Miller-Fisher Syndrome and Guillain-Barre Syndrome overlap syndrome in a patient post Oxford-AstraZeneca SARS-CoV-2 vaccination

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SUMMARY

We describe a patient who developed bilateral ocular motor nerve palsy, ataxia, facial diplegia and lower limb weakness 2 weeks post-Oxford-AstraZeneca SARS-CoV-2 vaccination, consistent with Miller-Fisher syndrome (MFS) and Guillain-Barre syndrome (GBS) overlap syndrome. Although some features of the patient’s presentation were typical of recently reported cases of a rare GBS variant post-Oxford-AstraZeneca vaccination, including severe facial weakness and a lack of respiratory involvement, to our knowledge this is the first reported case of MFS associated with SARS-CoV2 vaccination. While postvaccination GBS remains rare, it appears to have a favourable prognosis, and recognising this entity is therefore important for patient counselling and monitoring for potential complications.

BACKGROUND

Guillain-Barre syndrome (GBS) is a clinically heterogeneous immune-mediated condition involving the peripheral nervous system. It is the most common form of acute neuropathy with a lifetime risk for an individual of under 1 in 1000, and an incidence of one to two cases per 100 000 worldwide. Antecedent infections, such as Campylobacter jejuni and cytomegalovirus are commonly associated with GBS. A proposed pathophysiological mechanism is that of molecular mimicry whereby a foreign antigen evokes an immune response which cross reacts with gangliosides on the peripheral nerves. Miller-Fisher syndrome (MFS) is a rare variant of GBS characterised by the triad of ataxia, areflexia and ophthalmoplegia. Cases in which MFS coexists with other features of GBS, such as limb weakness and facial weakness, are considered MFS-GBS overlap syndromes.

An association between GBS and vaccination has previously been described. A meta-analysis of adverse event data following mass influenza A (H1N1) vaccination in 2009 revealed a small increased risk of GBS (relative risk 2.35, 95% CI 1.42 to 4.01). Interestingly, two recent case series have documented striking examples of the rare bifa- cial weakness with limb paraesthesia (BFP) variant of GBS within 3 weeks post Oxford-AstraZeneca vaccination. These patients developed profound facial weakness and lower limb symptoms but had a good prognosis with no respiratory complications. Reports of postvaccination MFS are less frequent and, as yet, no cases of MFS have been reported post-SARS-CoV-2 vaccine.

Here, we report the first case of a patient presenting with MFS-GBS overlap syndrome following their first dose of Oxford-AstraZeneca vaccination. The patient avoided respiratory complications and had a good outcome, supporting the evidence for a favourable prognosis in cases of GBS following SARS-CoV-2 vaccination. This case also highlights the need for vigilance for a broad spectrum of SARS-CoV-2 vaccine-associated GBS variants.

CASE PRESENTATION

A 63-year-old man presented 14 days following his first dose of Oxford-AstraZeneca vaccination. He experienced new-onset lower back pain 9 days post-vaccination, and then 5 days later developed severe bilateral facial weakness, unsteadiness, lower limb weakness and paraesthesia over a 48-hour period. There were no urinary or bowel symptoms. He had no medical history and was not on any regular medications. He lived with his wife at home and was independent.

Neurological examination revealed profound sensory ataxia, facial diplegia involving the forehead, proximal lower limb weakness, bilateral lower limb areflexia and impaired distal lower limb proprioception. He was unable to walk without assistance and had incomplete eye closure bilaterally.

On his second day of admission, he reported diplopia on lateral gaze bilaterally. Examination revealed impaired adduction, restriction upward gaze and intorsion with down gaze bilaterally, consistent with partial cranial nerve III palsies. Direct and consensual pupillary responses were normal. His bedside forced vital capacity was 2.2 L. His GBS disability rating score was 3.

INVESTIGATIONS

Testing for acquired causes of neuropathy was normal, including urea and electrolytes, liver function tests, glycated haemoglobin, B12/folate levels, thyroid function tests, serum protein electrophoresis and serum-free light chains, hepatitis B/C, HIV and syphilis serology, erythrocyte sedimentation rate, C reactive protein, antinuclear antibodies and antineutrophil cytoplasmic antibodies.

Cerebrospinal fluid (CSF) examination revealed albumin cytological dissociation with a markedly elevated protein of 2.99 g/L (normal range 0.15–0.45 g/L), 0/µL erythrocytes, 4/µL lymphocytes and 1/µL polymorphonuclear leukocytes. Serum
anti-GQ1b antibody was negative. Other potential causes for an elevated CSF protein were unrevealing with a negative CSF culture, normal CSF cytology and flow cytometry.

MRI of the brain with contrast demonstrated enhancement of the facial and oculomotor nerves bilaterally, consistent with the clinical examination findings of facial diplegia and partial bilateral cranial nerve III palsies (figure 1).

Nerve conduction studies (NCS) and needle electromyography (EMG) showed evidence of a long-standing axonal neuropathy with reduced motor and sensory amplitudes and length-dependent chronic neurogenic changes on EMG, but no acute abnormalities. NCS and EMG may not show changes early in the course of GBS and may be normal in cases of MFS.9–11

TREATMENT

Given the clinical features and temporal progression of symptoms, the patient was treated with a 5-day course of intravenous immunoglobulin (2g/kg). He also received supportive care with topical lubricating eye drops, eye patches and an indwelling catheter after developing urinary retention. He underwent 4 weeks of inpatient rehabilitation for persistent gait difficulties and lower limb weakness.

OUTCOME AND FOLLOW-UP

At 6 weeks after onset of symptoms, the patient’s ophthalmoplegia and ataxia have resolved and he has minimal residual bilateral facial weakness. He has persistent bilateral proximal lower limb weakness (Medical Research Council grade 4). The patient has returned home, is living independently and is ambulating with a four-wheel frame.

DISCUSSION

To our knowledge, this is the first to report the MFS-GBS overlap syndrome associated with the Oxford-AstraZeneca SARS-CoV2 vaccination. Recently, two case series have documented 12 patients who developed the rare BFP variant within 3 weeks following the first dose of the Oxford-AstraZeneca vaccination.7–8 Notably, patients in these series had a favourable prognosis, a lack of respiratory complications, and four had markedly elevated CSF protein (>1.9g/L).7–8 Our case exhibited strikingly similar clinical characteristics, raising the possibility of a similar pathogenic mechanism, but also had classical features of MFS with ataxia, ophthalmoplegia and areflexia.

Several cases of MFS have also been documented following SARS-CoV-2 infection.12 Interestingly, although the GQ1b antibody is observed in 85% of MFS cases,13 all documented cases of MFS post-SARS-CoV-2 infection have been anti-GQ1b negative,12 suggesting a novel immunopathogenic mechanism.12–14 The anti-GQ1b antibody was also negative in our case which may imply a shared mechanism to MFS post-SARS-CoV-2, perhaps via molecular mimicry of the SARS-CoV-2 spike protein.

We performed a literature review using PubMed, MEDLINE and Embase for all published postvaccination MFS cases up to 21 August 2021. The following keywords, [“Miller Fisher Syndrome”] AND [“vaccine” OR “post vaccination” OR “SARS-CoV-2” OR “COVID-19”], were used in the search strategy. There have been six prior case reports of postvaccination MFS: two following combined diphtheria, tetanus and pertussis vaccine (Tdap), one following seasonal influenza vaccine, one following combined Pneumovax and seasonal influenza vaccine, two following combined seasonal influenza and H1N1 vaccine, and one following H1N1 vaccine alone.11–15–20 All cases presented with symptoms from 5 days to 14 days postvaccination, except in one patient with HIV in which symptom onset was 40 days postseasonal influenza vaccine and 34 days post-H1N1 vaccine, respectively.17

All previous cases of post-vaccination MFS had raised CSF protein of under 1g/L. Our case is distinct in this respect, with a CSF protein of 2.99g/L, but consistent with cases of the BFP variant post-Oxford-AstraZeneca Vaccine.7 Three previous cases had positive anti-GQ1b antibodies, and in two cases, GQ1b status was not reported. All cases were treated with intravenous immunoglobulin (2

Patient’s perspective

I initially presented to my local general practitioner (GP) with severe back pain and was prescribed painkillers. Two days later, I revisited my GP as the pain had not diminished and had spread to my legs. He advised me to present to the hospital, given the increasing severity of symptoms. Fortunately, the neurological team ran a battery of tests and were quickly able to confirm I had GBS with the Miller-Fisher variant. Within less than 24 hours of admission, I was receiving the first of five doses of IVig treatments. I remained in hospital for a further week for observation then transferred to a rehabilitation facility for 3 weeks to work on the damage to my legs’ nerves, which is progressing well. During the whole process, I was kept informed of both what was being done and outcomes to expect, and this really took the stress out of the situation.

Learning points

➤ Guillain-Barre syndrome (GBS) is a heterogeneous autoimmune condition and can present with the overlap of rare clinical variants including bifacial weakness with limb paraesthesia and Miller-Fisher syndrome (MFS).
➤ The SARS-CoV-2 vaccination may be associated with atypical forms of GBS, including MFS–GBS overlap syndrome.
➤ SARS-CoV-2 vaccine-associated GBS may be associated with atypical investigation results, such as markedly elevated CSF protein and anti-GQ1b seronegativity, but appears to have a favourable prognosis.

Figure 1 T1 MRI brain postcontrast images. (A, top) bilateral oculomotor nerve enhancement. (B, bottom) Bilateral facial nerve enhancement.
immunoglobulin, except one case post-Tdap vaccine reported by Garg and Moudgil, which was treated with plasmapheresis. All cases of MFS following Tdap vaccine and one case post-seasonal influenza vaccine had complete resolution of symptoms; however, the prognosis of other cases may have been confounded by the variable duration of follow-up.

Our case highlights the wide clinical spectrum of GBS variants and the need for close surveillance for atypical complications of the SARS-CoV2 vaccination. Although a coincidental relationship with the Oxford-AstraZeneca vaccination cannot be excluded, we feel that the temporal onset of clinical symptoms, the presence of anti-GQ1b seronegativity and the coexistence of the rare BFP variant that has emerged as an associated clinical syndrome makes this unlikely. The patient avoided respiratory and autonomic complications and is making a promising recovery, supporting the existing evidence for a good prognosis in post-SARS-CoV2 vaccine GBS.

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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.

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