The emergence of cognitive COVID

The scale of the COVID-19 pandemic has impacted health care systems on a global level. As the pandemic moves into its second year, attention is beginning to turn towards the medium- and long-term consequences of the infection. High on the list of priorities is the issue of cognitive impairment, not only as a direct effect of neurotropic viral brain infiltration but also due to indirect factors associated with the pandemic, such as increased social isolation and mental health problems.

While associations between neurotropic respiratory viruses and brain changes have been documented since the 1918 influenza epidemic, the cognitive consequences of these changes have until now received very little attention. The increasing interest in both the spread of coronaviruses to the central nervous system (CNS) and the longer-term clinical presentations of infected individuals has led to a re-evaluation of the importance of cognitive changes.

A meta-analysis of 3,559 adult cases collectively drawn from the severe acute respiratory syndrome (SARS), Middle East respiratory syndrome (MERS) and COVID-19 epidemics identified memory impairment in one third of cases at hospital admission and in 19% of cases post-illness, with the latter notably also affecting younger adults. Initial studies indicate that cognitive dysfunction may extend beyond the acute stage of COVID-19 infection. A study of 18 patients with mild to moderate COVID-19 disease (not requiring intensive care unit admission) and a mean age of 42 years, examined a median of 85 days after recovery, found that over 75% had episodic memory, attention and concentration difficulties which were not associated with fatigue, depression, hospitalization, treatment, viremia or acute inflammation. These initial data indicate that cognitive changes may occur even after milder infections.

Given the scale of the pandemic and the implications for both working age adults and the older population at risk of dementia, these emerging data highlight the urgent need to better understand the mechanisms resulting in cognitive dysfunction, with a view to introducing interventions and public health strategies to combat these deleterious longer-term effects of the pandemic.

The effect of SARS-CoV-2 on cognition may relate to the vulnerability of various CNS cells to the virus and to its direct infiltration of the CNS. Viral attachment to host cells results from binding of the S1 subunit of the S protein, one of four structural proteins of the SARS-CoV-2 virion, to the angiotensin-converting enzyme 2 (ACE2) receptor on cell surfaces, with subsequent intracellular entry of the viral genome occurring after fusion of viral and host cell membranes. As such, the cellular tropism of SARS-CoV-2 relates to the expression of the ACE2 receptor. Outside the CNS, the receptor is expressed in alveoli, gut, kidney and epidermis, as well as vascular endothelial cells. Within the CNS, it is expressed in neurons, astrocytes, oligodendrocytes and endothelial cells. Regionally, high concentrations of the ACE2 receptor are found in the olfactory bulb, substantia nigra, middle temporal gyrus, and posterior cingulate gyrus.

Two direct mechanisms underpin the neurotropism of SARS-CoV-2 and its access to the CNS: a) retrograde axonal transport following invasion of peripheral olfactory neurons, and b) haematogenous breach of the blood-brain barrier following infection of this barrier or choroid plexus endothelial cells. The pathological effect of this direct viral infiltration is augmented by a brisk immune response and inflammation, with the associated cytokine storm further compromising the blood-brain barrier, by vasculopathy arising from disseminated intravascular coagulation, and by hypoxaemia.

The resultant clinical manifestations of this CNS pathology are multiple. They include inflammatory disorders (meningoencephalitis, acute disseminated encephalomyelitis), encephalopathies presenting with behavioural disturbances, seizures, and cerebrovascular disease (both thrombotic and haemorrhagic). The prevalence of CNS manifestations in severe infection is high: of 58 patients with acute respiratory distress syndrome, 69% had agitation and 65% had confusion, with a high proportion of those imaged showing magnetic resonance imaging (MRI) changes in the form of altered perfusion, ischaemic stroke and leptomeningeal enhancement.

The relative recency of the pandemic means that there are at present only limited data on the impact of COVID-19 infection on cognitive function beyond the acute illness. However, both direct and indirect effects of the infection indicate a likelihood of longer-term cognitive impairment. SARS-CoV-2 invasion of peripheral olfactory neurons, now recognized as one component of the virally-induced acute anosmia, permits trans-synaptic viral spread to cortical regions receiving primary and secondary input from the olfactory tract, notably the entorhinal cortex and the hippocampus. The involvement of these regions in episodic memory and spatial navigation raises the possibility of COVID-19 infection causing longer-term impairment in these cognitive domains. This will be amplified by indirect consequences of the infection in terms of other pathophysiological effects, notably virally-mediated vascular pathology and inflammatory responses, psychological trauma and need for critical care. Preliminary estimates of the prevalence and timescales of such effects can be gleaned from previous neuropsychological studies of long-term post-ventilation outcomes, with cognitive impairment observed in 78% of patients at one year and with memory problems persisting up to five years in around 50%, independent of psychological problems.

Finally, there is the potential risk that COVID-19 infection may cause long-term cognitive decline by accelerating the onset of neurodegenerative dementia. The severity of the infection is greater at higher ages, and the neural pathways along which SARS-CoV-2 may be transported overlap with those implicated at the onset of Parkinson’s and Alzheimer’s disease, such as the cognitively eloquent regions within the medial temporal lobe. This overlap in regional vulnerability may provide the anatomical basis for an interaction between SARS-CoV-2 and neurode-
Post-traumatic stress disorder in the aftermath of COVID-19 pandemic

Post-traumatic stress disorder (PTSD) is a potentially debilitating mental health disorder which affects an important minority of people exposed to events involving actual or threatened death, serious injury or sexual violence. The COVID-19 pandemic is unfortunately providing multiple opportunities for people to experience traumatic situations which may lead to PTSD.

Imagine the previously fit person who rapidly goes from an active lifestyle to a chemical induced coma, surviving only after weeks on a mechanical ventilator. Or the nurse who volunteers to join a rapidly assembled intensive care team with minimal preparatory training, and faces the stark reality that many of those cared for end up dying alone, with relatives being unable to visit the unit. These situations have a high potential to induce PTSD. Indeed, it has been reported that up to 20% of intensive care unit survivors go on to develop PTSD. On the other hand, there is evidence that repeated exposure to traumatic events in health care workers can lead to the development of PTSD even if the staff member cannot identify which specific traumatic event caused him/her to become unwell.

Whilst PTSD must follow trauma exposure, other factors substantially influence the likelihood of developing this condition. Comprehensive meta-analyses of risk factors for PTSD consistently find that the nature of the post-trauma environment is a more important predictor than pre-traumatic factors such as childhood adversity, or demographic factors such as gender or ethnicity. In particular, there is strong evidence that psychological stress experienced during the initial post-exposure period, as well as the availability and quality of post-trauma social support, are highly influential determinants. Whilst we know that social support is highly protective against the development of PTSD, social distancing restrictions are making it more difficult for people to access non-professional support, so that the onset of PTSD after trauma exposure may become more likely.

Another important risk factor for PTSD is moral injury, which is defined as the psychological distress, including feelings of deep shame and guilt, resulting from doing, or not preventing, events that someone believes are “wrong”. Many health care workers are likely to experience morally injurious events during this pandemic. Feeling unable to deliver high-quality care, or having to make hard choices about who will and who will not receive a given intervention due to shortage of available equipment, have become somewhat commonplace, especially when the rates of hospitalization are high. Moral injury is also a relevant concept outside of work environments, especially when people are concerned about having infected loved ones who have died. Moral injury is important as it can predispose people to developing PTSD as well as making it less likely that they will seek treatment if they do.

Within organizational settings, a number of approaches have been tried to prevent the onset of PTSD. Pre-employment, or pre-role, psychological health screening aims to identify higher risk individuals, so they can either not be employed in trauma-exposed roles or be provided with extra support to mitigate the risk. However, there is consistent evidence that this approach is ineffective. It may indeed be harmful, by providing employers with false reassurance that screened personnel are resilient to trauma and will not develop PTSD. Whilst health care managers understandably may wish to exclude vulnerable staff from dealing with the most severe COVID-19 patients, in order to protect their mental health, the reality is that the state of the current evidence base on screening is unsatisfactory and this practice cannot be recommended.

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