GRP78 Antibodies Are Associated With Blood-Brain Barrier Breakdown in Anti–Myelin Oligodendrocyte Glycoprotein Antibody–Associated Disorder

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Abstract

Background and Objectives
To analyze (1) the effect of immunoglobulin G (IgG) from patients with anti–myelin oligodendrocyte glycoprotein antibody (MOG-Ab)–associated disorder on the blood-brain barrier (BBB) endothelial cells and (2) the positivity of glucose-regulated protein 78 (GRP78) antibodies in MOG-Ab–associated disorders.

Methods
IgG was purified from sera with patients with MOG-Ab–associated disorder in the acute phase (acute MOG, n = 15), in the stable stage (stable MOG, n = 14), healthy controls (HCs, n = 9), and disease controls (DCs, n = 27). Human brain microvascular endothelial cells (BMECs) were incubated with IgG, and the number of nuclear NF-κB p65-positive cells in BMECs using high-content imaging system and the quantitative messenger RNA change in gene expression over the whole transcriptome using RNA-seq were analyzed. GRP78 antibodies from patient IgGs were detected by Western blotting.

Results
IgG in the acute MOG group significantly induced the nuclear translocation of NF-κB and increased the vascular cell adhesion molecule 1/intercellular adhesion molecule 1 expression/permeability of 10-kDa dextran compared with that from the stable MOG and HC/DC groups. RNA-seq and pathway analysis revealed that NF-κB signaling and oxidative stress (NQO1) play key roles. The NQO1 and Nrf2 protein amounts were significantly decreased after exposure to IgG in the acute MOG group. The rate of GRP78 antibody positivity in the acute MOG group (10/15, 67% [95% confidence interval, 38%–88%]) was significantly higher than that in the stable MOG group (5/14, 36% [13%–65%]), multiple sclerosis group (4/29, 14% [4%–32%]), the DCs (3/27, 11% [2%–29%]), or HCs (0/9, 0%). Removal of GRP78 antibodies from MOG-IgG reduced the effect on NF-κB nuclear translocation and increased permeability.

Discussion
GRP78 antibodies may be associated with BBB dysfunction in MOG-Ab–associated disorder.
Anti-myelin oligodendrocyte glycoprotein antibody (MOG-Ab)-associated disorder has been recently recognized as a new entity in the spectrum of inflammatory demyelinating diseases, differing from either multiple sclerosis (MS) or neuromyelitis optica spectrum disorder (NMOSD).1,4 Using live cell-based assays (CBAs), MOG antibodies bound to the cell surface are detected in patients with pediatric/adult acute disseminated encephalomyelitis (ADEM), AQP4 antibody–negative NMOSD, isolated optic neuritis, transverse myelitis, encephalomyelitis, or brainstem encephalitis.2 The clinical spectrum of MOG-Ab–associated disorder thus seems to be broader than that of AQP4 antibody–positive NMOSD,3,4 but what factors determine the varied clinical phenotypes of the disease remain unclear.

A previous histopathologic analysis of 9 cases with MOG-Ab–associated disorder described an MS pattern II lesion, characterized by active demyelination with marked infiltration of macrophages and T cells, along with deposition of immunoglobulin G (IgG)/complements5–9 suggesting Ab-mediated demyelination. Intrathecal transfer experiments of MOG-Ab from patients with MOG-Ab–associated disorder to experimental autoimmune encephalomyelitis (EAE) models induced pathologic changes, resembling an MS pattern II pathology, together with myelin-reactive T cells (myelin basic protein–specific T cells) and MOG-specific T cells.10 These findings suggest that MOG-Ab s exert a pathogenic effect when they enter the CNS space in cases of MOG-Ab–associated disorder.

Blood-brain barrier (BBB) breakdown is a key pathologic feature of MS and neuromyelitis optica (NMO).11–14 We previously reported the effect of glucose-regulated protein 78 (GRP 78) autoantibodies from patients with MS on human brain microvascular endothelial cell (BMEC) dysfunction.15,16 Both a history of preceding infectious prodrome in 50% of patients with MOG-Ab–associated disorder17 and the absence of oligoclonal IgG bands in the CSF18 suggest that anti-MOG antibodies are produced peripherally and reach the CNS following BBB breakdown, but the molecular mechanism regarding BBB disruption in MOG-Ab–associated disorder remains unclear.

In the present study, we analyzed the contribution of serum IgG from individual patients with MOG-Ab–associated disorder to BBB breakdown using human BMECs with the high-content imaging and RNA-seq. We also investigated the GRP78 antibody positivity of patients and the effect of this antibody on the BMEC activation in MOG-Ab–associated disorder.

**Methods**

**Patient Samples**

This research was approved by the ethics committees of the Medical Faculties of Yamaguchi Universities (IRB#: H22-137-6). We obtained written informed consent from each participant.

Sera were collected from 15 patients with MOG-Ab–associated disorder who had been diagnosed at Tohoku University or Yamaguchi University Hospital (Table). All the patients were positive for MOG-Abs according to a cell-based assay performed at Tohoku University. We included sera from MOG-Ab–associated disorder patients (15 sera in the acute phase [acute MOG, n = 15] and 14 sera in the stable stage [stable MOG, n = 14]) (Table). The 15 serum samples from acute MOG were collected during the acute phase within 14 days at the time of the symptom onset, and the 14 sera from stable MOG were collected in the clinical stable phase from patients, who had been treated with corticosteroids over 3 months (3–57 months) after the acute phase. IgGs from 9 healthy controls (HCs: men, n = 4; women, n = 5; mean age, 32.6 years) and 27 disease controls (DCs) were used as controls (amyotrophic lateral sclerosis, n = 17; cervical spondylosis, n = 2; multiple system atrophy, n = 3; spinocerebellar degeneration, n = 1; hereditary peripheral neuropathy, n = 2; progressive supranuclear palsy, n = 1; and normal pressure hydrocephalus, n = 1).

All sera were stored at −80°C until the experiments were performed. Sere were inactivated at 56°C for 30 minutes before the experiments. IgG was prepared from serum samples using a Melon Gel IgG Spin Purification Kit (Thermo Fisher Scientific).

**Patient Information**

We investigated the clinical phenotype, the age/sex, the relapse number, ΔEDSS score (change in the score on the Expanded Disability Status Scale [EDSS] between before and after relapse), presence of a preceding infection, IgG index (intrathecal IgG synthesis), Q Albumin (Q Alb, BBB
breakdown), and the presence of Gd-enhanced lesions on MRI in the 15 patients with acute MOG-Ab–associated disorder (Table).

**Immunohistochemistry of NF-κB, ICAM-1, NOQ1, or Nrf2 and the High-Content Imaging Assay**

TY10 cells derived from human adult BMECs and immortalized with temperature-sensitive SV40 large T antigen (tsA58) were used for experiments. Two days after the shift in temperature from 33 to 37°C, all experiments were performed. The cells were maintained on collagen type 1 and the culture dish in MCDB 131 medium containing IgG (500 μg/mL) from patients with MOG-Ab–associated disorder, DCS, or HCs after substitution for MCDB 131 medium for 1 hour for NF-κB p65 immunostaining and for 24 hours for intercellular adhesion molecule 1 (ICAM-1), NAD(P)H quinone dehydrogenase 1 (NQO1), Nrf2, and JC-1 immunostaining.

For NF-κB p65, NQO1, or Nrf2 staining, cells were fixed with 4% paraformaldehyde, incubated with 0.3% Triton X-100, and then blocked overnight in 5% fetal bovine serum/0.3% Triton X-100 in phosphate-buffered saline (PBS). The cells were incubated with each primary Abs (NF-κB p65 monoclonal antibody, NQO1 polyclonal antibody [Proteintech], or Nrf2 polyclonal antibody [Proteintech]), followed by the incubation with secondary Abs (Alexa Fluor 488 goat anti-rabbit IgG, Thermo Fisher Scientific).

For JC-1 staining, a final concentration of 1 μM of JC-1 Mitomycin Detection Kit (DOJINDO, Japan) was added to living TY10 cells in the dark at 37°C for 45 minutes. JC-1 dye shows fluorescence emission at 2 typical wavelengths: red fluorescent J-monomers. The red/green intensity ratio reflecting higher mitochondrial concentrations reflects higher mitochondrial potential and green fluorescent J-aggregates at higher mitochondrial concentrations reflecting higher mitochondrial potential and green fluorescent J-aggregates at high-content imaging. For high-content imaging, 5,000 cells per well were plated onto CELLSTAR 96-well plates (Greiner). After immunostaining for NF-κB p65 was performed, the images in the plate were captured using an In Cell Analyzer 2000 (GE Healthcare) at ×20 magnification with 4 fields of view per well (equivalent to almost 800 cells). The images were then analyzed with the IN Carta image analysis software (Cytiva) or the In Cell Analyzer software program (Cytiva). The data represent the mean value of 12 experiments for NF-κB p65 or 3 experiments for ICAM-1, NQO1, Nrf2, and JC-1.

**Paracellular Permeability of 10-kDa Dextran**

TY10 cells were cultured on 0.4-mm pore size 24-well collagen-coated Transwell culture inserts (Corning) on the luminal side for 3 days at 33°C and then 2 days at 37°C. TY10

### Table: Clinical Information of MOG Patients

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<th>Preceding infection</th>
<th>IgG index</th>
<th>Q Alb</th>
<th>Gd-MRI</th>
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**Abbreviations:** ADEM = acute disseminated encephalomyelitis; CE = cortical encephalitis; EDSS = Expanded Disability Status Scale; Gd-MRI = presence of Gd-enhanced spinal lesions on MRI; GRP Ab = presence of GRP78 autoantibodies in IgG from patients; M = myelitis; NE = not examined; ON = optic neuritis; preceding infection = presence of preceding infection; Pt Nos = patient numbers.

%NFκB, % of NF-κB p65 nuclear-positive cells; ∆EDSS score, change in the score on the EDSS from before to after relapse.
cells were exposed to individual IgG from patients with MOG-Ab–associated disorder (acute, n = 15; stable, n = 14), DCs (n = 27), or HCs (n = 9) (500 μg/mL) for 24 hours at 37°C. After the cells were washed, FITC-10-kDa dextran fluorescence (Sigma-Aldrich) was added to the luminal insert (concentration, 1 mg/mL). A total 100 μL of medium was then transferred from the abluminal chamber into 96-well black plates over 40 minutes. Fluorescence signals were calculated at 490/520 nm (absorption/emission) by using a FlexStation 3 Multi-Mode microplate reader (Molecular Devices).

Whole Transcriptome Analyses With RNA-Seq
TY10 cells were exposed to IgGs from 4 patients with MOG-Ab–associated disorder in the acute phase or 3 healthy individuals (500 μg/mL) for 12 hours at 37°C. TY10 cells without incubation with IgGs were used as controls.

The method for the whole transcriptome analysis with RNA-seq was previously described. In brief, total RNA was extracted from TY10 cells using the RNeasy Mini Kit (Qiagen), and messenger RNA was purified as described previously. Complementary DNA libraries were produced using a NEBNext Ultra II RNA Library Prep kit (New England Biolabs) and NEBNextplex Oligos for Illumina, as described previously. In this approach, messenger RNA was fragmented in NEBNext First Strand Synthesis Reaction Buffer at 94°C for 15 minutes in the presence of NEBNext Random Primers and was reverse transcribed with NEBNext Strand Synthesis Enzyme Mix. The library fragments were then concentrated, and index sequences were inserted during polymerase chain reaction amplification. The products were purified using AMPure XP beads (Beckman Coulter), and the quality of the library was confirmed with an Agilent 2200 TapeStation (D1000, Agilent Thermo Fisher). The libraries mixed to equal molecular amounts were sequencing on an Illumina NextSeq DNA sequencer with a 75-bp paired-end format to analyze the detected genes (Qiagen). The mapped read counts were normalized to transcripts per million, and the FC increased by >50% or decreased by >50% were calculated at 490/520 nm (absorption/emission) by using a FlexStation 3 Multi-Mode microplate reader (Molecular Devices).

Removal of GRP78 Autoantibodies From MOG-IgG by Immunoprecipitation
The method of GRP78 Ab removal was previously described. In brief, 500 μg/mL of MOG-IgG (patients 1 and 9) was incubated with 5 μg of human embryonic kidney cells 293T cell lysates with or without the overexpression of FLAG-tagged GRP78 (Origene) for 4 hours. After the antigen-antibody immune complexes of GRP78 had precipitated, 40 μL of Red Anti-FLAG M2 Affinity Gel beads (Sigma-Aldrich) was incubated for 2 hours. After centrifuging the sample, the supernatants (MOG-IgG with/without GRP78 antibodies) were used for the analysis. The GRP78 antibodies in both supernatants were detected by a Western blot analysis.

Statistical Analyses
All statistical analyses were performed using the Prism 7 software program (Graph Pad). A paired Student t test (2 sided) was used for single comparison analyses. For multiple comparison analyses, a 1-way analysis of variance was used with the Tukey multiple comparison test when the data were normally distributed or the nonparametric Kruskal-Wallis test when the data were not normally distributed. The Fisher exact probability test was used to calculate the significance of differences in the positivity of GRP78 antibodies among the acute MOG, stable MOG, DC, HC, and MS groups. To assess the association, Pearson correlation coefficients were used. *p < 0.05, **p < 0.01, and ***p < 0.001 were considered to have statistical significance.

Data Availability
Data will be shared on reasonable request. Please contact the corresponding author.

Results

IgGs From the Acute Phase of the Patients With MOG-Ab–Associated Disorder Activated BMECs
We first compared the proportion of the nuclear translocation of NF-κB p65 in BMECs among the acute MOG, stable MOG, DC, and HC groups (Figure 1, A–E). The proportion of NF-κB p65 nuclear-positive cells in the acute MOG group

Western Blotting
Detection of GRP78 Autoantibodies by Western Blotting
The method of Western blotting was previously described. In brief, 2 μg of human full length GRP78 recombinant protein (Abcam) was fractionated and electrophoretically transferred to the polyvinylidene difluoride (PVDF) membranes (Amersham). The GRP78 antibodies (Abcam) and individual IgG (5 μg/mL) from 15 patients with MOG-Ab–associated disorder (acute, n = 15; stable, n = 14), DCs (n = 27), MS (n = 29), HCs (n = 9), and 27 DCs diluted with PBS-T/5% milk were used as the primary antibodies. The PVDF membranes were incubated with each primary antibody for an hour, followed by the anti-human secondary fluorescent antibodies for an hour. For vascular cell adhesion molecule 1 (VCAM-1) and ICAM-1 staining, VCAM-1 antibodies (R&D Systems) and ICAM-1 (Santa Cruz) were used as the primary antibodies, respectively, followed by an anti-mouse secondary antibody for an hour. The bands were visualized using a chemiluminescence kit (Immunostar LD, Japan), and the relative density of each band was calculated with the Quantity One software program (Bio-Rad).

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Data Availability
Data will be shared on reasonable request. Please contact the corresponding author.
was significantly greater than in the DC group (Figure 1E). The proportion of NF-κB p65 nuclear-positive cells was significantly decreased between the acute and stable MOG in the same individual (Figure 1F). The permeability was found to be significantly increased after incubation with IgG from the acute MOG group compared with that from the stable MOG, DC, or HC groups (Figure 1G). In a time course study (6, 12, 18, and 24 hours), the 10-kDa dextran permeability in BMECs was increased after >12 hours of incubation with 2 IgGs from MOG patients (patients 1 and 9) (eFigure 1, links.lww.com/NXI/A602). Furthermore, a Western blot analysis showed that the VCAM-1 expression was significantly increased after exposure to IgG from the acute MOG/DC group compared with that from the HC group (Figure 2, D and E).

**Profiling of Altered Gene Expression in BBB Endothelial Cells After IgG Exposure From the Acute Phase of the Patients With MOG-Ab–Associated Disorder**

We selected IgG from 4 patients with MOG-Ab–associated disorder presenting with myelitis or cortical encephalitis and showing an elevated effect on the induction of NF-κB p65 nuclear translocation of TY10 (patients 1, 3, 5, and 9 in Table) for RNA-seq. A whole transcriptome analysis using RNA-seq in BMECs after exposure to IgG from the patients with MOG-Ab–associated disorder (n = 4) and HCs (n = 3) was performed to identify important signaling pathway. BMECs without exposure to IgGs were used as controls. In TY10 cells, over 32,000 genes were detected from approximately 30 million reads in each sample. Volcano plots (Figure 2A) and heat maps (eFigure 2, links.lww.com/NXI/A603) at the same
p values and FC revealed that 189 genes were significantly differentially expressed (FC > 1.5; p < 0.05), including 83 upregulated genes and 106 downregulated genes, between the patients with MOG-Ab–associated disorder (n = 4) and HC/control groups (n = 4) (eFigure 1, links.lww.com/NXI/A602). Next, Ingenuity Pathway Analysis (an IPA) was performed using the above upregulated or downregulated genes to determine the signaling pathway of BMECs after exposure to IgGs from the acute phase of patients with MOG-Ab–associated disorder. In the network analysis of the upregulated genes, NF-κB was detected in the center of the network analysis, and PLAAT4 and TNFRSF6B were observed as upstream molecules of NF-κB (Figure 3B). In the network analysis of downregulated genes, NQO1 and DNAJB1 were detected in the center of the network analysis, suggesting that oxidative stress had been induced (Figure 3C).

IgGs From the Acute Phase of the Patients With MOG-Ab–Associated Disorder Reduced Protective Proteins Against Oxidative Stress in BBB Endothelial Cells

To confirm the data suggested by RNA-seq, the change in amounts of protective proteins against oxidative stress after exposure to IgGs from the patients with MOG-Ab–associated disorder was observed. We selected 14 IgGs from the acute MOG group, 7 IgGs from the DC group (all samples from patients with amyotrophic lateral sclerosis), and 9 IgGs from the HC group for this purpose. High-content imaging system
revealed that the expression of NQO1 and Nrf2 protein in BMECs and the mitochondria membrane potential (red/green ratio using JC-1 dye) were significantly decreased after exposure to IgGs from the acute phase of the patients with MOG-Ab–associated disorder compared with that from DCs or HCs (Figure 4, A–E). Furthermore, a Western blot analysis showed that the amount of Nrf2 protein of TY10 cells was also decreased after exposure to IgGs from the acute phase of
Figure 4 Changes in the Protective Proteins Against Oxidative Stress After IgG Exposure From Acute MOG Antibody–Associated Disorders

(A, B) Immunostaining of human brain microvascular endothelial cells (TY10 cells) for NQO1 or Nrf2 (green), or JC-1 (green/red) after exposure to IgG (500 μg/mL) from patients with MOG-Ab–associated disorder in the acute phase (acute MOG), DCs, or HCs (upper, NQO1/lower, Nrf2). Images were captured by an In Cell Analyzer 2000. The JC-1 red/green intensity ratio reflects the mitochondria potential. Scale bar, 50 μm. (C–E) Scatter plots of the intensity of NQO1 or Nrf2, or JC-1 red/green intensity ratio in TY10 cells, as determined by high-content imaging after exposure to IgG from patients with MOG-Ab–associated disorder in the acute phase (acute MOG, n = 14), DCs (n = 9), and HCs (n = 7). The data were normalized to cultures that had not been exposed to human IgG and are shown from 3 independent experiments. The p values were determined by a 1-way ANOVA followed by the Tukey multiple comparison test (*p < 0.05 vs the DC or HC group followed by the Tukey multiple comparison test). (F) Western blotting of Nrf2 in TY10 after exposure to acute MOG-IgG (n = 15), DC-IgG (n = 7), or HC-IgG (n = 7) (500 μg/mL). (G) Scatter plots of quantification by Western blotting for Nrf2 in relation to actin. Each scatter plot reflects the combined densitometry data (mean ± standard error of the mean). The p values were determined by a 1-way ANOVA followed by the Tukey multiple comparison test (***p < 0.001 vs the DC or HC group followed by the Tukey multiple comparison test). (H) Effect of 2 MOG-IgG (Pts 1 and 9, 250 μg/mL) on 10-kDa dextran permeability on BMECs after incubation with and without bardoxolone methyl (MOG vs MOG + bardoxolone methyl), which has the effect of activating Nrf2 and inhibiting NF-κB activation. ANOVA = analysis of variance; BMEC = brain microvascular endothelial cell; DC = disease control; HC = healthy control; MOG = myelin oligodendrocyte glycoprotein; MOG-Ab = myelin oligodendrocyte glycoprotein antibody.

(A, B) Immunostaining of human brain microvascular endothelial cells (TY10 cells) for NQO1 or Nrf2 (green), or JC-1 (green/red) after exposure to IgG (500 μg/mL) from patients with MOG-Ab–associated disorder in the acute phase (acute MOG), DCs, or HCs (upper, NQO1/lower, Nrf2). Images were captured by an In Cell Analyzer 2000. The JC-1 red/green intensity ratio reflects the mitochondria potential. Scale bar, 50 μm. (C–E) Scatter plots of the intensity of NQO1 or Nrf2, or JC-1 red/green intensity ratio in TY10 cells, as determined by high-content imaging after exposure to IgG from patients with MOG-Ab–associated disorder in the acute phase (acute MOG, n = 14), DCs (n = 9), and HCs (n = 7). The data were normalized to cultures that had not been exposed to human IgG and are shown from 3 independent experiments. The p values were determined by a 1-way ANOVA followed by the Tukey multiple comparison test (*p < 0.05 vs the DC or HC group followed by the Tukey multiple comparison test). (F) Western blotting of Nrf2 in TY10 after exposure to acute MOG-IgG (n = 15), DC-IgG (n = 7), or HC-IgG (n = 7) (500 μg/mL). (G) Scatter plots of quantification by Western blotting for Nrf2 in relation to actin. Each scatter plot reflects the combined densitometry data (mean ± standard error of the mean). The p values were determined by a 1-way ANOVA followed by the Tukey multiple comparison test (***p < 0.001 vs the DC or HC group followed by the Tukey multiple comparison test). (H) Effect of 2 MOG-IgG (Pts 1 and 9, 250 μg/mL) on 10-kDa dextran permeability on BMECs after incubation with and without bardoxolone methyl (MOG vs MOG + bardoxolone methyl), which has the effect of activating Nrf2 and inhibiting NF-κB activation. ANOVA = analysis of variance; BMEC = brain microvascular endothelial cell; DC = disease control; HC = healthy control; MOG = myelin oligodendrocyte glycoprotein; MOG-Ab = myelin oligodendrocyte glycoprotein antibody.
Figure 5 Western Blotting of GRP78 Autoantibodies in IgG From Patients With MOG Antibody–Associated Disorders

(A) The results of Western blotting of individual IgG samples (5 μg/mL) from patients with acute and stable MOG, MS, DCs, and HCs, as determined using recombinant human GRP78 protein prepared from *Escherichia coli*. The arrowhead indicates an immunoreactive band corresponding to GRP78. Rabbit anti-GRP78 antibodies were used as the positive control. The rate of GRP78 antibody positivity: (1) the acute MOG group (10/15, 67% [95% CI 38%–88%]), (2) stable MOG group (5/14, 36% [95% CI 13%–65%]), (3) MS group (total MS: 4/29, 14% [95% CI 4%–32%]), (4) DC group (3 of 27, 11% [95% CI 2%–29%]), (5) HC group (0 of 9, 0% [95% CI 0%]), (6) secondary progressive MS group (3 of 10, 30% [95% CI 6%–65%]), (7) acute MS group (0 of 10, 0% [95% CI 0%], and (8) stable MS group (1 of 9, 11% [95% CI 0.3%–48%]). (B) Immunofluorescence labeling of TY10 cells with MOG-IgG (50 μg/mL) (green) and commercial anti-GRP78 antibodies (red) shows colocalization of the 2 proteins (merged in yellow). Scale bar, 50 μm. CI = confidence interval; DC = disease control; HC = healthy control; MOG = myelin oligodendrocyte glycoprotein.
patients with MOG-Ab–associated disorder compared with that from DCs or HCs (Figure 4, F and G). The effect of MOG-IgG (MOG1 and 9) on 10-kDa dextran permeability in BMECs was reversed after incubation with bardoxolone methyl, which has the effect of activating Nrf2 and inhibiting NF-xB activation.

**Detection of GRP78 Autoantibodies in MOG-Ab–Associated Disorder**

The GRP78 autoantibodies were detected in the IgG from patients with MOG-Ab–associated disorder by Western blotting. The rate of GRP78 antibody positivity observed in the acute MOG group (10/15, 67% [95% confidence interval {CI} 38%–88%]) was significantly higher than that in the stable MOG group (5/14, 36% [95% CI 13%–65%]), the MS group (total MS: 4/29, 14% [95% CI 4%–32%]), the DC group (3 of 27, 11% [95% CI 2%–29%]), or the HC group (0 of 9, 0% [95% CI 0%]) (Figure 5A). The rate of this antibody positivity in the acute MOG group (10/15, 67% [95% CI 38%–88%]) was also significantly higher than that in secondary progressive MS (3/10, 30% [95% CI 6%–65%]), acute MS (0/10, 0% [95% CI 0%]) and stable MS (1/9, 11% [95% CI 0.3%–48%]. Double immunostaining with commercial anti-GRP78 Abs and IgG from GRP78 antibody–positive patients with MOG-Ab–associated disorder demonstrated colocalization in BMECs (Figure S8).

**Removal of GRP78 Antibodies in MOG-Ab–Associated Disorder**

We selected GRP78 antibody–positive IgGs from 2 acute MOG patients who showed a ΔEDSS score of >5 and the highest effect on the induction of NF-xB p65 nuclear translocation of TY10 (patients 1 and 9 in Table). We prepared 2 MOG-IgGs (1 with and 1 without GRP78 antibodies) from these MOG patients using immunoprecipitation. Complete depletion of GRP78 antibodies was observed by a Western blot analysis (eFigure 3, links.lww.com/NNX1/A663). The removal of GRP78 antibodies from MOG-IgGs from these 2 patients with MOG-Ab–associated disorder resulted in a significant reduction in NF-xB nuclear translocation and decreased permeability of BMECs (Figure 6, A and B).

There was no significant association between the clinical phenotype/ΔEDSS score and positivity for GRP78 antibodies/number of NF-xB p65 nuclear-positive BMECs after MOG-IgG exposure (data not shown).

**Discussion**

MOG is specifically expressed in oligodendrocytes, not in BMECs, so MOG-Abs itself cannot bind and react to the BBB via immune activation.\(^2\) In the present study, we demonstrated that IgG derived from MOG-Ab–associated disorder activated BBB endothelial cells; fraction of TY10 with the NF-xB p65 nuclear translocation was significantly increased after the acute MOG group than that after the DC group and significantly lower in the same individual MOG patients between the acute and stable phases. The permeability of the monolayer TY10 cells was significantly higher in the acute MOG group than in the stable MOG/DC/HC group. The amount of VCAM-1 and ICAM-1 protein in the TY10 cells was also higher in the acute MOG group than that in the stable MOG and HC group. These results suggest that IgG from the acute-phase MOG patients induced endothelial activation and increased the BBB permeability. Whole RNA-seq and the pathway analysis revealed that the genes associated with NF-xB signal and oxidative stress were significantly changed after exposure to IgG from GRP78 antibody–positive patients with MOG-Ab–associated disorder compared with HCs. In addition, the amount of NQO1 and Nrf2 protein and the mitochondrial membrane potential were lower in the acute MOG group than in the DC and/or HC groups, and activation of Nrf2 reversed the increased permeability of TY10 cells after MOG-IgG exposure, suggesting that IgG from acute-phase MOG patients induced oxidative stress in TY10 cells. Furthermore, the positive rate of GRP78 autoantibodies in the acute MOG
group was significantly higher than that in the stable MOG, DC, MS, and HC groups (acute MOG 66% vs stable MS 36%, MS 14%, DCs 11%, and HCs 0%). Double immunostaining with commercial anti-GRP78 antibody and MOG-IgG demonstrated colocalization in TY cells, suggesting that MOG-IgG reacted with GRP78. Depletion of GRP78 antibodies from acute MOG-IgG reduced the effect on NF-κB p65 nuclear translocation and BMEC permeability. These results suggest that IgGs from patients with MOG-Ab–associated disorder induced the disruption of the BBB via the NF-κB signaling and the oxidative stress, and GRP78 antibodies were associated with BBB breakdown in patients with MOG-Ab–associated disorder. Unfortunately, we were unable to perform the in vivo experiments in which GRP78-IgG–seropositive and –seronegative myelin oligodendrocyte glycoprotein antibodies–associated disorder sera were administered peripherally to MOG-EAE animals to evaluate the relationship between GRP78 autoantibodies and BBB permeability in the present study.

We recently reported that GRP78 autoantibodies were a potential biomarker associated with the BBB breakdown in NMO.15 In addition, the positivity rate of GRP78 autoantibodies in the longitudinally extensive transverse myelitis phenotype of NMOSD was higher than that in the optic neuritis (ON) phenotype (longitudinally extensive transverse myelitis 71% vs ON 17%), and positivity of GRP78 antibodies was associated with an increased clinical severity (ΔEDSS score) in patients with NMOSD and BBB permeability in our in vitro BBB model.16 The present study showed that GRP78 antibodies were associated with MOG-Ab–associated disorder beyond NMOSD, suggesting that these antibodies may be able to be triggered in the activation of BMECs in the disease as well as NMOSD.

NQO1 is a cytosolic homodimeric flavoprotein that catalyzes the 2-electron reduction of quinones, and its production is induced under many stress conditions, including oxidative stress, to protect against cellular damage. Nrf2 plays an important role in the regulation of many antioxidant enzyme genes, such as NQO1.22 A reduction in the Nrf2 signal leads to the induction of oxidative stress via a decrease in NQO1 and increase in inflammatory signaling pathways, including that of NF-κB, resulting in dysfunction of the BBB endothelial cells and increased paracellular permeability.23 Bardoxolone methyl activates the Nrf2 system and protects the cells against oxidative stress through the inhibition of reactive oxygen species generation.24 BMEC permeability induced by MOG-IgG was decreased after incubation with bardoxolone methyl in the present study. We revealed the role of GRP78 antibody in MOG-Ab–associated disorder on the BBB dysfunction: the induction of inflammation via NF-κB and the increase in oxidative stress through Nrf2 on BBB endothelial cells.

We next evaluated clinical data using statistical analyses to address whether NF-κB nuclear translocation/permeability of BMECs and GRP78 antibodies was correlated with the clinical data in MOG-Ab–associated disorder. However, we failed to observe any significant association between NF-κB nuclear translocation/permeability/GRP78 antibodies and the clinical phenotype/disease activity/clinical data. Multiple potential factors, including the small sample size, low assay sensitivity by Western blotting, a lack of detection of GRP78 antibodies titer, a lack of information of pathogenic autoantibody epitopes, and a relatively poor understanding of clinical metric and the disease pathophysiology, may have contributed to the lack of a correlation. The establishment of a better assay for GRP78 autoantibodies, including an ELISA using pathogenic antibody epitopes and a CBA to determine the GRP78 antibody titer, will be needed to address these important questions.

A case report described 2 patients with MOG-Ab–associated encephalitis mimicking small-vessel CNS vasculitis: Brain biopsy samples of abnormal MRI scans from these 2 patients revealed prominent lymphocytic infiltration of the wall of the small vessels, including T and B lymphocytes with edema, perivascular demyelination, and reactive gliosis, without fibrinoid necrosis.25 The present study showed the upregulation of VCAM-1 in BBB endothelial cells after exposure to IgG from patients with MOG-Ab–associated disorder. The overexpression of VCAM-1 in the brain microvasculature may induce lymphocytic infiltration around the small vessels of the CNS.26 GRP78 antibodies may play a role in inducing the inflammation of small vessels of the brain in MOG-Ab–associated encephalitis via the upregulation of the VCAM-1 and ICAM-1 expression.

In conclusion, our data show that GRP78 autoantibodies play a role in increasing the permeability of BMECs, thus suggesting the potential ability to the entry of pathogenic MOG-Abs into the CNS in MOG-Ab–associated disorder. Further studies using better assay, including an ELISA and CBA, to determine the GRP78 antibody titer will be required to understand the association between the clinical phenotype of the disease and GRP78 antibodies. An in vivo study will be needed to confirm the association between GRP78 autoantibodies and BBB permeability.

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**References**


GRP78 Antibodies Are Associated With Blood-Brain Barrier Breakdown in Anti–Myelin Oligodendrocyte Glycoprotein Antibody–Associated Disorder

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