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## CHAPTER 4

# **Characterisation of Immune Response to Neurofilament Light in Experimental Autoimmune Encephalomyelitis**

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## Abstract

Autoimmunity to neuronal proteins occurs in several neurological syndromes where cellular and humoral responses are directed to surface, as well as intracellular antigens. Similar to myelin autoimmunity, pathogenic immune responses to neuroaxonal components such as neurofilaments may contribute to neurodegeneration in multiple sclerosis. We studied the immune response to the axonal protein neurofilament light (NF-L) in the experimental autoimmune encephalomyelitis (EAE) animal model of multiple sclerosis. To examine the association between T cells and axonal damage, pathology studies were performed on NF-L immunised mice. The interaction of T cells and axons was analysed by confocal microscopy of central nervous system tissues, and T-cell and antibody responses to immunodominant epitopes identified in Biozzi ABH (H2-A<sup>g7</sup>) and SJL/J (H2-A<sup>s</sup>) mice. These epitopes, algorithm-predicted peptides and encephalitogenic motifs within NF-L were screened for encephalitogenicity. Confocal microscopy revealed both CD4<sup>+</sup> and CD8<sup>+</sup> T cells alongside damaged axons in the lesions of NF-L immunised mice. CD4<sup>+</sup> T cells dominated the areas of axonal injury in the dorsal column of spastic mice in which the expression of granzyme B and perforin was detected. Identified NF-L epitopes induced mild neurological signs similar to the observed with the NF-L protein, yet distinct from those characteristic of neurological disease induced with myelin oligodendrocyte glycoprotein. Our data suggest that CD4<sup>+</sup> T cells are associated with spasticity, axonal damage and neurodegeneration in NF-L immunised mice. In addition, defined T-cell epitopes in the NF-L protein might be involved in the pathogenesis of the disease.

## Introduction

Multiple sclerosis (MS) is a chronic demyelinating and neurodegenerative disease of the central nervous system (CNS) widely considered to be due to aggressive, autoreactive T cells and antibodies to myelin (1-3). However, accumulating evidence shows that immune responses to neuronal and axonal proteins are also present in a wide range of neurodegenerative disorders including MS (4-6). That these responses may contribute to axonal and neuronal damage, pathological hallmarks of MS, is supported by observations that immunisation with neuronal antigens and transfer of antibodies directed to neuronal and axonal proteins induce neuronal damage in animals (7-10).

The lack of expression of molecules of the major histocompatibility complex (MHC) class II on neurons indicates that neurons cannot activate CD4<sup>+</sup> cells in an antigen specific manner. However, neurons constitutively express or readily up regulate expression of MHC class I during inflammation, indicating that neurons may become targets for CD8<sup>+</sup> T cells (11). Conceptually, both CD4<sup>+</sup> and CD8<sup>+</sup> T cells could mediate attack on axons and neurons, either by direct contact via antigen-independent interactions or as a result of collateral damage (12). Activated T cells in the CNS are reported to produce cytotoxic molecules as well as glutamate, nitric oxide and reactive oxygen species that could contribute to the damage and progressive neurodegeneration observed in MS and other neurodegenerative diseases in which inflammation has been described (13-15). In addition, activation of B cells might lead to the production of specific antibodies to neuronal antigens that could also contribute to the damage and progressive neurodegeneration (16).

To examine the mechanisms of autoimmunity to neurons we have developed a model of autoimmune induced axonal and neuronal damage following immunisation of mice with the neuronal cytoskeletal protein neurofilament light (NF-L) (10).

Whether T-cell responses to neuroaxonal components are pathogenic in MS is as yet unknown; although we have recently shown that NF-L is phagocytosed by MHC class II<sup>+</sup> microglia/macrophages in MS brain lesions (17), indicating a potential source by which autoreactive T cells could become reactivated in MS. In mice, we have shown that autoimmunity to NF-L causes spasticity and neurodegeneration and that axonal damage is a direct consequence of such responses (9, 10). Infiltration of CD3<sup>+</sup> T cells and B220<sup>+</sup> cells mainly localised in the dorsal column of the spinal cord of NF-L immunised mice (9). In addition, immunoglobulin deposits were observed into the axons in mice immunised with NF-L (10).

In the present study, we characterised the T-cell infiltrates in the CNS of spastic mice. Both CD4<sup>+</sup> and CD8<sup>+</sup> T cells were found in close association to axons, although CD4<sup>+</sup> T cells dominated the infiltrates in lesions. In addition, increased perforin expression and cells positive for granzyme B could be observed in the spinal cord of mice immunised with NF-L. Furthermore, NF-L peptides were screened for T-cell and B-cell responses in ABH (H2-A<sup>g7</sup>) and SJL/J (H2-A<sup>s</sup>) mice, and the pathogenic potential of these peptides and predicted binding motifs to H2-A<sup>g7</sup> present in the NF-L sequence were investigated.

In summary, our study reveals that T cells associated with the expression of cytotoxic molecules are present in lesions in the dorsal columns of spastic mice immunised with NF-L and we show, for the first time, that active immunisation with defined NF-L peptides induced neurological disease in ABH mice.

## Materials and methods

### *Mice*

Male and female 10-week-old Biozzi ABH (H-2<sup>dq1</sup>) and SJL/J (H-2<sup>s</sup>) mice were obtained from Harlan (Bicester, UK) and Charles River laboratories (Kent, UK) or were bred at QMUL (London, UK). All procedures were performed in accordance with the UK Animals (Scientific Procedures) Act (1986) and approved by the local ethics committee. All procedures were performed following Institutional ethical review in accordance to the United Kingdom Animals (Scientific Procedures) Act (1986) and European Union Directive 2010/63/EU. Animals were housed and monitored consistent with the principles of the ARRIVE guidelines as described previously (18).

### *Antigens*

Spinal cord homogenate (SCH) prepared from 60 ABH mice was lyophilised and reconstituted in PBS as described previously (18). Recombinant mouse NF-L (rmNF-L) was prepared as described previously (17). Overlapping 16-amino-acid peptides (Table 1) based on the mouse protein sequence (NCBI protein ID: NP035040) and myelin oligodendrocyte glycoprotein (MOG<sup>35-55</sup>) peptide (MEVGWYRSPFSRVVHLYRNGK) were synthesised as peptide amides (CONH<sub>2</sub>; Cambridge Research Biochemicals Ltd, UK).

### *Induction of experimental autoimmune encephalomyelitis*

Mice were injected subcutaneously with 1 mg SCH, 200 µg rmNF-L, 200 µg MOG<sup>35-55</sup> or rmNF-L and MOG<sup>35-55</sup> (1:1), or pools of 30 µg of each peptide emulsified with incomplete Freund's adjuvant (Difco Laboratories, UK) supplemented with 48 µg *Mycobacterium tuberculosis* and 6 µg *M. butyricum* (Difco Laboratories) on day 0 and day 7 as described previously (19). Control mice were immunised with complete Freund's adjuvant (CFA) only. All mice were injected with 200 ng *Bordetella pertussis*

toxin (Sigma, USA) intraperitoneally, immediately after immunisation and 24 h later.

To identify encephalitogenic epitopes, mice were immunised with rmNF-L, individual or pooled peptides. To optimise identification, sequences containing motifs that bind to or interact with H2-A<sup>g7</sup> were selected as described previously (19). The Rankpep server was additionally used to predict binding to H2-A<sup>g7</sup> (<http://imed.med.ucm.es/Tools/rankpep.html>).

Mice were monitored daily and scored according to a neurological scale: 0, normal; 0.5, partial loss of tail tone; 1, paralysis or spasticity of the tail; 2, impaired righting reflex; 3, paralysis or spastic paresis of one limb; 4, paralysis or spastic paresis of two limbs; and 5, moribund (10, 18). Mice were sacrificed by carbon dioxide inhalation and brains and spinal cords snap-frozen in liquid nitrogen or processed for pathology (10).

**Table 1. Sequences of mouse neurofilament-light peptides<sup>a</sup>**

Sequence	Amino Acids	Sequence	Amino Acids
1-16	SSFGYDPYFSTSYKRR	273-288	MQNAEWFKSRFTVLT
9-24	FSTSYKRRYVETPRVH	281-296	KSRFTVLTESAANKTD
17-32	YVETPRVHISSVRSYG	289-304	ESAAKNTDAVRAAKDE
25-40	ISSVRSGYSTARSAYS	297-312	AVRAAKDEVSESRLL
33-48	STARSAYSYSAPVSS	305-320	VSESRLLKAKTLEIE
41-56	SYSAPVSSLSVRRSY	313-328	KAKTLEIEACRGMNEA
49-64	SLSVRRSYSSSSGSLM	321-336	ACRGMNEALEKQLQEL
57-72	SSSSGSLMPSLENLDL	329-344	LEKQLQELEDKQNADI
65-80	PSLENLDLSQVAASIN	337-352	EDKQNADISAMQDTIN
73-88	SQVAASINDLKSIRTQ	345-360	SAMQDTINKLENELRS
81-96	DLKSIRTQEKALQDL	353-368	KLENELRSTKSEMARY
89-104	EKALQDLNDRFASFI	361-376	TKSEMARYLKEYQDLL
97-112	NDRFASFIERVHELEQ	369-384	LKEYQDLLNVKMALDI
105-120	ERVHELEQQNKVLEAE	377-392	NVKMALDIEIAAYRKL
113-128	QNKVLEAELLVLRQKH	385-400	EIAAYRKLLEGEETRL
121-136	LLVLRQKHSEPSRFRA	393-408	LEGEETRLSFTSVGSI
129-144	SEPSRFALYEQEIRD	401-416	SFTSVGSITSGYSQSS
137-152	LYEQEIRDLRLAEDA	409-424	TSGYSQSSQVFGRSAY
145-160	LRLAEDATNEKQALQ	417-432	QVFGRSAYSLQSSSY
153-168	TNEKQALQGEREGLEE	425-440	SGLQSSSYLMSARSFP
161-176	GEREGLEETLRNLQAR	433-448	LMSARSFPAYTSHVQ
169-184	TLRNLQARYEEVLSR	441-456	AYTSHVQEEQTEVEE
177-192	YEEVLSREDAEGRML	449-464	EEQTEVEETIEATKAE
185-200	EDAEGRLMEARKGADE	457-472	TIEATKAEAKDEPPS
193-208	EARKGADEAALARAE	465-480	EAKDEPPSEGEAE
201-216	AALARAELEKRIDSLM	473-488	EGEAEEEEKEKEEGEE
209-224	EKRIDSLMDEIAFLKK	481-496	KEKEEGEEEGAE
217-232	DEIAFLKKVHEEEIAE	489-504	EEGAEAAAEDSEDT
225-240	VHEEEIAELQAQIQYA	497-512	AKDESEDTKEEEGGE
233-248	LQAQIQYAQISVEMDV	505-520	KEEEEGGEGEEEDTKE
241-256	QISVEMDVSSKPDLSA	513-528	GEEEDTKESSEEEKKE
249-264	SSKPDLSAALKDIRAQ	521-536	SEEEKKEESAGEEQV
257-272	ALKDIRAQYEKLAAKN	529-544	ESAGEEQVAKKKD
265-280	YEKLAAKNMQNAEEWF		

<sup>a</sup>Mouse neurofilament-light sequence (NCBI reference sequence: NP035040).

### Immunohistochemistry

Sections (3 µm) from snap-frozen spinal cord tissues were fixed with acetone and incubated overnight at 4°C with mAb directed to CD4 (YTS 191.1.2), CD8 (YTS 169AG, ImmunoTools, Friesoythe, Germany), MHC I antigens (HM1091; Hycult Biotech, USA) or biotinylated MHC II (OX 6, a kind gift of Jack van Horssen, VU University Medical Center) diluted in antibody diluent (Immunologic, The Netherlands). After washing, endogenous peroxidase was blocked with 0.3% H<sub>2</sub>O<sub>2</sub> in PBS. Sections

stained for CD4, CD8 and MHC I were incubated with biotinylated rabbit anti-rat Ig (Dako, Denmark) for 1 h, followed by peroxidase-coupled avidin–biotin complex (ABC kit; Vector Laboratories, USA). Sections stained with biotinylated MHC II were incubated with streptavidin–horseradish peroxidase complex (Dako) for 1 h. All secondary antibodies were visualised with 3,3'-diaminobenzidine. Antibodies were prescreened on brain, liver, lung, spleen and tonsil tissues and isotype control mAb served as negative control. The percentage of CD4<sup>+</sup> and CD8<sup>+</sup> T cells were counted at 25 x objective at three levels of the spinal cord.

For immunofluorescence, sections were incubated with blocking solution (CleanVision IHC/ICC, Immunologic) containing 10% normal goat serum for 2 h, washed in PBS and incubated with mAb to NF-L (10H9), SMI32 (Covance, USA) or Neu-N (Merck Millipore, Germany) and CD3 (CD3-12; Serotec, UK), CD4 (YTS 191.1.2) or CD8 (YTS 169AG, ImmunoTools) overnight at 4°C. After washing in PBS, sections were incubated with goat anti-mouse IgG1 Alexa 594 or goat anti-rat IgG Alexa 488 (Invitrogen, UK) for 60 min at room temperature. Sections were viewed using confocal laser scanning microscopy (Leica DMI6000; The Netherlands). Image processing was performed using NIH Image J software (<http://rsb.info.nih.gov/ij/index.html>).

Granzyme B staining was performed on paraffin embedded sections (4 µm). In brief, sections were deparaffinised and rinsed in H<sub>2</sub>O. Subsequently, endogenous peroxidase was blocked as described above. After rinsing in PBS, antigen retrieval in Tris–ethylenediamine tetraacetic acid buffer (pH 9.0) was performed in a microwave followed by incubation with 10% normal goat serum. Sections were incubated overnight at 4°C with polyclonal rabbit anti-granzyme B (Abcam, UK) in 1% BSA. Subsequently, sections were washed in PBS and incubated with secondary antibody Envision anti-rabbit labelled with horseradish peroxidase (Dako) for 30 min and visualised with 3,3'-diaminobenzidine.

#### *Reverse transcriptase polymerase chain reaction*

Spinal cords from control and NF-L immunised ABH mice were dissolved in lysis buffer (NucleoSpin RNA/ Protein kit, Machery-Nagel GmbH, Germany) and homogenised with 1.4 mm ceramic beads (Precellys 24; Peqlab Biotechnologie GmbH, Erlangen, Germany) at 5,000 rpm for 15 seconds. Subsequently, RNA was isolated using NucleoSpin (Macherey-Nagel) according to the manufacturers recommendations. Purity was confirmed using 260:280 OD ratios (optical density, Nano-Drop 1000; Peqlab Biotechnologie GmbH). RT reactions were performed with the MMLV RT-kit and random hexanucleotide primers (Invitrogen) and gene expression was measured using Taq-Polymerase (Biomol GmbH, Germany).

Primers for perforin amplification (sense, 5'-CTGCCACTCGGTCAGAATG-3'; antisense, 5'-CGGAGGGTAGTCACATCCAT-3') were used at annealing temperature of 59°C, amplifying an 88-base-pair fragment. Expression levels of the reference gene hypoxanthine guanine phosphoribosyl transferase were used as control. Primers for hypoxanthine guanine phosphoribosyl transferase amplification (sense, 5'-GCTGGTGAAAAGGACCTCT-3'; antisense: 5'-CACAGGACTAGAACACCTGC-3') were used at an annealing temperature of 60°C, amplifying a 248-base-pair fragment, and primers for 18s RNA amplification (sense, 5'-CGGCTACCACATCCAAGGAA-3'; antisense, 5'-GCTGGAATTACCGCGGCT-3') were used at an annealing temperature of 60°C, amplifying a 187-base-pair fragment. Gene expression was performed using RT-PCR technology (Bio-Rad, Germany) with SYBR green (SensiMix™, Biotline, Germany), as published previously (20).

### *T-cell proliferation assays*

ABH and SJL/J mice were immunised with rmNF-L in CFA or CFA only. Spleen cells were collected 10 days after priming and single-cell suspensions ( $3 \times 10^5$ /mL) cultured in RPMI medium (Gibco, UK) with 5% FCS (Gibco), 2 mM L-glutamine, 100 U/mL penicillin, 100  $\mu$ g/mL streptomycin, 5 mM Hepes, and  $5 \times 10^{-5}$  M 2-mercaptoethanol (Gibco). Cells were incubated with NF-L peptides or rmNF-L for 72 h. Proliferation was determined by incorporation of [ $^3$ H]-thymidine (GE Healthcare, Sweden). Stimulation indices (SIs) were calculated as the proliferative response in the presence of antigen divided by the response in the absence of antigen. SI in the CFA control group ranged from 0.6 to 1.1. Positive stimulation was defined as SI > 1.5.

### *Enzyme-linked immunosorbent assay*

To identify B-cell epitopes in NF-L, Biozzi ABH and SJL/J mice were immunised with the rmNF-L protein in complete adjuvant and serum was collected on day 15 post immunisation. The mouse immune sera were tested for their reactivity to NF-L overlapping peptides. Nunclon plates (Nunc, Denmark) were coated overnight at 4°C with 10  $\mu$ g/mL mouse NF-L peptides or rmNF-L protein in carbonate buffer. Plates were washed twice in PBS and blocked for 1 h at 37°C with 2% BSA/PBS. After blocking, ABH or SJL/J immune sera, diluted 1:100 in 1% BSA/PBS, were added and incubated for 1 h at room temperature. After washing in PBS-Tween 0.1%, the plates were incubated for 1 h at room temperature with horseradish peroxidase-conjugated rabbit anti-mouse Ig (Dako). The reaction product was developed with TMB substrate (Thermo Fisher Scientific, UK) and stopped by the addition of 2 M hydrochloric acid. The absorbance was measured at 450 nm, using a Synergy HT microplate reader (Bio-Tek instruments, USA). Background values were obtained by the reactivity of immune serum on peptide uncoated wells. An absorbance above the mean plus three standard deviations of the background reactivity was taken as positive.

### *Statistical analysis*

For comparison of clinical EAE scores, significance between the groups was determined by nonparametric Mann-Whitney rank-sum test (SigmaStat, Systat Software Inc., USA). Data represent the mean  $\pm$  standard error of the mean (\*\*  $p < 0.01$ , \*  $p < 0.05$ ).

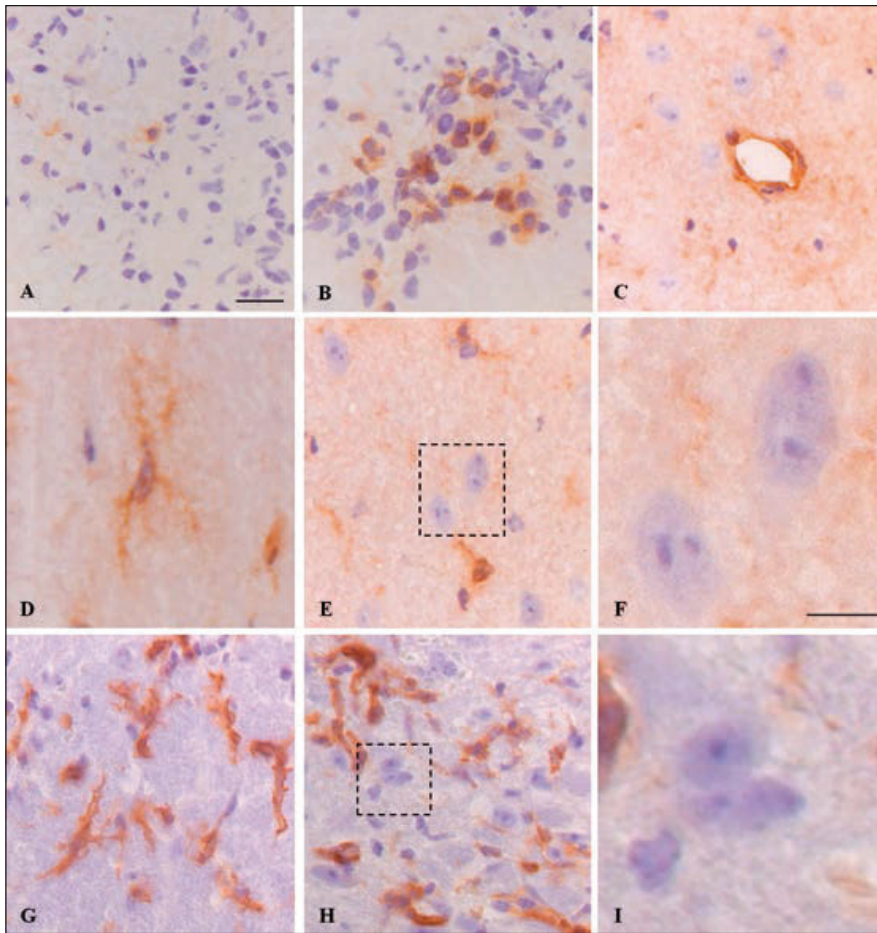
## **Results**

### *CD4<sup>+</sup> and CD8<sup>+</sup> T cells are associated with neuronal/axonal damage in spastic mice*

To examine the presence of T cells in the CNS during disease, CD4<sup>+</sup> or CD8<sup>+</sup> T cells within the lesions of NF-L immunised mice at the peak of disease were examined. Using immunohistochemistry, the ratio of CD4<sup>+</sup> and CD8<sup>+</sup> T cells in the lesions was determined (Fig. 1A, B, respectively). We observed that significantly more CD4<sup>+</sup> T cells were present within the lesions in a ratio of 9:1 CD4<sup>+</sup>:CD8<sup>+</sup> T cells (data not shown). To support a role for CD8<sup>+</sup> T cells we examined the expression of MHC class I in the spinal cord of diseased mice. In contrast to expression of MHC class I antigens on inflammatory cells in the lesion as well as perivascular cells (Fig. 1C) and microglia in normal appearing white matter (Fig. 1D), there was no evidence of MHC class I antigen expression on neurons or axons in the lesions (Fig. 1E, F) or in normal-appearing tissues (data not shown). Likewise, while MHC class II was expressed on inflammatory cells in the blood vessels and in the parenchyma,

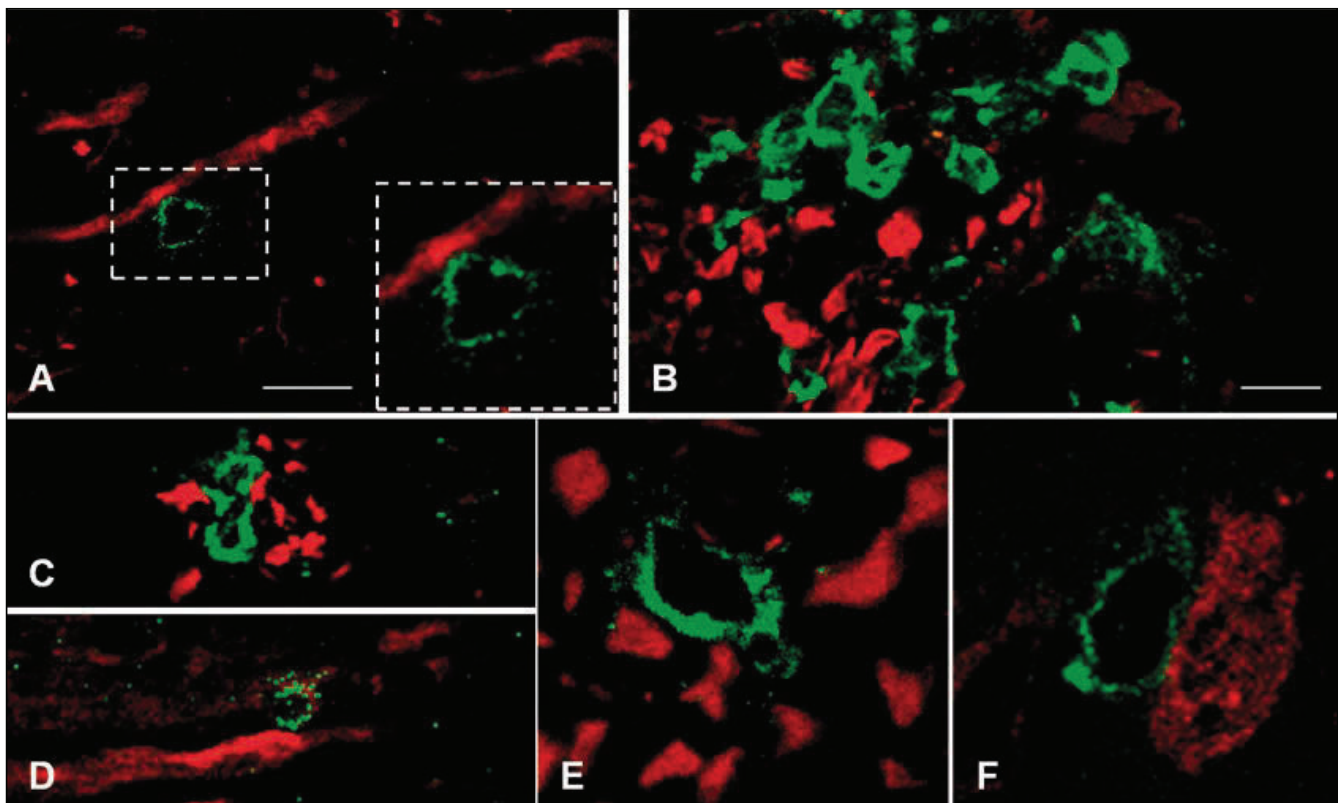


being present on ramified cells resembling microglia (Fig. 1G), no expression on neurons and axons was observed (Fig. 1H, I), as has been shown before in normal or pathological tissues during EAE (21).



**Figure 1. T cells in lesions of spastic mice.** Immunohistochemistry showing (A) CD8 and (B) CD4 T cells in the dorsal column of a spastic mouse. (C) to (F) Major histocompatibility complex (MHC) class I antigen staining expressed on (C) endothelial cells, and (D) microglia (E, F), but not on neurons. (G) to (I) MHC class II staining on ramified cells resembling microglia but not neurons (H, I). A to E, G to H, bar=10  $\mu\text{m}$ ; F and I, bar=5  $\mu\text{m}$ .

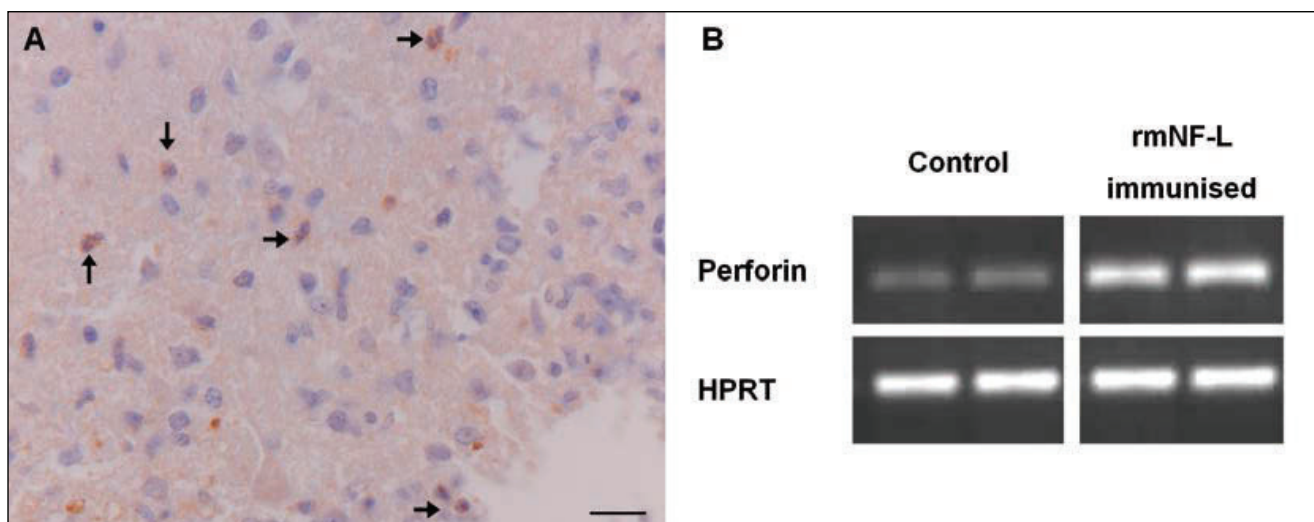
To examine the interaction of T cells and axons we used confocal microscopy of CNS tissues from spastic mice. We observed a close association of CD3<sup>+</sup> T cells aligning with possible damaged axons as indicated by SMI32 staining (Fig. 2A), in which there was a close association between the T cell and the axon (Fig. 2A, insert). In cross-sections of the lesion, CD4<sup>+</sup> T cells were observed within areas of axonal damage (Fig. 2B) and frequently observed between axons with increased immunoreactivity for SMI32 and neurofilament, indicating axonal damage (Fig. 2C, D). While CD8<sup>+</sup> T cells were also observed between thickened axons in the lateral funiculus and adjacent to Neu-N<sup>+</sup> neurons in the spinal cord (Fig. 2E, F), these were rare as compared with CD4<sup>+</sup> T cells.



**Figure 2. T cells associated with axonal damage in spastic mice.** Confocal images of (A) CD3 T cell (green) in contact with SMI32 axon (red). (B, C) CD4 T cells in area of axonal damage and (D) next to swollen NF-L axons. (E) CD8 T cells between NF-L axons and (F) Neu-N neurons in the spinal cord. A, E, F, scale bar=5  $\mu$ m; B, C, D, scale bar=25  $\mu$ m.

#### Cytotoxic molecules are present in the spinal cord of NF-L immunised mice

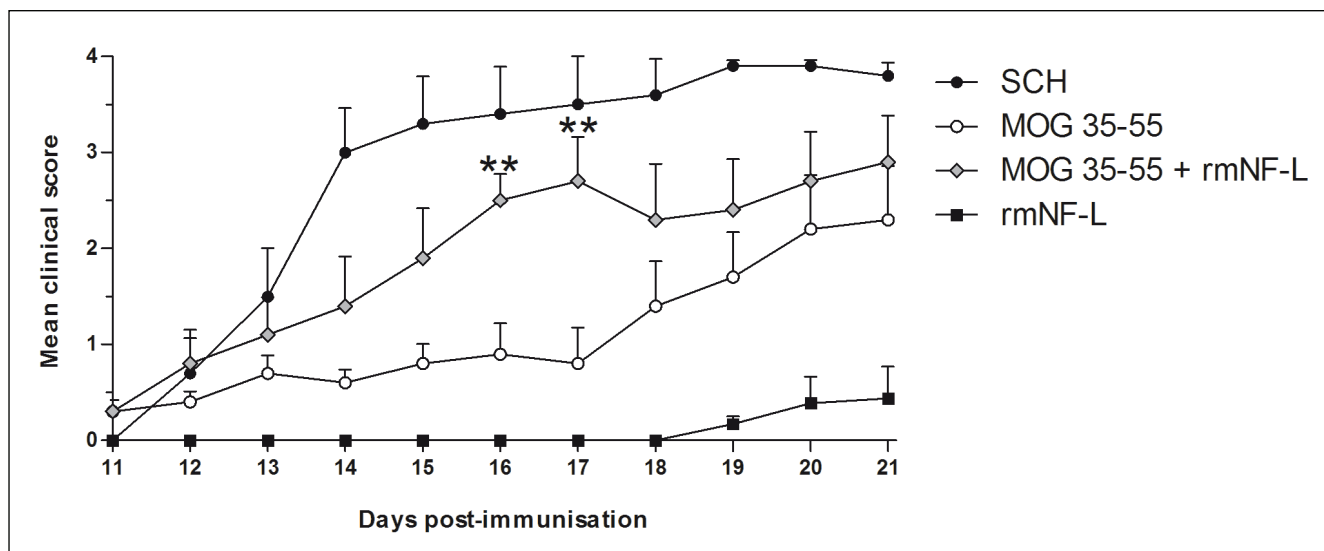
To examine the potential mechanisms of T cell-mediated axonal damage, the expression of the cytotoxic molecules granzyme B and perforin was examined in the spinal cord of spastic mice. Immunostaining revealed the expression of granzyme B in spinal cord lesions of mice immunised with NF-L (Fig. 3A). Immunodetection of perforin in mouse CNS was not possible due to lack of specific mAbs. Instead, RT-PCR on spinal cord tissues was performed and showed that perforin mRNA levels were increased in rmNF-L immunised mice compared with controls (Fig. 3B). In addition, granzyme B and perforin were expressed by peripheral CD4<sup>+</sup> T cells from NF-L immunised mice compared with CD8<sup>+</sup> T cells (data not shown).



**Figure 3. Production of cytotoxic molecules in spastic mice.** **A)** Granzyme B-positive staining (indicated by arrows) was performed on paraffin-embedded spinal cord sections: scale bar=15  $\mu$ m. **B)** Gel electrophoresis of PCR products show perforin expression is increased in spastic mice ( $n=4$  mice per group), hypoxanthine guanine phosphoribosyl transferase (HPRT) = internal control). See figure on the previous page.

#### Autoimmunity to NF-L exacerbates MOG<sup>35-55</sup> experimental autoimmune encephalomyelitis in ABH mice

To increase the likelihood of exposing NF-L protein in the CNS, ABH mice were co-immunised with MOG<sup>35-55</sup> to induce myelin damage (Fig. 4). While SCH induced classical EAE about 11 days after immunisation, co-immunisation of rmNF-L with MOG<sup>35-55</sup> induced significantly augmented disease in which spasticity was observed on days 16 and 17 ( $p=0.005$ ), indicating that myelin damage may first be necessary to expose neuronal antigens. Within this timeframe, mice immunised with rmNF-L only did not develop clinical disease as we have reported previously (10). However, by day 21, NF-L immunised mice did develop neurological signs of disease with a mean day of onset  $19.8 \pm 1.6$ , EAE score  $1.5 \pm 1.0$ , group score  $0.5 \pm 0.4$  and an EAE incidence of 50%.



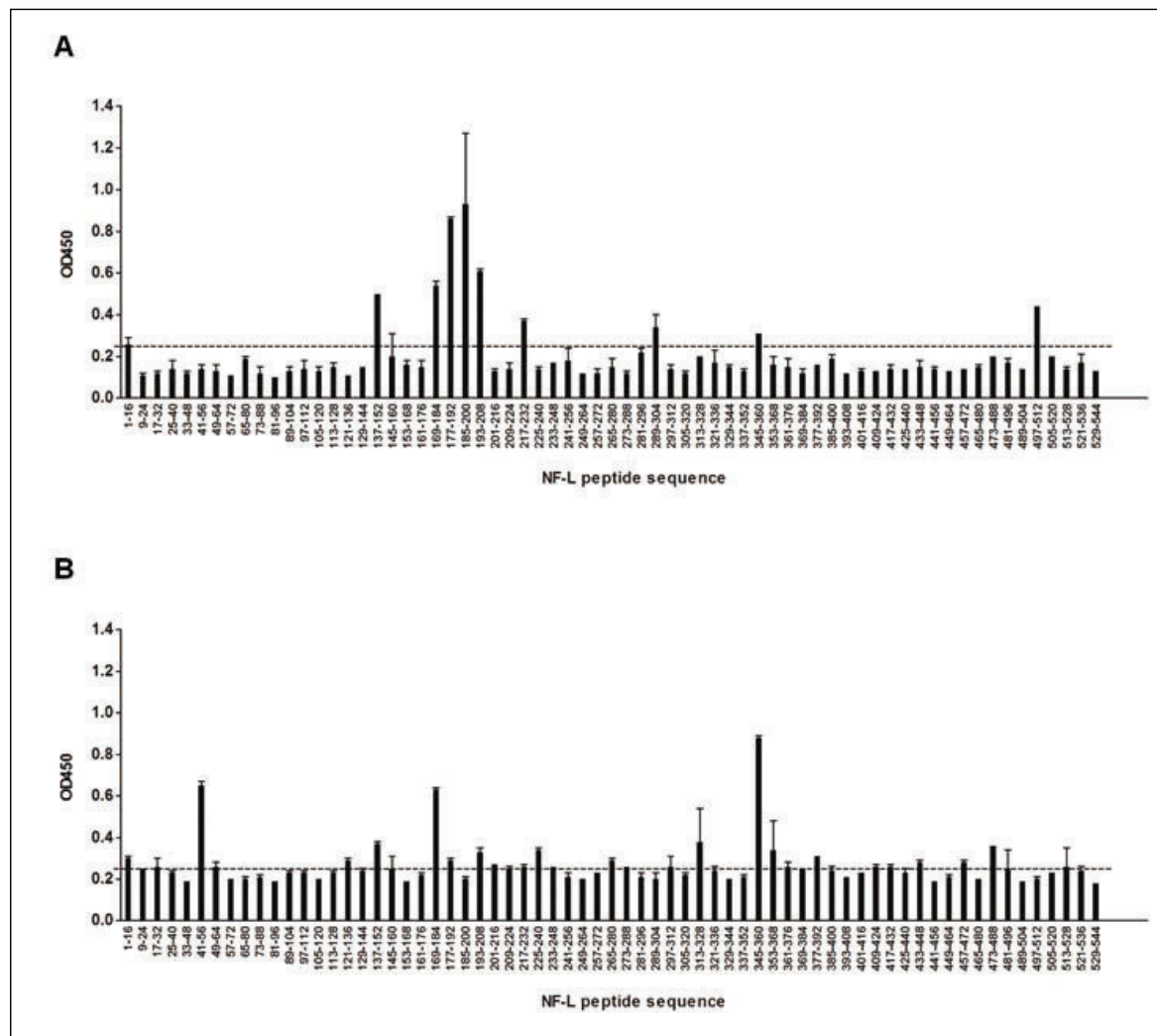
**Figure 4. Augmentation of MOG<sup>35-55</sup>-induced experimental autoimmune encephalomyelitis by neurofilament light.** ABH mice were immunised with SCH ( $n=8$ ), MOG<sup>35-55</sup> ( $n=9$ ), rmNF-L ( $n=9$ ) or rmNF-L and MOG<sup>35-55</sup> ( $n=9$ ). \*\*  $p<0.005$ ; Mann-Whitney U test.

#### Immunodominant B-cell and T-cell epitopes of neurofilament light in mice

To determine immunodominant epitopes in the NF-L protein, antibody and T-cell responses to synthetic 16-mer overlapping peptides spanning the mouse NF-L sequence (Table 1) were screened in ABH and SJL/J mice immunised with rmNF-L in CFA (19). Since we have seen before that immunisation with NF-L in Biozzi mice can result in antibody production and antibody binding to axons, it was of interest to screen the antibody responses to the overlapping peptides (10). Antibody reactivity to NF-L peptides in both strains of mice revealed an immunodominant region that corresponds to the overlapping region (NF-L amino acids 169 to 208) and the single region (amino acids 169 to 184) in ABH and SJL/J mice respectively (Fig. 5A, B). This region is located in the coil 1b domain of NF-L. In line with our observations, these motifs (amino acids 154 to 195) are likely to be linear B-cell epitopes according to the BCPred prediction algorithm (<http://ailab.cs.iastate>).

edu/bcpreds/). In SJL/J mice, NF-L specific antibodies also reacted to the single peptides amino acids 41 to 56 and amino acids 345 to 360 (Fig. 5B). In both strains, the mean absorbance values for antibody response to rmNF-L protein were within the optical density range of 1.21 to 1.34.

Immunodominant T-cell responses to two dominant regions, amino acids 241 to 288 and 345 to 368, were observed in ABH mice (Fig. 6A). In comparison, T-cell responses in SJL/J mice were observed to amino acids 241 to 288 (Fig. 6B). In both strains, the SIs were lower than responses to rmNF-L alone (SI=5 to 10). These results are in line with data using overlapping 15-mer peptides of NF-L in ABH mice (data not shown), indicating that immunodominant epitopes reside within the rod domain (linker 2 and coil 2B) of NF-L. No overlapping peptide induced significant proliferation in cells from CFA immunised mice.

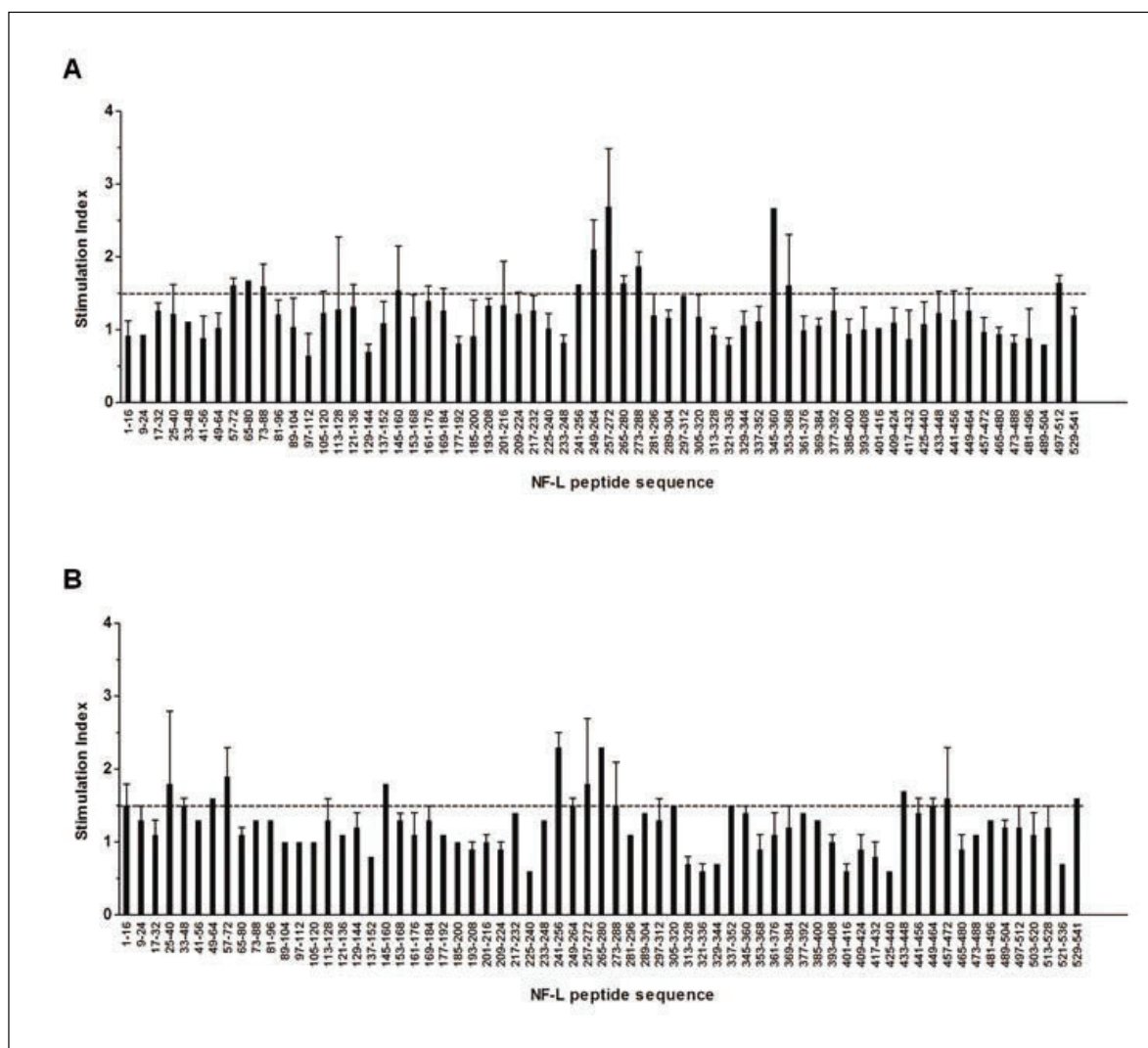


**Figure 5. B-cell responses to neurofilament light peptides in ABH and SJL/J mice.** **A)** ABH mice ( $n=4$ ) and **(B)** SJL/J mice ( $n=4$ ) were immunised with rmNF-L and serum antibody reactivity to NF-L peptides was measured on day 10. Mean absorbance ( $OD > 0.25$ ) was considered positive.

#### Pathogenicity of immunodominant epitopes

To determine the pathogenicity of immunodominant epitopes, ABH mice were immunised with individual NF-L peptides spanning the T-cell immunodominant region (amino acids 241 to 272) or with a pool of immunodominant B-cell epitopes. In contrast to SCH-induced EAE, immunisation of ABH mice with amino acids 241 to 256, 249 to 264, 257 to 272, 345 to 360 and 353 to 368 induced spasticity of

the tail and hind limbs in line with studies using rmNF-L (Table 2, panel A) (10). Immunisation with NF-L peptides also induced a delayed onset of clinical signs occurring 10 days later than SCH-induced EAE (Table 2, panel A). Rota-rod studies did not reveal additional signs of neurological disease (data not shown). SJL/J mice immunised with rmNF-L also developed very mild clinical disease of spasticity (data not shown). The pathogenic potential of NF-L peptides was also observed with some, but not all, peptides predicted to bind with H2-A<sup>g7</sup> (Table 2, panel B). No significant effect was observed after immunisation with the immunodominant B-cell epitopes (amino acids 169 to 208) (Supp. Fig. 1). Additionally, the capacity of NF-L-specific T cells to transfer spasticity was tested in ABH mice. Lymph node and spleen cells from rmNF-L-immunised ABH mice were collected 12 days after immunisation and stimulated for 10 days *in vitro* with rmNF-L. Activated cells were adoptively transferred into irradiated naïve recipients and followed until day 25. Mild signs of disease (score 0.5) were observed in two of seven mice revealing the pathogenic potential of T cells to NF-L (data not shown).



**Figure 6. T-cell responses to NF-L peptides in ABH and SJL/J mice.** **A)** ABH mice ( $n=4$ ) and **(B)** SJL/J mice ( $n=4$ ) were immunised with rmNF-L and T-cell proliferation to NF-L peptides was measured on day 10. Stimulation index  $>1.5$  was considered positive.

**Table 2. Pathogenic neurofilament light peptides ABH mice<sup>a</sup>**

Antigen/peptides	Number with EAE	Mean group Score <sup>b</sup>	Mean EAE Score <sup>c</sup>	Mean day of onset
<b>Panel A</b>				
SCH	4/4	4.0 ± 0.0*	4.0 ± 0.0	15.3 ± 0.3
rmNF-L	4/5	0.7 ± 0.3*	0.9 ± 0.2	31.0 ± 1.4
241-256	1/5	0.1 ± 0.1	0.5 ± N/A	21.0 ± N/A
249-264	5/5	0.9 ± 0.3*	0.9 ± 0.3	25.6 ± 6.4
257-272	3/5	0.4 ± 0.2	0.7 ± 0.2	29.3 ± 0.6
265-280	0/5	N/A	N/A	N/A
273-288	0/5	N/A	N/A	N/A
345-360	3/5	0.7 ± 0.3	1.2 ± 0.3	29.3 ± 0.3
353-368	3/5	0.5 ± 0.2	0.8 ± 0.2	25.7 ± 4.4
<b>Panel B</b>				
129-144	2/5	0.2 ± 0.1	0.5 ± 0.0	18.5 ± 6.5
297-312	2/5	0.2 ± 0.1	0.5 ± 0.0	30.5 ± 0.5
313-328	0/5	N/A	N/A	N/A
113 to 128, 281 to 296, 289 to 304, 321 to 336	3/4	0.9 ± 0.4	1.2 ± 0.3	28.7 ± 0.7
<b>Panel C</b>				
CFA	0/5	N/A	N/A	N/A

<sup>a</sup>Mice were injected with 200 µg SCH, rmNF-L or single immunodominant NF-L peptides in CFA (panel A) or with peptides in CFA predicted to bind H2-A9<sup>7</sup> (panel B) or with CFA alone (panel C).

<sup>b</sup>Mean ± SEM of the maximum EAE score of mice in the group.

<sup>c</sup>Mean ± SEM of the maximum EAE score of affected mice.

\*  $p < 0.05$ , compared with CFA immunised mice.

N/A, not applicable.

## Discussion

Accumulating evidence indicates that as well as myelin-specific T cells, neuronal-specific T cells may gain access to the CNS and contribute to neurodegeneration. Immune responses to neurons are reported in Rasmussen's encephalitis (22), Alzheimer's disease (23), Parkinson's disease (24) and paraneoplastic disorders (6), underscoring the potential role of pathogenic neuronal-reactive T cells in MS in which axonal damage correlates with disability. To examine the role of T cells in neurodegeneration, we have studied autoimmune-mediated neurodegeneration in mice immunised with NF-L in which spasticity and paralysis, clinical features characteristic of MS, are observed (9, 10). In previous studies, infiltration of CD3<sup>+</sup> and B220<sup>+</sup> cells and immunoglobulin deposits were observed in spinal cord lesions of NF-L immunised mice (10).

In this study, we aimed to further investigate the cellular and humoral immune response in the mouse model of NF-L-induced neurodegeneration. Immunohistochemistry revealed the association of both CD4<sup>+</sup> and CD8<sup>+</sup> T cells with neuronal damage and the expression of cytotoxic molecules in the CNS. We also identify immunodominant regions and encephalitogenic epitopes on the NF-L protein. Together, our data support the accumulating evidence that T cells play a role in neuronal and axonal damage in neurodegenerative disorders. Adoptive transfer experiments to test the ability of NF-L peptide-specific T cells to induce disease would be of interest in future studies.

A pre-requisite for antigen-specific CD8<sup>+</sup> cytotoxic damage of neurons is the expression of MHC class I molecules. Strong evidence exists of MHC class I on neurons, neurites and axons, revealing differential expression depending on neuronal subtype, development stage and the inflammatory stimulus. MHC class I expression is highly up regulated on neurons following interferon-gamma (IFN $\gamma$ )

treatment but not tumor necrosis factor alpha (TNF $\alpha$ ) (25). Intriguingly, CD8<sup>+</sup> T cells directed to NF-L generated from spastic mice produce high levels of IFN $\gamma$  (5), indicating that T cells to NF-L could trigger progressive neurodegeneration. Further support for T cell-mediated neuronal damage comes from the finding that, in EAE, myelin-reactive T cells are activated by neurofilament peptides (26). Such interaction between T cells and neurons may induce antigen-specific lysis of neurons (27) or dysfunction by impairing electrical signalling (28) and induction of rapid microtubule axonal destabilisation (29). In the spastic model it is unlikely that antigen-specific CD8<sup>+</sup> T-cell activation leads to neuronal damage since MHC class I was not present on neurons, either in the lesions or in normal appearing tissue. While it is unlikely that CD4<sup>+</sup> T cells induce neuronal death in an antigen-specific fashion, as neurons do not express MHC class II, pathogenic CD4<sup>+</sup> T cells expressing NKG2C could injury neurons and axons expressing HLA-E, as reported for oligodendrocytes in MS (30). T cell-mediated neuronal damage may alternatively occur via antigen-independent interactions involving Fas-Fas ligand, TRAIL, CD11a and CD40. In culture, polyclonally activated CD4<sup>+</sup> and CD8<sup>+</sup> T cells are cytotoxic to human neurons (27, 29), underscoring the potential pathogenic role of both CD4<sup>+</sup> and CD8<sup>+</sup> T cells. In line with these studies, our results show that granzyme B and perforin are present in the lesions of spastic mice, indicating that neuronal damage is more likely due to antigen-independent mechanisms. The predominance of CD4<sup>+</sup> T cells in the lesions of NF-L immunised mice may be due to selective recruitment or selective depletion of CD8<sup>+</sup> T cells (31). Moreover, the ratio between the T-cell subsets in the lesions in our model is similar to myelin-induced EAE (32). Further studies on the expression of cytotoxic molecules in T-cell subpopulations in the CNS of NF-L immunised mice will be necessary.

Our observations that co-immunisation with MOG and NF-L leads to exacerbated disease is pertinent to what might occur in MS during myelin damage, in which neurons may be more vulnerable to immune responses to neuronal antigens. In this way, the course of neuronal degeneration might be accelerated in the presence of NF-L-reactive T cells (5) and antibodies to NF in MS (33).

EAE studies have been instrumental in identifying the role of T cells and antibodies to myelin in disease. These have revealed important information about peptide:MHC:T-cell receptor interactions crucial for development of tolerance regimes using altered peptide ligands or tolerogenic routes for delivery of pathogenic peptides to modulate myelin-specific T cells (34, 35). These approaches have proved effective in chronic EAE models, preventing the clinical relapses, but do not control progressive disease (34) indicating that mechanisms other than myelin autoimmunity, such as autoimmunity to neuronal proteins, are involved in neurodegeneration in MS.

In ABH mice, immunisation with MOG and proteolipid protein readily induces neurological disease in which flaccid paralysis and preferential myelin loss is observed. In contrast, immunisation with NF-L protein induces a predominantly spastic disease in which neuronal damage is the primary pathological feature. To determine the pathogenic potential of NF-L peptide, we used a peptide mapping approach and identified a peptide core within the NF-L protein that contains similar elements to previously defined H2-A<sup>g7</sup> motifs within myelin basic protein, MOG and proteolipid protein (Table 3) (19). This supports many of the predictions made by computer modelling of peptide H2-A<sup>g7</sup> interactions (36).

**Table 3. Pathogenic peptides associated with neurological disease in ABH mice**

Protein and peptide residues	Amino acid sequence	Reference
PLP 54-76	DYEYLNVIHAFQYVIGASF	(19)
MBP 12-35	YLATASTMDHARHGFLPRHRDTSGI	(19)
MOG 1-23	GQFRVIGPGYP <del>IRAL</del> VGDEQED	(19)
alpha-B crystallin 52-61	F <del>FL</del> RAPSWI	(37)
NF-L 129-144	SEPSRF <del>RAL</del> YEQ <del>IE</del> IRD	This study
NF-L 257-272	ALKD <del>IRA</del> QY <del>EK</del> LAAKN	This study

PLP, proteolipid protein; MBP, myelin basic protein.

Similar to the finding following immunisation with myelin antigens (19), several encephalitogenic epitopes were identified in NF-L immunised ABH mice. One should note that when using overlapping peptides, inappropriate sequence alignment could mask encephalitogenic epitopes or induce peptide epitopes to become tolerogenic (38). In addition, neurofilament modulates oligodendrocyte proliferation and differentiation (39), thereby masking the possible pathogenic impact of these proteins. Peptide mapping approaches have identified immunodominant epitopes of myelin basic protein in mice and humans and prompted tolerance strategies to myelin basic protein-specific T cells in MS. Current studies are underway to identify T-cell responses to NF-L epitopes in MS patients who respond to the NF-L protein (5). These studies may reveal whether such responses are present in subtypes of MS or correlate with extent of cognitive changes or progressive disease. Such findings may be key to the development of personal therapeutic approaches using tolerance regimens.

In conclusion, we show that peptide epitopes of NF-L induce neurological disease and that potentially pathogenic CD4<sup>+</sup> T cells dominate the lesions of NF-L immunised mice, a model for immune-mediated neurodegeneration. Our data suggest that, similar to peptide therapies targeting myelin responses (40), immune therapies targeting neuronal-specific T cells could thus be beneficial in reducing neurodegeneration in inflammatory disorders such as MS.

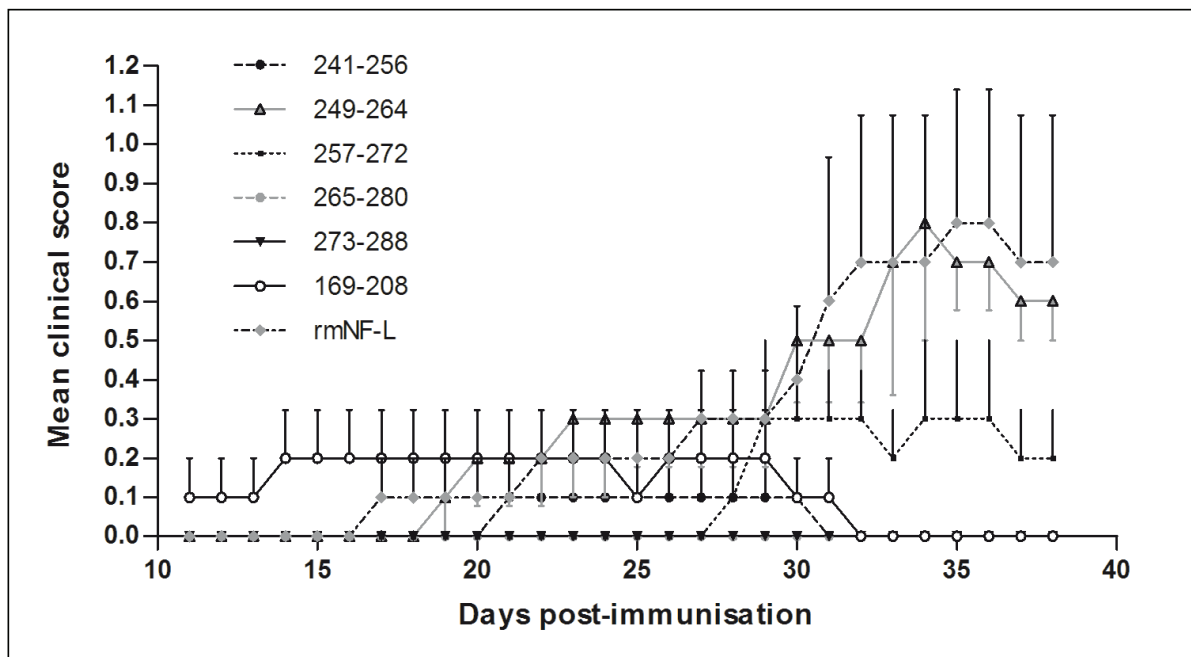
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### Consent

Written informed consent was obtained from the patient for the publication of this report and any accompanying images.





**Supplementary Figure 1. Pathogenicity of immunodominant NF-L epitopes.** ABH mice ( $n=5$  per group) were immunised with NF-L peptides spanning the immunodominant regions (amino acids 241 to 288 and 169 to 208). Mice were immunised subcutaneously with 200  $\mu\text{g}$  rmNF-L protein or NF-L peptides emulsified in complete Freund's adjuvant containing *Mycobacterium tuberculosis*. Plots show the mean  $\pm$  standard error of the mean daily clinical score.

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