor as SARS-CoV.

Most CoVs share a similar viral structure and infection pathway, and therefore the infection mechanisms previously found for other CoVs may also be applicable for SARS-CoV-2. A growing body of evidence shows that neurotropism is one common feature of CoVs. Therefore, it is urgent to make clear whether SARS-CoV-2 can gain access to the central nervous system (CNS) and induce neuronal injury leading to the acute respiratory distress.

2 | THE CLINICAL FEATURES OF SARS-CoV-2 INFECTION

SARS-CoV-2 causes acute, highly lethal pneumonia with clinical symptoms similar to those reported for SARS-CoV and MERS-CoV. Imaging examination revealed that most patients with fever, dry cough, and dyspnea showed bilateral ground-glass opacities on chest computerized tomography scans. However, different from SARS-CoV, SARS-CoV-2-infected patients rarely showed prominent upper respiratory tract signs and symptoms, indicating that the target cells of SARS-CoV-2 may be located in the lower airway.

Based on the first-hand evidence from Wuhan local hospitals, the common symptoms of COVID-19 were fever (83%-99%) and dry cough (59.4%-82%) at the onset of illness. However, the most characteristic symptom of patients is respiratory distress (~55%). Among the patients with dyspnea, more than half needed intensive care. About 46% to 65% of the patients in the intensive care worsened in a short period of time and died due to respiratory failure. Among the 36 cases in the intensive care reported by Wang et al., 11.1% received high-flow oxygen therapy, 41.7% received noninvasive ventilation, and 47.2% received invasive ventilation. These data suggest that most (about 89%) of the patients in need of intensive care could not breathe spontaneously.

It is now known that CoVs are not always confined to the respiratory tract and that they may also invade the CNS inducing neurological diseases. Such neuroinvasive propensity of CoVs has been documented almost for all the βCoVs, including SARS-CoV, MERS-CoV, HCoV-229E, HCoV-OC43, mouse hepatitis virus, and porcine hemagglutinating encephalomyelitis coronavirus (HEV).

With respect to the high similarity between SARS-CoV and SARS-CoV2, it remains to know whether the potential neuroinvasion of SARS-CoV-2 plays a role in the acute respiratory failure of patients with COVID-19.

3 | THE NEUROINVASIVE POTENTIAL OF SARS-CoV-2

It is believed that the tissue distributions of host receptors are generally consistent with the tropisms of viruses. The entry of SARS-CoV into human host cells is mediated mainly by a cellular receptor angiotensin-converting enzyme 2 (ACE2), which is expressed in human airway epithelia, lung parenchyma, vascular endothelia, kidney cells, and small intestine cells. Different from SARS-CoV, MERS-CoV enters human host cells mainly via dipeptidyl peptidase 4 (DPP4), which is present in the lower respiratory tract, kidney, small intestine, liver, and the cells of the immune system.

However, the presence of ACE2 or DPP4 solely is not sufficient to make host cells susceptible to infection. For example, some ACE2-expressing endothelial cells and human intestinal cell lines failed to be infected by SARS-CoV while some cells without a detectable expression level of ACE2, such as hepatocytes could also be infected by SARS-CoV. Likewise, the infection of SARS-CoV or MERS-CoV was also reported in the CNS, where the expression level of ACE2 or DPP4 is very low under normal conditions.

Early in 2002 and 2003, studies on the samples from patients with SARS have demonstrated the presence of SARS-CoV particles in the brain, where they were located almost exclusively in the neurons. Experimental studies using transgenic mice further revealed that either SARS-CoV or MERS-CoV, when given in transnasally, could enter the brain, possibly via the olfactory nerves, and thereafter rapidly spread to some specific brain areas including thalamus and brainstem. It is noteworthy that in the mice infected with low inoculum doses of MERS-CoV virus particles were detected only in the brain, but not in the lung, which indicates that the infection in the CNS was more important for the high mortality observed in the infected mice. Among the involved brain areas, the brainstem has been demonstrated to be heavily infected by SARS-CoV or MERS-CoV.

The exact route by which SARS-CoV or MERS-CoV enters the CNS is still not reported. However, hematogenous or lymphatic route seems impossible, especially in the early stage of infection, since almost no virus particle was detected in the nonneuronal cells in the infected brain areas. On the other hand, increasing evidence shows that CoVs may first invade peripheral nerve terminals, and then gain access to the CNS via a synapse-connected route.

The trans-synaptic transfer has been well documented for other CoVs, such as HEV679.10,18,19 and avian bronchitis virus.36,37 HEV 67N is the first CoV found to invade the porcine brain, and it shares more than 91% homology with HCoV-OC43.38,39 HEF first oronasally infects the nasal mucosa, tonsil, lung, and small intestine in suckling piglets, and then is delivered retrogradely via peripheral nerves to the medullary neurons in charge of peristaltic function of the digestive tract, resulting in the so-called vomiting diseases. The transfer of HEV 67N between neurons has been demonstrated by our previous ultrastructural studies to use the clathrin-coating-mediated endocytotic/exocytotic pathway.
Similarly, the trans-synaptic transfer has been reported for avian bronchitis virus. Intranasal inoculation in mice with avian influenza virus was reported to cause neural infection besides bronchitis or pneumonia. Of interest, viral antigens have been detected in the brainstem, where the infected regions included the nucleus of the solitary tract and nucleus ambiguus. The nucleus of the solitary tract provides sensory information from the mechanoreceptors and chemoreceptors in the lung and respiratory tracts, while the efferent fibers from the nucleus ambiguus and the nucleus of the solitary tract provide innervation to airway smooth muscle, glands, and blood vessels. Such neuroanatomical interconnections indicate that the death of infected animals or patients may be due to the dysfunction of the cardiorespiratory center in the brainstem. 

Taken together, the neuroinvasive propensity has been demonstrated as a common feature of CoVs. In light of the high similarity between SARS-CoV and SARS-CoV-2, it is quite likely that SARS-CoV-2 also possesses a similar potential. Based on an epidemiological survey on COVID-19, the median time from the first symptom to dyspnea was 5.0 days, to hospital admission was 7.0 days, and to the intensive care was 8.0 days. Therefore, the latency period may be enough for the virus to enter and destroy the medullary neurons. As a matter of fact, the previous studies mentioned above has reported that some patients infected with SARS-CoV-2 did show neurologic signs such as headache (about 8%), nausea and vomiting (1%). More recently, one study on 214 COVID-19 patients by Mao et al. further found that about 88% (78/88) among the severe patients displayed neurologic manifestations including acute cerebrovascular diseases and impaired consciousness. Therefore, awareness of the possible neuroinvasion may have a guiding significance for the prevention and treatment of the SARS-CoV-2-induced respiratory failure.

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REFERENCES
1. Glass WG, Subbarao K, Murphy B, Murphy PM. Mechanisms of host defense following severe acute respiratory syndrome-coronavirus (SARS-CoV) pulmonary infection of mice. J Immunol. 2004;173:4030-4039.
2. Huang C, Wang Y, Li X, et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. Lancet. 2020;395:497-506.
3. Yu F, Du L, Ojcius DM, Pan C, Jiang S. Measures for diagnosing and treating infections by a novel coronavirus responsible for a pneumonia outbreak originating in Wuhan, China. Microbes Infect. 2020. https://doi.org/10.1016/j.micinf.2020.01.003
4. Song Z, Xu Y, Bao L, et al. From SARS to MERS, thrusting coronaviruses into the spotlight. Viruses. 2019;11:59. https://doi.org/10.3390/v111010059
5. Lu R, Zhao X, Li J, et al. Genomic characterisation and epidemiology of 2019 novel coronavirus: implications for virus origins and receptor binding. Lancet. 2020;395:565-574.
6. Wan Y, Shang J, Graham R, Baric RS, Li F. Receptor recognition by novel coronavirus from Wuhan: an analysis based on decade-long structural studies of SARS. J Virol. 2020. https://doi.org/10.1128/JVI.00127-20
7. Yuan Y, Cao D, Zhang Y, et al. Cryo-EM structures of MERS-CoV and SARS-CoV spike glycoproteins reveal the dynamic receptor binding domains. Nat Commun. 2017;8:15092.
8. Hulswit RJ, de Haan CA, Bosch BJ. Coronavirus spike protein and tropism changes. Adv Virus Res. 2016;96:29-57.
9. Li YC, Bai WZ, Hirano N, Hayashida T, Hashikawa T. Coronavirus infection of rat dorsal root ganglia: ultrastructural characterization of viral replication, transfer, and the early response of satellite cells. Virus Res. 2012;163:628-635.
10. Wang D, Hu B, Hu C, et al. Clinical characteristics of 138 hospitalized patients with 2019 novel coronavirus-infected pneumonia in Wuhan, China. JAMA. 2020. https://doi.org/10.1001/jama.2020.1585
11. Khan S, Ali A, Siddique R, Nabi G. Novel coronavirus is putting the whole world on alert. J Hosp Infect. 2020. https://doi.org/10.1016/j.jhin.2020.01.019
12. Chen N, Zhou M, Dong X, et al. Epidemiological and clinical characteristics of 99 cases of 2019 novel coronavirus pneumonia in Wuhan, China: a descriptive study. Lancet. 2020;395(10223):507-513.
13. Li K, Wohlford-Lenane C, Perlman S, et al. Middle East respiratory syndrome coronavirus causes multiple organ damage and lethal disease in mice transgenic for human dipeptidyl peptidase 4. J Infect Dis. 2016;213:712-722.
14. Talbot PJ, Ekandé S, Cashman NR, Mounir S, Stewart JN. Neurotropism of human coronavirus 229E. Adv Exp Med Biol. 1993;342:339-346.
15. Dubé M, Le Coupéan A, Wong AHM, Rini JM, Desorges M, Talbot PJ. Axonal transport enables neuron-to-neuron propagation of human coronavirus OC43. J Virol. 2018;92. https://doi.org/10.1128/JVI.00404-18
16. Zhou X, Huang F, Xu L, et al. Hepatitis E virus infects neurons and brains. J Infect Dis. 2017;215(8):1197-1206.
17. Yu Y, Bai WZ, Hirano N, et al. Neurotropic virus traces suggesting a membranous-coating-mediated mechanism for transsynaptic communication. J Comp Neurol. 2013;521:203-212.
18. Mengeling WL, Bothe AD, Ritchie AE. Characteristics of a coronavirus (strain 67N) of pigs. Am J Vet Res. 1972;33(2):297-308.
19. Andries K, Pensaert MB. Immunofluorescence studies on the pathogenesis of hemagglutinating encephalomyelitis virus infection in pigs after oronasal inoculation. Am J Vet Res. 1980;41(9):1372-1378.
20. To KF, Lo AW. Exploring the pathogenesis of severe acute respiratory syndrome (SARS): the tissue distribution of the coronavirus (SARS-CoV) and its putative receptor, angiotensin-converting enzyme 2 (ACE2). J Pathol. 2004;203:740-743.
21. Tang JW, To KF, Lo AW, Sung JJ, Ng HK, Chan PK. Quantitative temporal-spatial distribution of severe acute respiratory syndrome-associated coronavirus (SARS-CoV) in post-mortem tissues. J Med Virol. 2007;79:1245-1253.
22. Kam YW, Okumura Y, Kido H, Ng LF, Bruzzone R, Altmeyer R. Cleavage of the SARS coronavirus spike glycoprotein by airway proteases enhances virus entry into human bronchial epithelial cells in vitro. PLoS One. 2009;4(11):e7870.
23. Donoghue M, Hsieh F, Baronas E, et al. A novel angiotensin-converting enzyme-related carboxypeptidase (ACE2) converts angiotensin I to angiotensin 1-9. Circ Res. 2000;87(5):E1-E9.
24. Harmer D, Gilbert M, Borman R, Clark KL. Quantitative mRNA expression profiling of ACE 2, a novel homologue of angiotensin converting enzyme. FEBS Lett. 2002;532:107-110.
25. Hamming I, Timens W, Bulthus ML, Lely AT, Navis G, van Goor H. Tissue distribution of ACE2 protein, the functional receptor for SARS coronavirus. A first step in understanding SARS pathogenesis. J Pathol. 2004;203(2):631-637.
26. Mattern T, Scholz W, Flad HD, Ulmer AJ. Expression of human coronavirus 229E in colonic cells in vitro. Adv Exp Med Biol. 1993;342:339-346.
27. Boonacker E, Van Noorden CJ. The multifunctional or moonlighting protein CD26/DPPIV. Eur J Cell Biol. 2003;82:53-73.
28. Chan PK, To KF, Lo AW, et al. Persistent infection of SARS coronavirus in colonic cells in vitro. J Med Virol. 2004;74:1-7.
29. Ding Y, Wang H, Shen H, et al. The clinical pathology of severe acute respiratory syndrome (SARS): a report from China. J Pathol. 2003;200:282-289.

30. Bernstein HG, Dobrowolny H, Keilhoff G, Steiner J. Dipeptidyl peptidase IV, which probably plays important roles in alzheimer disease (AD) pathology, is upregulated in AD brain neurons and associates with amyloid plaques. Neurochem Int. 2018;114:55-57.

31. Ding Y, He L, Zhang Q, et al. Organ distribution of severe acute respiratory syndrome (SARS) associated coronavirus (SARS-CoV) in SARS patients: implications for pathogenesis and virus transmission pathways. J Pathol. 2004;203:622-630.

32. Gu J, Gong E, Zhang B, et al. Multiple organ infection and the pathogenesis of SARS. J Exp Med. 2005;202(3):415-424.

33. Xu J, Zhong S, Liu J, et al. Detection of severe acute respiratory syndrome coronavirus in the brain: potential role of the chemokine mig in pathogenesis. Clin Infect Dis. 2005;41:1089-1096.

34. Netland J, Meyerholz DK, Moore S, Cassell M, Perlman S. Severe acute respiratory syndrome coronavirus infection causes neuronal death in the absence of encephalitis in mice transgenic for human ACE2. J Virol. 2008;82:7264-7275.

35. McCray PB Jr, Pewe L, Wohlford-Lenane C, et al. Lethal infection of K18-hACE2 mice infected with severe acute respiratory syndrome coronavirus. J Virol. 2007;81:813-821.

36. Matsuda K, Park CH, Sunden Y, et al. The vagus nerve is one route of transneural invasion for intranasally inoculated influenza virus in mice. Vet Pathol. 2004;41:101-107.

37. Chasey D, Alexander DJ. Morphogenesis of avian infectious bronchitis virus in primary chick kidney cells. Arch Virol. 1976;52:101-111.

38. González JM, Gomez-Puertas P, Cavanagh D, Gorbalenya AE, Enjuanes L. A comparative sequence analysis to revise the current taxonomy of the family coronaviridae. Arch Virol. 2003;148:2207-2235.

39. Li Z, He W, Lan Y, et al. The evidence of porcine hemagglutinating encephalomyelitis virus induced nonsuppurative encephalitis as the cause of death in piglets. Peer J. 2016;4:e2443.

40. Kalia M, Mesulam MM. Brain stem projections of sensory and motor components of the vagus complex in the cat: II. Laryngeal, tracheobronchial, pulmonary, cardiac, and gastrointestinal branches. J Comp Neurol. 1980;193:467-508.

41. Hadzieficndic S, Haxhiu MA. CNS innervation of vagal preganglionic neurons controlling peripheral airways: a transneuronal labeling study using pseudorabies virus. J Auton Nerv Syst. 1999;76:135-145.

42. Raux H, Flamand A, Blondel D. Interaction of the rabies virus P protein with the LC8 dynein light chain. J Virol. 2000;74:10212-10216.

43. Mao L, Wang M, Chen S, He Q, Chang J, Hong C, Zhou Y, Wang D, Li Y, Jin H, Hu B. Neurological Manifestations of Hospitalized Patients with COVID-19 in Wuhan, China: a retrospective case series study. MedRxiv. https://doi.org/10.1101/2020.02.22.20026500

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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