Demyelination as a result of an immune response in patients with COVID-19

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Abstract
The coronavirus disease of 2019 (COVID-19) is caused by the severe acute respiratory syndrome coronavirus-2 (SARS CoV-2), that already appeared as a global pandemic. Presentation of the disease often includes upper respiratory symptoms like dry cough, dyspnea, chest pain, and rhinorrhea that can develop to respiratory failure, needing intubation. Furthermore, the occurrence of acute and subacute neurological manifestations such as stroke, encephalitis, headache, and seizures are frequently stated in patients with COVID-19. One of the reported neurological complications of severe COVID-19 is the demolition of the myelin sheath. Indeed, the complex immunological dysfunction provides a substrate for the development of demyelination. Nevertheless, few published reports in the literature describe demyelination in subjects with COVID-19. In this short narrative review, we discuss probable pathological mechanisms that may trigger demyelination in patients with SARS-CoV-2 infection and summarize the clinical evidence, confirming SARS-CoV-2 condition as a risk factor for the destruction of myelin.

Keywords COVID-19 · SARS-CoV-2 · Demyelination · Immune response · Multiple sclerosis

Introduction
The outbreak of novel coronavirus disease known as severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2), first identified in Wuhan, China, in December 2019. Previously, other coronaviruses were recognized to cause severe respiratory disease in humans and were associated with epidemic prevalence, such as the severe acute respiratory syndrome coronavirus (SARS-CoV) and the Middle East respiratory syndrome coronavirus (MERS-CoV), both enhancing the risk of mortality [1]. The average diameter of the COVID virus is almost 100 nm and is spherical or oval in shape. The large spikes of membrane glycoproteins of the virus are located on the viral surface, and these negatively stained virus particles are expressed as typical crown-like shapes under electron microscopy [2]. The viral genome encodes several important structural proteins, including spike, membrane, envelope, and nucleocapsid proteins [3].

Similar to the previously identified SARS and MERS diseases, the main indications of COVID-19 are one or a combination of fever, fatigue, dry cough, and shortness of breath, followed by other symptoms like headache, nasal congestion, sore throat, myalgia, or arthralgia [4, 5]. However, extra respiratory complications such as neurological findings have been reported. Neurological manifestations, including headache, disturbed consciousness, paresthesia, and seizures are appeared in almost 36.4% (78/214) of patients suffering from COVID-19 [6]. On March 4, 2020, Beijing Ditan Hospital, for the first time, reported a patient with viral encephalitis instigated by a novel coronavirus attacking the central nervous system (CNS). This study illustrated the potential effect of COVID-19 in triggering damage to the nervous system [7]. According to accumulating evidence, the different types of Coronaviruses can induce neurological disorders such as polyneuropathy, encephalopathy, ischemic stroke, and demyelinating lesions, like Multiple sclerosis (MS) and Guillain-Barré syndrome (GBS) [8, 9]. The neurological problems could be presented delayed to respiratory signs [8], and severely affected patients are more vulnerable to neurological manifestations related to patients with mild or moderate disease [10]. Amongst neurological impairments, demolition of myelin appears as a visible complication in patients

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with severe COVID-19 [6]. MS is an acquired inflammatory immune-mediated disorder of the CNS, characterized by the focal demyelination associated with various degrees of neurodegeneration [11]. GBS, as another demyelinating disease, is an acute/subacute inflammatory polyneuropathy and immune-mediated disease triggered by a preceding bacterial or viral infection [12]. To support this assumption, the existence of demyelination in cases with COVID-19 has been reported, as well as particles of the SARS-CoV virus and genome sequences have been found in autopsy reports [13, 14]. Therefore, this review discusses the involvement of COVID-19 in triggering immunological host response and demyelination in central and peripheral nervous systems.

The entrance of SARS-CoV-2 virus into the CNS

Similar to other Coronaviruses, two possibilities appear for penetration of SARS-CoV-2 into the CNS, including (1) hematogenous dissemination of SARS-CoV-2 and rapidly spreading through the cerebral circulation, where the slower blood flow accelerates the capillary endothelium damages and makes the brain accessible, and (2) gaining entry through the cribriform plate and olfactory bulb [15]. Here, infectious agents like coronaviruses can spread through cranial nerves, including the olfactory, the trigeminal, the glosso-pharyngeal, and the vague nerves, via retrograde axonal transport [16]. Notably, the entrance of SARS-CoV-2 into the brain through the olfactory nerves infects neurons that control breathing [17]. Similar to SARS-CoV, the SARS-CoV-2 and a host receptor, Angiotensin-converting enzyme 2 (ACE2) have a stronger binding capacity [18]. ACE2, as a cardio-cerebral vascular protection factor, is capable of converting the angiotensin (Ang) I to Ang II that activates Ang II receptor type1 (AT1R), promoting the blood pressure, inflammation, and even neurodegeneration [19]. This receptor is present in human endothelial cells, neurons, respiratory epithelia, pulmonary parenchyma, small bowel cells, renal cells, and nerve cells, enabling the dispersal of SARS-CoV-2 into the host cells [20, 21]. Thus, the wide expression of the ACE2 receptor on nerve cells and capillary endothelium allows the direct access of SARS-CoV-2 to the CNS [22–24], where the virus binds to nerve cells via attachment between the spike glycoprotein and the existent ACE2 receptors [25]. On the other hand, the interaction of SARS-CoV-2 with ACE-2 receptors in the vascular endothelial cells may be conductive of the blood–brain-barrier (BBB) destruction, leading to the direct virus entry into the brain [26, 27]. In addition to ACE2, basigin (BSG; CD147) and neuropilin-1 (NRP1) act as docking receptors for SARS-CoV-2 virus, while several proteases including transmembrane protease serine 2 (TMPRSS2), cathepsin B and L (CatB/L), and furin are implicated in the virus entry to cell and replication [28]. NRPI, a transmembrane receptor lacking a cytosolic protein kinase domain, is highly expressed in the olfactory epithelium and improves the SARS-CoV-2 entry into the brain [29]. CD147 (also known as Basigin) is an alternative receptor for SARS-CoV-2 invasion into the brain, which is present on neural cells [30]. Furthermore, other endosomal cysteine proteases, namely CatB/L contribute in priming of the spike protein of SARS-CoV-2 virus [31]. Hoffmann et al. demonstrated that SARS-CoV-2 uses ACE2 for entry and needs the TMPRSS2 for S protein priming to efficiently enter the cell, and TMPRSS2 inhibitor can block virus entry to exert a treatment effect [32]. TMPRSS2 is expressed throughout olfactory epithelial cells and choroid plexus of both mouse and human [33, 34]. The other results indicated that TMPRSS2 and lysosomal cathepsins are capable of activating SARS-CoV-2 entry; moreover, furin has cumulative effects on this process [35].

Furthermore, another avenue for the accession of SARS-CoV-2 to the host cells is binding to integrins. These receptors are present on the surface of different cells in the body and contribute in triggering several signaling pathways [36].

Evidence of Covid-19 virus involvement in demyelination

Accumulating evidence confirms the SARS-CoV-2 viral infection as a risk factor of demyelination both in the peripheral and central nervous systems. In agreement with this idea, Zanin et al. reported a case of COVID-19 admitted for interstitial pneumonia and seizures. Brain MRI indicated newly diagnosed demyelination injuries. However, treatment with high-dose steroid resulted in respiratory and neurological recovery. They postulated that SARS-CoV-2-induced delayed immune response was a causative factor for the demolition of the myelin sheath [37].

In another report, Mehta et al. detected demyelinating lesions associated with the neurological damage in a case with COVID-19. The brain and spine MRI of the patient exhibited a new onset of multiple, non-enhancing demyelinating lesions. They assumed that following SARS-CoV-2 infection, the pro-inflammatory environment induced by the cytokine storm might be responsible for the activation of glial cells with subsequent demyelination [38]. Furthermore, Zoghi et al. reported a 21-year-old male with encephalomyelitis after experiencing intermittent 4 days vomiting and malaise that showed upper respiratory indications 2 weeks before this presentation. Brain MRI revealed bilateral posterior internal capsule lesions expanding to the ventral part of the pons and a marbled splenium hyperintensity pattern. In their study, Cervical and thoracic MRI showed the sizable transverse myelitis [39]. In another report, Gilles et al.
stated a 54-year-old woman affected to SARS-CoV-2 with brain lesions representing acute demyelination. The brain CT scan of the patient confirmed hypodense lesions, involving supratentorial white matter and pallidum bilaterally. A brain MRI showed lesions with restricted diffusion without any hemorrhage or enhancement. The thalamus, the striatum, and the posterior fossa were spared. A follow-up MRI displayed no new lesions, and the spinal cord MRI was without abnormalities [40].

In another case report, a 33-year-old male suffering from morbid obesity, diabetes mellitus, and hypoxemic showed a respiratory failure secondary to COVID-19 affection. MRI presented widespread foci of the demolition of the myelin with restricted diffusion, most located in the corpus callosum and the pericallosal white matter [41]. Furthermore, Karapanayiotides et al. described a 57-year-old male affected by COVID-19 along with particular localization and a concentric demyelination pattern. MRI documented the occurrence of hemosiderin deposits concurrent with a concentric demyelination pattern [42].

A recent study described a COVID-19 patient with isolated symmetrical demyelinating lesions of bilateral posterior internal capsules that presented stroke-like clinical demonstration. The lack of grey matter damage and absence of hemorrhage or cavitation confirmed demyelination, rather than encephalomyelitis [43].

In support of the involvement of SARS-CoV-2 infection in GBS and consequent demyelination, Bracaglia described a 66-year-old woman with acute demyelinating polynu-ritis related to asymptomatic SARS-CoV-2 infection. She was unable to walk, speech and swallow, as well as tendon reflexes were abolished. The patient showed normal vital signs, adverse medical history and no report of the previous month infection. They assumed GBS and conducted nerve conduction studies consistent with demyelinating polyneuropathy [44].

The immunopathological pathway resulting in COVID-19 mortality contains three stages, that progress rapidly from onset of disease to death over 16–18.5 days. Stage 1 is the initiation phase, with early signs of systemic inflammation, early induction of predominant chemokines, and decreased peripheral lymphocyte counts. Stage 2 is the amplification phase, with a massive production of inflammatory cells, including macrophages, monocytes, and neutrophils to intensify the immunopathological process. Stage 3 is the consummation phase. In this stage, the peripheral neutrophil counts further increase, while lymphopenia worsens with continuous enhancement of inflammatory mediators and cytokine storm, leading to the widespread organ damages [48].

Most patients with COVID-19 display normal or decreased leukocyte count [49–51], and lymphopenia [52]. Lymphopenia is a common feature in patients with severe COVID-19, but not in patients with a mild infection [53]. In this condition, the numbers of CD4+ T cells, CD8+ T cells, B cells, and natural killer (NK) cells subsets are extremely reduced in the lymph nodes and spleen of the patients (Fig. 1) [54]. Besides, in stage 2, the percentage of monocytes and macrophages is increased following viral infection, which leads to the release of a large amount of pro-inflammatory cytokines, referred to as cytokine storm [55, 56]. This phenomenon is an enhancement of serum levels of pro-inflammatory cytokines including IL-6 and IL-1β, as well as IL-2, IL-8, IL-17, G-CSF, GM-CSF, IP10, MCP1, MIP1α (also known as CCL3), and TNF, which has found in most patients with severe COVID-19 [57]. Accordingly, a dysfunctional and disharmonic immune response in severe COVID-19 cases initiates an extensive lung and systemic inflammation by triggering cytokine storm, that possibly worsens infection in the brain [58]. The cytokine storm targets the CNS because the released cytokines are able to cross the BBB by disrupting the integrity of this structure [59]. This process allows the penetration of monocytes, macrophages, and T-lymphocytes into the brain [60]. The cytokine storm induced by SARS-CoV-2 infection, at least in part caused by the toll-like receptors (TLRs) involvement in the immune responses in the CNS (Fig. 2). In the normal innate immune system, various pattern recognition receptors (PRRs) are expressed in macrophages, monocytes, dendritic cells, and neutrophils that recognize pathogen-associated molecular patterns (PAMPs), which are expressed by infectious agents. PRRs are the fundamental players in mediating innate immune response and comprise TLRs and nod-like receptors (NLRs) [61]. It has postulated that TLRs are the main PRRs, taking part in the induction of cytokine storm in the COVID-19-infected patients [61, 62]. Ten TLRs, TLR1-10 have been recognized in humans, while TLR1-13, with the exemption of TLR10 are present in rodents [63]. It is remarkable that IL-6 and TNF-α, the main cytokines that

**Immunological host response against SARS-CoV-2 infection**

It is worth noting that the severe form of COVID-19 causes not only the viral infection, but also leads to an abnormal and aggravated immunological host response, resulting in severe systemic damage [45]. SARS-CoV-2 infection is capable of triggering innate and adaptive immune responses. Uninhibited inflammatory innate responses and debilitated adaptive immune responses may cause harmful locally and systemically tissue damage [46]. Accordingly, unpredictable effects of acute viral infection on the host immune response is probably an important reason for nervous tissue damage [47].
participated in severe COVID-19 cases, are downstream of the TLR4 signaling pathway [64]. TLRs are expressed by monocytes, astrocytes, macrophages, and dendritic cells, which are the primary operators of the innate immune system and take part in host defense and the recognition of invading pathogens, suggesting the role of these receptors in MS pathophysiology [65]. Interestingly, these receptors identify viral particles and are involved in the modulation of the immune response in MS patients. Apparently, there may be an association between viral infections, such as coronavirus, and the growth of demyelination [65]. Two necessary proteins that activate these receptors are the cluster of differentiation 14 (CD14) and myeloid differential protein-2 (MD-2), triggering two internal signaling pathways, including the myeloid differentiating primary response gene 88 (MyD88)-dependent and the MyD88-independent pathways, which is known as toll/interleukin-1 receptor (TIR)-domain-containing adapter-inducing interferon-β (TRIF)-dependent signaling pathway (Fig. 2) [66]. With the exemption of TLR3, all other TLRs utilize the MyD88-dependent signaling pathway. TLR3 uses the MyD88-independent signaling pathway [67]. TLR4 is the only TLR that employs both the MyD88-dependent and the TRIF-dependent signaling pathways. TLRs trigger the internal signaling pathways to activate some transcription factors responsible for the generation of pro-inflammatory cytokines, like nuclear factor-kβ (NF-κβ), as well as interferon regulatory factors (IRF) that mediates the type I interferon (IFNs)-dependent antiviral response [62, 68]. Activation of TLRs initiates a cell signaling cascade, leading to the release of several interleukins, interferons, and other pro-inflammatory cytokines [62]. Activation of diverse TLRs can lead to a robust enhancement in cytokine release, inducing the cytokine storm in the CNS [69]. The main cytokines involved in the cytokine storm in the CNS are IL-6, interferons and IL-10 that result from astrocyte and microglia TLR signaling. Microglial cells and astrocytes are CNS immune cells and the cytokine storm that mostly produced by microglia can increase BBB permeability and may be responsible for neurological manifestations of COVID-19 [70, 71]. Some of the produced cytokines lead to the death of neurons and oligodendrocytes and, subsequently demyelination [71]. They also could increase glial activation and promote the expression of TLR in neural cells, exacerbating the cytokine storm and further inflammatory response [72]. Moreover, these signals are capable of attracting macrophages, NK cells, mast cells, and etc., which eventually may release reactive oxygen species (ROS) and reactive nitrogen species (RNS). An additional mechanism of CNS involvement and demyelination caused by autoimmune encephalitis in COVID-19 cases, a condition that results from extreme self-response and antigen-driven immune responses [74]. SARS-CoV-2 virus can elicit autoimmune responses through several mechanisms, including molecular mimicry between viral proteins and host self-antigens, epitope spreading due to T-cell-mediated damage by the virus, bystander activation of T cells by the action

![Diagram](https://via.placeholder.com/150)

**Fig. 1** Schematic representation of mechanisms that SARS-CoV-2 infection may result in demyelination. One crucial way is the decreased number of CD4+ T cells, CD8+ T cells, B cells, and natural killer (NK) cells. Furthermore, enormous cytokine release creates a pro-inflammatory condition that some of the produced cytokines lead to the death of neurons and oligodendrocytes, increase glial activation and promote the expression of TLR in neural cells. These signals are capable of attracting macrophages, NK, mast cells, and etc., which eventually may release reactive oxygen species (ROS) and reactive nitrogen species (RNS). Another mechanism is spontaneous or provoked autoimmune reactions and the generation of auto-antibodies against SARS-CoV-2 virus.
of virus-induced inflammatory cytokines, the presentation of cryptic antigens, poly-clonal B cell activation, and viral superantigens [75, 76]. The studies indicated the presence of high-level IgA and IgG antibodies (antineuronal antibodies) in the CSF and serum of patients, demonstrating that these antibodies can reach the brain [77, 78].

Thus, in patients with COVID-19, antibodies that are produced against SARS-CoV-2 attack antigens presented in human endothelial or neural cells (through the interrupted BBB), leading to cerebral edema and autoimmune encephalitis [25]. Given the evidence, SARS-CoV-2 infection may take part in proceeding MS and demyelination via several mechanisms, including reduction in the number of CD4+ T cells, CD8+ T cells, B cells, and NK cells, enormous cytokine release, creating a pro-inflammatory condition, and provoked autoimmune reactions (Fig. 1) [11].

**Conclusion**

This narrative review briefly described the mechanisms of CNS affection, altering the host immunological function, and induction of demyelination by SARS-CoV-2 infection, and summarized the evidence of the myelin sheath demolition induced by this viral infection. In summary, SARS-CoV-2 can enter the host cells in the brain and exert the neurologic symptoms via the binding to ACE2 and integrins presented on existing cells. Given the evidence, SARS-CoV-2 infection is able to cause MS and demyelination via several mechanisms. One important mechanism is the decreased number of CD4+ T cells, CD8+ T cells, B cells, and NK cells. Another mechanism is provoked autoimmune reactions, a condition which results from excessive self-response and antigen-driven immune responses. Thus, in patients with COVID-19, antibodies that are produced against SARS-CoV-2 attack antigens in human endothelial cells presented in cerebral vessels or neurons, leading to cerebral edema and autoimmune encephalitis. The most prominent type of mechanism is massive cytokines release by activating TLRs, which creates a pro-inflammatory environment. Some of the released cytokines induce neuronal and oligodendrocyte death and demyelination. They also could enhance glial activation and promote TLR expression in neural cells, intensifying the cytokine storm and inflammatory response.

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**Declarations**

**Conflict of interest** None.

**Ethical approval** The manuscript is not be submitted to more than one journal for simultaneous consideration. The submitted work is original and have not been published elsewhere in any form or language (partially or in full). A single study is not be split up into several parts to increase the quantity of submissions and is not submitted to various journals or to one journal over time (i.e. ‘salami-slicing/publishing’). Concurrent or secondary publication is sometimes justifiable, provided certain conditions are met. Examples include: translations or a manuscript that is intended for a different group of readers. Results are presented clearly, honestly, and without fabrication, falsification or inappropriate data manipulation (including image-based manipulation). Authors adhered to discipline-specific rules for acquiring, selecting and processing data. No data, text, or theories by others are presented as if they were the author’s own (‘plagiarism’). Proper acknowledgements to other works must be given (this includes material that is closely copied (near verbatim), summarized and/or paraphrased), quotation marks (to indicate words taken from another source) are used for verbatim copying of material, and permissions secured for material that is copyrighted.
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