Mutagenesis of the Aquaporin 4 Extracellular Domains Defines Restricted Binding Patterns of Pathogenic Neuromyelitis Optica IgG

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Background: Patient-derived aquaporin 4 (AQP4)-specific recombinant monoclonal antibodies (rAbs) cause neuromyelitis optica (NMO)-specific nervous system injury in animal models.

Results: AQP4-specific rAbs bind human AQP4 based on differential sensitivity to loop A and C mutations.

Conclusion: AQP4-specific rAbs derived from NMO patients recognize multiple conformational epitopes within the extracellular domains of human AQP4.

Significance: High resolution mapping of AQP4 autoantibody epitopes identifies target regions for potential blocking therapies.

Neuromyelitis optica-immunoglobulin G (NMO-IgG) binds to aquaporin-4 (AQP4) water channels in the central nervous system leading to immune-mediated injury. We have previously demonstrated that a high proportion of CSF plasma cells of NMO patients produce antibody to the extracellular domains of the AQP4 protein and that recombinant IgG (rAb) derived from these cells recapitulate pathogenic features of disease. We performed a comprehensive mutational analysis of the three extracellular loops of the M23 isoform of human AQP4 using both serial and single point mutations, and we evaluated the effects on binding of NMO AQP4-reactive rAbs by quantitative immunofluorescence. Whereas all NMO rAbs required conserved loop C (137-TP138 and Val150) and loop E (230-HW231) amino acids for binding, two broad patterns of NMO-IgG recognition could be distinguished based on differential sensitivity to loop A amino acid changes. Pattern 1 NMO rAbs were insensitive to loop A mutations and could be further discriminated by differential sensitivity to amino acid changes in loop C (148-Asn149 and His151) and loop E (Asn226 and Gln228). Alternatively, pattern 2 NMO rAbs showed significantly reduced binding following amino acid changes in loop A (63-EKP65 and Asp69) and loop C (Val141, His151, and Leu154). Amino acid substitutions at 137-TP138 altered loop C conformation and abolished the binding of all NMO rAbs and NMO-IgG, indicating the global importance of loop C conformation to the recognition of AQP4 by pathogenic NMO Abs. The generation of human NMO rAbs has allowed the first high resolution mapping of extracellular loop amino acids critical for NMO-IgG binding and identified regions of AQP4 extracellular structure that may represent prime targets for drug therapy.

Neuromyelitis optica (NMO)2 is a severe inflammatory demyelinating disorder of the central nervous system (CNS) that preferentially targets optic nerves and spinal cord leading to paralysis and vision loss. Although once considered a variant of multiple sclerosis, clues to the nature of disease were first revealed by the detection of an NMO-specific serum antibody (Ab) response, termed NMO-IgG, that immunostained the surface of microvessels, pia, subpia, and Virchow-Robin spaces in the CNS (1). NMO IgG was subsequently shown to recognize the water channel aquaporin-4 (AQP4), which is expressed on astrocytes in the CNS and is preferentially polarized to astrocytic end-foot processes (2). We showed that during disease exacerbation, CSF from NMO patients contained a dynamic population of expanded and clonally related plasmablasts that were enriched for cells producing Abs against the extracellular domains of AQP4 (3). More importantly, AQP4-specific recombinant Abs (rAbs) derived from these CSF plasmablasts, when administered in experimental animals, recapitulated the myriad pathological features of NMO lesions that include perivascular loss of astrocytes, terminal complement activation, granulocyte infiltration, and subsequent oligodendrocyte cell death and myelinolysis (3–5). The pathology produced in animal models parallels that observed with serum-derived NMO-IgG, indicating that CSF rAbs reproduce the disease-specific response of AQP4-seropositive patients (5–7).

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‡ The abbreviations used are: NMO, neuromyelitis optica; rAb, recombinant monoclonal Ab; AQP4, aquaporin 4; BN, blue native; OAPs, orthogonal array of particles; TIRF, total internal reflection fluorescence; TIRFM, TIRF microscope; NGS, normal goat serum; Ab, antibody.
Binding Patterns of AQP4-IgG

Although knowledge of NMO pathology has been greatly enhanced by the identification of NMO-IgG, many questions remain regarding the origin of this CNS B cell response, its variation across the NMO patient population, and the steps and signaling pathways leading from astrocyte destruction to demyelination. One critical factor that promotes Ab-mediated CNS tissue damage is the ability of the M23 isoform of AQP4 to assemble from tetramers into large supramolecular structures called orthogonal array of particles (OAPs). AQP4 is expressed in alternative isoforms termed long (M1) and short (M23). The M23 isoform promotes OAP formation, and the size of OAPs is determined by the ratio of M23:M1 isoforms (8, 9). A quantitative comparison of binding to the M1 and M23 isoforms shows that both serum NMO-IgG and most NMO CSF-derived rAbs bind with higher affinity to OAPs than to tetramers and that structural changes in the AQP4 epitope upon array assembly, and not bivalent binding of IgG, drive this increased affinity (10). OAP formation is also required for anti-AQP4 Abs to initiate complement-dependent cell lysis, which likely occurs through the promotion of a multivalent interaction of C1q molecules with cell surface-bound AQP4 Abs (10, 11).

Presently, it is not known whether the binding of Abs to specific epitopes on AQP4 may preferentially affect lesion formation and contribute to differences in disease severity. Studies to define the depth and specificity of the NMO serum IgG response have been somewhat disparate. In two separate studies using serum NMO-IgG to define AQP4 epitopes, Pisani et al. (12) identified two major conformational AQP4 epitopes in the extracellular domains of AQP4 by mutagenesis of amino acids within the extracellular domains, whereas Iorio et al. (13) reported a broader array of AQP4 epitopes displayed on denatured AQP4 monomers, tetramers, and OAPs. Recently, muta-
tion of Asp69 has been shown to alter loop A conformation and contribute to differences in disease severity. Studies of which may be nonpathogenic species directed against denatured AQP4 protein or protein fragments. In this study, we define the dominant epitopes recognized by a panel of AQP4-specific monoclonal recombinant Abs from CSF plasmablasts recovered from five NMO patients. Site-directed mutagenesis was used to generate seven 4- to 6-amino acid serial substitutions and 29-point mutations encompassing most amino acids (glycines excluded) within each of the three extracellular loops (A, C, and E) of the M23 isoform of human AQP4. The effects of each mutation on OAP array formation and AQP4 rAb binding were evaluated to define distinct and not previously appreciated patterns of antigen recognition.

EXPERIMENTAL PROCEDURES

Patients—CSF was obtained from five NMO-IgG seropositive patients 1–8 weeks after the onset of unilateral monosymptomatic optic neuritis or transverse myelitis as part of their standard clinical care. The clinical testing of each patient’s serum for NMO-IgG was performed at the Mayo Clinic Laboratories, Rochester, MN, using their standardized commercial clinical assay. Informed consent was obtained for all individuals prior to participation in this study.

CSF Cell Labeling and FACS—CSF cell collection, fluorescent labeling, and cell sorting of CD138+ CSF plasma cells and plasmablasts were performed as described (3). Cells were first selected in the size range of lymphocytes and plasma cells by forward and side light scattering. CD138+ cells were then identified and sorted into single wells of a 96-well PCR plate containing 20 μl of 1× RT buffer.

cDNA Synthesis and Amplification of VH and VL Chain Sequences—cDNA synthesis, nested PCR amplification, and purification of PCR products were performed as described (3, 15). Purified PCR products were sequenced at the University of Colorado Cancer Center DNA Sequencing Core. Sequences were analyzed and edited with 4Peaks software (Mek&Tosj.com) and then aligned to functional human immunoglobulin germline sequences using IMGT/V-QUEST.

Construction, Expression, and Purification of rAbs—rAbs were produced from NMO CSF plasma cell and plasmablast clonal populations using a dual vector transient transfection system. VH and VL PCR products were cloned into the expression vectors plgG1FLAG and pCEP4, respectively, as described. Final constructs were sequenced and verified. Constructs were co-transfected into HEK293 cells using Lipofectamine 2000 (Invitrogen). After transfection, the cells were grown for 6–7 days in DMEM + 10% fetal bovine serum (FBS); the supernant was harvested, fresh medium was added, and cells were propagated for 6–7 days. The cell culture supernatant was subsequently removed and combined with the previous collection. The cell culture supernatant was centrifuged at 10,000 × g to pellet cells and debris. Cell-free supernatants were passed twice through protein A-Sepharose (Sigma), and rAbs were eluted in 0.1 M glycine, 1 M NaCl, pH 3.0, and neutralized by the addition of 0.1 M Tris-HCl, pH 8.0. Recombinant IgG was subsequently exchanged and concentrated in storage buffer (PBS + 0.1% IgG/protease-free BSA) using Ultracel YM-30 microconcentrators (EMD Millipore, Darmstadt, Germany). Antibody integrity was confirmed by nondenaturing gel electrophoresis, and the IgG concentration was determined by a human IgG capture ELISA or by BCA colorimetric protein quantification (Pierce). The specificity of each rAb used in this study was confirmed either by FACS or by fluorescence immunostaining of mock- and AQP4-expressing cells (3).

Mutagenesis of M23-AQP4—Both 4–6 amino acid serial alanine or glycine substitutions and point mutations were introduced into M23 AQP4 extracellular loops using the PCR-based GeneArt site-directed mutagenesis system (Invitrogen) (Table 1). Partially overlapping sense and antisense primers were designed from the M23 AQP4 sequence with the desired nucleotide mutation(s) located near the center of each primer sequence. Methylation and mutagenesis reactions of AQP4 DNA were done using 20–25 ng of target AQP4 plasmid DNA according to the manufacturer’s instructions. AQP4 DNA mutations were confirmed by sequencing, and endonuclease-free plasmid DNA was purified for transfection into mammalian cells (Qiagen catalog no. 12362).

Plasmid DNA containing M23-AQP4 or mutated M23-AQP4 was then transfected with Lipofectamine 2000 into...
U-87MG or HEK293 EBNA cells growing on poly-L-ornithine-treated coverslips. Cells were incubated for ~24 h at 37 °C, rinsed in PBS, and fixed in 4% paraformaldehyde in PBS. Coverslips were stored at −20 °C until used.

**Generation of Stable U-87MG Cell Lines Expressing AQP4M23 and Mutated AQP4M23—**Plasmid DNA containing the AQP4-M23 cDNA sequence or the loop A mutant, ΔA3, cDNA sequence were linearized with PvuI for 1 h at 37 °C and then transfected into U-87MG astrocytoma cells with Lipofectamine 2000 per manufacturer’s protocol. At 24 h post-transfection, G418 was added at a concentration of 400 μg/ml, and cells were monitored for survival. After ~1 week, growing clonal populations were diluted in U-87MG selection media (DMEM high glucose media containing 10% FCS, 100 μg/ml penicillin/streptomycin, and 300 μg/ml G418) to less than 10 cells/ml, and 100 μl were deposited into a 96-well plate. Plates were monitored 1–2 weeks for clonal growth and then 5–6 individual wells were eluted and expanded into a T75 flask. G418 selection was maintained at 200 μg/ml, and then cells were passed and seeded onto coverslips to assess AQP4 expression. NMO rAbs were assayed for binding as described below.

**Immunostaining—**Cells on coverslips were blocked in 10% normal goat serum (NGS)/PBS for 30 min at room temperature. Individual AQP4-positive rAbs at 10–20 μg/ml in 5% NGS/PBS were then applied to coverslips and incubated overnight at 4 °C. The concentration of Ab represented concentrations at or below their apparent K_a values. A secondary control M23-transfected coverslip that was used to correct for background staining of the Alexa Fluor 488 anti-human secondary antibody alone was included and remained in blocking buffer overnight at 4 °C. Coverslips were washed four times in 1× PBS for 3–5 min each, fixed for 5 min with 4% paraformaldehyde, and then blocked in 5% NGS/PBS containing 0.1% Triton X-100. All coverslips were then incubated with 4 μg/ml of a rabbit polyclonal anti-human AQP4 Ab (Santa Cruz Biotechnology) in 2% NGS/PBS for 1 h at room temperature to normalize for transfection efficiency. Coverslips were washed and incubated with 1:600–1:1000 dilutions of both a goat anti-human Alexa Fluor 488 and goat anti-rabbit Alexa Fluor 594 Ab in 2% NGS/PBS for 1 h at room temperature. Cells were washed five times with 1× PBS for 3–5 min each and mounted onto clean microscope slides using Vectashield HardSet Mounting Medium with DAPI (Vector Laboratories).

**Image Analysis—**Immunostained coverslips were visualized on a Nikon E800 fluorescence microscope or on an Olympus IX81 confocal fluorescence microscope using either ×20 or ×40 objective lens. Images were acquired from 4 to 8 nonoverlapping visual fields per coverslip, and images were processed using AxioVision 4.8 software. For studies of Ab binding to AQP4 extracellular loop mutations, 2–5 independently transfected coverslips and 10–30 images were processed for quantitative analyses. The binding intensities of individual NMO rAbs (green fluorescence) and the C-terminal specific positive control anti-AQP4 rabbit polyclonal Ab (red fluorescence) were measured using Image software, and the average background binding of anti-human secondary Ab (green fluorescence) alone to coverslips was subtracted, which typically represented 0.1–3% of rAb binding levels. The average ratio of NMO rAb to control polyclonal Ab (green/red) was calculated for each rAb and wildtype M23 AQP4 to normalize for AQP4 expression and the ratio of the fraction antibody bound on mutant and wildtype M23 AQP4 was then computed and reported as % M23 binding ± S.E. using Equation 1,

\[
\% \text{ M23 Binding} = \frac{\text{Mutant (Green/Red)}}{\text{M23 (Green/Red)}} \times 100
\]  

(Eq. 1)

**Statistical Analysis—**The statistical effect of each mutation on binding to M23AQP4 was determined using an unpaired two-sided Student’s t test with unequal variances.

**BN-PAGE, SDS-PAGE, and TIRF Microscopy (TIRFM)—**Transfected U-87MG cells were collected 24–48 h post-transfection, washed in cold PBS, and lysed in native PAGE sample buffer (Invitrogen) containing 0.5% dodecyl-β-D-maltos. Samples were then electrophoresed through blue native gels, transferred onto PVDF membrane, and probed with a polyclonal anti-rabbit antibody against the intracellular C terminus of AQP4 Ab as described (10). Formation of OAPs was visualized by the characteristic ladder appearance (Fig. 1). Alternatively, OAP formation in transfected cells was visualized for some mutations by TIRF microscopy revealing a characteristic punctate staining at the cell surface (Fig. 1). The effects of serial and point mutations on tetramer and OAP formation are reported for each construct in Table 1.

For SDS-PAGE analysis of NMO rAb binding to AQP4 M23, low speed cell lysates were prepared from AQP4-transfected cells and quantified by the BCA protein assay (Pierce), and 20 μg of protein were resolved on a SDS-12.5% polyacrylamide gel. Proteins were electrophoretically transferred to nitrocellulose, disected into individual lanes, blocked in 3% BSA, and then incubated with 1–2 μg/ml of the indicated NMO CSF-derived human rAb or the polyclonal anti-rabbit intracellular AQP4 Ab. Nitrocellulose strips were then washed in TBS containing 0.1% Tween 20 (TBS/Tween 20), incubated with goat anti-human HRP- or goat anti-rabbit HRP-conjugated secondary antibody in 3% BSA, and then rewash in TBS/Tween 20. Binding was visualized by chemiluminescence (Supersignal West Pico kit, Pierce).

**Mapping of M23-AQP4-binding Sites—**The three-dimensional mapping of putative AQP4 epitopes was based on the crystal structure of human M1-AQP4 (Protein Data Bank code 3DG8) at 1.8 Å (16). Three-dimensional images of AQP4 monomers and tetramers were obtained and manipulated using Imol Viewer at the Protein Data Bank. False coloration of AQP4 extracellular domains were performed manually on exported images using Adobe Photoshop.

**RESULTS**

Monoclonal Anti-AQP4 NMO Antibodies Do Not Recognize Linear Epitopes on AQP4—FACS single-cell sorting was used to isolate single CD138⁺ antibody-secreting cells from the CSF of five NMO-IgG-positive patients. Sequencing of their PCR-amplified heavy- and light-chain variable regions identified clonal populations within each repertoire, and Abs were generated from each clone as described previously (3). NMO CSF-
derived rAbs specific for AQP4 failed to recognize denatured AQP4 protein monomers in SDS-PAGE protein immunoblots of lysates from AQP4-transfected CHOK1 cells (Fig. 2), indicating that CSF-derived Abs likely recognize conformational epitopes formed by the assembly of AQP4 tetramers and OAPs at the cell surface.

**Mutagenesis of AQP4 Extracellular Domains Identifies Two Patterns of AQP4 Recognition**—To define regions of the AQP4 extracellular domain important for Ab recognition, we first introduced a series of alanine and/or glycine substitutions into the three extracellular domains of human M23 AQP4 as listed in Table 1. This included three sets of nonoverlapping serial substitutions covering loop A (ΔA1-ΔA3) and loop C (ΔC1-ΔC3), and one serial substitution that covers loop E (ΔE1). Plasmid DNA of each mutation and M23 AQP4 were separately transfected into U-87MG glioblastoma or HEK-EBNA cell lines, and binding to AQP4 was assayed by a dual immunostaining protocol in which AQP4 was immunostained using a polyclonal Ab recognizing the intracellular AQP4 C terminus. The double staining of normal and mutated AQP4-transfected cells with ONO7-5 #53 rAb (green fluorescence) and the AQP4-positive control Ab (red fluorescence) are shown in Fig. 3A. Whereas binding of the positive control Ab is largely unaltered in cells expressing AQP4 serial extracellular loop substitutions, binding of rAb #53 is absent in cells expressing the ΔC1-ΔC3 and ΔE1 mutations. Other NMO rAbs displayed additional sensitivity to the serial substitutions ΔA2 and ΔA3 in loop A (Fig. 3B). As summarized in Table 2 for all AQP4-spe-
to AQP4 recognition, we assayed three pattern 2 NMO rAbs for binding to each of the loop A mutations and assayed three pattern 1 and three pattern 2 NMO rAbs for binding to each loop C and E. The point mutation W227G in loop E eliminated AQP4 plasma membrane expression and OAP formation preventing any assessment of the role of this amino acid in pattern 1 or pattern 2 rAb binding. Because the ΔA1 serial substitution did not alter binding of any AQP4-specific rAbs, point mutations were not generated in this region of loop A.

Single alanine substitutions within loop A sequence \( \text{TEK}^{230} \) identified the amino acids EKP at positions 63–65 as important contributors to pattern 2 Ab binding (Fig. 4). A modest but significant reduction in binding was observed at each of these positions suggesting that this trio of amino acids together help facilitate binding of pattern 2 rAbs to AQP4. ON07-5 #58, a human AQP4-specific rAb, was most affected by the K64A point mutation, which also represents a species-specific difference between human and mouse AQP4. In the region \(^{(66} \text{LPVD}^{66})\) covered by the loop A serial substitution ΔA3, the only residue impacting AQP4 binding was Asp\(^{69}\), which produced a significant loss of binding for all pattern 2 rAbs (Fig. 4).

Pisani et al. (14) recently reported that a D69H substitution affected loop A conformation and serum NMO-IgG binding. Loop C is the largest AQP4 extracellular domain and appears critical for rAb recognition as judged by the absence of rAb binding to M23 AQP4 mutant ΔC1. The crystal structure of human AQP4 reveals an \( \alpha \)-helix at positions \(^{139} \text{PSVV}^{142}\) within the N-terminal portion of loop C (16). To determine whether this helix might be important for antigen binding, we first generated two smaller point mutations (TP137–8 AA and SVV140–2AA). The TP137–8 AA mutation greatly reduced the binding of every AQP4-specific rAb assayed (Fig. 5). The effects of the SVV140–2AA mutation were mixed and varied by individual rAbs independent of their pattern 1 or pattern 2 classifications (data not shown). Single point mutations introduced at positions Ser\(^{140}\), Val\(^{141}\), and Val\(^{142}\) and across the region \(^{145} \text{LGVTM}^{149}\) further illustrated these Ab-specific positional effects, which ranged from modestly increased binding to AQP4 for some pattern 1 Abs (ON07-5 #53 at Val\(^{141}\) and ON10-1 #153 at Val\(^{147}\)) to moderately decreased binding for many other rAbs. Mutations causing the greatest disruption of individual rAb binding within this region included V142A for ON07-5 #10, T148A and M149A for ON07-5 #43, and M149A for ON10-1 #153. The Met\(^{149}\) amino acid is specific to human AQP4 reflecting the specificity of ON07-5 #43 Ab for human AQP4. The species specificity of ON10-1 #153, which was also significantly affected by this mutation, has not been determined. Experiments to independently evaluate the contributions of Thr\(^{139}\), Pro\(^{140}\), and Pro\(^{141}\) were not performed. Excluding the \(^{137} \text{TP}^{138}\) amino acids, which were necessary for the binding of all NMO rAbs, no other single mutation encompassed within the ΔC1 and ΔC2 mutations unambiguously distinguished pattern 1 from pattern 2 binding.

Point mutations at Val\(^{150}\) in loop C identified this amino acid as critical for AQP4 recognition by all NMO rAbs. When alanine was substituted for Val\(^{150}\), plasma membrane expression and OAP formation were abrogated thus preventing NMO rAb binding (Figs. 1 and 5C); however, a V150G substitution at this

### TABLE 1

**Mutations introduced into the extracellular domains of AQP4**

| Loop | AQP4 mutation* | Mutational changes* | Mutated amino acid positions* | Tetramer and OAP array formation* |
|------|----------------|---------------------|------------------------------|---------------------------------|
| Loop A | ΔA1 | TINW AAAG | 56–59 | OAP |
| | ΔA2 | TEKP AAGA | 62–65 | OAP |
| | ΔA3 | LPVD AAAAA | 66–69 | OAP |
| | T62A | AEKP AEKP | 62 | OAP |
| | E63A | TEKP TAKP | 63 | OAP |
| | K64A | TEAP TEAP | 64 | OAP |
| | P65A | TEKA TEKA | 65 | OAP |
| | L66A | LPVD APVD | 66 | OAP |
| | P67A | LPVD LAVD | 67 | OAP |
| | V68A | LPVD LAPD | 68 | OAP |
| | D69A | LPVD LPYA | 69 | OAP |
| Loop C | ΔC1 | TPPSVV AGAAGA | 137–142 | OAP |
| | ΔC2 | LGVTM AGAAA | 145–149 | NEG |
| | ΔC3 | VHGNLT AGAAGA | 150–155 | NEG |
| | TP137–8AA | TPPSVV AAPSVV | 137–138 | OAP |
| | SVV140–2AA | TPPSVV TTPAAAA | 140–142 | OAP |
| | S140A | TPPSVV TPAVV | 140 | OAP |
| | V141A | TPPSVV TPPSAV | 141 | OAP |
| | V142A | TPPSVV TPPSVVA | 142 | OAP |
| | L145A | LGVTM AGVTM | 145 | OAP |
| | V147A | LGVTM LGATM | 147 | OAP |
| | T148A | LGVTM LGVAM | 148 | OAP |
| | M149A | LGVTM LGVTA | 149 | OAP |
| | V150A | VHGNLT AGHGNLT | 150 | OAP |
| | V150G | VHGNLT AGHGNLT | 150 | OAP |
| | H151A | VHGNLT YAGNLT | 151 | OAP |
| | N153Q | VHGNLT VHQLQTL | 153 | OAP |
| | N153A | VHGNLT VHGLALT | 153 | OAP |
| | L154A | VHGNLT VHGPAT | 154 | OAP |
| | T155A | VHGNLT VHGLNA | 155 | OAP |

* The indicated amino acids located within the A, C, and E extracellular domains of the M23 isoform of human aquaporin 4 were mutated to alanine and or glycine residues as shown in column 2. An additional asparagine to glutamine mutation was also introduced at position 153.

* Amino acids are numbered according to the M1 isoform of hAQP4 as published by Ho et al. (16).

* The effects of amino acid mutations on the formation of tetramers and OAPs were assessed by BN-PAGE and by TIRF immunostaining (see Fig. 1).

Specific rAbs studied, both pattern 1 (loop A-independent) and pattern 2 rAbs (loop A-dependent) were recovered from each NMO patient. Using a U-87MG stable cell line expressing the ΔA3 AQP4 substitution mutation, we confirmed that all pattern 2 rAbs required intact extracellular loop A sequences (Fig. 3C). The necessity of individual regions within loops C and E (Table 1) remained unclear as the serial substitutions ΔC2, ΔC3, and ΔE1 eliminated plasma membrane expression and OAP formation as measured by BN-PAGE and TIRFM (see Fig. 1). Because of the uninformative nature of these large serial substitutions, point mutations that did not disrupt AQP4 plasma membrane expression and OAP assembly were required to generate a detailed map of individual amino acids contributing to epitope formation.

**Single Amino Acid Substitutions Identify Critical Residues Required for AQP4 Recognition**—We next introduced alanine and or glycine replacement mutations for most amino acid in loops A, C, and E, excluding native glycine residues (Table 1). To identify specific amino acids within each loop contributing
position maintained OAP formation, but significantly reduced binding for each of the six NMO rAbs assayed, with pattern 1 Abs having a somewhat greater sensitivity than pattern 2 rAbs. A second critical amino acid within the C-terminal portion of loop C that influenced the binding of both pattern 1 and 2 rAbs was Asn153, which is part of a consensus N-glycosylation motif \((153NLT155)\). A conservative N153Q substitution abolished or significantly reduced the binding of pattern 2 rAbs but not pattern 1 rAbs, whereas a second N153A substitution reduced the binding of all pattern 1 and 2 rAbs with the exception of ON07-5 #43, which was not affected by either substitution. The T155A mutation did not produce equivalent global reductions in pattern 2 binding indicating that the entire consensus sequence was not required.

Loop E \(\left(226NWENHW231\right)\) mutations revealed three residues (Trp\(^{227}\), His\(^{230}\), and Trp\(^{231}\)) that significantly affected the binding of each NMO rAb (Fig. 6). Unfortunately, the W227A substitution was uninformative as it abolished AQP4 plasma membrane and OAP formation (Table 1). The single point mutations H230G and W231G reduced the binding of all NMO rAbs without impacting OAP formation. ON07-5 #43 Ab represented a unique pattern 1 rAb with additional sensitivities to mutations N226A and E228A.

Loop C Amino Acids His\(^{151}\) and Leu\(^{154}\) Define a Subset of Pattern 2 Binding—The H151A and L154A mutations identify critical loop C amino acids that further delineate the heterogeneity of AQP4 recognition by AQP4-specific rAbs. Binding of both the pattern 1 rAb, ON07-5 #43, and pattern 2 rAb,
ON07-5 #10, was severely impaired by the H151A substitution. ON07-5 #10 binding was also significantly reduced by the H151A substitution as was the pattern 1 rAb ON10-1 #153, and ON07-5 #10, was severely impaired by the H151A substitution.

To determine whether amino acids necessary for AQP4 binding to M23 AQP4 indicating the global importance of loop C conformation for the binding of different classes of AQP4 autoantibodies. A

DISCUSSION

The binding of NMO patient-derived antibodies to the extracellular domains of AQP4 initiate an immune-mediated pathology that is characterized by activation of complement, astrocyte cytolysis, inflammation with granulocyte infiltration, and oligodendrocyte cell death leading ultimately to demyelination (3, 6, 17, 18). Although first described as a serum response, plasmablasts secreting anti-AQP4 Abs are clonally expanded within the CSF compartment and may be a significant source of AQP4-specific CNS Abs (3, 19). To study the CSF Ab response in NMO, we have used single-cell PCR to generate a panel of AQP4-specific monoclonal rAbs from expanded clones of plasmablasts present in CSF of NMO patients. The rAb products of these NMO patient-derived plasmablasts have reproduced CNS tissue pathology in multiple models of NMO lesion pathogenesis (3, 5, 20) demonstrating their ability to reproduce the pathogenicity of serum NMO-IgG. Hence, we used a panel of 12 CSF-derived AQP4-specific Abs from five NMO-IgG seropositive patients to evaluate the importance of each amino acid in the three extracellular loops of AQP4 for NMO-IgG recognition. To assess binding in its native state, cells were transfected with one of a series of 37 mutated AQP4 M23 constructs, and their binding was quantified by fluorescence microscopy with normalization for AQP4 expression using a second Ab.

In contrast to prior studies using polyclonal human serum containing a broad collection of antibodies with a wide range of affinities and epitopes, our use of AQP4-specific monoclonal rAbs allowed us to determine distinct amino acids important for the binding of different classes of AQP4 autoantibodies. A common feature of each NMO rAb was the importance of conformation and quaternary structure for AQP4 recognition. All CNS-derived AQP4-specific rAbs examined to date do not rec-
recognize AQP4 protein in denaturing SDS-PAGE immunoblots but require the expression of conformationally intact protein on the plasma membrane. In this study, all serial or single amino acid substitutions within AQP4 extracellular domains that abrogated M23 plasma membrane expression and OAP formation (Table 1) abolished AQP4 rAb binding relative to that of the C-terminal specific polyclonal control Ab. This included serial substitutions A9002C2, A9002C3, and A9002E1 as well as the single point mutations V150A in loop C and W227G in loop E. We hypothesize that these mutations are likely impairing AQP4 tetramer oligomerization and/or transport of AQP4 tetramers from the endoplasmic reticulum to the cell surface based on TIRFM (Fig. 1). Interestingly, although the A9002C2 mutant prevented plasma membrane expression and OAP formation, none of the four constituent amino acids (LVTM) produce the same inhibition when mutated individually. Together, these observations indicate the presence of important topology within the AQP4 extracellular domains that is necessary for oligomerization and/or transport to the cell surface.

While conformational membrane-expressed AQP4 was essential for NMO rAb binding, prior studies have reported serum NMO-IgG binding to AQP4 peptides, monomers, and higher order arrays (13, 19, 21). The distinction is due to the polyclonal nature of serum NMO-IgG, which contains many AQP4 autoantibodies directed against multiple targets as follows: conformational AQP4 protein, denatured AQP4 multimers and monomers, and AQP4 peptides. The relevance of AQP4 autoantibodies directed against denatured AQP4 or AQP4 pep-
tides in disease pathogenesis, however, remains unclear. Although NMO CSF rAbs have been shown to bind to AQP4-expressing cells or CNS astrocytes and mediate target destruction by *in vitro*, *ex vivo*, and *in vivo* methods (3, 5, 20), the ability of autoantibodies against denatured or linear AQP4 epitopes to bind plasma membrane AQP4 and activate antibody effector function has not been tested. In one study, a significant proportion of anti-AQP4 antibodies targeted linear epitopes localized in the intracellular domains of the protein (21), and in a second study, serum antibodies against loops A and E peptides lacked disease specificity (13). Although serum Abs to denatured linear AQP4 epitopes may be diagnostic for seropositive NMO (13), the uniform absence of such broad specificity to denatured epitopes in AQP4-specific rAbs derived from NMO CSF implies distinctions between global anti-AQP4 responses and the AQP4-specific B cells infiltrating the CNS. Pathogenicity is likely driven by conformationally dependent Abs binding to surface-exposed epitopes in tetramers that are stabilized or optimized in OAPs (10).

Our analysis revealed amino acids that were either critical for the binding of all AQP4-specific rAbs or defined unique epitopes specific to certain rAbs. Five AQP4 constructs greatly impaired NMO rAb recognition as follows: C loop mutants TP137–8AA and V150G (Fig. 8, purple), C loop mutant N153A (Fig. 8, turquoise), and E loop mutants H230G and W231G (Fig. 8, blue). The most remarkable feature distinguishing patterns of AQP4 antigen recognition is their differential sensitivity to mutations within the first extracellular domain, broadly categorized as either pattern 1 (loops C- and E-dependent) or pattern 2 (loops A-, C, and E-dependent) binding. Importantly, all the pattern 2 rAbs are uniquely sensitive to be mutagenesis of the same amino acids within the loop A domain (Fig. 8, highlighted in red). All pattern 2 rAbs are uniquely sensitive to both the ΔA2 and ΔA3 serial substitutions (Table 2 and Fig. 3D) and single amino acid mutations at Asp⁶⁶⁹ and the sequence E⁶³EK⁶⁶⁵. Although most pattern 1 and pattern 2 rAbs share the same sensitivity to the loop C mutants TP137–8AA, V150G, and N153A, a subset of pattern 2 rAbs, derived from different patients (Figs. 7 and 8) showed additional dependence on loop C amino acids His¹⁵¹ and Leu¹⁵⁴. This sub-pattern may be further defined by the amino acid Val¹⁴⁷, based on ON07-5 #10 rAb staining, but it has not been verified with other pattern 2 rAbs dependent on His¹⁵¹ and Leu¹⁵⁴ (Fig. 8). There are also multiple amino acids across loop C that have modest effects on the binding of individual rAbs, independent of their pattern classification, but do not delineate clear AQP4 epitopes, suggesting that subtle changes to the overall conformation of AQP4 extracellular domains influence AQP4-IgG binding. The epitopes of Abs classified as pattern 1 are less well defined than pattern 2 and show modest to strong sensitivities to unique amino acid substitutions across both loops C and E. The most unique pattern 1 Ab is ON07-5 #43, which displays a specific requirement for amino acids Thr¹⁴⁸, Met¹⁴⁹, His¹⁵¹, Asn²²⁶, and Glu²²⁸, and is the only rAb studied that is not affected by the N153A substitution.

The present analysis is limited by the inability to differentiate mutations that directly affect Ab contact from those that cause both global and local conformational changes. To preliminarily address this issue, we examined the three-dimensional structure of human AQP4 to investigate the juxtaposition of critical amino acids that likely define AQP4 epitope domains. Fig. 9 displays the crystal structure of human M1 AQP4 at 1.8 Å resolution (16) visualized at different angles to show the position of critical amino acids defined by this mutagenesis study. Because OAP formation drives high affinity binding of most CSF-derived rAbs and are formed from the M23 AQP4 isoform (10), a detailed examination of putative rAb-AQP4 interactions across OAPs is not feasible. Nevertheless, the following features are notable when viewing the three-dimensional structure of M1 AQP4 tetramers. First, amino acids projecting farthest outward from the extracellular surface form local structures comprised of critical residues required for NMO rAb binding (Fig. 9A). They include the loop A amino acids E⁶³EK⁶⁶⁵ that are critical for all pattern 2 rAb binding, and the loop C amino acid Asn¹⁵³ that is located on the surface of a horseshoe bend. This prominent bend within loop C also includes His¹⁵¹ and Leu¹⁵⁴.
that delineate a subset of pattern 2 Abs. Gly152 is also located within this bend, but its putative contribution to AQP4 recognition has not been investigated. Second, additional amino acids critical for NMO rAb binding are positioned in regions that likely dictate the correct three-dimensional spacing and conformation of extracellular loops (Fig. 9B). The loop C amino acids 137TPP139 create an almost 180° turn that positions loop C parallel to much of the extracellular surface of AQP4, whereas the other critical loop C amino acid Val150 creates an angle that directs the prominent loop C horseshoe (151HGNL154) perpendicular from the plane of the cell surface. The role of Asp69 in pattern 2 binding is not readily apparent from the three-dimensional models, but it has been reported to affect loop A conformation (14). Third, although loop E does not protrude outward from the cell surface as far as loops A and C, it runs parallel to a portion of loop C (145LGVTM149) forming a pocket that may be necessary for both pattern 1 and pattern 2 binding. It may also explain the unique sensitivity of individual rAbs to amino acid mutations in this region (Fig. 7C). Loop E also forms a protruding horseshoe bend that brings the loop C amino acids Thr148 and Met149 in close proximity (~7–8 Å) to the loop E amino acids Asn226, Trp227, and Glu228, positions that uniquely define the pattern 1 epitope represented by NMO rAb ON07-5 #43.

Finally, the three-dimensional structure of the AQP4 tetramer reveals important structural features that likely contribute to epitope recognition. The first and most striking feature is the juxtaposition of loop A amino acids 63EKP65 in one monomer to the 151HGNL154 amino acids of loop C in the adjacent monomer (~16 Å), thus aligning the two prominent extracellular bends in each loop critical to pattern 2 rAb binding (Fig. 9D).
The second feature is the location of loop E at the periphery of each monomer subunit, possibly providing access for rAb recognition of the pocket created by loops C and E (Fig. 9D).

The epitopes identified using NMO rAbs show some similarities to those identified previously using serum NMO-IgG, including the absence of binding to denatured AQP4 by protein...
Binding Patterns of AQP4-IgG

immunoblotting and the enhanced staining of AQP4 in cells forming supramolecular OAPs (22). Select mutagenesis of some human AQP4 extracellular amino acids based on their sequence homologies to other OAP-forming aquaporins also revealed two major conformational epitopes that partially overlapped with the critical residues differentiating both pattern 1 (loops C- and E-dependent) and 2 (loops A-, C-, and E-dependent) binding (12); they include the serial mutation of amino acids 61GTEK64 in loop A, 146GVTMV150 in loop C, and 227WE228 in loop E. Using our panel of NMO monoclonal rAbs, we have identified novel amino acids critical for the binding of all (137TP138 and Val150), most (Asn153), or subsets (Val141, His151, and Leu154) of AQP4-specific rAbs. Previous studies revealed consistently greater affinity of NMO-IgG binding to M23 than M1 AQP4 (10), and Fab fragments of NMO rAbs showed a similar preference of AQP4 isoform binding, indicating that structural changes in AQP4 upon array assembly, and not bivalent cross-linking of whole IgG, result in the greater binding affinity to OAPs. Whether AQP4 OAPs create novel surface epitopes or stabilize conformational epitopes on AQP4 tetramers remains to be determined.

In conclusion, NMO CSF-derived AQP4-specific B cells show distinct patterns of antigen recognition that can be classified into two broad categories based on their differential dependence on select loop A amino acids. Localization of the critical amino acids in the AQP4 crystal structure identifies pockets or domains both between and within monomers that align important loop A, loop C, and loop E amino acids and contribute to antigen recognition. These domains may be important for development of potential therapies designed to inhibit NMO-IgG interaction with AQP4 and subsequent CNS lesion formation.

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