A possible Guillain-Barré syndrome/transverse myelitis overlap syndrome after recent COVID-19

Riyadh Alrubaye,1 Vijayamala Bondugula,2 Vidya Baleguli,3 Rosemary Chofo4

SUMMARY
Neurological manifestations are common in SARS-CoV-2 infection, including life-threatening acute muscle weakness, due to neuromuscular disorders such as acute transverse myelitis (TM) and Guillain-Barré syndrome (GBS). These syndromes can rarely coexist and present as an overlap syndrome. Here, we report a patient who developed acute symmetrical proximal lower limb weakness 5 days after diagnosis of COVID-19. GBS was diagnosed due to the presence of motor signs, albumin-cytological dissociation in cerebrospinal fluid examination and axonal damage according to nerve condition tests. However, abnormal areas on MRI of the thoracic spine and lack of improvement with intravenous immunoglobulin supported a diagnosis of TM. Therefore, a possible overlap between GBS and TM was established. To our knowledge, this is the third case report of GBS/TM overlap syndrome after COVID-19. The patient’s full and rapid recovery with intravenous corticosteroids and plasmapheresis supports our diagnosis.

BACKGROUND
COVID-19, caused by the virus SARS-CoV-2, has quickly spread worldwide since December 2019, with more than 250 million confirmed cases reported by the WHO in November 2021. The pandemic has continued to hammer hospitals with increasing numbers of cases, with broad impacts in limiting access to certain procedures and imaging that are routinely performed under regular circumstances.

Initially, the main symptoms of COVID-19 reported in late 2019 were respiratory symptoms; however, a wide range of acute neurological manifestations associated with COVID-19 have been reported in 2020 and 2021, with both the peripheral and the central nervous systems potentially affected.2 3

Guillain-Barré syndrome (GBS) is an acute inflammatory demyelinating polyradiculopathy that classically presents with ascending weakness, areflexia/hyporeflexia and sensory involvement.4 GBS is usually attributed to the production of antibodies that attack myelin protein in response to various infections, including SARS-CoV-2 infection.5 6

Acute transverse myelitis (TM) is an inflammatory disease of the central nervous system that mostly affects the paediatric population and is also triggered by an immune-mediated response to infection. The classical diagnostic criteria include back pain with acute weakness of the limbs, sensory deficit and autonomic dysfunction, along with abnormal changes on MRI.7 There are very few case reports of TM in patients with COVID-19 as compared with such reports for GBS.2 3

Since TM and GBS are both immune-mediated inflammatory diseases, it is possible for the two disorders to be concurrent, which is referred to as GBS/TM overlap syndrome. Herein, we report a case of GBS/TM overlap syndrome after COVID-19, which was initially considered to be only GBS, highlighting the importance of a differential diagnosis of this rare comorbidity during the pandemic with potentially limited imaging resources.

CASE PRESENTATION
A 72-year-old man was diagnosed with COVID-19 in the emergency department. He was administered 700 mg (7 mg/kg) of intravenous bamlanivimab and was discharged with oral dexamethasone (6 mg daily). Five days later, he returned because of worsening generalised weakness, severe back and leg pain, constipation, urine retention and multiple ground-level falls. He denied any sensory loss or visual disturbances. On examination, the patient was alert and oriented to person, place and time. Vital signs were within normal limits, with blood oxygen saturation of 98% breathing ambient air. His neurological assessment was notable for anisocoria, the left larger than the right (from previous injury), with the left being non-reactive. Other cranial nerves were intact, including the facial nerve. Motor examination using the Medical Research Council Manual Muscle Testing scale showed normal muscle power in the upper extremities. However, he had bilateral proximal lower extremity weakness with power of 1/5 and bilateral distal lower extremity with power of 4/5. Deep tendon reflexes were hypoactive in the upper extremities and absent in the lower limbs. The patient had normal muscle tone and no tremor, and fine coordination in the hands was intact. His gait was not assessable, as he was unable to bear weight. Flexor plantar responses were present bilaterally. Neurosensory assessment showed intact light touch, temperature and vibration sensation in the hands and feet.

INVESTIGATIONS
Laboratory testing revealed no significant abnormalities, including normal antinuclear antibody and vitamin B12 levels. Chest radiograph showed bilateral interstitial infiltrates, which were unchanged from 5 days previously.

Cerebrospinal fluid (CSF) analysis showed albumin-cytological dissociation with elevated

Correspondence to
Dr Riyadh Alrubaye; ralrubaye@gmail.com

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protein of 211 mg/dL, white cell count of 2 cell/L, normal glucose concentration and no oligoclonal bands. Routine bacterial and fungal culture and Gram stain were negative. Serological tests of the CSF were negative for cytomegalovirus, Epstein-Barr virus, venereal disease research laboratory, herpes simplex virus and SARS-CoV-2, and the paraneoplastic and autoimmune CSF panel were normal.

Whole-spine MRI sagittal sections with gadolinium did not show signal changes in the spinal cord (figure 1A and B) owing to some technical difficulties. However, thoracic axial spinal MRI T2-weighted images showed abnormal T2-hyperintense signals within the central thoracic spinal cord, extending from T7-T8 inferiorly to T11 and T12 (figure 2A and B) with multiple old canal stenoses at the cervical and lumbar levels and lack of root enhancement.

Nerve conduction analysis was performed with some technical difficulties due to resource restrictions associated with the COVID-19 pandemic, which showed decreased conduction velocities and decreased amplitudes, suggesting acute motor axonal neuropathy.

**DIFFERENTIAL DIAGNOSIS**

The possible differential diagnoses for the 72-year-old man with acute proximal lower limb weakness and areflexia after COVID-19 include acute inflammatory demyelinating polyneuropathy and TM.

Initially, we considered GBS (possibly motor variants) as the principal diagnosis, as the patient had sudden onset of lower limb weakness and areflexia. By contrast, acute flaccid weakness, back pain, autonomic dysfunction and hyperintense T2 MRI signals from T7 to T12 indicated a possible diagnosis of TM.

We believed that the patient’s neuromuscular disorder was mostly secondary to COVID-19. We investigated whether bamlanivimab could be a factor; however, his symptoms started before he was administered bamlanivimab and we found little information regarding the adverse effects of bamlanivimab in the literature.

**TREATMENT**

Based on the patient’s clinical features, we initially decided to administer intravenous immunoglobulin (IVIG) for possible GBS. However, the possibility of TM concurrency was subsequently entertained based on the worsening of both motor functions and urinary retention according to abnormal spinal images. Intravenous corticosteroid was administered at a dose of 1000 mg daily for 5 days as of day 6 from the admission date (figure 3). Surprisingly, the patient showed significant improvement in right hip movement, with improvement in right lower proximal muscle power from 0 to 4; however, the power in his left hip muscles remained at 0. Based on this partial recovery,
the patient was administered plasmapheresis from day 12 for five alternative sessions, which remarkably improved his left hip movement (figure 3).

OUTCOME AND FOLLOW-UP
The patient was discharged on day 22 after significant recovery. The patient was able to walk after five sessions of plasmapheresis and was discharged without the need for rehabilitation. The patient’s back and leg pain resolved, and he went back to his routine life within 1 week of discharge.

DISCUSSION
The concurrency of GBS and TM (ie, GBS/TM overlap syndrome) represents a rare paradox that typically affects children and young adults more frequently than elderly patients.6-9 The clinical features of GBS/TM overlap syndrome can include GBS elements such as acute flaccid paralysis with hyporeflexia and TM elements such as pyramidal signs, sphincter dysfunction and sensory level.8 It is critical to note that the presence of both sensory level and sphincter dysfunction suggests the concurrency of TM in the overlap syndrome.8 Albumin-cytological dissociation (elevated CSF protein and normal cell count) with an increased IgG index is a non-specific CSF finding. However, two practical measures are useful to evaluate the concurrency of GBS and TM: abnormal electrophysiological findings and the presence of a central T2-hyperintense thoracic spinal cord signal in MRI.9,10

Based on a review of the literature, the majority of reported cases of GBS/TM overlap syndrome have been attributed to infections such as acute viral illness, influenza, Mycoplasma, Legionella, Bartonella and Zika virus.8,11,12 Two previous cases of overlap syndrome after SARS-CoV-2 infection have been reported: one was a paediatric case,13 and the other was in a 40-year-old man.14 In the latter case, the patient presented with upper respiratory symptoms followed by lower limbs weakness, with mild COVID-19 findings determined by high-resolution computed chest tomography despite a negative SARS-CoV-2 PCR test. The initial diagnosis was TM based on cervical spine MRI findings; however, lack of improvement on steroid therapy necessitated CSF analysis, which showed albuminocytological dissociation, and nerve conduction analysis, which showed acute sensory and motor axonal neuropathy, a variant form of GBS that eventually improved with plasmapheresis.14 Sporadic cases of vaccine-induced GBS/TM overlap syndrome have also been reported.15 Finally, immunotherapy-related neurological disorders account for 1%-5% of all immune medication side effects.16 Both TM and GBS have been separately reported in some case reports after administration of nivolumab or pembrolizumab but their TM and GBS have been separately reported in some case reports of GBS/TM overlap syndrome can include GBS elements such as acute flaccid paralysis with hyporeflexia and TM elements such as pyramidal signs, sphincter dysfunction and sensory level. It is critical to note that the presence of both sensory level and sphincter dysfunction suggests the concurrency of TM in the overlap syndrome. Albumin-cytological dissociation (elevated CSF protein and normal cell count) with an increased IgG index is a non-specific CSF finding. However, two practical measures are useful to evaluate the concurrency of GBS and TM: abnormal electrophysiological findings and the presence of a central T2-hyperintense thoracic spinal cord signal in MRI.

Intravenous steroid and plasmapheresis showed a better outcome in our case than intravenous immunoglobulin and steroids.

Due to limited cases and reviews, further study of this overlap syndrome may help to improve the diagnostic and therapeutic approach.

Patient’s perspective
Initially they diagnosed me with COVID and when I got home my legs just gave away and I had severe back pain which led me to come back to the hospital. I lost use of my hip and all my muscles just shut down. After steroid and plasma treatment I started to feel great. Am currently doing great. Am walking and working all day. I want to compliment the doctors and nurses for their help.

Learning points
► Guillain-Barré syndrome (GBS)/transverse myelitis (TM) overlap syndrome should be suspected in patients presenting with severe back pain, motor weakness, areflexia, sensory level and urine retention.
► TM concurrent with GBS caused by COVID-19 can be initially missed secondary to the limited MRI use during the pandemic. This can delay the patient’s recovery and increase the length of stay.
► Intravenous steroid and plasmapheresis showed a better outcome in our case than intravenous immunoglobulin and steroids.

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Case reports provide a valuable learning resource for the scientific community and can indicate areas of interest for future research. They should not be used in isolation to guide treatment choices or public health policy.

REFERENCES


