Immune-mediated neurological syndrome in SARS-CoV-2 infection: a review of literature on autoimmune encephalitis in COVID-19

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Abstract

Introduction The novel Coronavirus Disease 2019 (COVID-19) is an infection caused by the Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) which has been spreading rapidly amongst humans and causing a global pandemic. The notorious infection has shown to cause a wide spectrum of neurological syndrome, including autoimmune encephalitis.

Objective Here, we systematically review the literature on autoimmune encephalitis that developed in the background of SARS-CoV-2 infections and also the possible pathophysiological mechanisms of auto-immune mediated damage to the nervous system.

Methodology An exhaustive search was made in Medline/PubMed, Embase, Scopus and other medical databases, and 28 relevant published articles were selected according to the strict inclusion criteria.

Results Autoimmune encephalitis can occur via three possible proposed pathophysiological mechanism and can manifest during or after the acute infection period. It is more common in adult but can also occur in the paediatric patients. There were various spectra of autoantibody panels reported including antineuronal antibody, anti-gangliosides antibody and onconeural antibody. Majority of the patients responded well to the immunomodulating therapy and achieved good recovery.

Conclusion In conclusion, SARS-CoV-2 infection can induce various spectrum of autoimmune encephalitis. It is a major concern since there is very limited long-term study on the topic. Hence, this review aims to elucidate on the potential long-term complication of SARS-CoV-2 infection and hopefully to improve the management and prognosis of COVID-19.

Keywords COVID-19 · SARS-CoV-2 · Pandemic · Neurology · Autoimmune · Encephalitis

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Introduction

The Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) is a newly emerged zoonotic virus that causes the Coronavirus Disease 2019 (COVID-19) [1]. It was first reported in Wuhan, Hubei Province, China on 29 December 2019, where four cases of an acute respiratory syndrome of unknown aetiology were linked to a local Huanan South China Seafood Market and have been causing a global catastrophic since then [2].

SARS-CoV-2 is an enveloped, single-stranded positive-sense RNA genome virus, which has a spherical or oval shape and an average size of 100 nm in diameter. There are large club-shaped spikes of glycoprotein membrane on the viral surface, which makes the viral particles appear like a typical crown-like shape in an electron microscope [3]. Taxonomically, the SARS-CoV-2 is part of the SARS-related coronaviruses species that belong to the subgenus Sarbecovirus, along with the other subgenuses namely Embecovirus, Hibeceovirus, Merbecovirus and Nobecovirus which are part of the genus Beta-coronavirus [4]. These viruses are more common in animals worldwide. However, there are a few types, such as HCoV-HKU1, HCoV-OC43, HCoV-229E and HCoVNL63 that are known to affect the humans and cause a mild respiratory illness [5, 6]. And there are some that have caused a large-scale epidemic before in the last two decades, such as the case of the Severe Acute Respiratory Syndrome (SARS-CoV) in the year 2002 to 2003, where it has infected more than 8400 individuals with the mortality rate of 11% across 26 countries around the world [7, 8], and then the Middle-East Respiratory Syndrome (MERS-CoV) in the year 2012 to 2013 which has affected about 2500 people and caused the death of 850 people [9]. SARS-CoV-2 is the first coronavirus to have caused a global pandemic.

The SARS-CoV-2 infection results in a syndrome of various systemic and respiratory symptoms such as dry cough, breathing difficulty, fever and fatigue that sometimes can be critical causing severe pneumonia and cardiopulmonary failure that require specialized management in intensive care units [10, 11]. In addition to that, it is well documented that SARS-CoV-2 infection can affect the nervous system causing a wide range of neurological symptoms [11]. Autoimmune encephalitis, which is an immune-mediated inflammatory condition involving the brain tissue, is one of the syndromes that have been reported to occur in the background of SARS-CoV-2 infection. It has been postulated that the autoinflammatory condition was triggered in response to the viral infection in addition to any other environmental confounding factors in a genetically susceptible patient [12]. This condition will have a long-term implication on the patient who recovered from the SARS-CoV-2 infection. Therefore, in the light of the possibility of autoimmune disorders linked to SARS-CoV-2 infection, this review aims to summarize all the published reporting of autoimmune encephalitis that occurs in COVID-19 patients and hopefully provide useful information for the diagnosis and management of the condition.

Methods

An exhaustive search of scientific publications which consist of a case series and reports was conducted using the following online databases: Medline/PubMed, Embase, Scopus, Web of Science, as well as Google scholar. The search MeSH term keywords used were: {“Encephalitis”, OR “autoimmune encephalitis”, OR “limbic encephalitis”, OR “NMDA-encephalitis”} AND {“SARS-CoV-2”, OR “COVID-19”, OR “coronavirus”}. All relevant articles were analysed for the possible autoimmune encephalitis related to COVID-19. These articles were published from 1 January 2020 to 31 July 2021. There were 45 articles that were organized according to the search words. The reference list of these articles was also searched and analysed for additional reporting related to autoimmune encephalitis in COVID-19 disease which identifies additional 15 articles. A flow chart of the search process is shown in Fig. 1. These articles were then carefully filtered for their relevance based on selection criteria. Amongst the selection criteria that were considered: diagnosed case of COVID-19, confirmed case of autoimmune encephalitis either by the presence of the serum or cerebrospinal fluid autoantibodies or the typical radiological findings and have a clear description of the clinical cases. The occurrence of autoimmune encephalitis in COVID-19 is explored, and a brief review is then performed.

Results

We reviewed more than 60 scientific papers on autoimmune encephalitis that occurs in confirmed cases of SARS-CoV-2 infection that were published since 1 January 2020 to 31 July 2021 which include limited case series and case reports. We focused on the details on how the diagnosis of autoimmune encephalitis was made based on the serology and radiology investigations done, treatment given and the prognosis of each case. After considering the selection criteria, 26 case reports and 2 limited case series are included in the study. A brief summary of the selected articles is briefly described in Table 1.
**Pathophysiological mechanism of autoimmune encephalitis in SARS-CoV-2 infection**

The exact pathophysiological mechanism of autoimmune encephalitis in SARS-CoV-2 infections is still yet to be understood. One of the most plausible proposed mechanisms was a molecular mimicry in response to the SARS-CoV-2 infection which led to activation of host antibodies that cross-reacted and identified self antigens as foreign and cause damages to various system, including the central nervous system (CNS) [13]. There were various anti-neuronal autoantibodies that have been discovered to affect the nervous system in SARS-CoV-2 infected patients, namely the anti-N-methyl-D-aspartate-receptor (NMDAR) antibody [14–19], anti-contactin-associated protein-like-2 (CASPR2) antibodies [20], anti-myelin oligodendrocyte glycoprotein (MOG) antibody [21, 22], anti-glutamic acid decarboxylase (GAD) antibody [23], anti-GD1a antibody [24] and many others. This mechanism was also suggested to explain other neurological complications of SARS-CoV-2 infection, namely Guillain–Barré syndrome [25] and acute disseminated encephalomyelitis [26], which further support the theory of molecular mimicry as the pathophysiology of autoimmune encephalitis.

Another proposed mechanism was systemic hyperinflammation due to over activation of host immune system causing the release of large amounts of inflammatory cytokines as a response to SARS-CoV-2 infection in a phenomenon known as “cytokine storm”. These inflammatory mediators are transported throughout the body including the central nervous system and result in inflammatory damage or encephalitis [27]. This mechanism was further supported by the evidence from the laboratory finding on the serology and cerebral spinal fluid (CSF) analysis which shows elevated proinflammatory parameters such C-reactive protein and D-dimer [11, 28, 29] and also the inflammatory mediators, such as the IL-6 [14, 16, 29]. In addition to that, the electroencephalogram (EEG) findings from some of the patients reveal diffuse patterns which are indicative of extensive inflammation [16, 19–21, 26, 30–32].
| No | Article                          | Age, Gender | Neurological presentation                                                                 | Investigation results                                                                                                                                     | Treatment                  | Outcome                                      |
|----|---------------------------------|-------------|------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------|----------------------------------------------|
| 1  | Panariello A, et al. (2020) [14] | 23, male    | Occurred during acute infection before typical COVID-19 symptoms; psychomotor agitation, anxiety, thought disorganization, persecutory delusions, hallucinations and insomnia | MRI brain did not show significant finding  
EEG showed theta activity, unstable, non-reactive to visual stimuli and no significant asymmetries  
CSF analysis raised IL-6, no elevated protein and cell counts  
Anti-NMDAR antibodies positive in CSF | Corticosteroid; IVIG | Clinical conditions are ameliorating to date                                              |
| 2  | Bravo GA, et al. (2020) [15]    | 30, female  | Occurred during acute infection before typical COVID-19 symptoms; psychomotor agitation, paranoid ideation, dysarthria, visual hallucinations and seizure | MRI brain showed hyperintensities in the left hippocampus  
EEG showed epileptic discharges in the left frontotemporal region  
Repeated EEG showed delta brush pattern together with spike-and-wave discharges in anterior regions  
CSF showed high protein and pleocytosis (lymphocyte predominance)  
Anti-NMDAR antibodies positive in serum and CSF | Corticosteroid; IVIG; surgical removal of ovarian teratoma | Incomplete recovery; with cognitive sequelae and memory disorder |
| 3  | Monti G, et al. (2021) [16]     | 50, male    | Occurred during acute infection with no fever or respiratory symptoms; acute psychosis, later developed focal motor seizures and progressed into refractory status epilepticus | Serial MRI brain was normal  
EEG showed diffuse delta activity (delta brush pattern)  
CSF analysis raised IL-6 (also in serum), mild elevated protein and mild pleocytosis  
Anti-NMDAR antibodies positive in CSF | IVIG; PLEX; corticosteroid | Complete recovery                                                |
| 4  | Allahyari F, et al. (2021) [17] | 18, female  | Occurred before acute infection; lack of concentration, anhedonia and seizure               | CT brain showed brain oedema  
MRI brain normal  
EEG not available  
CSF showed high protein and pleocytosis (lymphocyte predominance)  
Anti-NMDAR antibody positive in CSF | Corticosteroids; IVIG | Complete recovery                                                |
| No | Article                        | Age, Gender | Neurological presentation                                                                 | Investigation results                                                                 | Treatment               | Outcome                           |
|----|--------------------------------|-------------|-------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------|-----------------------------------|
| 5  | Guilmot A, et al. (2021) [20]  | 80, male    | Occurred during acute infection with mild respiratory symptoms; short-term memory disturbances, visual hallucinations, anxiety and seizure | MRI brain normal<br>EEG show generalized slowing<br>CSF analysis elevated protein and positive oligoclonal band with normal cell counts<br>Anti CASPR2 antibody was positive in CSF and serum | Corticosteroid PLEX    | Complete recovery                  |
| 6  | Gaughan M, et al. (2021) [23] | 16, female  | Occurred during acute infection with no fever or respiratory symptoms; hallucinations and ritualistic behaviours, subsequently severe encephalopathy with akinetic mutism. She then developed bilateral limb rigidity with subtle high frequency tremor | MRI brain showed 2 tiny punctate T2/FLAIR hyperintensities in the centrum semi-ovale bilaterally, with no diffusion restriction or contrast enhancement<br>CSF analysis normal<br>EEG delta slowing, more prominent in the right hemisphere posteriorly<br>Anti-GAD antibody level was positive in serum but negative in CSF | IVIG; corticosteroid    | Complete recovery                  |
| 7  | Ayuso LL, et al. (2020) [24]  | 71, female  | Occurred after acute infection; unsteadiness, dizziness and severe truncal ataxia          | MRI brain showed hyperintense lesions in the caudal vermis and right flocculus, and contrast enhancement was observed in the floor of the fourth ventricle<br>EEG was normal<br>CSF was normal<br>Anti-GD1a IgG antibodies positive in serum | Corticosteroid           | Almost complete recovery with mild unsteadiness |
| 8  | Yousuf F, et al. (2021) [30]  | 60, male    | Occurred after acute infection; Altered mental status, memory impairment and aphasia      | PET scan of brain showed abnormal mixed brain hypometabolism and hypermetabolism (suggesting an early pattern of autoimmune encephalitis)<br>48 h of video EEG showed severe diffuse encephalopathy, no seizures or epileptiform discharges<br>CSF analysis elevated protein<br>Autoantibody panel negative | IVIG                    | Almost complete recovery           |
| No | Article | Age, Gender | Neurological presentation | Investigation results | Treatment | Outcome |
|----|---------|-------------|---------------------------|-----------------------|-----------|---------|
| 9  | Grimaldi S, et al. (2020) [31] | 72, male | Occurred during acute infection before typical COVID-19 symptoms; subacute cerebellar syndrome and myoclonus | MRI brain normal PET scan of brain showed a diffuse pattern compatible with encephalitis and cerebellitis EEG showed symmetric diffuse background slowing and reactive to stimulation without interictal paroxysm CSF analysis no elevated protein and cell counts | IVIG corticosteroid | Complete recovery |
|    |         |             |                           |                       |           |         |
| 10 | McAlpine LS, et al. (2021) [32] | 60, male | Occurred during acute infection before typical COVID-19 symptoms; confusion and cognitive decline concerning for encephalopathy | MRI brain showed scattered foci of FLAIR hyperintensity in the periventricular and subcortical white matter (likely sequelae of chronic small vessel disease) EEG showed mild to moderate generalized slowing without seizures or epileptiform patterns CSF analysis not available Autoantibody panel not available | None | Ongoing improvement |
| 11 | Oosthuizen K, et al. (2021) [33] | 52, male | Occurred during acute infection with no fever or respiratory symptoms; progressive gait instability with other cerebellar signs, including nystagmus, dysarthria and truncal ataxia | MRI brain showed features consistent with brainstem encephalitis EEG not available CSF analysis revealed pleocytosis with normal protein. PCR for SARS-CoV-2 RNA was positive Anti-amphiphysin antibody positive in the serum | Corticosteroid | Almost complete walking independently with dysarthria, broad-based gait |
| No | Article                        | Age, Gender | Neurological presentation                                                                 | Investigation results                                                                 | Treatment  | Outcome                                |
|----|--------------------------------|-------------|------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|------------|----------------------------------------|
| 12 | Zambreanu L, et al. (2021) [36] | 66, female  | Occurred during acute infection with mild respiratory symptoms; confusion, memory deficits and seizure | MRI brain showed T2 hyperintensities in limbic lobes, upper pons and medial thalami, without gadolinium enhancement (consistent with limbic encephalitis) | Corticosteroid, IVIG | Incomplete recovery; ongoing cognitive impairment |
|    |                                |             |                                                                                         | EEG not available                                                                     |            |                                        |
|    |                                |             |                                                                                         | CSF analysis elevated protein                                                          |            |                                        |
|    |                                |             |                                                                                         | Autoantibody panel negative                                                           |            |                                        |
| 13 | Pilotto A, et al. (2020) [37]  | 68, male    | Occurred during acute infection with severe respiratory symptoms; altered mental state, status epilepticus and dysarthria | MRI consistent with limbic encephalitis                                               | None       | Complete recovery                      |
|    |                                |             |                                                                                         | EEG was abnormal, but no detail                                                       |            |                                        |
|    |                                |             |                                                                                         | CSF analysis elevated protein and pleocytosis                                         |            |                                        |
|    |                                |             |                                                                                         | Autoantibody panel negative                                                           |            |                                        |
| 14 | Hamill A (2021) [38]           | 59, female  | Occurred during acute infection before typical COVID-19 symptoms; altered mental status and syncopal attack | MRI brain showed diffuse abnormal hyperintensities over right hippocampus (consistent with limbic encephalitis) | Not available | Still in ICU                            |
|    |                                |             |                                                                                         | EEG normal                                                                           |            |                                        |
|    |                                |             |                                                                                         | CSF analysis elevated protein                                                          |            |                                        |
|    |                                |             |                                                                                         | Autoantibody panel not available                                                     |            |                                        |
| 15 | Chenna V, et al. (2021) [39]   | 58, male    | Occurred after acute infection; altered sensorium (during his admission for COVID-19 and was intubated) | MRI brain showed bilateral cerebellar, parietooccipital hyperintensities on T2/FLAIR mode | IVIG       | Complete recovery, able to walk        |
|    |                                |             |                                                                                         | EEG not available                                                                     |            |                                        |
|    |                                |             |                                                                                         | CSF analysis normal                                                                   |            |                                        |
|    |                                |             |                                                                                         | Autoantibody panel not available                                                     |            |                                        |
| 16 | Khoo A, et al. (2020) [40]     | 65, female  | Occurred after acute infection; generalized myoclonus, ocular flutter with convergence spasm and acquired hyperkplexia | MRI brain normal                                                                      | Corticosteroid | Incomplete recovery; walking with stick ongoing fine myoclonus |
|    |                                |             |                                                                                         | EEG not available                                                                     |            |                                        |
|    |                                |             |                                                                                         | CSF analysis normal                                                                   |            |                                        |
|    |                                |             |                                                                                         | Autoantibody panel not available                                                     |            |                                        |
| No | Article | Age, Gender | Neurological presentation | Investigation results | Treatment | Outcome |
|----|---------|-------------|---------------------------|-----------------------|-----------|---------|
| 17 | Dono F, et al. (2021) [41] | 81, male | Occurred after acute infection; altered mental status, fever, then reduced consciousness with recurrent myoclonic jerk | MRI brain showed multiple hyperintense areas in T2-FLAIR and axial DWI in the bilateral parietal cortex, left temporal cortex and right cingulate cortex with no contrast enhancement after gadolinium injection | Corticosteroids; IVIG | Minimal recovery; myoclonic jerk stops, then the patient succumbs to hospital acquired infection |
| 18 | Pizzanelli C, et al. (2021) [42] | 74, female | Occurred after acute infection; subacute confusion and focal motor seizures with impaired awareness | MRI brain showed bilateral T2/FLAIR hyperintensities in both hippocampi (consistent with limbic encephalitis) | Corticosteroid; antiseizure medications | Almost complete recovery with slight verbal deficits |
| 19 | Hosseini AA, et al. (2020) [43] | 79, female | Occurred during acute infection with fever and no respiratory symptoms; acute confusion, dysphasia and seizure | MRI brain showed partial diffusion restriction in limbic system (suggestive of limbic encephalitis) | Antiseizure medications | Poor recovery with impaired verbal fluency, delayed recall memory |
| 20 | Zuhorn F, et al. (2020) [44] | 54, male | Occurred during acute infection with no fever or respiratory symptoms; altered mental state, disoriented and stupor prior to respiratory failure and intubated | MRI brain showed signal alterations at the claustrum/external capsule region, showing reduced diffusion | None | Almost complete recovery with mild cognitive impairment |
| No | Article | Age, Gender | Neurological presentation | Investigation results | Treatment | Outcome |
|----|---------|-------------|---------------------------|-----------------------|-----------|---------|
| 21 | Bhagat, et al. (2021) [45] | 54, male | Occurred during acute infection before typical COVID-19 symptoms; headache, loss of consciousness and seizure | MRI brain showed increased signal in DWI without ADC correlation in the posterior aspect of right medial temporal lobe and para-hippocampal gyrus with associated T2-weighted-FLAIR signal hyperintensity EEG showed lateralized periodic discharges and focal delta slowing over right posterior quadrant CSF analysis normal Autoantibody panel negative | Antiseizure medications | Complete recovery |
| 22 | McAlpine LS, et al. (2021) [46] | 30, male | Occurred during acute infection after typical COVID-19 symptoms; bizarre delusion, sleep disturbance and hallucination | MRI brain normal EEG was normal CSF analysis was normal Anti-neural autoantibodies test on CSF revealed a novel immunostaining pattern | IVIG | Complete recovery |
| 23 | Ayatollahi P, et al. (2021) [47] | 18, female | Occurred during acute infection before typical COVID-19 symptoms; Drowsiness, confusion and seizure | MRI brain showed signal hyperintensities on FLAIR and T2-weighted sequences in the claustrum bilaterally and extended to the external capsules and anterior insular cortex, sparing mesial temporal structures EEG showed intermittent non-epileptiform abnormalities over the both frontocentrottemporal regions CSF analysis no elevated protein and cell counts Autoantibody panel negative | Corticosteroid; antiseizure medications | Almost complete recovery with persistent recent memory deficit |
| No | Article | Age, Gender | Neurological presentation | Investigation results | Treatment | Outcome |
|----|---------|-------------|--------------------------|----------------------|-----------|---------|
| 24 | Burr T, et al. (2021) [18] | 23-month girl | Occurred during acute infection with fever and no respiratory symptoms; communication and movement disorders, multiple seizures and gradual worsening of encephalopathy | MRI brain was normal EEG not available CSF analysis no elevated protein and cell counts Anti-NMDAR antibodies positive in serum and CSF | IVIG | Complete recovery |
| 25 | Sarigecili E, et al. (2021) [19] | 7-year-old boy | Occurred during acute infection with no fever or respiratory symptoms; unsteady gait, then progressed to choreiform movements, tongue protrusion, bruxism and lip smacking | MRI brain was normal Awake and sleep EEGs were encephalopathic with widespread delta waves CSF analysis no elevated protein and cell counts Anti-NMDAR antibody positive in CSF | IVIG corticosteroid | Almost complete recovery with mild ataxia |
| 26 | Vraka K, et al. (2021) [21] | 13-month girl | Occurred during acute infection before typical COVID-19 symptoms; impaired consciousness, seizures with decorticated posturing | MRI brain showed bilateral widespread high signal abnormalities over white matter, including the splenium of the corpus callosum with associated diffusion restriction and high signal in the thalami and pons EEG showed diffuse slow-wave background activity (encephalopathy), but no epileptiform discharge CSF analysis not available Anti-MOG antibody detected in serum | Corticosteroid | Complete recovery |

| No | Article | Age, Gender | Neurological presentation | Investigation results | Treatment | Outcome |
|----|---------|-------------|--------------------------|----------------------|-----------|---------|
| 10-year-old girl | Occurred during acute infection with fever and no respiratory symptoms; headache, fluctuating sensorium and right-hand weakness | MRI brain showed asymmetric bilateral high-signal lesions in the basal ganglia and the subcortical white matter in the frontal and temporal lobes, with involvement of the left internal capsule and left hippocampus EEG normal CSF analysis was normal Autoantibody panel negative | Not available | Incomplete recovery; neglect right upper limb impairment of verbal memory |
Table 1 (continued)

| No | Article                                      | Age, Gender | Neurological presentation                                                                 | Investigation results                                                                 | Treatment                  | Outcome                                         |
|----|----------------------------------------------|-------------|------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|---------------------------|------------------------------------------------|
| 27 | Ahsan N, et al. (2021) [22]                  | 7-year-old girl | Occurred during acute infection with fever and no respiratory symptoms; aphasia, encephalopathy, status epilepticus and prolonged Todd’s paralysis | MRI brain revealed peri Rolandic and posterior parietal lobe restricted diffusion and cortical oedema | Antiseizure medications; IVIG | Almost complete recovery with mild dysarthria   |
|    |                                              |             |                                                                                          | EEG showed cerebral slowing with left focal slowing                                     |                           |                                                 |
|    |                                              |             |                                                                                          | CSF analysis not available                                                              |                           |                                                 |
|    |                                              |             |                                                                                          | Anti-MOG antibody detected in serum                                                     |                           |                                                 |
|    |                                              |             |                                                                                          | MRI brain showed multifocal hyperintense T2-FLAIR signals in bilateral subcortical and periventricular white matter without contrast enhancement | IVIG corticosteroid           | Complete recovery                               |
|    |                                              |             |                                                                                          | EEG showed diffuse slowing consistent with encephalopathy but no seizures or epileptiform activity |                           |                                                 |
|    |                                              |             |                                                                                          | CSF analysis not available                                                              |                           |                                                 |
|    |                                              |             |                                                                                          | Autoantibody panel not available                                                        |                           |                                                 |

ADC apparent diffusion coefficient, CASPR2 contactin-associated protein-like-2, CSF cerebral spinal fluid, DWC diffusion weighted imaging, EEG electroencephalogram, FLAIR fluid-attenuated inversion recovery, GAD glutamic acid decarboxylase, IL-6 interleukin-6, IVIG intravenous immunoglobulin G, MOG myelin oligodendrocyte glycoprotein, MRI magnetic resonance imaging, NMADR N-methyl-D-aspartate-receptor, PCR polymerase chain reaction, PET positron emission tomography, PLEX plasma exchange, RNA ribonucleic acid, T2/T2-weighted image.
The third proposed pathophysiological mechanism was direct invasion of the virus into the central nervous system that led to the development of encephalitis. Since the earlier times of the pandemic, it has been postulated that the SARS-CoV-2 virus could enter the brain parenchyma through either the transsynaptic propagation or the hematogenous invasion [11]. Oosthuizen K, et al. [33] reported a case of middle-aged man with SARS-CoV-2 infection presented with progressive ataxia who was found to have SARS-CoV-2 particles detected via polymerase chain reaction (PCR) in the CSF. The patient was also noted to have positive anti-amphiphysin antibody in the serum, which is an onconeural antibody that is usually associated with paraneoplastic encephalitis despite no malignancy found upon extensive full body screening. This may show that viral infections can trigger antibodies against cell surface antigens as it was well documented in the case of Herpes Simplex virus (HSV) encephalitis which can trigger anti-NMDAR encephalitis through uncovering epitopes resulting in antibody responses [34]. However, Siahaan YM, et al. [35] suggested that direct invasion of SARS-CoV-2 virus into the CNS is less likely to be the main mechanism of encephalitis in SARS-CoV-2 infection. This is because there are very limited cases where SARS-CoV-2 infected patients with encephalitis had positive PCR against SARS-CoV-2 detected in the CSF.

Autoimmune encephalitis in SARS-CoV-2 infection

From the cases that we reviewed in this article, 24 (80%) of the patients were adults with the majority (58%) of them aged more than 55 years old [20, 24, 30–32, 36–43]. Six (20%) of the patients were from paediatric age, defined as 12 years and younger [18, 21–23, 26]. Amongst the most commonly reported neurological presentation of autoimmune encephalitis in SARS-CoV-2 infection are altered mental status [21, 30, 32, 36–39, 41–44, 47], pyramidal motor agitation [14, 15, 20], memory deficit [20, 30, 36], some form of psychogenic disturbance like hallucinations or delusions [14–16, 23, 46], ataxia [19, 24, 26, 31, 33] and seizure [15–18, 20–22, 36, 37, 42, 43, 45, 47]. Based on the typical findings on magnetic resonance imaging scan of the brain, six of the patients had limbic encephalitis [36–38, 42, 43], three had cerebellitis [24, 31, 39] and one had brain stem encephalitis [33]. There were 11 patients reported to develop autoantibodies either in the serum or/and in the cerebrospinal fluid (CSF) sample. Six of these autoantibodies were anti-NMDAR antibody [14–19], two were anti-MOG antibody [21, 22], and one anti-GAD [23], anti-GD1a [24] and anti-CASPR2 antibodies [20] each. Grimaldi S, et al. [31] reported a case of an elderly man who presented subacute cerebellar syndrome with myoclonus which occurs after the acute SARS-CoV-2 infection. His positron emission tomography scan of brain showed a diffuse pattern compatible with encephalitis and cerebellitis, and his CSF analysis revealed no elevated protein and normal cell counts, but there were high titres of autoantibodies directed against the nuclei of Purkinje cells, striatal and hippocampal neurons found. He was successfully treated with high-dose corticosteroid and intravenous (IV) immunoglobulin G and had complete recovery. McAlpine LS, et al. [32] on the other hand reported a novel immunostaining pattern from the anti-neuronal autoantibodies test on the CSF in a young male patient presented with bizarre delusion and hallucination. The patient was successfully treated with IV immunoglobulin G and achieved complete recovery.

In this review, majority (77%) of the patients had the autoimmune encephalitis occurred during or concurrent with the acute infection of the SARS-CoV-2 which is termed “para-infectious presentation” [14–16, 18–23, 26, 31–33, 36–39, 44–47]. From this, 30% of the patients presented with neurological manifestations before developing fever and respiratory symptoms [14, 15, 21, 26, 31, 32, 38, 45, 47]. Six (20%) of the cases occurred after the acute infection [24, 30, 40–43], and only one (3%) occurred before the acute infection which was reported by Allahyari F, et al. [17] in a young female with anti-NMDAR antibody positive in CSF. She was successfully treated with high-dose corticosteroid and IV immunoglobulin G with complete recovery.

There were 18 patients that were treated with high-dose corticosteroid, which is either oral prednisolone (1 mg/kg, max 60 mg) daily or pulsed intravenous (IV) methylprednisolone 1 g daily [14–17, 19–21, 23, 24, 26, 31, 33, 36, 37, 40–42, 47]. On the other hand, 15 patients were treated with IV immunoglobulin G [14–19, 22, 23, 26, 30, 31, 36, 39, 41, 46]. There were two patients that were treated with just antiseizure medication [43, 45], and three patients were not given any treatment at all [32, 37, 44]. There were nine patients that received a combination of high-dose corticosteroid and IV immunoglobulin G [14, 15, 17, 19, 23, 26, 31, 36, 41], and two were subjected into plasma exchange [16, 20]. Monti G et al. [16] reported a young patient with anti-NMDAR antibodies positive in the CSF which was treated with a combination of plasma exchange, IV immunoglobulin G and corticosteroid and had complete recovery. Guilmet A., et al. [20] on the other hand reported an elderly patient with positive anti-CASPR2 antibody in both CSF and serum which was successfully treated with high-dose corticosteroid and plasma exchange and had complete recovery.

The prognosis of autoimmune encephalitis in SARS-CoV-2 infection based on all the cases in our study was generally good where 20 (67%) of the patients have complete or almost complete recovery [16–24, 26, 30, 31, 33, 37, 39, 42–47]. There were five (17%) patients that have incomplete recovery [15, 21, 36, 37, 40], and only two (7%) patients had poor recovery, which was the case of limbic
encephalitis reported by Hosseini AA, et al. [43] and a case of post-SARS-CoV-2 infection autoimmune encephalitis presenting as new-onset refractory status epilepticus which was reported by Dono F, et al. [41].

Autoimmune encephalitis in SARS-CoV-2 infection in paediatric

Out of the 30 patients that were reviewed in this article, there were six paediatric patients defined as age 12 years and younger. Vraka K, et al. [21] reported a case of an anti-MOG antibody positive in the serum of a 13-month-old girl. The young girl was successfully treated with high-dose corticosteroid and achieved complete recovery. Another similar case was reported in a 7-year-old girl by Ahsan N, et al. [22] who also had anti-MOG antibody positive in the serum. She was treated with IV immunoglobulin G and had a good recovery with only mild dysarthria. Burr T, et al. [18] and Sarigecili E, et al. [19] on the other hand reported a 23-month-old girl and a 7-year-old boy with anti-NMDAR IgG antibody which were both treated with a combination of high-dose corticosteroid and IV immunoglobulin G and had a complete and almost complete recovery. It is worth to mention that not all of the paediatric patients have autoantibody positive in the serum or CSF. Vraka K, et al. [21] reported another case in their article on a 10-year-old girl who presented with right-hand weakness and fluctuating sensorium. She has normal CSF analysis findings and negative autoantibody panel. She was not given any immunomodulating therapy and still has neglect over her right hand and impairment of her verbal memory. Another case was reported by McLendon LA, et al. [26] in a 17-month-old girl who developed upper limb weakness and ataxia, which was also treated with a combination of high-dose corticosteroid and IV immunoglobulin G and had complete and almost complete recovery. However, the authors did not include the CSF analysis and the autoantibody panel test results in the report.

Discussion

SARS-CoV-2 belongs to the genus of beta-coronavirus which is a zoonotic disease that can infect both humans and animals. Since it was first reported in December 2019, the infection has spread rapidly across country borders and caused massive global pandemic. Up to date, there are more than 200 million reported confirmed cases of COVID-19, and more than four million deaths have taken place from more than 150 countries around the world [48]. Infection by SARS-CoV-2 causes a typical clinical syndrome such as fever, cough, shortness of breath, myalgia and fatigue. Other clinical presentations that are considered non-typical are diarrhoea, anorexia, conjunctival congestion, nausea and vomiting. About 20 to 30% of patients may progress to more critical conditions with acute respiratory distress syndrome, septic shock, disseminated intravascular coagulation, acute heart failure and acute kidney injury [49]. Another peculiarity about SARS-CoV-2 infection is the propensity to affect the nervous system. In our previous studies, we reviewed multiple clinical evidences that show SARS-CoV-2 involvement on the nervous system and raised a major concern on the possibility of future autoimmune complication involving the nervous system [11]. There have been multiple cases that have been reported on the possible autoimmune occurrence in SARS-CoV-2 infection. Therefore, here we studied the link between autoimmune encephalitis in SARS-CoV-2 infection and the prognosis of the condition.

Similar to what have been observed in autoimmune disease involving the nervous system induced by other viral infections like HSV which itself is more common to cause encephalitis [34], autoimmune encephalitis can occur after the resolution or more commonly during the acute phase of SARS-CoV-2 infection. It can happen in any spectrum of disease severity, and some can even occur without any of the typical symptoms of COVID-19. Apart from that, there were various types of serum and CSF autoantibodies related to encephalitis that have been reported, namely anti-gangliosides antibody (such as anti-GAD antibody [23] and anti-GD1a antibody [24]), anti-neuronal antibodies (anti-NMDAR antibody [14–19] and anti-CASPR2 antibody [20]) and onconeural antibodies (anti-amphiphysin antibody [33]). The clinical significance of the anti-amphiphysin antibodies in the case reported by Oosthuizen K, et al. [33] was unclear given that no malignancy was identified in the patient. However, the RNA of SARS-CoV-2 was detected in the CSF sample of the patient, which could indicate direct invasion of the virus to the CNS causing encephalitis that may have induced responses against amphiphysin, which is an intracellular synaptic antigen. However, the association still warrants future studies for further explanation and to explore the associations between SARS-CoV-2 infection and immune-mediated CNS disease.

The possibility of immune-mediated neurology disease in SARS-CoV-2 infection leading to future disabilities is still a major concern. This is because the unknown complications can occur years after the recovery from acute infection, especially when evidence have shown other types of coronaviruses that are linked to central nervous system dysfunction such as multiple sclerosis [50]. Furthermore, there were cases where antibody against MOG, which is a myelin protein that exclusively expressed on the surface of CNS myelin sheaths, was reported amongst the paediatric patients [21, 22]. Antibody against MOG has been associated with various autoimmune inflammatory demyelinating diseases, such as multiple sclerosis and acute disseminated encephalomyelitis (ADEM). ADEM is an autoimmune...
demyelinating disease that commonly affects the brainstem and spinal cord and typically presents as a monophasic disorder associated with multifocal neurologic symptoms and encephalopathy [22]. Therefore, more reporting of cases and studies are deemed necessary to illuminate the link and potential reversibility of the neurological syndrome, and the overall impact of the infection in promoting other neurological diseases in the future.

Limitations of the study
This study has limitations, where most of the articles analysed are case reports and limited case series. Apart from that, this article mainly uses descriptive analysis to review and summarize the clinical cases.

Conclusion
In conclusion, primary data on COVID-19 patients with autoimmune encephalitis remains limited. Therefore, this review aims to illuminate the occurrence of autoimmune encephalitis in SARS-CoV-2 infection based on the evidence presented from the previous published case reporting and also attempt to educate clinicians around the world regarding the conditions in order to improve the prognosis of COVID-19 by delivering the appropriate treatment promptly.

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Declarations

Ethics approval and consent to participate
None.

Conflict of interest
The authors declare no competing interests.

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