COVID-19-related refractory status epilepticus with the presence of SARS-CoV-2 (RNA) in the CSF: a case report

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Abstract

Amongst the neurologic complications of COVID-19 disease, very few reports have shown the presence of the virus in the cerebrospinal fluid (CSF). Seizure and rarely status epilepticus can be associated with COVID-19 disease. Here we present a 73-year-old male with prior history of stroke who has never experienced seizure before. He had no systemic presentation of COVID-19 disease. The presenting symptoms were two consecutive generalized tonic-clonic seizures that after initial resolution turned into a nonconvulsive status epilepticus despite antiepileptic treatment (a presentation similar to NORSE (new-onset refractory status epilepticus)). There was no new lesion in the brain magnetic resonance imaging (MRI). The CSF analysis only showed an increased protein levels and positive reverse transcription polymerase chain reaction (RT-PCR) of 2019-nCoV. Patient recovered partially after anesthetic, IVIG, steroid, and remdesivir. To our knowledge, this is the first report of a refractory status epilepticus with the presence of SARS-CoV-2 ribonucleic acid (RNA) in the CSF.

Keywords COVID-19 · Status epilepticus · CSF

Introduction

Seizure has been reported as a life-threatening neurological complication of severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) [1, 2]. Potential mechanisms have been speculated to cause seizure in the setting of COVID-19 including encephalitis, cytokine storm, hypoxia, and metabolic derangement [3]. Here we report a case of COVID-19-related refractory status epilepticus with the presence of SARS-CoV-2 ribonucleic acid (RNA) in the cerebrospinal fluid (CSF).

Case report

The patient is a 73-year-old man with a history of a previous left occipital lobe ischemic stroke in 2017. Although he had a mild right hemiparesis as a sequela of the stroke, he never had a seizure. He presented with two seizures on the day of admission. The seizures were focal motor (right versive seizure without loss of awareness) followed by secondary generalized tonic-clonic seizures. In the postictal phase, he was moderately confused with a gaze dominancy to the left and partial Broca-type dysphasia and increased weakness in the right arm (Todd’s paresis). There was no systemic presentation of an infection (no fever nor any respiratory symptoms) and no meningeal irritation signs like headache or neck stiffness. The patient received loading and maintenance dose of levetiracetam. The Todd’s paresis resolved, and the level of consciousness improved during the next 12 h; then he developed aphasia and fluctuations in awareness once more (aphasic seizure with loss of awareness). The brain computed tomography scan and magnetic resonance imaging (MRI) failed to show any new lesions in the brain. However, there was a slight cortical hyperintensity in diffusion-weighted (DWI) MRI without hypointensity in the apparent diffusion coefficient (ADC-map) images located in the left temporal cortex (Fig. 1). The routine lab data was normal apart from the elevation of serum leukocyte (13,000 cells/mm³). Serum and urine toxicology analyses were negative. The electroencephalography (EEG) showed a focal ictal pattern in the left temporal leads
(maximum potential: F7) with a slow background rhythm consistent with a focal nonconvulsive status epilepticus (Fig. 2a). Valproate loading dose helped the patient to recover to his normal baseline cognitive and language state. Unfortunately, after 12 h, there was a recurrence of symptoms despite optimal antiepileptic serum levels. After intravenous

Fig. 1  

(a) Previous stroke in the left PCA territory with bilateral periventricular white matter lesions. (b, c) Left temporal cortex flair and DWI hyperintensity

Fig. 2  

(a) There was very frequent rhythmic (more than 2.5 Hz) left temporal sharp compatible with EEG ictal pattern (starting from: F7>T3, T5). (b, c) There was no posterior dominant rhythm. There was continuous left temporal sharp/periodic lateralized epileptiform discharge (PLED)/EEG ictal pattern (maximum potential: T3, T5) and independent frequent right occipital sharp/PLED (maximum potential O2) and a continuous right temporo-occipital rhythmic and irregular delta slowing (maximum potential: O2, T6) which we assumed compatible with focal nonconvulsive status epilepticus. (d) After tapering the anesthetics, the background rhythm consisted of a low amplitude irregular delta, theta activity without epileptiform discharges.  

* F frontal, T temporal, O occipital
In the context of COVID-19 disease, patients can be susceptible to have seizures by multiple mechanisms. Patients with established epilepsy can experience seizure breakthrough. On the other hand, new onset seizures can occur in patients who have metabolic derangement due to systemic complications of COVID-19. Seizures can be due to structural insult in the brain which can be a new lesion after COVID infection as a result of an ischemic, inflammatory, or a potential direct viral encephalitis [3].

Our patient’s MRI showed that there was no new ischemic or inflammatory lesion in the brain, and the cortical findings could be justified by the frequent epileptic activity in that region. Given that there was no hypoxia or other metabolic derangement in this patient, one can assume that the reason for the new onset seizures can be associated to a direct viral insult orsecondary to inflammatory response in the CNS (compatible with an encephalitis process). Unlike previous reports of COVID-19-associated meningitis or hemorrhagic encephalitis with positive RT-PCR of viral RNA in the CSF, our patient did not have the cellular response in CSF nor the typical hemorrhagic lesions in MRI [5, 6]. Although the high CSF protein levels could be suggestive of inflammation, the normal IgG index failed to prove intrathecal IgG synthesis. We did not have access to more sophisticated laboratory tests like CSF interleukin-6, neopterin, and β2-microglobulin, and our assessment of the inflammatory response was not thorough.

Given that the old ischemic stroke was located in the left occipital region, and the current seizures come from the left temporal region, it is possible that there is a new irritated lesion due to viral invasion or inflammatory reactions that serves as the epileptogenic zone. There was no systemic condition that could explain the aggressive course of the epileptic activity, and we did not have an exact explanation to why the seizures turn into refractory status epilepticus. Status epilepticus has been reported in patients with COVID-19 [9]. In some reports, the NORSE (new-onset refractory status epilepticus) and anti-NMDA associated with COVID-19 were encountered long after the disease receded, which is expectable in a post-infectious inflammatory process [10, 11]. Nonetheless, this was not the case in our patient who experienced the status epilepticus during the acute phase of the viral infection. We assumed that similar to what occurs in NORSE, an ongoing inflammatory cascade could play a role in our patient’s condition. Therefore, we started an immunomodulatory treatment in combination with antiviral drug, which resulted in improvement in patient’s condition.

In retrospect, probably because of the delay in starting immunomodulatory and antiviral treatment, status epilepticus became refractory. This was in part due to lack of systemic and respiratory findings in this patient who had a pure neurological COVID-19 presentation. We propose that during the pandemic, all patients suspected to NORSE should be investigated for COVID-19 disease and have a PCR test. In addition, it may be beneficial to apply a strict anti-COVID-19 and immunomodulatory treatment in patients who do not have severe respiratory and systemic complications but rather rare neurologic symptoms since they may easily get aggressive and be potentially life-threatening.
Data Availability  All data not published within this article will be made available by request from any qualified investigator.

Declarations

Ethics approval  Ethical approval for this study was obtained from the Iran National Committee for ethics in biomedical research (IR.TUMS.VCR.REC.1399.520). Written informed consent for publication of this case report was obtained from his legal guardian.

Conflict of interest  The authors declare no competing interests.

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