CASE REPORT

COVID-19 leukoencephalopathy with subacute magnetic resonance imaging findings of vasculitis and demyelination

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Abstract
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) commonly results in a respiratory illness in symptomatic patients; however, those critically ill can develop a leukoencephalopathy. We describe two patients who had novel subacute MRI findings in the context of coronavirus disease 2019 (COVID-19) leukoencephalopathy, which we hypothesize could implicate a potent small-vessel vasculitis, ischemic demyelination and the presence of prolonged ischemia. Recent evidence of the direct neuroinvasiness of SARS-CoV-2 leading to ischemia and vascular damage supports this hypothesis.

Keywords COVID-19 · Coronavirus · SARS-CoV-2 · Leukoencephalopathy · Demyelination · Vasculitis · MRI

Introduction
A wide spectrum of neurological disease is recognized in COVID-19 from mild to severe presentations (Mao et al. 2020; Xiong et al. 2021). We report two patients with severe respiratory failure and poor neurological status with novel subacute radiological findings that yield insight into the pathogenesis of COVID-19 leukoencephalopathy.

Case series
Both patients were symptomatic and tested positive for SARS-CoV-2 PCR on nasopharyngeal swabs. Patient 1 had 2 weeks of illness, and patient 2 had 1 week of illness at home. Both required ICU admission and intubation for severe type-1 respiratory failure secondary to COVID-19 pneumonia, IV antibiotics and hemodialysis for acute kidney injury. Neither received extra-corporeal membrane oxygenation nor experimental treatment. Table 1 provides a detailed summary of the clinical presentation, laboratory and imaging investigations and treatment regime for both patients.

Patient 1 was a 61-year-old woman with type-2 diabetes and raised inflammatory markers. After 3 weeks, she was extubated with poor neurological status; she was non-verbal and showed little movement. Brain CT on day 18 and MRI on day 24 demonstrated numerous lesions in the deep cerebral, callosal and cerebellar white matter demonstrating diffusion restriction (Fig. 1). Gadolinium was contraindicated due to renal failure.

Lumbar puncture was normal including negative tests for SARS-CoV-2 RNA RT-PCR, oligoclonal bands, JC virus and antibodies to myelin-oligodendrocyte-glycoprotein and aquaporin-4 (Table 1). EEG supported a moderate encephalopathy. An unusual post-viral acute disseminated encephalomyelitis (ADEM) was hypothesized based on the clinico-radiological findings in particular the splenial and middle cerebellar peduncle involvement, despite the normal CSF. She was treated with a 3-day course of IV methylprednisolone and a tapering course of oral prednisolone. There was clinical response, and by day 34 she could sit, but her neurologic status remained highly variable. Contrast-enhanced MRI at day 46 demonstrated incomplete ring and punctate enhancement in the existing lesions and the presence of microbleeds and gliosis (Fig. 2). In view of potentially active disease, she had 5 days of plasma exchange. MRI at
day 73 showed resolution of contrast enhancement with no new lesions (not shown). She improved, and by day 172 she could stand and complete some activities of daily living.

Patient 2 was a 62-year-old male ex-smoker. His admission CRP was high, highlighting a pro-inflammatory phenotype. He had poor neurological status on extubation and remained intubated. Brain CT on day 30 demonstrated widespread cerebral and callosal white matter hypoattenuation with swelling and a cerebellar white matter lesion topographically reminiscent of patient 1 (Fig. 3). He received 5 days of IV methylprednisolone for a co-existent organizing pneumonia and a tapering course of oral prednisolone.

On day 43, he developed intermittent facial twitching but remained intubated. Brain MRI was performed on day 77, delayed due to his significant cardiorespiratory instability and performed without contrast due to renal failure. This confirmed extensive leukoencephalopathy with deep cerebral, callosal and cerebellar white matter lesions, larger lesions demonstrated ring-shaped diffusion restriction (Fig. 3). EEG off sedation showed moderate-severe encephalopathy. He failed to improve neurologically, and repeat MRI on day 98 showed ongoing diffusion restriction with low apparent diffusion coefficient values (ADC) within the same lesions and microbleeds (Fig. 3). He had spontaneous eye opening but no other movements. Lumbar puncture was normal including negative SARS-CoV-2 RNA RT-PCR (Table 1). He was treated with another 3 days of IV methylprednisolone and 5 days of plasma exchange.
without improvement. As the prospect of meaningful levels of recovery was not possible, he was converted to a palliative management plan.

Discussion

What has become clear to clinicians around the world is that SARS-CoV-2 is a disease unlike any other. Acute leukoencephalopathy has been described; however, there are few reports on its imaging evolution (Agarwal et al. 2020b; Lang et al. 2020; Radmanesh et al. 2020b). This is of clinical importance as leukoencephalopathy and cerebral microbleeds are associated with critical illness and higher mortality.

We report incomplete-ring enhancement in our first patient, which could suggest ADEM where breakdown of the blood–brain-barrier facing the white matter occurs at the leading edge of demyelination. However, incomplete-ring enhancement is also common in primary cerebral angitis, and vasculitis was also considered a differential in our first case (Boulouis et al. 2017). A case report of an ADEM-like illness following mild SARS-CoV-2 infection with typical brain, optic nerve and cord lesions demonstrated an incomplete-ring enhancing brain lesion, but no microbleeds or diffusivity changes were described. In addition, the clinical presentation was in contrast to our critically ill patient with poor neurological status; the latter is the emerging phenotype of COVID-19 leukoencephalopathy (Novi et al. 2020). Of note, reports in the COVID-19 pandemic of other parainfectious immune-mediated neurological conditions such as cytotoxic lesions of the corpus callosum and Guillain-Barré syndrome remain relatively sparse considering the worldwide prevalence (Xiong et al. 2021). However, that does not discount ADEM as a possibility in our case where a clinical response was seen following immunotherapy.

Microbleeds and diffusion restriction are hallmarks of COVID-19 leukoencephalopathy seen in both of our cases, and small-vessel vasculitis has been postulated as the cause of such changes (Agarwal et al. 2020a; Lang et al. 2020). In our second patient, we report novel diffusion restriction with low ADC values persisting over 3 months into admission. This exceeds the expected temporal evolution for diffusion restriction in infarcts or demyelination, which usually normalizes on follow-up. The persistent diffusion restriction in this case of COVID-19 leukoencephalopathy could be secondary to a prolonged ischemic insult in the context of vessel wall inflammation, a phenomenon seen specifically in vasculitis and not occlusive or embolic infarcts (Stanley et al. 2017). However, inflammatory etiologies such as ADEM can rarely show ongoing diffusion restriction. The lack of clinical response to immunotherapy in our second patient could lend support to a dominant vascular etiology although ultimately the high burden, and severity of lesions may be responsible for his poor outcome.

Leptomeningeal enhancement is a common COVID-19 neuroimaging finding (Kremer et al. 2020). Whilst the etiology is currently unknown in COVID-19, it is a well-recognized feature of cerebral vasculitis, not just meningitis (Boulouis et al. 2017). Arterial wall enhancement and thickening have also recently been described in COVID-19 lending support to a vasculitis (Lersy et al. 2020). Vessel wall or angiographic imaging was not possible in our cases due to renal failure and patient instability. The significant
callosal involvement in both our cases goes against embolic infarcts, small vessel disease or a hypoxic-ischemic etiology.

Both our cases depicted lesions showing ring-shaped diffusion restriction, previously described in two cases (Toledano-Massiah et al. 2020). This feature is reminiscent of cytotoxic edema at the leading edge of demyelination. Post-hypoxic necrotizing leukoencephalopathy has been postulated in some COVID-19 cases where there is confluent white matter signal change and diffusion restriction, and in this condition, it is thought to reflect widespread demyelination secondary to ischemia (Radmanesh et al. 2020a). A recent neuropathological study of COVID-19 described vascular and ADEM-like findings and has raised demyelination as secondary to an ischemic insult rather than a primary phenomenon (Reichard et al. 2020).

There is growing evidence for SARS-CoV-2 neuroinvasion, and recently, infection has been shown directly within infarcts in human post-mortem brain studies, within in vitro human brain organoids and in vivo mice models associated with neuronal death and remodeling of the vasculature (Song et al. 2021). The absence of SARS-CoV-2 in the CSF in our cases and the literature does not imply a lack of neuroinvasion but could be due to low PCR sensitivity or CSF penetrance. The constellation of neuroimaging findings in our cases could hypothetically represent the sequelae of SARS-CoV-2 neuroinvasion, virally mediated small-vessel vasculitis and subsequent ischemic demyelination with a leading spreading edge (ring-shaped diffusion restriction, incomplete-ring enhancement) and prolonged ischemia (persistent diffusion restriction).

The clinical picture in critically ill patients remains complex confounded by profound hypoxia and a cytokine storm, and the contribution of thrombotic microangiopathy cannot be discounted. Serial neuroimaging with MRI in critically ill patients with COVID-19 pneumonitis remains clinically challenging.

Fig. 2 Follow-up MRI. The axial (A) and coronal (B) T1-weighted images post-gadolinium demonstrate incomplete-ring contrast enhancement around 2 of the larger callosal and periventricular lesions (white arrowheads) as well as numerous foci of punctate contrast enhancement which numbered 18 in total. A follow-up scan depicts maturation with gliosis on T2-weighted images (C). Susceptibility-weighted imaging shows punctate microbleeds centred on the T2 lesions (D) most extensive in the splenium of the corpus callosum (white arrows). Isodiffusivity was seen on ADC maps with mild T2 shine-through on diffusion-weighted images (not shown).
Conclusion

The constellation of neuroimaging findings in our two cases of COVID-19 leukoencephalopathy and response to immunotherapy in one case supports either vasculitic or demyelinating immune-mediated processes; however, in light of recent evidence of SARS-CoV-2 neuroinvasion and vascular damage, we hypothesize that there is in fact a virally induced potent small-vessel vasculitis and possible ischemic demyelination. Our report aims to help clinicians recognize COVID-19 leukoencephalopathy, the first step in understanding this unique entity and developing therapeutic strategies.

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Declarations

Conflict of interest The authors declare no competing interests.

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