Emotion processing in the minimally conscious state

As a newly described condition distinct from coma or the vegetative state, minimally conscious state (MCS) is characterised by a threshold level of consciousness, and diagnostic criteria have recently been proposed.1 In MCS, cognitively mediated behaviour occurs inconsistently, but is reproducible or sustained enough to be differentiated from reflexive behaviour. It is clinically essential to distinguish this condition from persistent vegetative state (PVS), due to a potentially more favourable outcome.1 So far, whether patients in MCS can process emotion is unknown.

Cortical processing has been described in PVS using auditory and visual functional paradigms with positron emission tomography.2 3 However, to date hardly any functional imaging studies are available in patients in MCS.4 We used fMRI to assess brain activity induced by an emotional stimulus in a patient in MCS.

A 17 year old man was riding his bicycle when he was hit by a train. The accident resulted in head trauma and immediate coma, progressing to MCS over the course of 4 months, when he was admitted to our institution. This research protocol was approved by the Institutional Ethics Committee. At the time of the fMRI study, 5 months after the accident, the patient localised noxious stimuli, had spontaneous eye opening, detectable sleep/wake cycles, sustained visual fixation, and contingent smiling, thus meeting criteria for MCS. A structural MRI study showed mild cortical atrophy and dilated ventricles. Auditory evoked potentials showed decreased conduction velocities at brainstem level. The patient increased his level of awareness 2.5 months after the functional study was conducted. Auditory evoked potentials after recovery were within normal range, while MRI showed much less ventricle dilatation. Six months after recovering full consciousness, he was able to chat normally and feed himself. Currently we are retesting the patient with the same paradigm.

Non-familiar voice ν silence and mother’s voice ν non-familiar voice recognition were tested in an fMRI block design with 30 seconds per epoch. The patient listened to his mother reading a story, followed 30 seconds later by an age matched voice reading the same story, for 30 seconds with silence epochs in between. Blood oxygen level dependent images were acquired using a T2 weighted gradient echo planar sequence on a General Electric Signa CVI, 1.5T system with real time image processing of multislice and multi-phase images during patient stimulation and rest periods. The Medx 3.4 Sensor System was used to carry out fMRI post-processing, including motion correction and Gaussian smoothing. An uncorrected significance threshold of P<0.001 was used because amygdala and insula activation was expected, owing to emotional voice processing. Activated clusters were localised following co-registration with an anatomical T1-IR volume.

Subtraction of the phrases read by the age matched voice from silence was the control experiment, showing a significant focus of activation in the transverse and superior temporal gyr, which spread to the planum temporale; more anterior activation was found in the superior (right) and inferior (left) insula (fig 1A). The subtraction of the mother’s phrases from the age matched voice disclosed a strong activation of the amygdala and insula spreading to the inferior frontal gyrus; there was also weaker activation of the transverse temporal gyrus, temporal operculum, and planum temporale (fig 1B,C). Activation was lower on the right hemisphere in both comparisons, non-familiar voice ν silence and familiar voice ν non-familiar.

To the best of our knowledge, our results provide for the first time anatomical evidence for the response of an MCS patient to a familiar voice, in which both amygdala and insula appear to play a major role.

The activation pattern of the control experiment agrees with previous studies.5 Our results showed that the mother’s voice activates the extended amygdala, an emotion related structure, and a directly connected area such as the insula, perhaps acting jointly as limbic integration cortex. Although residual cerebral activity was unequivocal in our case, representing fragmentary cognitive processing, it should not be assumed that it depicts a fully integrated system required for normal levels of awareness; however, our findings highlight the legal and ethical implications of careless bedside chatter. Whether functional imaging represents a reliable method to evaluate neural processing in MCS patients, in whom cognitive output is extremely difficult to assess, remains to be seen.

References


Figure 1 Brain areas of activation produced by non-familiar voice subtracted from silence in coronal view (control experiment, A). Brain areas of activation produced by mother’s voice subtracted from non-familiar voice in coronal view (B), and in axial view (C)
Neurosyphilis presenting with gummatous oculomotor nerve palsy

Although epidemiological studies suggest that the incidence of primary syphilis is rising, neurosyphilis remains an uncommon manifestation of *Treponema pallidum* infection. In addition, the MRI appearances of this treatable neurological condition are not well known. Many patients with neurosyphilis are asymptomatic, but manifestations include subacute basal meningitis, a meningovascular syndrome of small deep cerebral and cranial nerve infarctions, chronic gummatous inflammation with focal intracranial mass lesions, chronic compartmental dementia of general paresis, and chronic sensory-ataxic myelopathy of tabes dorsalis. We report a case in which a meningeal form of neurosyphilis presented with rapid evolution of a pupil-involving oculomotor nerve palsy to highlight the clinical, CSF, and MRI features and good response to treatment.

Case report

The patient was a 54 year old right handed homosexual man with a history of syphilis of unknown stage, treated with penicillin 25 years previously. He was well until 6 weeks prior to admission when he sustained minor head trauma in an automobile accident, followed by intermittent headaches, fatigue, photophobia, and anorexia. Four days before admission he developed worsening and persistent drooping of the right eyelid and double vision. On examination, his mental status was remarkable only for psychomotor slowing. The right pupil was round but enlarged at 6 mm and sluggishly constricted to 5 mm with direct and consensual light stimulation as well as near vision. The left pupil was round and 4 mm and constricted briskly to light. The right eye showed a moderate ptosis of the upper lid, and the globe was deviated laterally in primary gaze with markedly impaired adduction and elevation. In the left eye, ptosis was absent and ocular motility was normal. Other cranial nerve, sensory, motor, and reflex functions and gait were normal with the exception of a slight decrease in vibration and position sense in the feet. There were no signs of meningial irritation. Head computed tomography (CT) and CT angiography revealed no blood in the subarachnoid space nor evidence of intracranial aneurysm. MRI of the head (fig 1) showed a sphenoid contrast-enhancing lesion at the root of the right oculomotor nerve, which extended towards the cavernous sinus. Incidentally noted were right cerebellar and right frontal developmental venous anomalies. CSF examination revealed normal opening pressure at lumbar puncture, 344 white blood cells (WBCs) (97% lymphocytes), 14 red blood cells (RBCs), protein of 167 mg%, and glucose of 39 mg%. CSF VDRL and serum RPR titres were unchanged. At 6 months, no additional improvement in oculomotor nerve functions was seen but fatigue had subsided. Repeat MRI 7 months after hospital admission showed complete resolution of the oculomotor nerve abnormality.

Discussion

Neurosyphilis is known to cause oculomotor nerve palsies either in the meningovascular phase, due to small vessel vasculitis with resultant nerve infarction, or in granulomatous basal meningitis, due to inflammation of the nerve or its investiture; however, the literature on syphilitic mass lesions around the oculomotor nerve is sparse. Vogl et al. reported a case of oculomotor nerve palsy associated with MR findings similar to ours that also resolved with penicillin treatment. Standaert et al. described an enhancing penicillin-responsive lesion based in the interpeduncular cistern that compressed the ventral midbrain. The oculomotor nerve lesion in our patient was isoointense to adjacent brain on T1 and T2 sequences, with brisk enhancement after intravenous injection of gadolinium contrast. We believe the lesion was a manifestation of meningeal syphilis in the form of an oculomotor nerve gumma. A gumma is a focally accentuated, exuberant granulomatous response of the meninges, typically with sparse treponemal organisms. Nonetheless, treatment of the underlying infection quietes the inflammatory process and can, as in our patient, lead to significant reversal of neurological deficit. We add our case to the growing literature on MR correlates of neurosyphilis and encourage a search for neurosyphilis when an unexplained mass lesion is present in the basal subarachnoid space. Neurosyphilis, albeit rare, still deserves inclusion among eminently treatable causes of a rapidly developing oculomotor nerve palsy.

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References


High dose cyclophosphamide for severe refractory myasthenia gravis

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Drachman et al published the beneficial effects of high dose cyclophosphamide in three patients with severe refractory myasthenia. We recount our experience of three myasthenic patients treated in a similar way.

### Materials and methods

All patients participated in studies approved by the Drexel University College of Medicine and signed informed consent. These three patients, with severe (class IVb) refractory MG includes all patients treated. Patients received cyclophosphamide 50 mg/kg (adjusted ideal body weight)/day over four consecutive days. Patients received antibacterial, antiviral, and antifungal prophylaxis. Haemorrhagic cystitis prophylaxis included Mesna and forced diuresis. Packed red cells and platelets were transfused to maintain haemoglobin >8.5 g/dL and platelets >10 x 10^9/L. Patients received filgrastim (G-CSF) (5 μg/kg/day) starting day 10 until their absolute neutrophil count (ANC) reached 10 x 10^9/L for two consecutive days.

### Results

Patient 1 was diagnosed with seronegative MG at 30 years of age by a positive tensilon test and a decremental response on repetitive stimulation. Initial treatment included pyridostigmine and plasmapheresis, but worsening symptoms prompted thymectomies at 12 and 18 months later. Her thymic pathology revealed thymic hyperplasia. Additional treatment with only transient responses included low dose oral cyclophosphamide, intravenous Ig, azathioprine, methylprednisolone, and continued pyridostigmine with plasmapheresis. She required 27 intubations between initial diagnosis and immunosuppressive treatment at 41 years of age.

Patient 2, previously reported, suffered from both seronegative MG and chronic inflammatory demyelinating polyneuropathy (CIDP). He presented at 47 years of age with fluctuating double vision, ptosis, dysphagia, arm weakness, and breathing difficulties. Testing revealed a decremental response on repetitive stimulation. Pyridostigmine was initiated. Thymectomy revealed a 75 g lipoma. His MG resulted in two intubations. After thymectomy, to control symptoms, prednisone (25–40 mg daily) was required. At 54 years of age, CIDP was diagnosed. Despite steroids (plasmapheresis, intravenous Ig, azathioprine, and pyridostigmine) he continued with symptoms of double vision, dysphagia, and dysphasia with a continued decremental response to repetitive stimulation. At 56 years of age, he underwent high dose cyclophosphamide without stem cell rescue.

Patient 3 was diagnosed with antibody positive MG at 12 years of age, initially treated with pyridostigmine. She received her first thymectomy at age 18 years and continued on pyridostigmine and occasional steroids. By 36 years of age, she was steroid dependent. Between ages 38 and 41 years she required 11 intubations and only transiently responded to intravenous Ig and plasmapheresis. A second thymectomy was performed at age 39 and cyclosporine (CsA) was initiated. She continued on prednisone 25 mg qod, scheduled intravenous Ig every 3–4 weeks, and intermittent plasmapheresis. The CsA and Cellept were maintained but poorly tolerated. At 41 years of age, she underwent high dose cyclophosphamide without stem cell rescue.

### Treatment course

Patient 1 had 13 days of neutropenia, required three of packed red cells and three platelet transfusions. Patient 2 had 9 days of neutropenia, required two units of packed red cells, and three platelet transfusions. Patient 3 had 11 days of neutropenia, required five units of packed red cells, and two platelet transfusions. Patients 1 and 3 experienced MG flares requiring intravenous Ig and plasmapheresis, but neither required intubation.

### Neurological follow up

Patient 1, intubated 27 times before treatment, required a single intubation during 48 months of follow up. To control less severe exacerbations, during the first 40 months after immunosuppressive treatment, oral cyclophosphamide was necessary. She continues scheduled plasmapheresis and pyridostigmine. No other immunomodulatory medications are prescribed.

### Discussion

The patients discussed have all suffered from severe refractory MG, which requires multiple intubations. All underwent thymectomy: patients 1 and 3 repeat thymectomies. Patient 2 had an early and sustained response to treatment. Patients 1 and 3 had multiple exacerbations. As this treatment targets IgG production, exacerbations following treatment are expected. Patient 1, who required 27 intubations before treatment and only once since, and who has in the past 6 months stopped oral cyclophosphamide, may yet to enjoy the maximum benefit of this treatment. Patient 3, one year after treatment, has an improving activity level. The intervals between exacerbations are increasing: 5, 8, and 11 weeks. It is 26 weeks since her last exacerbation.

Recently, Drachman et al published a single institution case series of three patients with refractory MG who were also treated with high dose cyclophosphamide. In this series, one patient had AChR antibody negative MuSK antibody positive myasthenia. Their mean disease duration was 10.3 (range: 3–15) years; one required intubation and median follow up was 24 (range: 7–40) months. In comparison, in the three patients described here, two had antibody negative myasthenia and the mean disease duration was 16.3 (range: 9–29) years. All required multiple intubations: 27, 2, and 11, and our median follow up is 25 (range: 13–48) months. During follow up, patient 3’s serum AChR levels remained detectable and did not correlate with her clinical course. Drachman et al reported a decline in antibody levels in their patients treated in a similar way, although AChR antibody titres and MuSK antibodies persisted in their patients even after 2 years. This suggests that long term remissions in MG may be possible even without achieving complete immunosuppression. High dose cyclophosphamide has the potential to significantly reduce symptoms and increase quality of life among people with MG refractory compared to conventional treatment. Long term follow up is necessary to evaluate the duration effect and time to maximum benefit. High dose cyclophosphamide treatment warrants further study as a treatment for severe refractory MG.

### Table 1

<table>
<thead>
<tr>
<th>Age/sex</th>
<th>Patient 1 41/female</th>
<th>Patient 2 56/male</th>
<th>Patient 3 41/female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration of MG (y)</td>
<td>11</td>
<td>9</td>
<td>29</td>
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<tr>
<td>MG severity class</td>
<td>IVB</td>
<td>IVB</td>
<td>IVB</td>
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<tr>
<td>AChR positivity</td>
<td>Undetectable</td>
<td>Undetectable</td>
<td>Detectable</td>
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<tr>
<td>Pyridostigmine</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Thymectomy(ies)</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>iv Ig (no of infusions)</td>
<td>1</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Prednisone</td>
<td>10–100 mg qd, duration 3 years</td>
<td>40–100 mg qd, duration 7 years</td>
<td>10–60 mg qd, duration 4 years</td>
</tr>
<tr>
<td>Plasmapheresis</td>
<td>2/1</td>
<td>14</td>
<td>16</td>
</tr>
<tr>
<td>(no. of procedures)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azathioprine</td>
<td>50 mg/d, duration 7 months limited by nausea/vomiting</td>
<td>200 mg qd, duration 2 months limited by nausea/vomiting</td>
<td>50–150 mg qd, duration 15 months</td>
</tr>
<tr>
<td>Oral cyclophosphamide</td>
<td>100 mg qd, 28 months</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>50–125 mg bid, duration 3 months</td>
<td>250–500 mg qd, duration 7 months</td>
<td></td>
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<td>Cellcept</td>
<td></td>
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</tr>
</tbody>
</table>

MG: myasthenia gravis; iv, intravenous; Ig, immunoglobulin; qd, four times daily; bid, twice daily.
Acute head drop after cervical hyperflexion injury

Head drop is familiar to neurorogists, but not widely appreciated by neurosurgeons. There are multiple causes of this condition in which the patient is unable to hold their head up because of weakness of the neck extensor musculature. It predominantly results from primary muscle pathologies in the neck extensor muscles, with occasional evidence supporting a neurogenic aetiology.1 2 I describe three patients in whom acute head drop closely followed cervical hyperflexion injury, and suggest that the cause is bilateral traction neurapraxia of the cervical extensors or more cervical dorsal rami.

Patient A was an 84 year old man who enjoyed excellent health prior to falling backwards, striking his occiput on a wall and sustaining forced flexion of the cervical spine. He complained of posterior cervical pain but, when seen in casualty for closure of an occipital laceration, was found to be neurologically intact. Cervical x-rays showed only degenerative disease in the mid-lower cervical spine and loss of lordosis. Over 2 weeks the pain in his neck resolved, but he became aware of a difficulty holding his head up as the day progressed and, later, of aching in his neck extensor muscles. He was referred to neurosurgery as a possible case of delayed instability. Cervical x-rays demonstrated 5° of forward angulation at C4/C5, which did not change with neck flexion, but were otherwise unchanged. He remained neurologically intact but had progression of the angulation and development of neck pain. Cervical x-rays showed 9° of neck revealed normal soft tissue anatomy. A neurological opinion confirmed the normal examination, other than head tilt. There was no evidence of inflammatory, auto-immune, or neoplastic disease. There were no features of Parkinson's disease or amyotrophic lateral sclerosis (ALS).

Electroneuromyography (EMG) studies of the neck muscles were normal 3 weeks after injury were normal in the ventral muscles, but there were typical features of acute partial denervation in the neck extensors bilaterally, particularly in a band in the mid-to-lower cervical spine with more normal EMGs above and below this. However, electrophysiological examination of the limbs was normal also and consistent with an asymptomatic peripheral neuropathy. The patient declined muscle or nerve biopsy.

In view of patient A's course and the evidence in patient B of acute denervation that might recover, patient B was managed expectantly. The Philadelphia collar was used to maintain range of neck movement and encourage use of the neck extensor muscles. He was given a Philadelphia collar, which was worn by day once he became aware of head tilt. The patient was discharged to recover to normal over 4 months, including recovery of the spinal alignment at C5/6, and the Philadelphia collar was withdrawn. There has been no recurrence of head drop.

Patient C was similar to patient B. He suffered a whiplash injury in an RTA and developed head drop and angulation at C5/6 on cervical x-rays 2 weeks later. Investigation and management mirrored patient B. He had an asymmetry of his neck extensor muscle EMG, which suggests partial denervation, but otherwise was normal clinically, biochemically, and electrophysiologically. We did not suggest muscle or nerve biopsy as it was clear he would be managed conservatively. Physiotherapy and external bracing, patient C made a complete recovery in 2 months, including recovery of spinal alignment at C5/6. There was no recurrence of head drop.

Although there are reports of head drop in conditions predominantly affecting neural rather than muscular elements,1 2 Umapathi et al.3 cite Braun et al.4 who treat refractory torticollis by sectioning multiple cervical dorsal rami without generating significant functional deficits, as evidence that focal denervation of neck extensor muscles is unlikely to cause head ptosis. This surgical denervation, however, is unlikely and the denervated muscles are unlikely to be grossly abnormal because of secondary changes resulting from the underlying condition. The cat neck extensor muscle biventer cervicis (analogous to human semispinalis capitis) has tendinous insertions defining serially arranged compartments, each receiving segmental innervation from a cervical dorsal ramus. The muscle only generates useful tension if all compartments are co-activated; stimulated compartments act as weak springs in series and dissipate force out of the muscle.5 There is some evidence for similar architecture in human neck extensors: they receive innervation from several cervical dorsal rami and have tendinous inscriptions producing several at least partially serial compartments.6 Denervation of one compartment bilaterally would produce significant weakness and fatigability in such compartmentalised muscles. Additionally, the deeper muscles only traverse one motion segment and are innervated by one posterior primary root. Segmental denervation of either type of muscle would lead to angulation at a motion segment, limited in degree by intact joints, ligaments, and disc space.

Whiplash injury can cause neurapraxia of cranial nerve XI, XII, and branches of the cervical plexus,7 8 and there are other reports of traction neuropathies in the neck.9 In the present cases, the close temporal relationship of the head drop to a forced flexion injury and the EMG findings suggesting acute denervation of neck extensor muscles are consistent with a neurogenic mechanism. Although dystonia of neck flexor muscles can produce head drop, these patients could easily lift their chins and there was no evidence of ventral neck hypertonia on clinical examination. In addition, in patients B and C, there were normal EMG findings in the ventral neck muscles but abnormal findings in the neck extensors.

Neurapraxia of dorsal primary rami would be expected to recover in 21 months without intervention in patients B and C. Equally, muscle tearing would recover in time, but it is inconceivable that sufficient fibres would have been torn to produce head drop without also producing soft tissue abnormalities on MRI scanning. This is not the case. Only two of the cases were investigated to exclude primary neuromuscular disorders. These were excluded in patient C. Although patient B had evidence of a pre-existing peripheral neuropathy, this may simply have made him more prone to traction neurapraxia after whiplash and his eventual recovery is consistent with the proposed mechanism.

It is unclear why this syndrome has not been described before. Perhaps most whiplash injuries produce insufficient neurapraxia to provoke head drop unless patient factors adversely affect the transmission of forces to the nerves or their susceptibility to injury. In non-predisposed individuals, sufficiently severe injuries might instead produce fractures/dislocations, whose management masks signs of a concomitant neurapraxia. Less severe injuries might provoke a head drop, which is either not recognised or recovers quickly and never requires secondary referral. Furthermore, although motor deficits may be rare after whiplash, sensory symptoms may be components of the patient's symptoms in a case of “typical” whiplash syndrome. There is support for this
We report a case of acute disseminated encephalomyelitis (ADEM) temporally associated with Campylobacter gastroenteritis in a previously fit man. A Medline search using the keywords “ADEM”, “demyelination”, and “campylobacter” revealed no previous reports of ADEM associated with Campylobacter infection in isolation.

A 24 year old man presented to his general practitioner on day 8 of history of non-bloody diarrhoea associated with fevers and sweats. His past medical history was unremarkable. He drank 6 units of alcohol per week and smoked occasionally. His general practitioner prescribed loperamide for symptomatic relief. Campylobacter species was later isolated from stool samples. By day 5 of his illness, his diarrhoea had settled and he had become constipated. However, he remained febrile and developed nausea and vomiting. His general practitioner prescribed erythromycin but he tolerated only two doses because of nausea.

Fourteen days into the illness he was admitted to hospital complaining of headache, fever, and sweats. Examination revealed a temperature of 38.4°C, pulse of 65 beats/min and normal blood pressure. Rectal examination revealed hard faeces but no tenderness or focal neurological signs. His haemoglobin was 15.3 g/dl, leucocyte count was 13.3 x10^9/l (87.1% neutrophils) and C-reactive protein was 12.8 mg/l. Two days after admission (day 16 of illness), his family reported a change in his personality and he complained of slurring of speech, intermittent diplopia, and difficulty in walking. Examination revealed mild dysaesthesia, left sided facial weakness, mild left pyramidal limb weakness, and decreased sensation in the left leg. Tendon reflexes were brisk but plantar responses were flexor. His gait was ataxic. Cranial CT scan showed no significant abnormalities. Lumbar puncture revealed an opening pressure of 160 mm CSF, total cell count of 34/mm^3 with a white cell count of 20/mm^3 (100% lymphocytes), total protein of 541 mg/l, glucose of 3.2 mmol/l, and negative oligoclonal banding. No organisms were seen and PCR was negative for enteroviruses and herpes virus. An EEG showed mild excess of generalised slow wave activity. Cranial MRI scan was performed on a 1.5 T Siemens magnetic system. T2 weighted imaging of the brain showed multiple high signal foci in the supra- and infra-temporal compartments involving the cortex, white matter, and deep grey matter. One lesion in the right peri-trigonal white matter showed slight enhancement following intravenous gadolinium diethylenetriaminepenta-acetic acid (gadolinium DTPA) injection (fig 1). The abnormalities were consistent with ADEM. The patient was initially treated with aciclovir 1000 mg four times daily, ampicillin 2 g 4 times daily and ciprofloxacin 500 mg twice daily, but was subsequently given intravenous methylprednisolone 1 g daily for 3 days after the diagnosis of ADEM was made. Aciclovir and ampicillin were discontinued when the negative laboratory results were available but ciprofloxacin was continued for 7 days. One day after treatment with methylprednisolone he noticed an improvement in his speech and gait, and after 7 days of starting treatment he had no ataxia and was discharged home. He appeared to have made a full recovery when he was reviewed at 6 weeks and has since remained asymptomatic.

ADEM is an acute monophasic immune mediated inflammatory demyelinating disease of the central nervous system. It is an uncommon but a serious condition with mortality rates estimated between 10–30%. In the majority of cases ADEM develops after systemic viral infections but commonly measles, mumps, rubella, influenza A and B, herpes simplex, Epstein-Barr virus, varicella, and vaccinia. It has also been reported following bacterial infections with Mycoplasma pneumoniae, Chlamydia, Legionella, and Streptococcus, or following immunisations for rabies, diphtheria/tetanus/pertussis, smallpox, measles and Japanese B encephalitis. The pathogenesis of ADEM is poorly understood. However, the evidence suggests that activated T cells, which recognise amino acid sequences shared between microbial epitopes and myelin antigens, attack central nervous system structures alone or in synergy with antibodies. Viral or bacterial superantigens could likewise trigger autoimmune T cells with similar results.

The diagnosis of ADEM is usually made clinically with the aid of MRI scanning. Lumbar puncture finding can be normal or show mononuclear pleocytosis and mild protein elevation. There are few diagnostic criteria based on evidence based treatment regimens, but treatment is usually instituted with high dose glucocorticoids. Plasmapheresis and intravenous immunoglobulin have also been used. Campylobacter gastroenteritis is the most common cause of acute gastroenteritis in the UK, accounting for over 56 000 cases in 2000. Its incidence has risen progressively over the past 2 decades. In the majority of cases, the illness self terminates within a few days with no long term consequences. It is estimated that approximately 1/1000 reported campylobacteriosis cases leads to Guillain-Barré syndrome, and around 33% of Guillain-Barré syndrome cases in a year may be triggered by campylobacteriosis. Huber et al reported a case of combined ADEM and acute motor axonal neuropathy following Campylobacter jejuni infection and hepatitis A. Immunisation of the ADEM group may be triggered by campylobacteriosis. Nasralla et al reported a case of postinfectious encephalomyelitis in a patient with Campylobacter jejuni enteritis. Cranial MRI scanning showed a combination of predomi- nantly grey matter involvement with concomi- tant focal areas of subcortical white matter lesions with no parenchymal abnormalities, on which the authors felt to be different from the pattern of signal abnormalities seen in patients with ADEM. The MRI abnormalities in our case were in keeping with ADEM and the subsequent clinical course as well as improvement on steroids were consistent with ADEM. In our case, the paucity of reported cases of Campylobacter infection and Guillain-Barré syndrome is the predominant focal areas of subcortical white matter lesions with no parenchymal abnormalities, in which the authors felt to be different from the pattern of signal abnormalities seen in patients with ADEM. The MRI abnormalities in our case were in keeping with ADEM although, as with the case reported by Huber et al, the amount of enhancement was mini- mal, indicating that the majority of the lesions were not acute. The paucity of reported cases of ADEM following Campylobacter infection is surprising given the association between Campylobacter jejuni infection and Guillain-Barré syndrome and the pathogen- esis of the latter. In these cases, Campylobacter jejuni induces humeral and cellular immune responses due to molecular mimicry and specific lipopolysaccaride epitopes on the infecting agent and target epitopes on the
surface components of the peripheral nerves, resulting in myelin destruction and axonal degeneration.7 Furthermore, patients with ADEM often have peripheral nervous system involvement and there have been occasional cases of ADEM associated with Guillain-Barré syndrome. Our patient did not have any clinical features suggestive of peripheral nervous system involvement. However, nerve conduction studies were not performed and a degree of sub-clinical neuropathy cannot therefore be excluded.

We describe the first identifiable case of ADEM temporally associated with Campylobacter gastroenteritis alone. Our patient made an excellent recovery associated with therapy with high dose methylprednisolone.

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