MOG transmembrane and cytoplasmic domains contain highly stimulatory T-cell epitopes in MS

ABSTRACT

Objective: Recently, we reported that the 218 amino acid murine full-length myelin oligodendrocyte glycoprotein (MOG) contains novel T-cell epitopes p119-132, p181-195, and p186-200, located within its transmembrane and cytoplasmic domains, and that p119-132 is its immunodominant encephalitogenic T-cell epitope in mice. Here, we investigated whether the corresponding human MOG sequences contain T-cell epitopes in patients with multiple sclerosis (MS) and healthy controls (HC).

Methods: Peripheral blood T cells from patients with MS and HC were examined for proliferation to MOG p119-130, p181-195, p186-200, and p35-55 by fluorescence-activated cell sorting analysis using carboxyfluorescein diacetate succinimidyl ester dilution assay. Intracellular production of proinflammatory cytokines was analyzed by flow cytometry.

Results: MOG p119-130, p181-195, and p186-200 elicited significantly greater T-cell responses than p35-55 in patients with MS. T cells from patients with MS proliferated significantly more strongly to MOG p119-130 and p186-200 than did T cells from HC. Further, MOG p119-130-specific T cells exhibited Th17 polarization, suggesting this T-cell epitope may be relevant to MS pathogenesis.

Conclusions: Transmembrane and cytoplasmic MOG domains contain potent T-cell epitopes in MS. Recognition of these determinants is important when evaluating T-cell responses to MOG in MS and may have implications for development of myelin antigen-based therapeutics. Neurology.org/nn © 2014 American Academy of Neurology

GLOSSARY

aa = amino acid(s); AQP4 = aquaporin-4; CFSE = carboxyfluorescein diacetate succinimidyl ester; EAE = experimental autoimmune encephalitis; HC = healthy controls; Ig = immunoglobulin; MBP = myelin basic protein; MHC = major histocompatibility complex; MOG = myelin oligodendrocyte glycoprotein; MS = multiple sclerosis; NMO = neuromyelitis optica; PLP = proteolipid protein; PBMC = peripheral blood mononuclear cells; UCSF = University of California at San Francisco.

Evidence indicates that T cells specific for myelin autoantigens have an important role in the pathogenesis of multiple sclerosis (MS).1 Although several myelin antigens exist, investigations of T-cell reactivity in MS have focused attention on myelin basic protein (MBP) and proteolipid protein (PLP), which account for approximately 80% of myelin protein,2 as well as myelin oligodendrocyte glycoprotein (MOG). Although intact MOG protein accounts for only 0.05%–0.1% of total myelin proteins,2 it was initially reported to induce more potent T-cell responses than other myelin antigens in patients with MS.3 Subsequent studies in MS have concentrated primarily on T-cell recognition of the 117 amino acid (aa) N-terminal extracellular immunoglobulin (Ig) “variable-like” domain of MOG but have not consistently observed T-cell responses.4 However, native full-length MOG is 218 aa and contains transmembrane and cytoplasmic domains.5 In our companion manuscript, we identified 3 T-cell MOG determinants in mice, MOG p119-132, located within the transmembrane region, which induced potent clinical and histologic experimental autoimmune encephalitis (EAE), and MOG p181-195 and p186-200,

See companion article
2 discrete T-cell epitopes within the cytoplasmic domain. Upon recall to immunization of mice with full-length MOG, these T-cell epitopes were recognized more frequently than MOG p35-55, indicating they are dominant epitopes. We therefore questioned whether T cells in patients with MS recognize the corresponding peptide sequences of human MOG.

**METHODS** Patients. Twelve Caucasian patients with MS (66% female, mean age [SD]: 43.2 [12.9] years, mean disease duration [SD]: 5.5 [6.2] years, mean Expanded Disability Status Scale score [SD]: 1.8 [1.0]) and 12 Caucasian healthy controls (HC) (42% female, mean age [SD]: 40.5 [8.8] years) were recruited from the University of California at San Francisco (UCSF) MS Center. Out of 12 patients with MS, 10 had not received disease-modifying therapies prior to the study and 2 were treated with rituximab. Statistical significance between patients with MS and HC occurred regardless of whether the 2 patients treated with rituximab were included in T-cell proliferation and cytokine analysis. None of the patients had received steroids within 2 months preceding blood draws. Blood was collected by venipuncture.

Standard protocol approvals, registrations, and patient consents. This study was approved by the UCSF Committee on Human Research (Protocol 10-00650). Written informed consent was obtained from participants prior to enrollment.

Peptides. Human MOG p119-130 (FYWVSPGVLYLL), MOG p181-195 (TLFIVYPVLGPLVL), and p186-200 (VPVPLGPLVALICYN) were synthesized by Genemed Synthesis Inc. (San Antonio, TX). Human MOG p119-130 (FYWVSPGVLYLL), p181-195, and p186-200 were purchased from AnaSpec, Inc. (Fremont, CA).

Lymphocyte culture and proliferation assay. Human peripheral blood mononuclear cells (PBMC) were isolated by density gradient centrifugation over Ficoll (Ficoll-Paque PLUS; GE Healthcare, Milwaukee, WI). T-cell proliferation was evaluated as previously described. PBMC were stained with 0.5 μM 5,6-carboxyfluorescein diacetate succinimidyl ester (CFSE) (Invitrogen, Carlsbad, CA). After 10 days of culture with antigens, T-cell proliferation was examined by flow cytometric evaluation of CFSE dilution. Proliferation was expressed as the cell division index (defined as the number of CFSE<sup>−</sup> T cells cultured with antigen/number of CFSE<sup>+</sup> T cells without antigen). Culture medium consisted of X-VIVO 15 (Lonza, Walkersville, MD) supplemented with penicillin (100 U/mL) and streptomycin (0.1 mg/mL), Mouse monoclonal anti–HLA-DR (clone G46-6; BD Biosciences, San Jose, CA), anti–HLA-DQ (clone HG-38; Abcam, Cambridge, MA), anti–HLA-DP (clone B7/21; Abcam), and isotype control (clone G155-178; BD Biosciences) were used to evaluate inhibition of T-cell proliferation. Antibodies were added to CFSE-stained PBMC cultures 1 hour prior to antigen addition.

Flow cytometry analysis. Human single-cell suspensions were stained with antibodies against CD3 and CD4 (eBioscience, San Diego, CA and BD Biosciences). Intracellular cytokine production by CD4<sup>+</sup> T cells was analyzed by monitoring the expression of interferon-γ and interleukin-17 (1:100; eBioscience) after phorbol 12-myristate 13-acetate (50 ng/mL) and ionomycin (500 ng/mL; both from Sigma–Aldrich, St. Louis, MO) stimulation for 5 hours in the presence of GolgiPlug (BD Biosciences). Data were acquired on a FACSCanto flow cytometer (BD Biosciences).

**RESULTS** Peripheral blood CD4<sup>+</sup> T cells from patients with MS and HC were examined for their capability to respond to human MOG peptides p119-130, p181-195, and p186-200. Proliferation was measured by CFSE dilution assay. As shown in figure 1A, CD4<sup>+</sup> T cells from untreated patients with MS and HC proliferated in response to human MOG p119-130, p181-195, and p186-200 more strongly than to MOG p35-55. Further, T-cell proliferation to p119-130, p181-195, and p186-200 was significantly greater than to p35-55 (p = 0.0051, p = 0.0304, and p = 0.0007, respectively) in patients with MS. Proliferative responses to MOG p119-130 and p186-200 were most robust, and, in comparison, T cells from patients with MS proliferated significantly more to these determinants than T cells from HC. In contrast, no significant differences in proliferation to p181-195 were observed between patients with MS and HC.

In MS, the pathogenic involvement of CD4<sup>+</sup> T cells is supported by the association between disease susceptibility and major histocompatibility complex (MHC)-II genes. Using MHC II–blocking antibodies, we observed that T-cell proliferation to MOG p119-130 and p186-200 was inhibited by anti–HLA-DR but was not significantly inhibited by anti–HLA-DQ or anti–HLA-DP. These results demonstrated that HLA-DR molecules serve as restriction elements for MOG peptides. The frequency of Th17 cells and ratio of Th17 to Th1 cells were greater in response to MOG p119-130– and p186-200–specific T cells (figure 1B). Th1 and Th17 cells represent 2 proinflammatory T-cell subsets that may participate in MS pathogenesis. Thus, we examined proliferating MOG peptide–specific T cells for production of proinflammatory cytokines. The frequency of Th17 cells and ratio of Th17 to Th1 cells were greater in response to MOG p119-130 than to the other MOG T-cell determinants examined (figure 2). Further, Th17 p119-130–specific T cells were statistically more frequent in patients with MS than HC. In contrast, the frequency of MOG p186-200–specific Th1 cells appeared to be higher in patients with MS than HC, but it did not reach statistical significance. Collectively, these results clearly demonstrate that the T-cell determinants within the transmembrane and cytoplasmic domains of MOG, shared in mice and humans, are highly stimulatory in patients with MS.

**DISCUSSION** In 1993, when MOG protein was discovered as a CNS autoantigen, it was reported to induce more potent T-cell responses than MBP, PLP, or myelin-associated glycoprotein (MAG). However,
most subsequent studies utilized peptides corresponding to the extracellular MOG domain (MOG1-125) or recombinant MOG1-125 itself and did not consistently observe T-cell reactivity. These findings raised the possibility that T-cell epitopes exist in the other domains. In contrast to many studies that evaluated T-cell proliferation to MOG in MS, we utilized the CFSE dilution assay, which is a sensitive and robust method for measuring
In our study, the 3 transmembrane or cytoplasmic determinants examined elicited stronger T-cell proliferative responses than human MOG p35-55. These observations suggest that MOG p119-130, identified in our accompanying manuscript as an encephalitogenic murine T-cell determinant, may be pathologically relevant in both EAE and MS.

Of interest, a small number of reports that did evaluate T-cell reactivity to peptides corresponding to the intracellular domain of MOG observed similar, or lower, responses in patients with MS than HC. Of note, MOG-specific IgG antibodies may define a subset of AQP4-seronegative patients with NMO. Two recent studies suggested that MOG-specific IgG antibodies may define a subset of aquaporin-4 (AQP4)-seronegative patients with neuromyelitis optica (NMO), considered an astrocytopathy. IgG is a T-cell-dependent antibody subclass.

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Of interest, a small number of reports that did evaluate T-cell reactivity to peptides corresponding to the intracellular domain of MOG observed similar, or lower, responses in patients with MS than HC. Here, we observed not only that MOG p119-130 and p186-200 elicited more vigorous T-cell responses than p35-55 but also that these 2 novel T-cell epitopes elicited stronger proliferative responses in patients with MS than HC. In addition, p119-130–specific T cells exhibited significant proinflammatory Th17 polarization. Our demonstration that immunodominant T-cell epitopes exist within the C-terminal 100 residues of intact MOG in mice and that they correspond to determinants in MS should encourage reevaluation of T-cell recognition of MOG in patients with MS.

Two recent studies suggested that MOG-specific IgG antibodies may define a subset of aquaporin-4 (AQP4)-seronegative patients with neuromyelitis optica (NMO), considered an astrocytopathy. IgG is a T-cell-dependent antibody subclass. While it should be recognized that astroglial histopathology is necessary...
to confirm that this entity indeed represents NMO, it may also be important to evaluate T-cell responses to the immunodominant determinants of MOG in patients with NMO. It is interesting that a murine model containing T cells and B cells that selectively target MOG, a protein expressed on oligodendrocytes and not astrocytes, leads to opticospinal EAE.12 Thus, one can speculate that MOG IgG+ AQP4-seronegative NMO could represent a unique form of opticospinal MS.

While myelin antigen-based tolerigenic therapies, including altered peptide ligands, DNA vaccination, intravenous peptide administration, and oral tolerance, have not clearly proven efficacious in phase II and III MS trials, interest in development of therapies that target autoantigen-specific immune cells remains high.13 Of interest, a majority of such studies have tested therapies targeting immune recognition of MBP alone,13 and those investigations that did include MOG concentrated on T-cell epitopes within its extracellular domain.14 Our observations that the transmembrane and cytoplasmic domains of MOG contain highly stimulatory T-cell epitopes in patients with MS underscore the importance of including these domains should myelin antigen-based therapies be further advanced in MS. Further, recognition of these T-cell epitopes of MOG is clearly important when evaluating how established and novel MS therapeutic influence T-cell recognition of CNS autoantigens.

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