MOG antibody–positive, benign, unilateral, cerebral cortical encephalitis with epilepsy

ABSTRACT

Objective: To describe the features of adult patients with benign, unilateral cerebral cortical encephalitis positive for the myelin oligodendrocyte glycoprotein (MOG) antibody.

Methods: In this retrospective, cross-sectional study, after we encountered an index case of MOG antibody–positive unilateral cortical encephalitis with epileptic seizure, we tested for MOG antibody using our in-house, cell-based assay in a cohort of 24 consecutive adult patients with steroid-responsive encephalitis of unknown etiology seen at Tohoku University Hospital (2008–2014). We then analyzed the findings in MOG antibody–positive cases.

Results: Three more patients, as well as the index case, were MOG antibody–positive, and all were adult men (median age 37 years, range 23–39 years). The main symptom was generalized epileptic seizure with or without abnormal behavior or consciousness disturbance. Two patients also developed unilateral benign optic neuritis (before or after seizure). In all patients, brain MRI demonstrated unilateral cerebral cortical fluid-attenuated inversion recovery hyperintense lesions, which were swollen and corresponded to hyperperfusion on SPECT. CSF studies showed moderate mononuclear pleocytosis with some polymorphonuclear cells and mildly elevated total protein levels, but myelin basic protein was not elevated. A screening of encephalitis-associated autoantibodies, including aquaporin-4, glutamate receptor, and voltage-gated potassium channel antibodies, was negative. All patients received antiepilepsy drugs and fully recovered after high-dose methylprednisolone, and the unilateral cortical MRI lesions subsequently disappeared. No patient experienced relapse.

Conclusions: These MOG antibody–positive cases represent unique benign unilateral cortical encephalitis with epileptic seizure. The pathology may be autoimmune, although the findings differ from MOG antibody–associated demyelination and Rasmussen and other known immune-mediated encephalitides.

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GLOSSARY

ADEM = acute disseminated encephalomyelitis; AMPA = α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid receptor; AQP4 = aquaporin-4; CASPR2 = contactin-associated protein 2; CBA = cell-based assay; CBC = complete blood count; CFF = critical flicker frequency; DWI = diffusion-weighted imaging; EAE = experimental autoimmune encephalomyelitis; FLAIR = fluid-attenuated inversion recovery; GABAB = γ-aminobutyric acid receptor type B receptor; GAD = glutamic acid decarboxylase; GdT1WI = gadolinium enhancement on T1-weighted imaging; HIMP = high-dose IV methylprednisolone; HSV = herpes simplex virus; IgG = immunoglobulin G; IL-6 = interleukin-6; LETM = longitudinally extensive transverse myelitis; LGI1 = leucine-rich glioma-inactivated protein 1; MBP = myelin basic protein; MOG = myelin oligodendrocyte glycoprotein; MS = multiple sclerosis; NMDAR = NMDA receptor; NMOSD = neuromyelitis optica spectrum disorder; ON = optic neuritis; PSL = prednisolone; RE = Rasmussen encephalitis; T2WI = T2-weighted imaging; Tg = thyroglobulin; TPO = thyroid peroxidase; VA = visual acuity; VEP = visual evoked potential.

Myelin oligodendrocyte glycoprotein (MOG) is a myelin protein expressed at the outermost lamellae of the myelin sheath in the CNS.1–5 MOG is also used as an immunogen for experimental autoimmune encephalomyelitis (EAE).2–5 EAE studies have suggested that MOG
antibodies play a direct pathogenetic role in the animal model of inflammatory demyelinating disease, although previous studies designed to detect MOG antibody with the ELISA or Western blotting in human inflammatory demyelinating diseases have failed to reveal any characteristic findings in patients. However, recent studies have demonstrated that conformation-sensitive MOG antibody can be detected by cell-based assays (CBAs) in patients without multiple sclerosis (MS), such as those with pediatric acute disseminated encephalomyelitis (ADEM), aquaporin-4 (AQP4)–immunoglobulin G (IgG)–negative neuromyelitis optica spectrum disorders (NMOSD), optic neuritis (ON), and longitudinally extensive transverse myelitis (LETM). These findings suggest that the MOG antibody may serve as a biomarker to define a spectrum of inflammatory demyelinating diseases, and extensive studies of MOG antibody–positive cases may identify new clinical phenotypes directly or indirectly associated with this myelin antibody.

In the present study, we encountered an index case of MOG antibody–positive

| Table 1: Clinical features of 4 patients with unilateral cortical encephalitis positive for the MOG antibody |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Patient 1 (index case)** | **Patient 2** | **Patient 3** | **Patient 4** |
| **Sex/age at onset, y** | M/39 | M/36 | M/23 | M/38 |
| **Clinical manifestations** | | | | |
| **Encephalopathy** | Generalized tonic seizure with Todd palsy | Generalized tonic seizure | Involuntary movement of the left hand, generalized tonic seizure initially involved the right hand | Generalized tonic seizure initially involved the right hand |
| **Disturbance of consciousness** | Delirium | Only in seizure | Disorientation | Only in seizure |
| **Abnormal behavior and psychiatric symptoms** | Paranoia, hallucination, anorexia | — | — | Agitation, violent behavior |
| **Other focal brain symptoms** | — | — | — | Aphasias and right hemiparesis |
| **Optic neuritis** | + (R, 7 months before seizure) | + (R, after seizure) | — | — |
| **Myelitis** | — | — | — | — |
| **Brain MRI** | FLAIR hyperintensity | R frontoparietal cortex | R frontoparietal cortex | R parietal cortex | L hemisphere cortex |
| **CSF at acute phase** | | | | |
| **Cell counts/μL (mono:poly)** | 29 (23:6) | 63 (62:1) | 101 (51:50) | 311 (129:182) |
| **Protein, mg/dL** | 35 | 38 | 86 | 53 |
| **Oligoclonal IgG bands** | No data | — | — | — |
| **MBP, pg/mL** | No data | <31.3 | <31.3 | <31.3 |
| **MOG antibody titer** | | | | |
| **Serum (onset)** | 1:512 | 1:2,048 | 1:256 | 1:1,024 |
| **Serum (remission)** | 1:16 | 1:128 | — | — |
| **CSF** | 1:32 | 1:4 | 1:16 | No sample |
| **Treatment** | HIMP, PSL, CBZ, LTG | HIMP, PSL, CBZ | DEX, PSL, CBZ | HIMP, PSL, CBZ |
| **Response to treatments** | Full recovery after HIMP | Full recovery after HIMP | Full recovery after DEX | Full recovery after HIMP |
| **Relapse** | No | No | No | No |
| **Duration of follow-up, mo** | 30 | 40 | 23 | 72 |

Abbreviations: ADEM = acute disseminated encephalomyelitis; CBZ = carbamazepine; DEX = dexamethasone; FLAIR = fluid-attenuated inversion recovery; HIMP = high-dose IV methylprednisolone; IgG = immunoglobulin G; LTG = lamotrigine; MBP = myelin basic protein; MOG = myelin oligodendrocyte glycoprotein; PSL = prednisolone.
benign unilateral cerebral cortical encephalitis manifesting with generalized epileptic seizure and then investigated the presence of MOG antibody in an adult cohort of patients with steroid-responsive encephalitis of unknown etiology to identify any unique features of encephalitis in MOG antibody–positive cases.

METHODS Patients, sera, and CSF. We encountered an adult patient (index case, case 1) with unique benign unilateral cerebral cortical encephalitis manifesting with generalized epileptic seizure and seropositivity for MOG antibody in 2014. To explore any other cases with similar features, we identified 24 consecutive patients diagnosed with steroid-responsive encephalitis of unknown etiology seen at Tohoku University Hospital from 2008 to 2014. The patients were older than 20 years and were followed for more than 19 months. We defined steroid-responsive encephalitis of unknown etiology as cases with encephalopathy (epileptic seizure, abnormal behavior, disturbance of consciousness, or focal brain symptoms) that responded to corticosteroid therapy and could not be explained by fever, systemic illnesses, or postictal symptoms. Additional criteria included abnormal brain MRI and CSF findings during the acute phase that were compatible with encephalitis and not indicative of alternative CNS diseases. Sera and CSF were collected during the acute phases and were stored at −80°C. In some cases, sera obtained during remission phases were also stored.

Assays for autoantibodies. We conducted live CBA for MOG antibody based on our previous reports with modification (we used anti-human IgG1 as the secondary antibody to avoid nonspecific binding8,10). Briefly, full-length MOG-expressing or MOG-nonexpressing stable cell lines were incubated with a 1:16 dilution of serum and then incubated with a 1:400 dilution of Alexa Fluor 488 mouse anti-human IgG1 antibody (A10631; Thermo Fisher Scientific, Rockford, IL). After cell immunostaining, 2 investigators (R.O. and T.T.), who were blinded to patients’ data, judged MOG antibody positivity by comparing the staining results of MOG-expressing and MOG-nonexpressing cells. In MOG antibody–positive samples, the antibody titers were calculated by consecutive twofold dilutions to ascertain the maximum dilution with positive staining. Simultaneously, M23-AQP4 antibody in the serum was tested by live CBA using Alexa Fluor 488 goat anti-human IgG (A11008, Thermo Fisher Scientific) as the secondary antibody. Anti-NMDA receptor (NMDAR) antibody, anti-α-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid receptor (AMPA) antibody, anti-leucine-rich glioma-inactivated protein 1 (LG1) antibody, anti-contactin-associated protein 2 (CASPR2) antibody, and anti-γ-aminobutyric acid receptor type B receptor (GABA_B) antibody in the CSF were tested by indirect immunofluorescence using commercially available kits (Euroimmun, Lübeck, Germany).

Standard protocol approvals, registrations, and patient consents. This study was approved by the institutional ethics committee, and all patients provided written informed consent.

RESULTS In addition to the index case, 3 other patients were found to be seropositive for MOG antibody. The clinical characteristics of these 4 patients are summarized in table 1. All of these patients were male, and the median onset age was 34 years (range 23–39). All of the patients experienced epileptic seizures, 3 showed abnormal behavior, 2 patients had ON (figure 1), and 1 patient had dysuria, but no patient had myelopathy. In CSF examinations, the median cell count was 83/μL (range 29–311), and the median protein concentration was 46 mg/dL (range 35–86). Myelin basic protein (MBP) in CSF was not elevated in any of the 3 patients (cases 1, 2, and 3) whose CSF samples were available, although
they were positive for MOG antibody in the CSF. On brain MRI examination, all 4 cases showed unilateral hemispheric cortical hyperintense lesions in fluid-attenuated recovery (FLAIR) imaging (figure 2A and figure 3, A–C, F–H, J–R, and T). None of the MOG antibody–negative patients in our cohort of encephalitis patients showed such FLAIR-hyperintense cortical lesions. The 4 MOG antibody–positive patients were treated with high-dose IV corticosteroids and antiepilepsy drugs, and they fully recovered. We also screened for other encephalitis-related autoantibodies, including AQP4, NMDAR, AMPA, LGI1, CASPR2, and GABA_B antibodies, but negative results were obtained for all of the cases.

Case presentation. Case 1 (index case). A 38-year-old man developed right eye pain and visual loss. His visual acuity (VA) was 30/200, and the critical flicker frequency (CFF) was 16.4 Hz (normal >35) in his right eye. He presented with a right relative afferent pupillary defect and color vision defect in the right eye. Fundus examination revealed optic disc swelling in the right eye. Regarding visual evoked potentials (VEP), the amplitude of P100 was reduced with prolonged latency in the right eye (120.6 ms). This patient was diagnosed with idiopathic ON (figure 1, A and B) and treated with high-dose IV methylprednisolone (HIMP) (1,000 mg/d for 3 days) followed by an oral prednisolone (PSL) taper. His visual symptoms greatly improved soon after HIMP. Seven months later, the patient acutely developed loss of consciousness and generalized tonic seizure and was admitted to our hospital. On admission, he was alert, but his left hand was weak due to Todd palsy. Complete blood cell counts (CBC) and biochemistry were normal. A CSF study showed mild pleocytosis and elevated interleukin-6 (IL-6; 72.6 pg/mL, normal <4.0 pg/mL). Glutamic acid decarboxylase (GAD) antibody, thyroid peroxidase (TPO) antibody, and thyroglobulin (Tg) antibody results were negative in the serum, but the MOG antibody test was positive in the serum (1:512) and in the CSF (1:32) (table 1). EEG showed rhythmic slow waves in the right cerebral hemisphere but no epileptic discharge in the interictal stage. Brain MRI scanned on the day of epilepsy onset showed FLAIR hyperintensity in the right hemispheric cortical region, and the cortical layer was mildly swollen (figure 3, A–C) but did not show hyperperfusion on brain single photon emission computed tomography (SPECT). However, the FLAIR hyperintensities in the cortical regions disappeared after more than 2 years (E, J, O, T).
not show gadolinium enhancement on T1-weighted imaging (GdT1WI). Slight hyperintensity in the regions of diffusion-weighted imaging (DWI) and T2-weighted imaging (T2WI) were seen but were less evident than in the FLAIR image (figure 2, A–F). Brain SPECT showed hyperperfusion in the region (figure 3D). Whole-body PET-CT scans showed no malignancy or inflammation. We started carbamazepine (400 mg/d) and lamotrigine (25 mg/d), but 1 week later, the patient developed delirium, paranoia, hallucination, and anorexia. His symptoms worsened despite risperidone administration. Subsequently, we made a presumptive diagnosis of autoimmune encephalitis to start HIMP therapy, and his symptoms disappeared within a few days. Eight weeks after admission, he was asymptomatic and therefore discharged. He continued oral prednisolone (15 mg/d, then gradually tapered to 4 mg/d in 18 months), carbamazepine, and lamotrigine and experienced no relapse thereafter. At 26 months after discharge, his serum MOG antibody titer had decreased substantially (1:16). Brain MRI showed no residual lesions 30 months after discharge (figure 3E).

**Case 2.** A 36-year-old man let out a strange noise and lost consciousness for several minutes, resulting in a one-car accident when he was driving a car. He was admitted to a local hospital that day and was treated with carbamazepine (400 mg/d). Brain MRI taken on admission showed a FLAIR-hyperintense area in the right parietal cortex. After admission, he twice developed a generalized tonic seizure, right eye pain with visual loss, and dysuria. He was then transferred to our hospital. Neurologic examination showed impaired right VA and dysuria but no signs of meningeal irritation. His VA was normal, but visual field testing showed central scotoma. The CFF was 25 Hz, and VEP revealed prolonged P100 latency (128.4 ms). CBC and blood biochemistry results were normal, while the MOG antibody test was positive in the serum (1:2,048) and in the CSF (1:4) (table 1). The GAD antibody, TPO antibody, and Tg antibody tests were negative in the serum. The EEG examination revealed rhythmic slow waves in the right hemisphere, especially in the right parietal region, but no epileptic discharge was seen in the interictal state. Brain MRI scanned 1 month after the onset of epilepsy showed FLAIR hyperintensity in the right hemispheric cortical region (figure 3, K–N). Abnormalities in the region in DWI, T2WI, and GdT1WI were equivocal. Whole-body CT showed no malignancy or inflammation. VEP was not examined. Tests for cytomegalovirus antigen in the blood and *Mycobacterium tuberculosis* (QuantiFERON) were negative, and PCR for herpes simplex virus (HSV), gram stains, and culture results were negative in the CSF. However, because we could not rule out CNS infectious disease, we initially treated the patient with IV ceftriaxone, isoniazid, ethambutol, acyclovir, fluconazole, and dexamethasone (33 mg/d). His symptoms disappeared soon after the treatment, and we suspected autoimmune encephalitis rather than CNS infection. Four weeks after admission, he was discharged with no symptoms, but oral prednisolone (15 mg/d, gradually tapered off in 1 year) and carbamazepine (600 mg/d) were continued. Eighteen months later, he had not experienced a relapse. At 40 months after discharge, the MOG antibody titer was reduced (1:128) and brain MRI showed no residual lesions (figure 3J).

**Case 3.** A 23-year-old man with involuntary movement of the left hand was diagnosed with epilepsy and was treated with carbamazepine (400 mg/d) with no apparent effect. One month later, he developed a generalized tonic seizure that lasted for 1 hour. The following month, he was admitted to our hospital for a severe headache. But he did not complain of visual impairment. Upon neurologic examination, he was disoriented without neck stiffness. Although the CBC and blood biochemistry results were normal, the MOG antibody test was positive in the serum (1:256) and in the CSF (1:16) (table 1). The GAD antibody, TPO antibody, and Tg antibody tests were negative in the serum. The EEG examination revealed rhythmic slow waves in the right hemisphere, especially in the right parietal region, but no epileptic discharge was seen in the interictal state. Brain MRI scanned 1 month after the onset of epilepsy showed FLAIR hyperintensity in the right hemispheric cortical region (figure 3, K–N). Abnormalities in the region in DWI, T2WI, and GdT1WI were equivocal. Whole-body CT showed no malignancy or inflammation. VEP was not examined. Tests for cytomegalovirus antigen in the blood and *Mycobacterium tuberculosis* (QuantiFERON) were negative, and PCR for herpes simplex virus (HSV), gram stains, and culture results were negative in the CSF. However, because we could not rule out CNS infectious disease, we initially treated the patient with IV ceftriaxone, isoniazid, ethambutol, acyclovir, fluconazole, and dexamethasone (33 mg/d). His symptoms disappeared soon after the treatment, and we suspected autoimmune encephalitis rather than CNS infection. Four weeks after admission, he was discharged with no symptoms, but oral prednisolone (15 mg/d, gradually tapered off in 1 year) and carbamazepine (400 mg/d) was continued, and he did not experience any relapse. At 40 months after discharge, the MOG antibody titer was reduced (1:128) and brain MRI showed no residual lesions (figure 3J).
Table 2  Differential diagnosis of autoimmune or immune-mediated encephalopathy

<table>
<thead>
<tr>
<th>Age, y</th>
<th>Sex</th>
<th>Clinical manifestations</th>
<th>Autoantibodies</th>
<th>Brain MRI findings</th>
<th>Treatments</th>
<th>Prognosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present cases</td>
<td>23–39</td>
<td>All male</td>
<td>Seizure in all, abnormal behavior, psychiatric symptoms or other focal brain symptoms in some cases, no cancer</td>
<td>MOG antibody</td>
<td>FLAIR-hyperintense unilateral cerebral cortical lesions, no atrophy</td>
<td>HIMP, AED</td>
</tr>
<tr>
<td>ADEM(^{16})</td>
<td>Children and young adults (&lt;40 y)</td>
<td>Male 60%</td>
<td>Altered consciousness and behavior, weakness, ataxia, cranial nerve palsy, seizure, fever, headache, vomiting, and meningeal signs</td>
<td>MOG antibody 50%</td>
<td>T2/FLAIR multiple, large lesions (&gt;2 cm) in the white matter, basal ganglia, brainstem, cerebellum, and spinal cord</td>
<td>HIMP, IVlg, PLEX</td>
</tr>
<tr>
<td>Rasmussen encephalitis(^{17})</td>
<td>Infancy to adulthood (mean 6 y)</td>
<td>No predominance</td>
<td>Hemiparesis, seizure, and unilateral movement disorder (athetosis/dystonia)</td>
<td>GluR antibody in some cases</td>
<td>Unilateral hemispheric focal cortical lesions with atrophy</td>
<td>AED, HIMP, IVlg, PLEX, surgery</td>
</tr>
<tr>
<td>NMDAR encephalitis(^{18,19})</td>
<td>8 months–85 years (mean 21 y)</td>
<td>Female dominant</td>
<td>Psychiatric disorder, movement disorder, autonomic instability, and central hypoventilation, ovarian teratoma usually found in young women</td>
<td>NMDAR GluN1 subunit antibody</td>
<td>Abnormal, mainly medial temporal lobe lesions in half of cases</td>
<td>HIMP, IVlg, PLEX, rituximab</td>
</tr>
<tr>
<td>VGKC-Ab-associated encephalopathy(^{16,20})</td>
<td>47–77 years (median 63)</td>
<td>Male dominant</td>
<td>Amnesia, seizure, psychiatric disturbance, autonomic dysfunction, and neuromyotonia</td>
<td>LGI1, CASPR2 antibodies</td>
<td>T2/FLAIR hyperintensity in the medial temporal lobe, hippocampus, or amygdala in half of cases</td>
<td>HIMP, PLEX, IVlg</td>
</tr>
<tr>
<td>GAD-Ab-associated limbic encephalitis(^{21})</td>
<td>17–66 years (median 23)</td>
<td>Female dominant</td>
<td>Epilepsy, mild cognitive disorder</td>
<td>GAD antibody (&gt;1,000 U/mL)</td>
<td>High signal on T2 in the medial temporal lobe or hippocampus</td>
<td>HIMP, PLEX</td>
</tr>
<tr>
<td>Other autoimmune limbic encephalitis(^{18})</td>
<td>Hu antibody: 50-63 years; Ma2 antibody: 22-70 years; AMPA receptor antibody: 23-81 years; GABA(_{A}) receptor antibody: 16-77 years</td>
<td>Male dominant: Hu, Ma2, GABA(_{A}), AMPA receptor antibodies, female dominant: AMPA receptor antibody</td>
<td>Confusion, working memory deficit, mood change, seizure associated with tumors (ex: small-cell lung carcinoma, testicular seminoma, thymoma)</td>
<td>Hu, Ma2, AMPA receptor, GABA(_{A}) receptor antibodies</td>
<td>FLAIR hyperintensities in the medial temporal lobe</td>
<td>HIMP, IVlg, PLEX, tumor treatment</td>
</tr>
<tr>
<td>Hashimoto encephalopathy(^{22})</td>
<td>23–83 years (mean 60 years)</td>
<td>Female dominant</td>
<td>Seizure, myoclonus, hallucination, or stroke-episodes</td>
<td>Thyroid peroxidase, thyroglobulin antibodies, α-enolase antibody</td>
<td>Normal or nonspecific abnormalities</td>
<td>HIMP</td>
</tr>
<tr>
<td>NMOSD(^{10,24,25})</td>
<td>AQ4 antibody-positive: around 40 years on average, 1%-2% in children, MOG antibody-positive 3-70 years (median: 37.5 years)</td>
<td>AQ4 antibody-positive: female dominant (60%-90%), MOG antibody-positive: no female preponderance</td>
<td>Optic neuritis, acute myelitis, hiccups or noise postera and vomiting, acute brainstem syndrome, symptomatic narcolepsy or acute diencephalic clinical syndrome, symptomatic cerebral syndrome</td>
<td>AQ4 antibody, MOG antibody</td>
<td>AQ4 antibody-positive: dorsal medulla/area postrema, peri-ependymal brainstem, subcortical or deep white matter, corpus callosum, long cortico-spinal tract lesions, MOG antibody positive: ADEM, brainstem lesions</td>
<td>HIMP, PLEX, IVlg, cyclophosphamide in acute attacks, immunosuppressants for relapse prevention</td>
</tr>
</tbody>
</table>

Abbreviations: ADEM = acute disseminated encephalomyelitis; AED = antiepileptic drug; AQ4 = aquaporin-4; DWI = diffusion-weighted imaging; FLAIR = fluid-attenuated inversion recovery; GAD = glutamic acid decarboxylase; HIMP = high-dose IV methylprednisolone; IVlg = IV immunoglobulin; MOG = myelin oligodendrocyte glycoprotein; NMDAR = NMDA receptor; NMOSD = neuromyelitis optica spectrum disorders; PLEX = plasma exchange; VGKC = voltage-gated potassium channel.

a diagnostic workup, he was discharged 1 week later (carbamazepine was continued). However, 35 months after the first admission, he developed a generalized tonic seizure, which initially involved the right hand. He also had aphasia and right hemiparesis and was readmitted to our hospital. Neurologic examination revealed delirium, emotional incontinence, aphasia, and mild right hemiparesis. He did not show ocular symptoms during the course of disease. The CBC and blood biochemistry were normal, but the MOG antibody test was positive in the serum (1:1,024) (table 1). Serum GAD antibody, TPO antibody, and Tg antibody tests were negative. Brain MRI scanned 4 days after the second episode of epilepsy showed FLAIR hyperintensity in the left hemispheric cortical region (figure 3, P–R). Brain SPECT demonstrated hyperperfusion in the region (figure 3S). Whole-body PET-CT findings showed no malignancy.
or swollen lymph nodes. VEP was not done. PCR for HSV, gram stains, and culture results were negative in the CSF. After admission, his symptoms became worse (agitation and violent behavior) despite the administration of sedatives. We suspected autoimmune encephalitis and started HIMP, after which time he became asymptomatic and was discharged. He continued carbamazepine (300 mg/d) and experienced no relapse. At 84 months after discharge, he was MOG antibody-negative. A brain MRI taken 72 months after discharge was normal (figure 3T).

**DISCUSSION** In the index case (case 1), we tested for MOG antibody because the patient had unilateral benign ON rather than unilateral cortical encephalitis with epileptic seizure. Then, we tested for MOG antibody in our cohort of 24 consecutive adult cases of corticosteroid-responsive encephalitis of unknown etiology and identified 3 additional patients with MOG antibody positivity. Unexpectedly, these 3 MOG antibody–positive patients also had unilateral cortical encephalitis with epileptic seizure as seen in the index case, and there were no cases of unilateral cortical encephalitis with epileptic seizure without MOG antibody positivity in our cohort. The unilateral cortical lesions best depicted by FLAIR images were unique and appeared distinct from brain lesions previously described in MOG antibody–positive diseases including ADEM.13

The unilateral cortical lesions in our cases 1–4 needed to be differentiated from seizure-induced brain MRI abnormalities.14 Such brain MRI abnormalities induced by epileptic seizure are localized in the cortical/subcortical regions, hippocampus, basal ganglia, white matter, or corpus callosum, and they are readily visible on DWI due to cytotoxic changes.15,16 However, the MRI findings in our cases were much more clearly seen in FLAIR images than in DWI and ADC findings (figure 1, A–F). Moreover, pleocytosis in the CSF and a favorable response to HIMP suggested that the unique unilateral cortical lesions were inflammatory, and hyperperfusion on SPECT corresponding to the cortical FLAIR hyperintensity supported the inflammatory nature and epileptogenicity of the swollen cortical lesions in the acute phase.

We also ruled out a variety of autoimmune-mediated or immune-mediated encephalitides (table 2) before we concluded that the unilateral cortical encephalitis with epileptic seizure in our cases was unique. Rasmussen encephalitis (RE) is described as unilateral cerebral cortical encephalitis, similar to that observed in our patients. However, RE is clinically characterized by focal epilepsy, progressive hemiplegia, and cognitive decline with unilateral hemispheric focal cortical atrophy in the chronic stage, and corticosteroid and other anti-inflammatory therapies are only partially effective.17 Our cases did not share these features of RE or fulfill the diagnostic criteria. The lesion distribution in our 4 patients was also dissimilar to the brain MRI abnormalities in cases of encephalitis with seizure associated with NMDAR antibody, VGKC antibody, GAD antibody, and antithyroid antibodies,18–22 and our patients were negative for those autoantibodies. Likewise, the clinical and neuroimaging features of our cases were distinct from limbic encephalitides with positivity for GAD, LGI1, GABAB, or AMPA antibodies18 and from the brain syndrome previously described in NMOSD.10,23,24

FLAIR-hyperintense lesions localized at the cerebral cortex or sulcus, similar to the findings observed in the present cases, can develop in various CNS diseases including meningitis, subarachnoid hemorrhage, leptomeningeal metastasis, acute infarction, and moyamoya disease.25 In a review of such MRI abnormalities, the left tempo-ro-occipital cortical FLAIR-hyperintense lesions in a 23-year-old man with the diagnosis of meningitis appeared to be similar to the brain MRI findings in our cases. More recently, Numa et al.26 reported a case of a 37-year-old woman who was diagnosed with ADEM when she was 4 years old and developed ON followed by recurrent ADEM 33 years later. She was MOG antibody–positive, and brain MRI showed unique cortical FLAIR-hyperintense lesions in the left temporal and frontal lobes. Thus, unilateral cortical encephalitis in MOG antibody–positive patients, as in the 4 cases in our study, may have been previously unnoted as a distinct phenotype. The relationship between MOG antibody and the unilateral cerebral cortical encephalitis observed in our cases remains unclear. Two of our patients had benign unilateral ON, in which MOG antibody is often detected, while cases 3 and 4 lacked such characteristics of CNS diseases such as ON, LETM, NMOSD, or ADEM.27–29 Thus, unilateral cerebral cortical encephalitis may be another characteristic manifestation of MOG antibody–positive patients. Although some cases of MOG antibody–associated diseases fulfill the diagnostic criteria of seronegative NMOSD,23 the spectrum of MOG antibody–associated diseases is obviously wider than NMOSD. In the near future, MOG antibody–associated diseases may be recognized as a distinct clinical entity of inflammatory demyelinating diseases of the CNS.30

There is some evidence to support the pathogenic potential of MOG antibody. Experimental studies have shown that MOG antibody can affect oligodendrocytes and myelins.31,32 In addition, in a few brain-biopsied cases of tumefactive brain lesions with MOG antibody positivity, pathologic examinations revealed active inflammatory demyelination with deposition of immunoglobulins and complement33,34 or MS type II pathology.35,36 Moreover, we recently reported high...
positive encephalitis. These findings suggest that MOG antibody may directly contribute to inflammatory demyelination in anti–myelin antibody–associated CNS diseases. However, in 3 of our MOG antibody–positive cases whose CSF-MBP levels were measured during the acute phase, there was no elevation in CSF-MBP despite the extensive cortical involvement and CSF pleocytosis. Thus, it is also possible that MOG antibody itself may not be directly associated with the unilateral cerebral cortical encephalitis with epileptic seizure in our patients and that another autoimmune disorder coexisting with MOG antibody positivity might be responsible for the encephalitis. In fact, MOG antibody can be detected in some patients with other autoantibody–associated encephalitides such as NMDAR antibody–positive encephalitis. In addition, a pathogenic autoimmune disorder coexisting with MOG antibody positivity might be responsible for the encephalitis. In fact, MOG antibody can be detected in some patients with other autoantibody–associated encephalitides such as NMDAR antibody–positive encephalitis. Another autoantibody that coexists with MOG antibody may possibly develop later in the disease course of cases 3 and 4. Therefore, an unknown autoantibody might be associated with the unilateral cerebral cortical encephalitis with epileptic seizure in a fraction of MOG antibody–positive cases although we need to perform immunohistochemistry or immunofluorescence with rodent brain tissue slices and the sera and CSF from the patients as an attempt to see whether there are any antibody reactivities to the cerebral cortical tissues.

Our study is retrospective and has some limitations. Because our patient cohort was small and derived from a single university hospital, the results should be verified in prospective, larger-scale, multicenter studies. In addition, we analyzed only adult patients in the present study, and it is important to determine whether MOG antibody–positive unilateral cerebral cortical encephalitis with epileptic seizure also occurs in children. Therefore, at this point, it is premature to discuss the frequency of MOG antibody–positive unilateral cerebral cortical encephalitis in corticosteroid-responsive encephalitis of unknown etiology. However, since we experienced 6 cases of NMDAR antibody–associated encephalitis and 1 with VGKC antibody–associated encephalitis during the same period (2008–2014), unilateral encephalitis with MOG antibody may not be so uncommon.

Taken together, we report a form of benign unilateral cerebral cortical encephalitis with epileptic seizure in 4 adult patients with MOG antibody positivity. The pathogenesis of this condition appears to be immune-mediated or autoantibody-mediated, although the clinical, MRI, and laboratory features differ from those in previously described MOG antibody–associated CNS diseases and known autoantibody-mediated encephalitides. Another autoantibody that coexists with MOG antibody may be responsible for this type of encephalitis.

**AUTHOR CONTRIBUTIONS**

R.O. analyzed the data and wrote the paper, substantial contribution to the study conception, acquisition, analysis, and interpretation of data for the work, writing the manuscript, drafting and correction of all versions of the manuscript including figures, tables, and references, completion of the work to be submitted, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. L.N. substantial contribution to the conception and design of the work, as well as supervision of the acquisition, analysis, and interpretation of data for the work, revised several versions of the manuscript critically for important intellectual content, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. T.T. substantial contribution to the conception and design of the work, as well as supervision of the acquisition, analysis, and interpretation of data for the work, revised several versions of the manuscript critically for important intellectual content, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. K.K. contribution to the plan of the work, acquisition, analysis, interpretation of data for the work, and drafting the original manuscript related to the case, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. T.A. acquisition, analysis, and interpretation of data for the work, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. Y.T. acquisition, analysis, and interpretation of data for the work, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. D.K.S. acquisition, analysis, and interpretation of data for the work, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. H.K. substantial contribution to the conception and design of the work, as well as supervision of the acquisition, analysis, and interpretation of data for the work, revised several versions of manuscript critically for important intellectual content, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. M.A. substantial contribution to the conception and design of the work, as well as supervision of the acquisition, analysis, and interpretation of data for the work, provided final approval of the version to be published, agreed to be accountable for all aspects of the work. Y.H. substantial contribution to the conception and design of the work, as well as supervision of the acquisition, analysis, and interpretation of data for the work, revised several versions of the manuscript critically for important intellectual content, final responsibility and approval of the version to be published, agreed to be accountable for all aspects of the work.

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REFERENCES


MOG antibody–positive, benign, unilateral, cerebral cortical encephalitis with epilepsy

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