Amyloid-like Behavior of Site-Specifically Citrullinated Myelin Oligodendrocyte Protein (MOG) Peptide Fragments inside EBV-Infected B-Cells Influences Their Cytotoxicity and Autoimmunogenicity

Can Araman,*† Miriam E. van Gent,‡ Nico J. Meeuwenoord,‡ Nicole Heijmans,§ Mikkel H. S. Marqvorsen,‖ Ward Doelman,† Bart W. Faber,† Bert A. ’t Hart,*§⊥ and Sander I. Van Kasteren*⊥

1Leiden Institute of Chemistry and Institute for Chemical Immunology, Leiden University, Einsteinweg 55, 2333 CC Leiden, The Netherlands
2Leiden Institute of Chemistry and Department of Bioorganic Synthesis, Leiden University, Einsteinweg 55, 2333 CC Leiden, The Netherlands
3Department of Immunobiology, Biomedical Primate Research Centre, 2288 GJ Rijswijk, The Netherlands
4Department of Parasitology, Biomedical Primate Research Centre, 2288 GJ Rijswijk, The Netherlands
5Department of Neuroscience, University of Groningen, University Medical Centre, 9700 AB Groningen, The Netherlands

Supporting Information

ABSTRACT: Multiple sclerosis (MS) is an autoimmune disorder manifested via chronic inflammation, demyelination, and neurodegeneration inside the central nervous system. The progressive phase of MS is characterized by neurodegeneration, but unlike classical neurodegenerative diseases, amyloid-like aggregation of self-proteins has not been documented. There is evidence that citrullination protects an immunodominant peptide of human myelin oligodendrocyte glycoprotein (MOG35-56) against destructive processing in Epstein-Barr virus-infected B-lymphocytes (EBV-BLCs) in marmosets and causes exacerbation of ongoing MS-like encephalopathies in mice. Here we collected evidence that citrullination of MOG can also lead to amyloid-like behavior shifting the disease pathogenesis toward neurodegeneration. We observed that an immunodominant MOG peptide, MOG35-56, displays amyloid-like behavior upon site-specific citrullination at positions 41, 46, and/or 52. These amyloid aggregates are shown to be toxic to the EBV-BLCs and to dendritic cells at concentrations favored for antigen presentation, suggesting a role of amyloid-like aggregation in the pathogenesis of progressive MS.

Multiple sclerosis (MS) is a group of autoimmune-driven neuroinflammatory disorders that are pathologically characterized by myelin sheath loss (demyelination) and axonal damage. The disease starts in ±85% of the patients with alternating episodes of neurological defects (relapse) and recovery (remission), so-called relapsing–remitting MS (RRMS), which in ±60% of MS patients converts to secondary progressive disease with chronic progression. Approximately 10–15% of MS patients show chronic progression from disease onset, a course called primary progressive multiple sclerosis (PPMS). While there are several therapies available for the treatment of RRMS, few therapeutic options yet exist for PPMS. This is due to their fundamentally different pathophysiology. Recently, anti-B-cell antibody therapy was approved for the treatment of PPMS, which ties in with the hypothesis that—aside from genetic predispositions and well-characterized environmental factors such as smoking and sun exposure (vitamin D)—viral infections play a role in MS pathogenesis, especially in PPMS. In particular, Epstein-Barr virus (EBV) infection, which has a selective tropism for B-lymphocytes (BLCs), has been considered to be a prerequisite for MS pathogenesis. One of the presumed effects of viral infection upon MS emergence is the qualitative alteration of the protein composition of myelin, especially myelin oligodendrocyte glycoprotein (MOG). Studies in mouse and non-human primate models of experimental autoimmune encephalomyelitis (EAE) revealed that MOG is an essential myelin component for the experimental induction of T- and B-cell autoimmunity in chronic progressive disease. Moreover,
MOG is expressed as a homodimer on the surface of oligodendrocytes, the myelin-forming glial cells of the central nervous system (CNS), and on the outermost lamellae of the myelin sheaths that wrap around axons, forming a protective layer that is essential for fast pulse conduction and trophic support of energy-demanding axons. The exact biological function of MOG is not known, but it is evident that the N-linked glycan at position 31 interacts with the C-type lectin receptor DC-SIGN, which is expressed on microglial cells and on antigen-presenting cells within the brain-draining cervical lymph nodes (CLNs). As ligands of DC-SIGN are known to suppress maturation of dendritic cells to a full immunogenic state, we previously posited that MOG may have a role in the avoidance of autoreactive T-cell activation and neuroinflammation.

In addition to the findings described above, there is solid evidence for a link between the chronic inflammation via the production of reactive oxygen/nitrogen species (ROS/RNS) and the dysregulation of the ionic balance, especially due to an increase in intracellular Ca2+ ion concentrations. The latter event in turn upregulates the activation of peptidyl arginine deiminases (PADs), which then convert arginines (Arg) into citrullines (Cit). Citrullinated self-proteins have been shown to become antigenic, as described by Toes and co-workers (Arg 41 and 46, respectively). Moreover, myelin basic protein (MBP), a major component of the myelin sheath, has been found to be progressively hypercitrullinated (~45% citrullinated MBP in MS and ~100% in the acute neuroinflammatory disorder Marburg’s disease). More recently, Woodroffe and colleagues showed that glial fibrillary acidic protein (GFAP) can be detected as an additional, aberrantly citrullinated protein in MS. Finally, the MS relevant murine T-cell epitope of MOG, MOG35-55, has been reported to exacerbate EAE in mice if citrullinated at position 41, a contact residue for T-cell receptors (TCRs). Moreover, in this study, an exacerbation of disease progression was observed when T-cells specific for citrullinated MOG35-55 were transferred to mice with ongoing, native peptide-induced EAE. Taken together, these data suggested a critical role for citrullination of MOG and other components of the myelin sheath in MS progression.

One essential question that remains unanswered is whether neurodegeneration or peripheral inflammation is the primary cause of PPMS, as neuronal atrophy occurs away from immune cell-containing zones in this disease. Neuronal damage may therefore not be due to direct immune cell–sheath contact. As many other neurodegenerative diseases are characterized by post-translational modification (PTM) of proteins leading to aggregation and accumulation of aggregates within brain tissue (e.g., in transmissible spongiform encephalopathies (TSEs); prion protein, prP), Alzheimer’s disease (amyloid-β and tau protein), and Parkinson’s disease (α-synuclein)], we hypothesized that PPMS may also have an aggregation-induced component to its disease pathology. However, unlike for these classical neurodegenerative diseases, a misfolded autoantigen has not been documented for progressive MS. Comparing the structures of misfolded prP models with the β-sheet rich crystal structure of native MOG protein (depicted in Figure 1), we hypothesized that MOG or a PTM variant of MOG could be the species forming amyloid fibrils in PPMS due to potential structural changes (i.e., conversion from β-sheet to cross-β-sheet) upon citrullination, which are ordered structures of misfolding proteins usually described by seeding nucleation or nucleation-dependent polymerization models.

Here we test the hypothesis that PPMS carries an amyloid component by assessing the aggregation characteristics of MOG as well as the key MOG-derived immunodominant peptides containing the relevant T-cell epitopes in vitro. These characteristics were investigated in vitro using a thioflavin T (ThT) assay and in vivo using a novel bioorthogonal chemistry-based approach. We found that the recombinant MOG protein was non-immunogenic but that the peptide fragments indeed displayed amyloid behavior (Figure 1). Moreover, they displayed this behavior only upon citrullination and were shown to be toxic to phagocytes, such as bone marrow-derived dendritic cells (BMDCs) and the BLCs in which they were generated. Furthermore, amyloid-like aggregation altered

![Figure 1. Strategy and proposed mechanism for amyloid aggregation-driven cytotoxicity of MOG.](image)
the antigenicity of citrullinated MOG peptides in a concentration-dependent manner, resulting in high cytotoxicity at physiological concentrations. This study provides the first evidence that amyloid fibril formation may play a role in PPMS pathogenesis and disease progression.

**MATERIALS AND METHODS**

**Fmoc SPPS, Peptide Analysis, and Purification.** Peptides were prepared either manually or on automated synthesizer (PTI Tribute UV-IR synthesizer, Gyros Protein Technologies). For manual synthesis, Fmoc deprotection was achieved with 20% piperidine in dimethylformamide (DMF) using two cycles of 3 and 7 min. All amino acids (2.5 equiv) were coupled using HCU (2.38 equiv) and DIEPA (5 equiv) for 30 mins, unless stated otherwise. For automated peptide synthesis, 5 equiv of each amino acid (aa) on a 100–200 μmol scale and 60 equiv of each aa on a 50 μmol scale with respect to the resin loading were used. An equimolar quantity of HCU was used as an activator. Coupling cycles of 1 h were utilized, and unreacted amines were capped after each cycle using a solution of 500 μL of acetic anhydride, 250 μL of DIPEA, and 4.25 mL of DMF for 5 min at room temperature twice. Peptides were globally deprotected and cleaved from dried resin with a mixture of TFA, trisopropylsilane (TIS), and water [92.5:5:2.5 (v/v)] for at least 3 h at room temperature. Precipitation of crude peptides was achieved by addition of cold diethyl ether. Isolated peptides were lyophilized overnight. Peptides were characterized using electrospray ionization mass spectrometry (ESI-MS) on a Thermo Finnigan LCQ Advantage Max LC-MS instrument with a Surveyor PDA plus UV detector or an Agilent 6120 Thermo Finnigan LCQ Advantage Max LC-MS instrument. On both systems, a Gemini-NX C18 column (Phenomenex, 3 μm, 110 Å, 50 mm × 4.6 mm) was used. Peptides were analyzed using a gradient from 10 to 90% or from 10 to 50% of B over 10–20 min (eluent A, H2O; eluent B, acetonitrile; eluent C, 1% TFA in H2O; eluent D, methanol). Peptide masses were calculated from the experimental mass to charge (m/z) ratios from all of the observed protonation states of the peptides using the Xcalibur Qual Browser or LC-MS_6120B software. Lyophilized peptides were dissolved in a mixture of tert-butanol, acetonitrile (ACN), and water [1:1:1 (v/v)] unless stated otherwise and purified using a preparative HPLC system (Gilson Inc.) with a Gemini-NX C18 semipreparative column (Phenomenex, 5 μm, 110 Å, 250 mm × 10.0 mm) with a linear gradient from 20 to 30% B in 30 min (eluent A, 0.2% TFA in H2O; eluent B, ACN). Fractions containing the peptides were pooled and lyophilized overnight.

**Biophysical Characterization. Circular Dichroism (CD) Spectroscopy.** Circular dichroism spectroscopy was performed at room temperature using a Jasco J-815 CD spectrometer with a 1 mm path-length cell and a bandwidth of 2.0 nm. The proteins were prepared either in 20 mM Tris buffer (pH 7.5) or in 20 mM NaOAc buffer (pH 5.0). Further analysis was performed via addition of 2–4 mM SDS (β-sheet enhancer) or 50% TFE (α-helix enhancer) with a final peptide concentration of 0.1–0.2 mg/mL. Spectra were recorded from 260 to 190 nm at an interval of 1 nm. Each spectrum was the average of five scans and blank subtraction.

**ThT Fluorescence Aggregation Assay.** Aggregation assays were modified from those described by Araman et al. The experiments were performed in 96-well plates at 37 °C using the Infinite M1000 Pro Tecan plate reader with an excitation wavelength of 444 nm and an emission wavelength of 485 nm with a bandwidth of 10 nm. A mixture of 199 μL of peptide (200 μM to 2 μM) and 1 μL of ThT was measured in each well at a 20000 μm z-position over time with a kinetic interval of 10 min.

**Transmission Electron Microscopy (TEM).** TEM imaging was conducted on a JEOL 1010 instrument operated with an accelerating voltage of 70 kV. Samples were diluted to final concentrations of 1–5 μM. A single drop (approximately 1 μL) was applied on the grid (PELCO Center-Marked Grids, 75 mesh, 3.0 mm outside diameter, copper) and drained with the corner of a tissue. The grid was left to dry for 1–3 h prior to measurement.

**Dynamic Light Scattering (DLS).** The hydrodynamic diameter (R_int) of amyloid fibrils of MOG peptides 5 and 7 was assessed by dynamic light scattering (DLS) spectroscopy on a Zetasizer Nano S (Malvern Instruments, Malvern, U.K.) using PMMA small-volume cuvettes (VWR international, Leuven, Belgium). Hereby, the working concentration was ~20 μM in 20 mM NaOAc (pH 5).

**Cell Biology. B3Z T-Cell Hybridoma Culture.** Mouse B3Z T-cell hybridomas were cultured as described previously.

**B-Cell Culture.** Human EBV-infected lymphoblastoid cell line BSM (ECACC 88052032) was purchased from Sigma-Aldrich and cultured in polystyrene tissue culture flasks in RPMI 1640 (GIBCO), 2 mM glutamine (GIBCO, catalog no. 35050-038), 10% FCS (TIC Europ, catalog no. 61024-C05), and 100 units/mL penicillin/streptomycin (GIBCO, catalog no. 15140-122) in a 5% CO2 atmosphere at 37 °C and diluted every 3–4 days at a concentration of ~0.5 × 106 cells/mL.

**Bone Marrow Dendritic Cell Culture.** Bone marrow (BM) was isolated from 8–12-week-old C57BL/6 mice (strain C57Bl/6NHsd, H-2b haplotype; Envigo Inc., Huntingdon, U.K.) as described previously with some modifications; 8–12-week-old mice were kept under specific pathogen-free (SPF) conditions in our own facilities. Femurs and tibiae of female or male mice were removed and cleaned from the surrounding muscle tissue. Then, intact bones were left in 70% ethanol for 2–5 min for disinfection and washed with sterile phosphate-buffered saline (PBS). Next, both ends were cut with scissors, and bone marrow was flushed from femurs and tibiae with prewarmed IMDM (Sigma) using a 20 mL syringe with a 0.45 mm × 23 mm hypodermic needle (G26, Terumo Europe, NN-2623R) through a 70 μm cell strainer (Falcon). Isolated BM cells were centrifuged for 5 min at 350g and resuspended in 10 mL of IMDM supplemented with 8% heat-inactivated fetal calf serum (FCS, Sigma), 2 mM Glutamax (GIBCO), 20 μM 2-mercaptoethanol (Gibco), 50 IU/mL penicillin, and 50 μg/mL streptomycin in the presence of 20 mg/mL rmGM-CSF (Peprotech). The cell suspension was diluted to ~0.5–1 × 106 cells/mL in complete medium, and cells were incubated in non-adhesive Petri dishes (Sarstedt) at 37 °C and 5% CO2 for 6–7 days.

**Cell Viability Assay with 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium Bromide (MTT).** Cell viability of BMDCs and B3Zs was tested via the MTT assay. For this purpose, the following protocols were used. BMDCs and B3Zs were seeded in a 96-well plate at a density of 2.5 × 104 cells/well and treated with varying amounts of peptides 1, 5, and 7 in IMDM (20, 10, 5, and 3 μM) incubated overnight at 37 °C and 5% CO2. Medium with (control) and medium without 2.5 × 104 cells (blank) were used as controls. After incubation for
24 h, cells were centrifuged (350g for 5 min at 4 °C), the supernatant was discarded, and 100 μL of 0.5 mg/mL MTT [diluted 1:10 (v/v) from a 5 mg/mL MTT stock solution in PBS] was added to each well. Upon incubation for 3–4 h at 37 °C and formation of intracellular formazan crystals, the supernatant was removed and the solubilizing solution (DMSO, 100 μL) was added. The plate was incubated for a further 30 min at 37 °C, and the absorbance was measured at 540 nm (A540) as well as 570 nm (Tecan M1000 plate reader). The following equation was used to assess cell viability:

\[
\% \text{ cell viability} = \frac{A_{540}(\text{sample}) - A_{540}(\text{blank})}{A_{540}(\text{control}) - A_{540}(\text{blank})}
\]

**T-Cell Culture Assay.** Mononuclear cells (MNC) isolated from axillary lymph nodes (ALN) were obtained from two adult common marmosets (*Callithrix jacchus*) that had been enrolled in earlier EAE studies. The cells have been stored deep-frozen directly after being isolated. All marmosets were acquired from the purpose-bred colony at the Biomedical Primate Research Centre and declared in good health by veterinary staff prior to enrollment in the study. Health checks included physical examination and tests for hematological, serological, and microbiological abnormalities. The used EAE model was induced with a synthetic peptide representing residues 34–56 of human myelin oligodendrocyte glycoprotein (MOG34–56; Cambridge Research Biochemicals Ltd., Cleveland, U.K.; sequence of GMEVGWYRPPFSRVVHLYR-56; Cambridge Research Biochemicals Ltd., Cleve- land, U.K.; sequence of GMEVGWYRPPFSRVVHLYR-NGKD) emulsified with IFA (Difco Laboratories, Detroit, MI) as previously described.68 As antigen-presenting cells, EBV-infected marmoset B-lymphoblastoid cells (BLCs) were used. BLC lines were generated by infection of blood MNC with human Epstein-Barr virus propagated in marmoset cell line B95-8.70 Briefly, PBMCs were cultured with supernatants from the EBV-producing B95-8 cell line for 1.5 h at 37 °C. Following incubation, cells were diluted at a 1:1 ratio with RPMI-1640 (GIBCO) containing 1 μg/mL phytohemeagglutinin (PHA) and supplemented with 10% fetal calf serum (FCS), 1-α-α-α-α-α-α-glutamine (GIBCO), and penicillin/streptomycin as antibiotics. Following successful transformation, cells were maintained in culture medium without PHA until further use. For proliferation experiments, marmoset EBV-BLCs were maintained in culture medium without PHA until further use. For proliferation experiments, marmoset EBV-BLCs were maintained in culture medium without PHA until further use. For proliferation experiments, marmoset EBV-BLCs were maintained in culture medium without PHA until further use.

**RESULTS**

Citrullination of MOG35–55 Peptides Alters Their Structural Behavior. We first analyzed the aggregation properties of the extracellular portion of recombinant MOG (aa 1–125, hereafter termed rMOG) and its pericellullated counterpart, cit-rMOG, using a ThT assay. ThT is a fluorogenic dye, which becomes fluorescent upon binding to cross-β-sheet structures, such as those found in amyloid aggregates.34 In brief, proteins were dissolved in 20 mM NaOAc buffer (pH 5.0). Upon addition of ThT and incubation under gentle agitation, the fluorescence emission was recorded at 485 nm over a period of at least 15 h to monitor the propensity of the proteins toward amyloid-like aggregation. rMOG and cit-rMOG were assessed for aggregation under these conditions, but neither protein showed any aggregation.
(Figure S1). We thus hypothesized that it was not rMOG that was the aggregating species but a fragment produced by proteolytic processing. We focused on the immuno-dominant peptide 35–55 and synthesized eight peptides spanning aa
35–55 of the native MOG sequence carrying TCR contact residues (aa 41–48) and site-specific citrulline residues at the respective (aa 41, 46, and/or 52) positions (Scheme 1 and Table 1, 1–8). Elongated variants (aa 31–55) were also prepared, as were bioorthogonal variants carrying propargylglycine (Pg) as a bioorthogonal handle at position 31 that would allow in-cell visualization (Scheme 1 and Table 1, 9–12). All peptides were synthesized in good yields (5.3–34.9%) and very good purity (>95%; determined by integrating the peak areas on analytical HPLC measurements (Table 1)). All MOG35–55 peptides were analyzed by liquid chromatography and mass spectrometry (LC–MS) (Figures S2–S9) and
circular dichroism (CD) spectroscopy (Figure 2a–c and Figure S10). In our hands, in the presence of SDS, nonmicellar \( \alpha \)-helical structure upon addition of 50% 2,2,2-trifluoroethanol (TFE), a known stabilizer of \( \alpha \)-helical conformations in proteins and peptides\(^{36} \) (Figure 2a–c and Figure S10). In our hands, in the presence of SDS, nonmicellar \( \beta \)-sheet enhancing \( \beta \)-sheet rich structures (Figure 2b and Figure S10C). Peptides 2, 3, 6, and 7, however, seemed to adopt a random coil structure under the same conditions (Figure S10A,B,D for 2, 3, and 6 and Figure 2c for 7). Moreover, 5 and 8 adopted \( \alpha \)-helical structures. This indicates that the peptides have distinct structures depending on the external solution conditions.

**Citrullinated MOG Peptides Show Amyloid-like Aggregation.** Next, we asked whether the peptides were prone to amyloid-like aggregation. To investigate amyloid-like aggregation using the aforementioned ThT fluorescence assay, peptides (200 \( \mu M \), Figure 2 and Figure S11) were dissolved in buffers with a pH range of 3.0–7.0. Noncitrullinated peptide 1 did not show any aggregation at any pH or at any of the concentrations tested (Figure S11 and Figure 2d). Citrullinated peptides 2–7, on the other hand, showed pH-dependent aggregation characteristics with the strongest aggregation observed at pH 4.0 or 5.0, which is the pH range found within the B-cell lysosomal pathway (Figure S11). At all other pH values tested, either no increase or only a slight (<2-fold) increase in fluorescence intensity was observed, suggesting that the lysosome-like pH value of 4.0–5.0 was essential for amyloid-like behavior of the citrullinated MOG peptides. Note that peptide 8 showed aggregation behavior different from those of all of the other peptides tested aggregating at pH 6 (~3-fold compared to pH 5) and pH 7 (~4-fold compared to pH 5) the fastest. On the basis of the results presented above, we subsequently analyzed the concentrations at which amyloid-like aggregation was observed for the citrullinated MOG\(^{35–55} \) variants 2–8. Each of the citrullinated peptides displayed distinct aggregation behavior, underlining the hypothesis that specific arginines are responsible for characteristics of amyloid aggregation for MOG\(^{35–55} \). Peptide 2 aggregated only at concentrations of >4 \( \mu M \) (Figure S12A). Peptides 3, 5, and 8 on the other hand did aggregate at concentrations of >4 \( \mu M \) but not at a final concentration of 1 \( \mu M \) (Figure S12B,E and Figure 2). Peptides 6 and 7 showed aggregation at all concentrations tested (Figure S12D and Figure 2f). Peptide aggregation at lower concentrations is regarded a hallmark of nucleation-dependent aggregation and is thus a strong indication to prove our hypothesis. The most prominent aggregation behavior at low concentrations (<10 \( \mu M \)) was observed with peptides 5 and 7 (plotted data, Figure 2g,h and Figure S25). Hence, these peptides were used for further analysis.

**Citrullinated Peptides Induce Seeded Nucleation and Show a Heterogeneous Fibril Morphology.** A further characteristic of amyloid behavior is the observation of seeded nucleation,\(^{37} \) in which amyloid precipitation can be accelerated by the addition of amyloid seeds. In the lag phase, only negligible amounts of self-propagated amyloid fibrils (also called seeds or nuclei) are formed, whereas upon their formation, an exponential increase in the level of fibril growth can be explored, which ends up reaching a plateau phase. Peptides 1 (Arg41, Arg46, and Arg52), 5 (Cit41, Cit46, and Arg52), and 7 (Arg41, Cit46, and Cit52) were therefore dissolized in 20 mM NaOAc (pH 5.0) at a concentration of 0.1

---

**Figure 3.** Seeded aggregation of peptides 5 and 7. Peptides 1, 5, and 7 were subjected to aggregation in the presence of seeds from either peptide 5 (depicted as “seed 5”) or 7 (depicted as “seed 7”) with different amounts (3, 30, and 300 ng in 200 \( \mu L \), depicted in corresponding diagrams in brackets). For aggregation-prone peptides 5 (a and b) and 7 (c and d), aggregation curves were fitted according to eq E1 in the Supporting Information (depicted as “fit”) to determine the change in \( t_{1/2} \) as a measure of aggregation kinetics. All data were recorded at an excitation wavelength of 444 ± 9 nm and an emission wavelength of 485 ± 9 nm. All samples were used at a pH of 5.0 and a concentration of 40 \( \mu M \).
mg/mL (40 μM) and incubated for at least 24 h at 37 °C under constant agitation to produce seeds for the experiment. We then separated the supernatant and pellet via centrifugation (20000g for 30 min) and analyzed by CD spectroscopy (Figure S13). The seeds were then homogenized in 20 mM NaOAc buffer (the same amount of buffer that was used for the incubation reaction) to an estimated final concentration of 0.1 mg/mL for further use in seeded nucleation assays.

The seed homogenate of peptide 5 formed a β-sheet-like structure, while the supernatant contained only random coil structures (Figure S13A). However, the minimum of the spectrum was observed at a wavelength (222 nm) higher than that normally observed for β-sheets (218 nm), which has also been observed for certain Aβ peptides38 and scrapie PrP (PrPSc).39 The seed homogenate of peptide 7 seems to reach a transition state-like conformation with characteristics of both α-helices and β-sheets (Figure S13B), and similar to that of 5, the peptide in the supernatant fraction of 7 was mainly random coil. As expected, no seeds for peptide 1 could be collected.

The collected seeds (3–300 ng per experiment) were then incubated with solutions of peptides 1, 5, and 7 (40 μM, 200 μL) to determine whether an acceleration of aggregation kinetics was observed (Figure 3 and Figure S14). This would confirm the amyloid characteristic of small amounts (sub-micrograms) of already aggregated species being able to act as seeds and drive faster amyloid aggregation by interacting with template peptides in solution. Indeed, faster aggregation kinetics for peptides 5 and 7 but not for 1 were observed. Sigmoidal curves were interpolated (eq E1 and Tables S1 and S2) and used to determine the time shift of aggregation at a relative intensity of 0.9-fold with all peptides starting to aggregate at this value. For seeded aggregation of peptide 1, under all conditions, no aggregation was observed (Figure S14A). For peptide 5, however, a positive time shift (thus faster aggregation) was observed. The most pronounced time shift was observed upon seeding with 0.3 μg of peptide 7 (Δt1/2 = 2.9 h (Figure 3b)). In addition, seeding with 0.03 μg of 5 resulted in significantly faster aggregation [Δt1/2 = 1.1 h (Figure 3a)], indicating a synergistic effect of both seeds with peptide 5.

Likewise, peptide 7 showed a prominent shift in time upon seeding with 0.003 μg of 5 (Figure 3c) (Δt1/2 = 3.5 h). Additionally, seeding with 0.03 μg of peptide 7 resulted in significantly faster aggregation [Δt1/2 = 1.2 h (Figure 3d)]. This indicates a synergistic effect of both seeds with 7. It is therefore likely that peptides 5 and 7 act as a substrate for either seed leading to faster aggregation kinetics. This kind of seeded aggregation is also observed for other amyloids, as has been shown for Aβ peptides, PrP, and α-syn.40-42 TEM was next used to obtain information about the morphology of the fibrils (Figure 4a,b). Both fibrils showed different morphologies. Peptide 7 formed needle-like, long (>100 nm) assemblies of fibrils with partially amorphous structures (Figure 4b), while fibrils of peptide 5 showed shorter (<100 nm) assemblies with tight packing corresponding to the higher susceptibility to aggregation over peptide 7 in vitro (Figure 4a). Additionally, DLS measurements were performed to determine the size of the fibrils (Figure 4c,d). The fibrils have hydrodynamic radii (Rhydration) of 78 ± 3.1 and 175 ± 2.5 nm for peptide 5 and 7 seeds, respectively. The Rhydration values are larger than expected for amyloid-like structures, which typically have a diameter of ~10 nm in an oligomeric state.43 Therefore, we concluded that the seeds of peptides 5 and 7 can be considered as mixed fibrils with amyloid-like structural characteristics as reported previously.44,45

**Citrullinated Peptides Aggregate inside BLCs.** We next asked whether this aggregation of citrullinated peptides also occurred within EBV-infected BLCs. To date, visualization of protein and peptide aggregation inside cells is a challenging task as fluorescence microscopy with thioflavin S (ThS), a derivative of ThT, is difficult due to aspecific binding to all amyloid proteins.46 Super-resolution fluorescence microscopy47 and cryo-electron microscopy48 have recently been used but are yet too complex for routine analysis. Having recently shown that bioorthogonal modifications affect peptide behavior only minimally inside antigen-presenting cells,49,50 we opted for a bioorthogonal approach to visualize MOG—peptide aggregation in the EBV-infected BLCs. Bioorthogonal reactions are chemoselective reactions in living systems, which do not interfere with other biochemical processes.51,52 Four new peptides that are N-terminally extended (aa 31−55) harboring an N→Pg mutation at position 31 were generated (Scheme 1 and Table 1, peptides 10−12). Position 31 was chosen to avoid any interference with the immunorelevant epitope 35−55 by using three additional amino acids of the native MOG sequence as linkers (aa 32−34). Aside from the traceable Pg, the peptides were synthesized without Cit (10) or double citrulline substitutions at critical positions (positions 41 and 46 and positions 46 and 52 for 11 and 12, respectively), and a nonbioorthogonal variant of noncitrullinated 10, peptide 9, was additionally generated. Neither 9 nor 10 aggregated (Figures S9, S17, and S20), and the aggregation behavior of 11 and 12 was not affected by the four-amino acid extension or by the presence of Pg compared to 5 and 7 (Figures S15−S20), confirming their suitability for tracing amyloid aggregation in cells. EBV-infected BLCs were therefore incubated under serum-free conditions with 10, 11, and 12 (0, 6.2, and 25 μM depicted in Figure 5; 3.1 and 12.5 μM depicted in Figures S21 and 22) for 48 h. After incubation, cells were centrifuged onto glass slides and fixed with PFA. The peptides were visualized

![Figure 4. Morphology of amyloid fibrils of 5 and 7.](image)
by the copper-catalyzed azide alkyne cycloaddition (CuAAC)53,54 reaction with Alexa647-azide (Thermo Fisher). Cells were co-stained with DAPI (blue channel) and LC3 (green channel), as this protein—which becomes post-translationally lipidated during autophagy as a marker of autophagic activity—is recruited to phagosomes containing aggregates (Figure 5). Peptides 11 and 12 in the absence of cells showed large aggregates, while peptide 10 had formed no aggregates. When peptides 11 and 12 were incubated with EBV-infected BLCs, spherical structures of different sizes (small spheres for 11 and large agglomerates for 12) were found, contained within LC3 positive vesicles, indicating intracellularly aggregated peptide. To our surprise, non-citrullinated peptide 10 also aggregated, once it was taken up by BLCs. This may be due to citrullination of peptide 10 by elevated levels of PAD enzymes present in EBV-infected B-cells.18 Earlier work from our group showed that EBV infection of B-cells increases the level of degradation of MOG35−55 in those cells. Thus, the aggregation behavior might not be restricted to B-cells but is likely a consequence of elevated PAD2 or PAD4 levels.71

Citrullination Converts Antigenic MOG Peptides into Cytotoxic Aggregates in Distinct Cell Types. We next tested whether amyloid aggregation of the peptides resulted in them becoming toxic to cells. We aimed to assess the toxicity of the peptides toward the core autoimmune process, which is the antigen-mediated cross-talk of EBV-infected BLCs with autoaggressive T-cells. In a first experiment, lethally irradiated marmoset EBV-BLCs (10³ per milliliter) were incubated for 1 h with titrating doses of peptides 10–12 or MOG14–36 as an irrelevant peptide (Figure 6a). Subsequently, a mixed lymphocyte population harvested from ALN of a marmoset immunized with MOG34–56 was added and cultured for 72 h. Proliferation of the lymphocytes was assayed via the incorporation of [³H]thymidine during the final 18 h. Figure 6a shows a dose-dependent stimulation of proliferation by the nonmodified peptide 10, while both citrullinated peptides (11 and 12) induced dose-dependent inhibition of proliferation even beneath the background response against antigens expressed by the EBV-BLCs. Peptide 10 showed stimulation of T-cell proliferation at all concentrations tested. To assess the targets of the citrullinated peptides, we stained the EBV-BLCs with the CellTrace dye, prior to the incubation with peptides and lymphocytes. After 24 h, cells were harvested, stained with Annexin V, a cellular probe binding phosphatidylserine, a marker of apoptotic and dead cells, and analyzed via fluorescence-activated cell sorting (FACS). Figure 6b shows that Annexin V staining, marking cytotoxicity, stains the CellTrace™ and CellTrace™+ fraction of cells incubated with the aggregation-prone peptides 11 and 12. As expected, for 11 and 12, an increased level of apoptosis was observed at concentrations of >1 μM/mL (Figure 6b, white circles), whereas peptide 10 did not show any cytotoxic effects in EBV-BLCs or in a co-culture with the lymphocytes.

Furthermore, the toxicity of MOG35−55 peptides 5 and 7 was assessed toward murine bone marrow-derived dendritic cells (mBMDCs), which have the capacity to phagocytose aggregates, and to the T-cell hybridoma cell line B3Z55 as a nonphagocytosing cell line (Figure 6c). Four different concentrations of peptides were tested (40, 20, 10, and 3 μM). Peptides 5 and 7 were toxic to mBMDCs [cell viability down to 70% for 5 and 62% for 7 (Figure 6c)] when compared to a nontreated control (Figure 6c, control). Both 5 and 7 showed cytotoxicity at concentrations of ≤10 μM. Additionally, 5 showed significant cytotoxicity at 40 μM. As expected, no decrease in cell viability was measured when B3Zs were incubated with increasing concentrations of peptides 5 and 7.

■ DISCUSSION

MOG has been accepted as a key autoantigen in MS. Nevertheless, the pathophysiological role of MOG remains complex. The response of T-cells to MOG epitopes depends on the genetic background of the species (human, marmoset, and mouse) as well as on environmental factors (especially

Figure 5. Monitoring the uptake and aggregation of bioorthogonal, site-specific citrullinated MOG peptides via confocal microscopy. Human EBV-infected BLCs were incubated for 48 h with either no peptide (0 μM) or 6.2 or 25 μM peptide 10, peptide 11, or peptide 12 (highlighted in yellow). Cells were fixed with 4% PFA and processed for immunofluorescence with the following primary antibodies. The nucleus was stained with DAPI (blue), and LC3 was used as an autophagosome marker (green). The bioorthogonal peptides were stained using CuAAC chemistry with azide Alexa-647 (Thermo Fisher). The scale bar is 25 μm (white bar).
EBV infection) and post-translational modifications (e.g., N-glycosylation and glucosylation). In the marmoset model, two key epitopes were identified, an MHC class II restricted epitope of inflammatory T-cells (MOG_{24-36}) and an MHC class Ib restricted epitope of cytotoxic T-cells (MOG_{40-48}). The latter pathway is particularly interesting for MS, as it leads to lesions in brain gray matter, which have the strongest impact on the disease.

A key finding underlining this complexity has been that the antigenicity of the MOG_{40-48} epitope is dependent on pathogenic post-translational citrullination. We have shown that citrullination of a peptide containing the MOG_{40-48} epitope at position 46 alters its proteolytic sensitivity to the endolysosomal serine protease cathepsin G in BLCs. Interestingly, it was shown in EBV-infected BLCs that expression levels of PAD2 and PAD4 are increased and that the autophagic flux is activated in infected cells, providing a protection mechanism for the peptide against rapid destruction. This mechanism putatively explains the requisite involvement of EBV-infected BLCs in the activation of lymphocytes that are not subjected to co-culturing as controls (black circles). Cultured cells were harvested and stained for Annexin V as a marker of late apoptotic/dead cells. The final analysis was done utilizing FACS. (c) Cytotoxicity assays with citrullinated MOG peptides in BMDCs. \( p < 0.05 \), \( ** p < 0.005 \), \( *** p < 0.0005 \), and \( **** p < 0.00005 \). n.s., not significant. The experiment was conducted twice and with three biological replicates. Group mean values were analyzed by one-way analysis of variance with the Bonferroni post hoc significant difference test using GraphPad Prism 6.0. Data are represented as means ± the standard deviation.

Figure 6. Cytotoxicity of MOG derived peptides (a and b) 10–12 and (c) 5 and 7 in EBV-BLC co-cultures with lymph node cells from EAE marmosets and in mBMDCs. (a) Marmoset EBV-induced BLCs were lethally irradiated and incubated for 1 h with titrating concentrations of peptide 10, 11, or 12 or an irrelevant peptide. Subsequently, lymph node or spleen cells from marmosets immunized with MOG_{34-56} were added. The responses of T-cells to the peptides were assayed by proliferation and are expressed as the stimulation index per culture condition. The experiment was conducted six times (marmosets) and with three biological replicates. Data are presented as means ± standard error of the mean. (b) To test which cell type is targeted by the peptides, EBV-BLCs (from two marmosets, M1 and M2) were incubated with CellTrace dye before incubation with peptide (white circles) and a mixture with the spleen/lymph node cells (red triangles). Lymphocytes that are not subjected to co-culturing were used as controls (black circles). Cultured cells were harvested and stained for Annexin V as a marker of late apoptotic/dead cells. The final analysis was done utilizing FACS. (c) Cytotoxicity assays with citrullinated MOG peptides in BMDCs. \( * p < 0.05 \), \( ** p < 0.005 \), \( *** p < 0.0005 \), and \( **** p < 0.00005 \). n.s., not significant. The experiment was conducted twice and with three biological replicates. Group mean values were analyzed by one-way analysis of variance with the Bonferroni post hoc significant difference test using GraphPad Prism 6.0. Data are represented as means ± the standard deviation.
autoreactive CD8+CD56+ T-cells, which drive disease progression in the marmoset EAE model.61 On the other hand, earlier in vivo experiments in mice showed that the two arginine residues in the MOG40–48 epitope are TRC contact residues of the pro-inflammatory CD4+ T-cells.23 The study presented here shows that this may also be the case in the marmoset EAE model, as T-cells from marmosets immunized with MOG34–56 are not stimulated (proliferation) by EBV-infected BLCs presenting citrullinated MOG34–56. In this study, we synthesized and characterized site-specifically citrullinated peptides as well as three bioorthogonal variants thereof to analyze processing and presentation of MOG-derived epitopes and their contribution to neurodegeneration. We showed that site-specific citrullination of MOG35–55 peptides resulted in amyloid-like aggregation behavior with formation of heterogeneous fibrils. This aggregation behavior was strongest at pH 5.0 (Figure S11), highlighting the potential involvement of intracellular vesicular proteolysis, like that found in the autophagy pathway.62 Moreover, above peptides aggregated at low concentrations (≤10 μM), the best and the fastest renders this concentration the critical aggregation concentration. One possible explanation for high concentrations (≥160 μM) not yielding to higher/faster aggregation susceptibility is that those are not within the critical aggregation concentration range and therefore might slow aggregation as shown for other proteins.63 Furthermore, Hevehan et al. reported refolding of aggregated lysozyme at high concentrations. These findings might explain the altered aggregation behavior of aforementioned MOG35–55 variants at high concentrations.64 Two doubly citrullinated peptides, 5 and 7, also showed characteristics of seeded nucleation,65 thereby supporting the hypothesis that these peptides display amyloid behavior. Collectively, these findings may shed new light on the role of EBV-infected BLCs and the immunodominant MOG34–56 peptide in the pathogenesis of MS and the role of EBV-infected BLCs and the immunodominant MOG34–56 peptide in the pathogenesis of MS.

This finding was further supported when using bioorthogonal variants of MOG35–55 in EBV-infected BLCs. Peptides were shown to aggregate within 48 h in these cells in a concentration-dependent manner (the higher the concentration, the larger the number of aggregates). Additionally, we could show that the autophagy pathway was stimulated, when using concentrations of ≤6.2 μM (Figure 5). We hypothesized that above this concentration, the aggregates are formed already before uptake in the cells and hence only a small portion of the peptides aggregate intracellularly and enter the autophagic pathway; instead, the aggregates may be taken up via alternative routes such as phagocytosis, endocytosis, or micropinocytosis as observed for amyloid β (Aβ)-40 and -42.65,66

To provide further evidence for a link between the amyloid behavior of these citrullinated MOG fragments and PPMS, the aggregates showed enhanced cytotoxicity toward phagocytes (EBV-BLCs and mBMDCs) in a concentration-dependent manner (≥10 μM for BLCs and ≤10 μM for mBMDCs).

In summary, we were able to show for the first time that citrullination is a protection mechanism against fast, physiological degradation of MOG peptides harboring T-cell epitopes and enhances antigenicity but promotes amyloid aggregation as a potential neurotoxic mechanism. These results provide evidence for the theory that a pathogenic mechanism initiated by EBV infection is one of the driving forces behind autoimmunity in MS. Our future work is directed toward live cell imaging amyloid-like aggregation, antigen processing, and presentation events in APCs incubated with MOG-derived peptides in marmoset EAE.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.8b00852.

Tables S1 and S2, eq E1, Figures S1–S25, and supporting materials (PDF)

**AUTHOR INFORMATION**

Corresponding Authors

*E-mail: m.caraman@lic.leidenuniv.nl.

*E-mail: hart@bprc.nl.

*E-mail: s.i.van.kasteren@chem.leidenuniv.nl.

**ORCID**

Can Araman: 0000-0002-6961-5607

Sander I. Van Kasteren: 0000-0003-3733-181X

**Funding**

C.A. was supported by The Netherlands Organization for Scientific Research (NWO)-CW-ECHO Grant program 2016. M.H.S.M. was supported by a Lundbeck Postdoc Abbroad grant. S.I.v.K. acknowledges the European Research Council (ERC-2014-StG-639005) for financial support.

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

The authors thank Sanne Vonk for the assistance with MTT assays and G. E. M. Lemmers for the technical support with TEM. Additionally, the authors acknowledge A. Kross for granting us the access to the JASCO J-815 CD spectrometer, the Infinite M1000 Pro Tecan plate reader, and the dynamic light scattering (DLS) spectrometer Zetasizer Nano S.

**REFERENCES**


DOI: 10.1021/acs.biochem.8b00852

Biochemistry 2019, 58, 763–775