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Mannan-conjugated myelin peptides prime non-pathogenic Th1 and Th17 cells and ameliorate experimental autoimmune encephalomyelitis

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1

2 **Short running title:** Peptide-specific immunotherapy in EAE

3

4 **Keywords:** EAE/MS, dendritic cells, T cells, anergy/suppression/tolerance, CNS

5 **Abstract**

6 Antigen presenting cells are critical for regulating immune responses. We
7 tested mannan-peptide conjugates for targeting myelin peptides to APC to induce T
8 cell tolerance and resistance to experimental autoimmune encephalomyelitis (EAE).
9 Myelin peptides conjugated to mannan in oxidized (OM) or reduced (RM) forms
10 protected mice against EAE in prophylactic and therapeutic protocols, with OM-
11 conjugated peptides giving best results. Protection was peptide-specific and
12 associated with reduced antigen-specific T cell proliferation, but not alterations in
13 Th1, Th17 or Treg cell differentiation or T cell apoptosis compared to EAE controls.
14 OM-MOG-loaded bone marrow-derived DC showed up-regulated expression of co-
15 stimulatory molecules, reduced PD-L1 expression and enhanced CD40-inducible IL-12
16 and IL-23 production, features consistent with immunogenic DC. OM-MOG induced
17 active T cell tolerance because i.d. administration or passive transfer of OM-MOG-
18 loaded DC suppressed ongoing EAE, while OM-MOG-vaccinated mice did not reduce
19 the proliferation of transferred MOG-specific T cells. As *in vivo*, MOG-specific T cells
20 cultured with OM-MOG-loaded DC showed reduced proliferation and equal Th1 and
21 Th17 cell differentiation as those with MOG-loaded DC, but surprisingly cytokine
22 production was unresponsive to CD40 engagement. Impaired effector T cell function
23 was further evidenced in spinal cord sections from OM-MOG-vaccinated EAE mice,
24 where markedly reduced numbers of CD3⁺ T cells were present, restricted to
25 leptomeninges and exceptional parenchymal lesions. Our results show that mannan-
26 conjugated myelin peptides protect mice against EAE through the expansion of
27 antigen-specific Th1 and Th17 cells with impaired proliferation responses and APC-
28 induced co-stimulatory signals that are required for licensing them to become fully
29 pathogenic T cells.

30

32 Introduction

33 Multiple sclerosis (MS) is a chronic inflammatory disease of the CNS that
34 shows autoimmune features and causes demyelination, early axonal injury and
35 progressive neurological impairment. Current treatments induce non-specific immune
36 or T cell suppression but are only partially effective and are often associated with
37 adverse effects. Selective treatments that target the immune cells involved in the
38 pathological processes are needed. One approach is to identify disease-associated
39 auto-antigens and to induce antigen-specific T cell tolerance. Several experimental
40 strategies are effective in switching antigen-specific pro-inflammatory IFN- γ -
41 producing Th1 cell responses to IL-4-producing Th2 cell responses or inducing
42 regulatory T cells or anergy, including oral (Chen, Kuchroo, Inobe, Hafler, and Weiner,
43 1994) and inhalation (Burkhart, Liu, Anderton, Metzler, and Wraith, 1999) tolerance
44 or altered peptide ligands (APL) which compete for Ag-specific T cell receptors and
45 induce differential signaling through the T cell receptor (TCR) resulting in tolerance
46 (Vergelli, Hemmer, Utz, Vogt, Kalbus, Tranquill, Conlon, Ling, Steinman, McFarland,
47 and Martin, 1996), (Evavold and Allen, 1991). However, these approaches have not
48 translated well into the clinic due to lack of efficacy or severe adverse effects such as
49 hypersensitivity responses (Hafler, Kent, Pietrusewicz, Khoury, Weiner, and Fukaura,
50 1997), (Bielekova, Goodwin, Richert, Cortese, Kondo, Afshar, Gran, Eaton, Antel,
51 Frank, McFarland, and Martin, 2000), (Kappos, Comi, Panitch, Oger, Antel, Conlon,
52 and Steinman, 2000). Another approach is to harness the immune regulatory
53 properties of professional antigen-presenting cells (APC), particularly dendritic cells
54 (DC), which are critical not only for the induction of adaptive immune responses to
55 foreign antigens but also for the maintenance of immune tolerance to self (Steinman,
56 2008).

57 C-type lectin receptors such as DEC205 and the mannose receptor (MR,
58 CD206) recognize glycosylated self and nonself-antigens, are highly expressed on APC
59 and have been used successfully for targeting peptide antigens to APC for
60 presentation to T cells and modulating immune responses (McGreal, Miller, and
61 Gordon, 2005). In one approach, *in vivo* targeting of antigens selectively to steady
62 state (immature) DC by fusing them to an antibody against the DEC205 endocytosis
63 receptor (Mahnke, Guo, Lee, Sepulveda, Swain, Nussenzweig, and Steinman, 2000)
64 induced peripheral T cell tolerance in mice (Hawiger, Inaba, Dorsett, Guo, Mahnke,
65 Rivera, Ravetch, Steinman, and Nussenzweig, 2001). The observation that
66 simultaneous activation of CD40 with FGK 45 agonistic CD40 antibody changed the
67 outcome from tolerance to prolonged T cell activation and immunity supported the
68 concept of immature DC being involved in the induction of tolerance (Hawiger *et al.*,
69 2001). Mice treated with anti-DEC-205 antibody fused to myelin self-antigens myelin
70 oligodendrocyte glycoprotein (MOG) (Hawiger, Masilamani, Bettelli, Kuchroo, and
71 Nussenzweig, 2004) or proteolipid protein (PLP) (Stern, Keskin, Kato, Waldner,
72 Schallenberg, Anderson, von, Kretschmer, and Strominger, 2010) peptides developed
73 impaired T cell responses to antigen and showed resistance to EAE induction. In other
74 approaches, peptide antigens were targeted to the MR which is expressed at high
75 levels on APC and captures and presents soluble ligands with selectivity for heavily
76 glycosylated proteins on the surface of yeasts, bacteria and parasites (Sallusto, Cella,
77 Danieli, and Lanzavecchia, 1995). Peptides that are mannosylated with added sugar

78 units (Engering, Cella, Fluitsma, Brockhaus, Hoefsmit, Lanzavecchia, and Pieters,
79 1997), (Tan, Mommaas, Drijfhout, Jordens, Onderwater, Verwoerd, Mulder, van der
80 Heiden, Scheidegger, Oomen, Ottenhoff, Tulp, Neefjes, and Koning, 1997) or
81 chemically conjugated to the mannan polysaccharide (Apostolopoulos, Pietersz,
82 Gordon, Martinez-Pomares, and McKenzie, 2000) show greatly enhanced
83 presentation by major histocompatibility complex (MHC) class II and I to T cells.
84 However, mannosylated peptides did not promote immune responses *in vivo*.
85 Instead, immunization of mice with a mannosylated myelin autoantigen, PLP₁₃₉₋₁₅₁ in
86 the presence of complete adjuvant containing *Mycobacterium tuberculosis*, showed
87 reduced T cell proliferation responses, impaired delayed-type hypersensitivity (DTH)
88 responses and protected mice against the induction of EAE following immunization
89 with PLP₁₃₉₋₁₅₁ (Luca, Kel, van, Wouter, Koning, and Nagelkerken, 2005), (Kel,
90 Oldenampsen, Luca, Drijfhout, Koning, and Nagelkerken, 2007), (Kel, Slutter,
91 Drijfhout, Koning, and Nagelkerken, 2008). These findings again support the
92 participation of APC in mediating tolerance to mannosylated self-antigens and
93 indicate that targeting antigens to MR may be a powerful strategy to suppress
94 autoimmune responses.

95 Mannan is a yeast polysaccharide that acts as a pathogen-associated
96 molecular pattern (PAMP) and at high concentrations stimulates the activation of
97 macrophages (Tada, Nemoto, Shimauchi, Watanabe, Mikami, Matsumoto, Ohno,
98 Tamura, Shibata, Akashi, Miyake, Sugawara, and Takada, 2002), induces the
99 phenotypic maturation of DC in a Toll-like receptor 4-dependent manner (Sheng,
100 Pouniotis, Wright, Tang, Lazoura, Pietersz, and Apostolopoulos, 2006) and enhances
101 antigen presentation and immune responses (Apostolopoulos *et al.*, 2000). Our
102 studies have shown that conjugation of the human tumour antigen mucin 1 (MUC1)
103 to mannan in its oxidized (OM) or reduced (RM) forms leads to its efficient
104 presentation by MHC class I or MHC class II and the induction of T1 or T2 immune
105 responses respectively, with OM-MUC1 giving the best IFN- γ -producing cytotoxic T
106 cell responses and protection against tumour formation (Apostolopoulos *et al.*, 2000),
107 (Apostolopoulos, Pietersz, Loveland, Sandrin, and McKenzie, 1995). In view of these
108 results, in this study we investigated the potential of conjugating mannan to self-
109 antigens as a possible strategy for diverting myelin-specific T cell responses towards
110 an immunomodulatory profile and reducing the susceptibility of mice to EAE. We
111 show that mannan-conjugated myelin antigens induced peptide-specific T cell
112 tolerance and strongly ameliorated the clinical signs of EAE when administered to
113 mice in prophylactic (vaccination) and therapeutic protocols. Surprisingly, however,
114 tolerance was not associated with immune deviation of the effector T cell response or
115 the induction of regulatory T cells but with the efficient induction of antigen-specific
116 Th1 and Th17 cells that were anergic to re-stimulation with cognate antigen and
117 showed marked reduction of encephalitogenic potential.

118

119 **Materials and Methods**

120 *Mice*

121 C57BL/6 (CD45.2), SJL/J and C57BL/6-Tg(Tcra2D2, Tcrb2D2)1Kuch/J (2D2)
122 mice were purchased from the Jackson Laboratory. C57BL/6 expressing EGFP under
123 the actin promoter, TgN(act-EGFP)OsbC14-Y01-FM131, (TgEGFP) were kindly

124 provided by Masaru Okabe (Osaka University). CD45.1 congenic 2D2 C57BL/6 mice
125 were kindly provided by Burkhard Becher and Melanie Greter (University of Zurich).
126 Mice were kept under specific pathogen-free conditions in the experimental animal
127 unit of the Hellenic Pasteur Institute. All animal procedures were performed to
128 minimize suffering and conformed to the principles of the three Rs (replacement,
129 refinement and reduction) following the guidelines of the EU directive for animal
130 research 2010/63/EU. Experimentation licences were provided by the General
131 Secretariat of Agricultural Economy and Veterinary Medicine of the Greek State
132 according to the presidential directive 160/91. The reporting of the animal
133 experiments in this study follows the ARRIVE guidelines.

134

135 *Synthesis of myelin peptides*

136 Peptides (murine MOG₃₅₋₅₅, PLP₁₃₉₋₁₅₁, PLP₁₇₈₋₁₉₁, myelin basic protein (MBP)
137 ₈₃₋₉₉) were synthesized by Fmoc/tBu methodology using the acid sensitive 2-
138 chlorotriyl chloride (CLTR-Cl) resin (0.6-1.0 mmolCl⁻/g) and N^α-Fmoc (9-
139 fluorenylmethyloxycarbonyl)-protected amino acids (Tselios, Probert, Daliani,
140 Matsoukas, Troganis, Gerathanassis, Mavromoustakos, Moore, and Matsoukas,
141 1999). The final products were further purified using semi-preparative reverse
142 phase-high performance liquid chromatography (RP-HPLC). The purity of peptides
143 was >95% as determined by analytical RP-HPLC and electron spray ionization-mass
144 spectrometry. All peptides for conjugation with mannan were synthesized using (Lys-
145 Gly)₅ at the N-terminus which acts as a linker between the peptide and mannan. The
146 synthesis of the polypeptide [(Lys-Gly)₅-(Glu, Ala, Tyr, Lys) (POL) was based on the
147 synthesis of GA (Sela and Mozes, 2004) and involved the polymerization of five
148 benzotriazolyl esters derived from alanine, γ-tert-butyl-glutamate, N^ε-
149 butyloxycarbonyl-lysine, O-tert-butyl-tyrosine and N^α-butyloxycarbonyl-[(N^ε-
150 butyloxycarbonyl-lysine)-glycine]₅. The side chain-protected units were combined in
151 an average molar fraction 1 for [(Lys-Gly)₅], 1.51 for (Glu), 4.95 for (Ala), 1 for (Tyr),
152 3.54 for (Lys) in a ratio of 4.95. The 1-hydroxybenzotriazole (5.85 mmol) and N,N'-
153 diisopropylcarbodiimide (4.29 mmol) were added as coupling reagents in
154 dimethylformamide solvent. The mixture was left to react for 72 h at room
155 temperature, the solvent was removed on a rotary evaporator and the obtained oily
156 product was precipitated from water as an amorphous pale yellow solid. The linear
157 protected polypeptide was treated with 90% trifluoroacetic acid (TFA) in
158 dichloromethane in the presence of 0.3% triethylsilane, anisole and H₂O as
159 scavengers for 5 hours at room temperature. The solvents were removed on a rotary
160 evaporator and the obtained oily product was precipitated from cold dry diethyl
161 ether as amorphous light yellow solid. The crude peptide product was further
162 purified by semi-preparative RP-HPLC: (column: Nucleosil C18, 5 μm, 4.6x250 mm),
163 eluents: A, 0.08% TFA/H₂O, B, 0.08% TFA/acetonitrile, gradual gradient: from 10% to
164 60% B in 45 min, flow rate: 3 ml/min, detection 230 nm, 254 nm, 277 nm. All the
165 fractions between 13-15 min were collected, lyophilized and passed through a pre-
166 packed column Sephadex G-25 Medium to remove the low molecular weight
167 contaminants (Mr < 1000).

168

169 *Conjugation of peptides to mannan*

170 Peptide-mannan conjugation was achieved as previously described

171 (Apostolopoulos, Pietersz, and McKenzie, 1996). Briefly, mannan (poly-mannose
172 from *Saccharomyces cerevisiae*; Sigma-Aldrich Ltd) in phosphate buffer (pH 6.0) was
173 oxidized to polyaldehydes by treating with sodium periodate. The conjugation of
174 peptides to oxidized mannan (OM) was achieved via Schiff base formation between
175 the free amino groups of Lys and the aldehydes of OM in bicarbonate buffer (pH
176 9.0). Reduction of the free aldehydes and Schiff base to alcohols and amines
177 respectively, to form reduced mannan (RM), was achieved by treating the OM-
178 peptide complex with sodium borohydride. Conjugates were analysed for
179 conjugation efficiency by capillary electrophoresis as previously reported (Tselios,
180 Lamari, Karathanasopoulou, Katsara, Apostolopoulos, Pietersz, Matsoukas, and
181 Karamanos, 2005).

182

183 *Administration of mannan-peptide conjugates to mice*

184 In a prophylactic vaccination protocol, groups of female C57BL/6 or SJL/J
185 mice (6-8 weeks old) were injected intradermally (i.d.) on the flanks with 100 µl PBS
186 containing OM-MOG₃₅₋₅₅ (OM-MOG), RM-MOG₃₅₋₅₅ (RM-MOG), OM-PLP₁₇₈₋₁₉₁, OM-
187 POL, RM-POL or an unconjugated mixture of OM and MOG₃₅₋₅₅ (MOG) (in the
188 C57BL/6 strain), and OM-PLP₁₃₉₋₁₅₁ or OM-MBP₈₃₋₉₉ (in the SJL/J strain) (all 30 µg
189 peptide equivalent/injection and 700 µg mannan equivalent/injection). As controls,
190 age-matched groups of mice were vaccinated with OM or RM (700 µg),
191 unconjugated peptide (30 µg) or PBS vehicle. Three consecutive injections were
192 performed at 15-day intervals. Immunization for the induction of EAE was performed
193 15 days after the last i.d. injection. In therapeutic administration protocols, groups of
194 female C57BL/6 mice (6-10 weeks old) were injected i.d. with 100 µl PBS containing
195 peptide conjugates or controls, as above, at the time of immunization (day 0) and 7
196 later, or after the onset of clinical signs, as indicated.

197

198 *EAE induction*

199 MOG-EAE was induced in 6-8 week-old female C57BL/6 mice (17-18 week-old
200 bone marrow chimeric mice) by subcutaneous (s.c.) tail-base injection of 30 µg of
201 murine MOG₃₅₋₅₅ in 100 µl PBS emulsified in an equal volume of complete Freund's
202 adjuvant (CFA), 15 days after the third vaccine injection. Mice received
203 intraperitoneal (i.p.) injections of 200 ng of *Bordetella pertussis* toxin (PTx) (Sigma-
204 Aldrich) at the time of immunization and 48 h later. In some experiments an
205 agonistic rat IgG antibody to mouse CD40 (FGK45) (Rolink, Melchers, and Andersson,
206 1996) kindly provided by Antonius Rolink, University of Basel) was administered i.p.
207 (90 µg / mouse) from day 2 post-immunization and thereafter twice-weekly for 2
208 weeks (total of 5 injections). PLP-EAE was induced in female SJL/J mice by s.c. tail-
209 base injection of 150 µg of PLP₁₃₉₋₁₅₁ emulsified in CFA, 15 days after the third
210 vaccine injection, without the administration of PTx. CFA used in all experiments was
211 supplemented with 400 µg/injection of H37Ra *Mycobacterium tuberculosis* (Difco).
212 Mice were monitored daily for the clinical signs of EAE according to the following
213 scores: 0, normal; 1, limp tail; 2, hind limb weakness; 3, hind limb paralysis; 4,
214 forelimb paralysis; and 5, moribund or dead (0.5 gradations represent intermediate
215 scores). Moribund animals were euthanized and given a clinical score of 5 for the
216 remaining days of the experiment. All mice were allowed free access to food and
217 water throughout the experiments.

218

219 *TgEGFP bone marrow chimeric mice*

220 Bone marrow cells were flushed from tibia and femur bones of naïve female
221 TgEGFP mice and red blood cells were lysed using ammonium chloride potassium
222 (ACK) buffer. Washed cells were transplanted i.v. to lethally irradiated female
223 C57BL/6 recipients (5-6 weeks old) (5×10^6 / mouse). Efficiency of bone marrow
224 engraftment was determined after 6 weeks of recovery by FACS analysis for GFP
225 expression in peripheral blood cells. Mice showing $>70\%$ EGFP⁺ blood cells were used
226 for vaccination with OM-MOG or PBS control and induction of MOG-EAE and were
227 sacrificed for immunohistochemical analysis of TgEGFP cell distribution in peripheral
228 (gut, lung) and CNS tissues, and spinal cord lesion development (see below) in tissues
229 taken at the peak of clinical disease in the control group.

230

231 *Bone marrow-derived DC culture and in vivo transfer*

232 Bone marrow-derived DC were isolated as previously described (Lutz,
233 Kukutsch, Ogilvie, Rossner, Koch, Romani, and Schuler, 1999). Bone marrow cells
234 were isolated from naïve C57BL/6 mice as described above and cultured at 3×10^6
235 cells per plate in 10 ml of RPMI 1640 supplemented with 10% heat inactivated FBS
236 for 9 days. Cells were treated with 20 ng/ml of recombinant GM-CSF (Sigma) on days
237 0, 3 and 6 to stimulate differentiation. Adherent cells were harvested on day 9 and
238 loaded *in vitro* with OM-MOG, MOG, OM or PBS (10 µg/ml peptide equivalent and
239 233 µg/ml mannan equivalent). After 24 h, the phenotype of the CD11c⁺ cells was
240 evaluated by staining with fluorochrome-labeled antibodies for cell surface markers
241 (MHC class II (anti-I-A/I-E), CD8a, CD80, CD86, CD40) and intracellular staining for
242 PD-L1 using a FACSCalibur cytometer and CellQuest software (BD). Production of IL-
243 23 and IL-12p70 was measured in DC supernatants by ELISA (see below). As a
244 positive control for DC maturation, cells were stimulated with LPS (1 µg/ml) derived
245 from *E. coli* (Sigma). Cells were used in DC-T cell co-culture assays or for adoptive
246 transfer into mice. For *in vivo* experiments, peptide-loaded DC (1×10^6 cells/mouse)
247 were transferred i.v. to C57BL/6 recipient mice on day 8 post-immunization for
248 MOG-EAE induction, just before the onset of clinical signs, and mice were scored
249 daily for clinical signs of EAE.

250

251 *T cell proliferation and death assays*

252 Splenocytes or draining lymph node (DLN) cells were isolated from
253 immunized C57BL/6 mice and cultured for 72 h in RPMI 1640 (Invitrogen Life
254 Technologies) supplemented with 10% heat-inactivated FCS, 50 µM 2-β
255 mercaptoethanol (Sigma), and increasing concentrations of MOG peptide. Cells were
256 stimulated in triplicate at 2×10^6 cells / ml in round-bottom 96-well plates (Costar).
257 Cells were pulsed with 1 µCi / 5×10^5 cells [³H]-thymidine (Amersham Radiochemicals)
258 for the last 16 h of culture. [³H]-Thymidine incorporation was measured by liquid
259 scintillation counting (Wallac). Results are expressed as the stimulation index (SI)
260 calculated from the radioactivity counts per minute (cpm) of cells cultured in the
261 presence of peptide divided by cpm of cells cultured in medium alone.

262 To measure T cell proliferation *in vivo*, splenocytes and DLN cells were
263 isolated from CD45.1 congenic MOG₃₅₋₅₅-specific T cell receptor (2D2) transgenic
264 donor mice (Bettelli, Pagany, Weiner, Linington, Sobel, and Kuchroo, 2003) and

265 labeled with 5-(and 6-) carboxyfluorescein diacetate, succinimidyl ester (CFSE)
266 (CFDA-SE; Molecular Probes). Cells were washed and resuspended at a concentration
267 of 10^7 /ml in PBS. CFSE was added at a final concentration of 5 μ M and incubated for
268 5 min at RT. The reaction was stopped by washing the cells with RPMI 1640 (Life
269 Technologies) containing 10% FCS. Cells (10×10^6 cells/mouse) were injected i.v. in
270 the tail vein of vaccinated, CD45.2 recipient mice on day 2 post-immunization for
271 MOG-EAE induction. On day 7 of EAE DLN cells were isolated and analyzed by flow
272 cytometry.

273 Measurement of CD4⁺ T cells undergoing apoptotic cell death after prolonged
274 stimulation with peptide-loaded DC in DC-T cell co-cultures (assayed at days 6, 7, 9,
275 12 of culture) was made using the FITC Annexin V apoptosis detection kit (BD
276 Pharmingen).

277

278 *Cell phenotyping and cytokine production*

279 For intracellular cytokine staining mononuclear cells stimulated for 3h with
280 PMA and ionomycin in the presence of Brefeldin A, were fixed in 2%
281 paraformaldehyde solution in PBS for 10 min at room temperature and
282 permeabilized with 0.5% wt/vol saponin prior to staining for surface markers and
283 intracellular cytokines using fluorochrome-labeled antibodies (anti-CD4, clone L3T4;
284 anti-CD45.1, clone A20; anti-CD11c, clone HL3; anti-CD8a, clone 53-6.7; anti-CD40,
285 clone 3/23; anti-CD80, clone 16-10A1; anti-CD86, clone GL1; anti-I-A/I-E (MHC class
286 II), clone 2G9; anti-IL-17, clone TC11-18H10; anti-IFN- γ , clone XMG1.2; anti-FoxP3,
287 clone; FJK-16s; anti-PD-L1 (CD274), clone MIH5; IgG1 isotype control, clone R3-34;
288 IgG2a isotype control, clone B39-4 all from BD Biosciences). Data was acquired and
289 analyzed with a FACSCalibur cytometer and CellQuest software (BD) and with
290 FlowJo, version 10.0.6 (Tree Star). The production of IL-23 and IL-12p70 by DC was
291 measured in cell supernatants by mouse ELISA Ready-SET-Go kits (e-Bioscience)
292 (sensitivity for IL-12p70 at 15 pg/ml and for IL-23 at 8 pg/ml).

293

294 *DC-T cell antigen presentation assays*

295 Splenocytes and DLN cells were isolated from 2D2 mice and co-cultured at $2 \times$
296 10^5 cells/well in 96 well plates with peptide-loaded DC at different ratios. All
297 combinations of responder and stimulator cells were cultured in triplicate for 72 h.
298 Cell proliferation was measured by [³H]-thymidine incorporation and cytokine
299 production was measured by intracellular cytokine staining as described above. In
300 experiments with CD40 costimulation, cells were incubated for 72 h in the absence
301 or presence of agonistic anti-CD40 antibody FGK45 (10 μ g/ml).

302 For T helper cell polarization experiments, splenocytes and peripheral LN
303 cells were isolated from 2D2 mice and co-cultured with peptide-loaded, *in vitro*
304 matured, bone marrow-derived DC. The polarization conditions were: Th1 culture;
305 IL-12 (10 ng/ml, Peprotech), anti-IL4 (10 μ g/ml, R&D systems): Th17 culture; IL-6 (50
306 ng/ml, Peprotech), IL-23 (10 ng/ml, eBioscience), TGF- β (5 ng/ml, Peprotech), anti-
307 IFN- γ (10 μ g/ml, R&D systems): Treg culture; TGF- β (10 ng/ml, Peprotech), anti-IFN- γ
308 (10 μ g/ml, R&D systems), anti-IL-4 (10 μ g/ml, R&D systems). On day 3 of culture, Th1
309 and Treg polarizing cultures were supplemented with IL-2 (10 ng/ml, Peprotech) and
310 Th17 polarizing cultures were supplemented with IL-23 (10 ng/ml). Cells were
311 harvested on day 5 for FACS analysis of lineage markers.

312

313

314 *Histopathological analysis*

315 Mice were transcardially perfused with ice-cold 4% paraformaldehyde in PBS
316 under deep anaesthesia. CNS tissues were post-fixed in the same fixative for 3 h at
317 4°C and processed for standard histopathological analysis. Inflammation was
318 visualized by staining with H&E and demyelination was demonstrated by Luxol Fast
319 Blue/periodic acid-Schiff staining. Quantification of inflammation and demyelination
320 was done in a blinded manner. Inflammation in the spinal cord was determined by
321 absolute true quantification; the numbers representing inflammatory
322 infiltrates/mm² of tissue. In the brain a semi-quantitative scoring of inflammation
323 was used in which, 0.5 means single perivascular infiltrates and 1 means multiple
324 inflammatory infiltrates. Demyelination was also evaluated by semi-quantitative
325 scoring as follows: 0.5: single perivascular sleeves of demyelination, 1: ubiquitous
326 perivascular or subpial demyelination, 2: confluent demyelinated plaques, 3:
327 profound focal demyelination, involving about 1/2 of the spinal cord white matter at
328 least in one spinal cord segment, 4: extensive demyelination, for instance complete
329 demyelination of spinal cord white matter at least in one segment of the spinal cord.

330

331 *Immunohistochemistry*

332 Immunohistochemistry was performed on paraffin sections (4 µm) to evaluate tissue
333 distribution of TgEGFP bone marrow-derived immune cells, CD3⁺ T cells as well as
334 production of the p22phox subunit of NADPH oxidase and iNOS by inflammatory
335 macrophages. Antigen retrieval in paraffin sections was performed in a food steamer
336 in citrate buffer (pH 6) for 40 min. The primary Abs used were polyclonal rabbit anti-
337 GFP IgG (1/200; Molecular Probes; A11122), monoclonal rabbit anti-CD3 (1/2000;
338 Neomarkers; RM-9107), polyclonal rabbit anti-p22phox (1/100; Santa Cruz Biotech;
339 sc-20781) and polyclonal rabbit anti-rat iNOS (1/375; Chemicon; AB1631) followed
340 by biotinylated secondary anti-IgG Ab (1/500; Vector laboratories). An avidin-biotin
341 complex was used for detection of the biotinylated Abs and immune complexes
342 were visualized by incubation with 3,3'-diaminobenzidine tetrachloride (DAB) (both
343 from Vector laboratories).

344

345 *Statistical analysis*

346 All statistical analyses were performed with Sigma Stat 3.5, Sigma Plot 11 and
347 Microsoft Excel. All data are given as mean ± standard error of the mean (SEM).
348 Student's t test and Kruskal-Wallis test were used. Results were considered
349 statistically significant when $p < 0.05$.

350

351

352

353 **Results**

354 *Administration of OM- or RM-MOG prophylactically or therapeutically protects mice* 355 *against MOG-EAE*

356 To examine the effect of the H-2^b binding MOG₃₅₋₅₅ peptide (MOG) conjugated
357 to oxidized mannan (OM-MOG) or reduced mannan (RM-MOG) upon the
358 development of MOG-EAE, we delivered peptide conjugates to C57BL/6 (H-2^b) mice in

359 a prophylactic vaccine protocol prior to the induction of EAE by immunization with
360 MOG emulsified in CFA and administration of *Bordetella pertussis* toxin
361 (MOG/CFA/PTx). Three intradermal (i.d.) injections of OM- or RM-MOG, but not
362 unconjugated OM, RM, MOG or a mixture of unconjugated OM and MOG (OM/MOG)
363 in dilute soluble form and spaced at 15 day intervals prior to immunization with
364 MOG/CFA/PTx, protected mice from the subsequent development of EAE compared
365 to PBS (Fig. 1a, Supplementary Fig. 1a), with OM-MOG giving increased protection
366 when compared to RM-MOG (Fig. 1a).

367 We next tested the effects of mannan-conjugated peptides when
368 administered in therapeutic protocols in the presence of adjuvants and PTx. In one
369 approach peptide conjugates were injected i.d. on the day of EAE induction with
370 MOG/CFA/PTx and again seven days later. OM-MOG and RM-MOG, but not
371 unconjugated OM, RM, or MOG, showed protective effects compared to PBS. With
372 this protocol protection by OM-MOG and RM-MOG showed statistically similar levels
373 of protection up to the last time point tested (Fig. 1b). In a second approach we
374 injected OM-MOG after the onset of MOG-EAE, when mice have reached at least
375 clinical score 2 and found that OM-MOG rapidly reduced the severity of ongoing
376 disease, with the clinical condition of the experimental animals progressively
377 improving upon each injection (Fig. 1c).

378 In conclusion, i.d. administration of OM-MOG strongly protected mice against
379 clinical MOG-EAE in all prophylactic and therapeutic treatment protocols tested,
380 even in the presence of PTx and activated encephalitogenic T cells, and OM-
381 conjugated peptides were chosen for subsequent experiments investigating the
382 mechanism of tolerance induced by mannan-conjugated self-antigens.

383

384 *Vaccination with OM-MOG and RM-MOG protects mice against EAE neuropathology*

385 To determine whether amelioration of the clinical signs of MOG-EAE by OM-
386 and RM-MOG was associated by less severe neuropathology, vaccinated mice were
387 sacrificed 24 days after EAE induction for neuropathological analysis of spinal cord
388 and brain tissues (Fig. 1d and e). PBS-treated mice showed substantial mononuclear
389 cell infiltration and extensive demyelination in the spinal cord. In contrast, mice
390 vaccinated with OM-MOG showed reduced inflammatory cell infiltration and little or
391 no demyelination in the spinal cord. RM-MOG-vaccinated mice showed reduced
392 neuropathology compared to PBS-treated mice. Mice vaccinated with unconjugated
393 MOG, OM or RM showed equivalent spinal cord pathology to PBS-treated animals
394 (Fig. 1d). A quantitative assessment of demyelination and inflammation in each
395 individual mouse confirmed that CNS pathology in OM-MOG-vaccinated mice was
396 significantly reduced compared to PBS-treated mice (Fig. 1e).

397 We conclude that the administration OM-MOG as a prophylactic vaccine in
398 mice inhibits the accumulation and infiltration of immune cells into the CNS
399 parenchyma during EAE and the development of inflammatory and demyelinating
400 lesions.

401 *Protection from EAE by mannan-conjugated peptides is peptide-specific*

402 To determine whether protection by mannan-peptide conjugates can apply
403 to other CNS antigens and also whether it depends on peptide specificity, we next
404 performed peptide criss cross experiments where we complemented the MOG-EAE

405 model with another EAE model in SJL/J mice (H-2^S) in which disease is induced by
406 immunization with the H-2^S binding peptide, proteolipid protein 139-151 (PLP)
407 (McRae, Kennedy, Tan, Dal Canto, Picha, and Miller, 1992). OM was conjugated to
408 PLP (OM-PLP) or another H-2^S binding peptide, myelin basic protein 83-99, which
409 itself is capable of inducing EAE when used to immunize SJL/J mice (Miller, Karpus,
410 and Davidson, 2010), as peptide control. Groups of SJL/J mice were vaccinated, as
411 above, with OM-PLP₁₃₉₋₁₅₁, OM-MBP₈₃₋₉₉ or PBS. Fifteen days after the last injection,
412 EAE was induced by immunization with PLP₁₃₉₋₁₅₁ in CFA (PLP/CFA). Control mice
413 vaccinated with PBS developed acute severe clinical signs typical of PLP-EAE (Fig. 2a).
414 As in the MOG-EAE model, mice vaccinated with mannan-conjugated to cognate
415 peptide, OM-PLP₁₃₉₋₁₅₁, but not irrelevant peptide OM-MBP₈₃₋₉₉, were strongly
416 protected against PLP-EAE (Fig. 2a). In the crossover experiment using the MOG-EAE
417 model, OM was conjugated to another H-2^b-binding peptide, PLP 178-191 which
418 itself is capable of inducing EAE in C57BL/6 mice (Tompkins, Padilla, Dal Canto, Ting,
419 Van, and Miller, 2002), as peptide control. Groups of C57BL/6 mice were vaccinated
420 with OM-MOG₃₅₋₅₅, OM-PLP₁₇₈₋₁₉₁ or PBS and fifteen days after the last injection EAE
421 was induced by immunization with MOG/CFA/PTx. As predicted, mice vaccinated
422 with OM-MOG₃₅₋₅₅, but not OM-PLP₁₇₈₋₁₉₁, were strongly protected against MOG-EAE
423 (Fig. 2b).

424 In a second approach we used the MOG-EAE model to test the prophylactic
425 efficacy of OM and RM conjugated to a polypeptide mixture of synthesized randomly
426 from four amino acids (L-Glutamic acid, L-Lysine, L-Alanine and L-Tyrosine) based on
427 GA [30] (Glu, Ala, Tyr, Lys) (POL), which has been shown to induce T cell tolerance in
428 MOG-EAE and is used in the treatment of MS (Sela *et al.*, 2004). Groups of C57BL/6
429 mice were vaccinated, as above, with OM- and RM-POL conjugates or unconjugated
430 OM, RM or POL with or without the [(Lys-Gly)₅] linker. Fifteen days after the last
431 injection, mice were immunized with MOG/CFA/PTx to induce EAE. Under these
432 conditions all mice, including those injected with POL, developed EAE with
433 equivalent severity and timing as PBS-injected controls (Supplementary Fig. 1b).

434 We conclude that the induction of T cell tolerance by mannan-conjugated
435 peptides applies in different EAE models and is peptide-specific, at least during the
436 time frame of our experiments, applying only to the peptide used to immunize mice
437 for the induction of EAE.

438

439 *Antigen-specific responses of T cells exposed to OM-MOG show reduced proliferation*
440 *but efficient expansion of Th1, Th17 and Treg populations*

441 To determine whether the therapeutic effects of OM-MOG in MOG-EAE were
442 associated with alteration of antigen-specific immune responses, we compared T cell
443 responses to MOG immunization in mice that had been previously treated with the
444 different vaccination components. Mice were immunized with MOG/CFA/PTx 15
445 days after the last vaccine injection and DLN and/or splenocytes were isolated at
446 different time points prior to and after the onset of clinical signs in the PBS-treated
447 group, re-stimulated with MOG peptide *in vitro* and analyzed for proliferation and
448 effector T cell cytokine production. Time points analyzed were pre-onset (days 7 &
449 10 post-immunization), peak of EAE (day 15) and chronic EAE (day 25). Antigen-
450 specific proliferation responses were measured in splenocytes isolated 25 days post-
451 immunization with MOG/CFA/PTx and restimulated with MOG₃₅₋₅₅ peptide *in vitro*.

452 Lymphocytes from mice vaccinated with OM-MOG or RM-MOG, but not OM, RM, or
453 MOG showed reduced antigen-induced proliferation responses compared to PBS
454 (Fig. 3a, and data not shown). Similar results were obtained using DLN cells isolated
455 from vaccinated mice 10 days post-immunization with MOG/CFA and restimulated
456 with MOG₃₅₋₅₅ peptide *in vitro* (data not shown). The percentages of total CD4⁺ T cells
457 isolated from the DLN and spleens of mice from different vaccination groups were
458 similar following immunization (Supplementary Fig. 2).

459 We next examined whether T cells isolated from naïve MOG₃₅₋₅₅-specific T
460 cell receptor (2D2) transgenic donor mice (Bettelli *et al.*, 2003), and transferred into
461 recipient mice that had been previously vaccinated with OM-MOG, would acquire
462 tolerance at a time when OM-MOG was no longer present in the recipient mouse.
463 Splenocytes were isolated from 2D2 CD45.1 donor mice, labelled *ex vivo* with CFSE
464 and adoptively transferred into vaccinated CD45.2 recipient mice, 17 days after the
465 last i.d. injection of OM-MOG and 2 days after immunization with MOG/CFA/PTx.
466 Analysis of fluorescence in the transferred CFSE-labeled 2D2 CD45.1 T cells isolated
467 from DLN 7 days post-immunization showed that, unlike the endogenous cells, there
468 were no differences in the proliferation of exogenously administered cells between
469 mice vaccinated with OM-MOG, OM, MOG or PBS (Fig. 3b). This result indicates that
470 the presentation of OM-MOG to T cells is required for the induction of long-lasting
471 reduction in antigen-specific T cell proliferation capacity and that tolerance is not
472 transferred in the absence of OM-MOG.

473 Unexpectedly, the expansion of effector T helper cell populations was
474 unaffected in OM-MOG-vaccinated mice, as Th1 and Th17 populations were equal to
475 those in control mice. Thus, CD4⁺IFN γ ⁺ and CD4⁺IL-17⁺ DLN (day 7) (Fig. 4c) and
476 CD4⁺IL-17⁺ DLN (Fig. 4d, left panel) and splenocytes (Fig. 4d, right panel) (days 10 and
477 15) showed no differences from control mice or other vaccination groups as
478 measured by intracellular cytokine staining. Also, no IL-4 was detectable upon
479 restimulation of cells from any of the vaccinated groups (data not shown), indicating
480 that there was no overt emergence of Th2 cells in OM-MOG-vaccinated mice. In one
481 experiment, in which DLN were isolated on day 13 post-immunization with
482 MOG/CFA/PTx from mice that were treated with OM-MOG or OM on days 0 and 7
483 post-immunization (therapeutic protocol), the production of IL-10 in CD4⁺ T cells
484 was reduced in OM-MOG-treated mice compared to OM-treated controls (Fig. 3e).

485 The induction of regulatory T cells is a critical mechanism of tolerance in EAE
486 where they limit encephalitogenic T cell expansion and function (Kohm, Carpentier,
487 Anger, and Miller, 2002). We measured the induction of regulatory T cell populations
488 in mice that were vaccinated with the various peptide conjugates and subsequently
489 immunized with MOG/CFA. DLN were isolated 7 days post-immunization and CD4⁺ T
490 cells were analyzed for Foxp3 expression, the signature transcription factor of
491 regulatory T cells. The proportions of CD4⁺ Foxp3⁺ T cells were not significantly
492 altered in OM-MOG-vaccinated mice compared to PBS-, OM- and MOG-vaccinated
493 control groups (Fig. 3f).

494 In conclusion, OM-MOG exposure of T cells induces lasting peripheral T cell
495 tolerance that is associated with reduced antigen-specific proliferation responses but
496 not altered expansion of effector Th1 and Th17 T cells, immune deviation or the
497 induction of regulatory T cell populations.

498

499 *OM-MOG-loaded DC up-regulate maturation markers, Th1 and Th17 polarising*
500 *cytokines, down-regulate PD-L1 and are sufficient to transfer tolerance into EAE*
501 *mice*

502 The MR is expressed by APC, mainly macrophages and DC (Sallusto *et al.*,
503 1995), (Linehan, Martinez-Pomares, Stahl, and Gordon, 1999) and mediates innate
504 activation signals from PAMPs such as mannan (Tada *et al.*, 2002). To investigate
505 whether murine DC can present OM-MOG to T cells and participate in the induction
506 of tolerance in vaccinated mice we cultured bone marrow-derived DC, loaded them
507 with the vaccine components *ex vivo* and used them for antigen presentation assays
508 *in vitro* and adoptive transfer experiments in mice. Peptide-loaded DC were first
509 analysed for the expression of DC maturation markers. CD11c⁺CD8⁻-gated DC, loaded
510 with unconjugated OM or OM-MOG (OM-MOG DC), showed increased expression
511 CD40, CD80 and CD86 compared to those loaded with MOG and PBS, showing levels
512 of CD40 and CD80 at least as high as in LPS-treated DC (1 µg/ml), while proportions
513 of MHC class II-expressing cells were equivalent in the peptide-loaded and control
514 populations (Fig. 4a). These results confirm that OM, in either unconjugated or
515 peptide-conjugated form, induces phenotypic maturation of *in vitro* DC. In contrast,
516 CD11c⁺CD8⁻-gated DC loaded with MOG (MOG DC) did not show upregulation of
517 activation markers compared to control PBS DC confirming that unconjugated
518 peptide does not induce phenotypic maturation of DC (Fig. 4a).

519 We next compared effector cytokine secretion by OM-MOG and MOG DC in
520 the absence or presence of CD40 costimulation by FGK 45 agonistic anti-CD40
521 antibody (Rolink *et al.*, 1996). CD40 is a costimulatory receptor on APC that is
522 triggered by CD40 ligand produced by strongly activated CD4⁺ T cells. It boosts
523 immune responses through the induction of T cell polarizing cytokines and is
524 essential for licensing DC to induce development of functional effector cells (Albert,
525 Jegathesan, and Darnell, 2001), (Fujii, Liu, Smith, Bonito, and Steinman, 2004),
526 (Sporri and Reis e Sousa, 2005), (Iezzi, Sonderegger, Ampenberger, Schmitz,
527 Marsland, and Kopf, 2009) and for the development of MOG-EAE (Becher, Durell,
528 Miga, Hickey, and Noelle, 2001), (Iezzi *et al.*, 2009). OM-MOG DC showed increased
529 production of the Th1 and Th17 polarising cytokines, IL-12 and IL-23 respectively,
530 compared to MOG DC and production in both cultures was increased by CD40
531 engagement with FGK 45 (Fig. 4b). We next tested whether OM-MOG-induced
532 tolerance could be associated with altered production of programmed death ligand-
533 1 (PD-L1), an APC-expressed ligand that negatively regulates TCR signalling via PD-1
534 on T cells and actively terminates antigen-specific T cell responses including those
535 that induce EAE (Carter, Leach, Azoitei, Cui, Pelker, Jussif, Benoit, Ireland, Luxenberg,
536 Askew, Milarski, Groves, Brown, Carito, Percival, Carreno, Collins, and Marusic,
537 2007). Interestingly, we found production of PD-L1 to be decreased in CD11c⁺ OM-
538 MOG DC compared to MOG-, OM- and PBS-loaded DC (Fig. 4e).

539 To directly evaluate whether *in vitro* grown, OM-MOG-loaded DC showing
540 expression of maturation markers and up-regulated Th1 and Th17 polarising
541 cytokines, are sufficient to actively induce tolerance in mice with on-going EAE, we
542 adoptively transferred MOG-, OM-MOG-, OM- or PBS-loaded DC into non-vaccinated
543 recipient mice, 8 days after EAE induction by immunization with MOG/CFA/Ptx.
544 Previous studies have shown that bone marrow-derived DC from Lewis rats with EAE
545 (Xiao, Huang, Yang, Xu, and Link, 2001) or CD11c⁺CD11b⁺ DC isolated from MOG-

546 tolerized mice (Li, Zhang, Chen, Xu, Fitzgerald, Zhao, and Rostami, 2008) are
547 sufficient to transfer peptide-specific resistance to EAE in naïve recipients, a property
548 associated with their immature status. While none of the transferred DC populations
549 prevented disease onset, both MOG DC and OM-MOG DC, and not OM DC or PBS
550 DC, reduced the severity of clinical symptoms during the chronic phase of disease
551 compared to PBS DC (Fig. 4c). Together these results show that immature MOG DC
552 and phenotypically mature OM-MOG DC are both competent and sufficient to
553 actively transfer tolerance and reduce clinical symptoms in mice with on-going EAE,
554 although possibly by different mechanisms.

555

556 *OM-MOG DC induce normal maturation of MOG-specific Th1 and Th17*
557 *responses but reduced T cell proliferation and CD40 costimulation responses*

558 To further assess the ability of OM-MOG DC to present cognate antigen to T
559 cells and to investigate the cellular mechanism of OM-MOG-induced tolerance we
560 performed *in vitro* antigen presentation assays between peptide-loaded DC and
561 MOG-specific 2D2 T cells in the absence of presence of FGK 45 , and measured T cell
562 responses as readout. Bone marrow-derived DC were loaded with peptide
563 conjugates *ex vivo*, co-cultured with lymphocytes from 2D2 transgenic mice under
564 neutral, and in some experiments Th1, Th17 and Treg polarising conditions and T cell
565 proliferation and cytokine responses to MOG were measured. Both MOG and OM-
566 MOG DC induced robust proliferation responses in 2D2 T cells and these responses
567 were further enhanced by the presence of FGK 45. However, proliferation induced
568 by OM-MOG DC was significantly lower than that induced by MOG-loaded DC at two
569 DC-T cell ratios tested, 1:1 (data not shown) and 1:5 (Fig. 5a). MOG and OM-MOG DC
570 also efficiently and equally induced the maturation of Th1 and Th17 effector T cell
571 populations under neutral (Fig. 5b), as well as Th1, Th17 and Treg polarizing
572 (Supplementary Fig. 3), conditions, an effect in line with the observation of normal
573 priming of CD4⁺IL-17⁺ and CD4⁺IFN γ ⁺ T cells in response to MOG immunization in
574 OM-MOG-vaccinated mice (see Fig. 3). Interestingly however, while FGK 45 boosted
575 cytokine production induced by MOG DC, it had no effect on IFN γ or IL-17 production
576 induced by OM-MOG DC (Fig. 5b) suggesting that OM-MOG-stimulated T cells,
577 besides showing reduced antigen-specific proliferation, also show anergy to CD40
578 co-stimulation of effector cytokine production.

579 To investigate whether the reduced proliferation of MOG-specific CD4⁺ T cells
580 observed *in vitro* and *in vivo* was associated with increased T cell death, we assessed
581 cell surface annexin V staining in the DC-T cell co-cultures after prolonged
582 stimulation of 2D2 T cells with peptide-loaded DC. The translocation of annexin V to
583 the cell surface of CD4⁺ T cells is an early event in apoptotic death occurring after cell
584 activation. CD4⁺ T cell surface annexin V staining was not increased in OM-MOG or
585 MOG DC cultures compared to PBS and OM DC cultures when measured at several
586 culture time points (representative culture at day 7 postactivation; Fig. 5c). The
587 finding that OM-MOG DC do not increase apoptosis in MOG-specific T cells indicates
588 that the reduced proliferation of MOG-specific lymphoblasts seen in OM-MOG-
589 vaccinated mice and OM-MOG DC-T cell co-cultures is not due to increased cell
590 death.

591 Since cell types other than DC express MR, including macrophages, some
592 endothelial cells and perivascular microglia (Linehan *et al.*, 1999), we next

593 investigated whether CD40 co-stimulation of APC *in vivo* could change the outcome
594 of EAE resistance in OM-MOG-vaccinated mice. We administered FGK45 (90 µg/
595 injection) i.p. twice-weekly starting 17 days after the last i.d. injection of OM-MOG
596 and 2 days after immunization with MOG/CFA/PTx for the induction of EAE. FGK45
597 administration caused a temporary delay but subsequent enhancement of clinical
598 signs in control PBS-vaccinated mice compared to non-treated controls (Fig. 5d). In
599 contrast, FGK45 did not alter the susceptibility of OM-MOG vaccinated mice to
600 disease. Both FGK45-treated and non-treated OM-MOG-vaccinated mice remained
601 disease-free up to the last time-point studied (Fig. 5d).

602 To investigate whether T cells exposed to OM-MOG DC showed anergy we
603 repeated the proliferation assay in the presence of exogenous IL-2. The proliferation
604 of 2D2 T cells stimulated with OM-MOG DC was significantly increased by IL-2, up to
605 the level shown by control cells cultured with MOG DC. The proliferation of 2D2 T
606 cells stimulated with MOG DC was not further stimulated by IL-2 (Fig. 5e)

607 We conclude that OM-MOG DC efficiently provide the initial activation signals
608 for antigen-specific Th1 and Th17 T cell responses but that these T cells show
609 reduced proliferation responses to MOG and effector cytokine production is not
610 boosted in response to CD40 co-stimulation. This, together with the additional
611 finding that exogenous IL-2 reversed the unresponsiveness of T cells to antigen
612 stimulation shows that OM-MOG induces differential TCR signaling in MOG-specific T
613 cells that results in selective functional anergy.

614
615 *CD3⁺ T cells infiltrate spinal cord leptomeninges and activate inflammatory*
616 *macrophages in small lesions but do not induce clinical signs in OM-MOG-vaccinated*
617 *mice*

618 To understand why the antigen-specific Th1 and Th17 cells that are primed in
619 OM-MOG-vaccinated mice are not able to initiate EAE we next investigated whether
620 T cells are able to infiltrate the parenchymal tissue of the spinal cord and trigger
621 downstream effector mechanisms such as macrophage recruitment and activation.
622 First we attempted to isolate mononuclear cells from the spinal cords of OM-MOG-
623 and PBS-vaccinated EAE mice at the peak of disease in the PBS group (with clinical
624 scores 3.5-4) and to measure the proportions of CD4⁺ T cells. In PBS-vaccinated mice
625 CNS-infiltrating mononuclear cells were readily isolated and contained
626 approximately 15% CD4⁺ T lymphocytes as measured by flow cytometry (data not
627 shown). However, only very low numbers of CNS-infiltrating cells could be recovered
628 from OM-MOG-vaccinated mice and flow cytometry analysis of these cells was not
629 possible.

630 We next investigated the tissue distribution of activated MOG-specific T cells
631 and the extent of immune cell infiltration into the spinal cord during EAE in OM-
632 MOG-vaccinated mice by generating bone marrow chimeric mice. We reconstituted
633 lethally-irradiated C57BL/6 wild-type mice with bone marrow cells isolated from
634 double transgenic TgEGFP x 2D2 mice. Chimeric mice showing ≥70% reconstitution of
635 blood leukocytes by GFP⁺ cells, as measured by flow cytometry, were vaccinated and
636 immunized for MOG-EAE. Immunohistochemical analysis of gut, lung and spinal cord
637 taken at the peak of EAE in the PBS-treated group using anti-GFP antibodies showed
638 bone marrow-derived cells distributed in the lamina propria and submucosa of the
639 gut, Peyer's patches of the ileum, lung tissue and large confluent lesions in the white

640 matter of the spinal cord (Supplementary Fig. 4). Immunostaining of serial sections
641 with anti-CD3 antibody did not reveal any T cells in gut or lung but showed
642 numerous T cells distributed throughout the spinal cord lesions (Fig. 6a). OM-MOG-
643 vaccinated mice showed a similar distribution as controls of GFP-immunoreactive
644 cells in the gut and lung and no CD3⁺ T cells were detected in these tissues. In the
645 spinal cord, consistent with the histopathological analysis (Fig. 1d), the numbers of
646 recruited GFP-immunoreactive (Supplementary Fig. 4) and CD3-immunoreactive cells
647 (Fig. 6a) were markedly reduced compared to EAE controls. Both CD3⁺ T cells and
648 GFP-immunoreactive immune cell infiltrates were restricted to the leptomeninges or
649 occasional small compact lesions in the white matter (Fig. 6a). Interestingly, these
650 lesions were detected in mice that displayed no clinical signs on any day of follow-
651 up.

652 To address whether the reduced antigen-specific T cell proliferation
653 measured in secondary lymphoid organs of OM-MOG-vaccinated mice following
654 MOG immunization, and in 2D2 T cells co-cultured with OM-MOG DC, might be
655 responsible for reduced immune cell infiltration of the spinal cord in OM-MOG-
656 vaccinated mice during EAE we performed double immunofluorescence staining
657 using antibodies to CD3 and Ki67, a marker of cell proliferation. In PBS-treated mice,
658 a small proportion of CD3⁺ T cells distributed throughout the tissue lesions and
659 leptomeninges showed Ki67-immunoreactivity (Fig. 6b), indicating that proliferation
660 of T cells occurs in white matter lesions during EAE. This is consistent with the
661 findings of Wekerle and colleagues that T cells enter the brain in the "migratory"
662 phenotype, encounter their specific antigen at APC in meninges and perivascular
663 spaces, proliferate and acquire the ability to pass the astrocytic glia limitans and
664 invade the CNS parenchyma (Lodygin, Odoardi, Schlager, Korner, Kitz, Nosov, van
665 den Brandt, Reichardt, Haberl, and Flugel, 2013), (Mues, Bartholomaeus, Thestrup,
666 Griesbeck, Wekerle, Kawakami, and Krishnamoorthy, 2013). In OM-MOG-treated
667 mice, similarly small proportions of CD3⁺ T cells showed Ki67-immunoreactivity (Fig.
668 6b) although, as mentioned above, the overall numbers of T cells were markedly
669 reduced compared to EAE controls, and were differentially distributed, being
670 detected only in leptomeninges (Fig. 6a). No double-labeled CD3+Ki67+ cells were
671 detected in the parenchymal lesions in OM-MOG-vaccinated mice.

672 We further investigated whether T cells in OM-MOG-vaccinated mice can
673 trigger downstream effector mechanisms such as macrophage recruitment and
674 activation in the spinal cord, by analyzing the accumulation of p22phox, which is an
675 essential subunit of NADPH oxidases, and iNOS, in sections from mice that had been
676 vaccinated with OM-MOG or PBS, as shown in Fig. 1d. In the active MOG-EAE model
677 used in this study, tissue injury is associated with massive infiltration of the tissue by
678 CD3⁺ T cells and the presence of numerous macrophage-like cells showing p22phox
679 and iNOS expression (Schuh, Wimmer, Hametner, Haider, Van Dam, Liblau, Smith,
680 Probert, Binder, Bauer, Bradl, Mahad, and Lassmann, 2014). Here, PBS-vaccinated
681 mice showed typical spinal cord lesions of MOG-EAE, with numerous CD3⁺ T cells and
682 p22phox- and iNOS-immunoreactive macrophage-like cells (Fig. 6b, first and second
683 columns). Surprisingly, OM-MOG-vaccinated mice also showed sparse infiltration of
684 the spinal cord by CD3⁺ T cells and p22phox- and iNOS-immunoreactive macrophage-
685 like cells which were much fewer in number and, as mentioned above, limited to

686 leptomeninges or to exceptional compact white matter lesions (Fig. 6b, third and
687 fourth columns).

688

689 Overall, these data show that antigen-specific CD3⁺ T cells traffic to the spinal cord in
690 OM-MOG-vaccinated mice during the development of EAE, and locally activate
691 macrophage-like cells to produce reactive oxygen and nitrogen species, although
692 numbers are greatly reduced and mainly restricted to leptomeninges compared to
693 the large confluent white matter infiltrates typical of EAE in PBS-treated control
694 mice.

695

696

697 Discussion

698
699 In this study we describe a method for targeting myelin peptide antigens to APC and
700 inducing robust peptide-specific T cell tolerance in mice, protecting them against the
701 development of EAE when administered as i.d. injection in prophylactic (vaccination)
702 or therapeutic protocols and in the presence of strong immune stimulants such as
703 PTx and agonistic anti-CD40 antibody. EAE-inducing myelin peptide epitopes were
704 synthesized with a (Lys-Gly)⁵ linker and chemically conjugated to the polysaccharide
705 mannan, a ligand for the MR, in its oxidized or reduced form. Specifically, we
706 conjugated mannan to H-2^b binding (MOG₃₅₋₅₅) and H-2^s binding (PLP₁₃₉₋₁₅₁) myelin
707 peptides and show that they protect mice against the induction of EAE in two
708 different models, a chronic form induced in C57BL/6J (H-2^b) mice by immunization
709 with MOG/CFA/PTx and a relapse-remitting form induced in SJL/J (H-2^s) mice by
710 immunization with PLP/CFA, respectively. Tolerance in both models was peptide
711 specific and, as further studied in the MOG-EAE model, was associated with reduced
712 antigen-specific T cell proliferation but not changes in differentiation of IFN- γ -
713 producing Th1, IL-17-producing Th17 cells or regulatory T cells. However, we
714 identified resistance in antigen-specific Th1 and Th17 towards CD40-mediated co-
715 stimulatory signals from APC. This dissociation between proliferation and cytokine
716 production in T cell responses indicates that the presentation of OM-MOG by APC
717 results in differential signaling through the TCR on MOG-specific T cells compared to
718 presentation of MOG, and that this results in partial T cell anergy. Indeed, CD3⁺ T
719 cells and activated macrophages accumulated in the leptomeninges of the spinal
720 cord in OM-MOG-vaccinated mice after immunization with MOG, suggesting that
721 MOG-specific T cells recognize target tissue and initiate inflammation. However,
722 markedly reduced numbers of T cells infiltrated the CNS parenchyma to form
723 inflammatory lesions compared to PBS-vaccinated EAE controls, and mice that
724 developed lesions did not necessarily develop clinical signs of EAE. Overall, our
725 results suggest that OM-MOG induces the expansion of Th1 and Th17 T cells that
726 show impaired proliferation responses to antigen and APC-induced co-stimulatory
727 signals that are required for licensing them to become fully pathogenic T cells.

728
729 Peripheral tolerance can be induced by via several mechanisms including T cell
730 deletion or anergy, the induction of regulatory T cells and immune deviation (Tisch,
731 2010). To gain insight into the mechanism of OM-MOG-induced tolerance we
732 monitored T cell responses to MOG stimulation *in vivo* in conjugate-vaccinated mice
733 and *in vitro* using DC-T cell co-cultures. We found no evidence for deletion of
734 antigen-specific T cells in the periphery of OM-MOG-vaccinated mice, as judging
735 from the normal differentiation of IFN- γ - and IL-17-producing CD4⁺ T cells and FoxP3⁺
736 regulatory T cells isolated from secondary lymphoid organs following immunization
737 of mice with MOG. This was supported by data from DC-T cells cultures showing that
738 OM-MOG DC and MOG DC stimulated equal production of IFN- γ and IL-17 by CD4⁺
739 2D2 T cells, and that OM-MOG DC did not induce increased cell surface expression of
740 the apoptosis marker annexin V compared to MOG, OM or PBS DC. Furthermore, we
741 found no evidence for immune deviation in antigen-specific T helper cell populations
742 and no alteration in regulatory T cell populations. We did, however, detect reduced
743 antigen-specific T cell proliferation measured both in secondary lymphoid organs of

744 OM-MOG-vaccinated mice following MOG immunization, and in 2D2 T cells co-
745 cultured with OM-MOG DC, as well as failure of OM-MOG DC-delivered co-
746 stimulatory signals to up-regulate T cell cytokine production. The uncoupling of
747 cytokine production from proliferation in T cells in the presence of competent APC
748 has been described previously in response to APL (Evavold *et al.*, 1991) and during
749 oral tolerance (Whitacre, Gienapp, Orosz, and Bitar, 1991), (Chen, Inobe, Kuchroo,
750 Baron, Janeway, Jr., and Weiner, 1996), (Karpus, Kennedy, Smith, and Miller, 1996)
751 and is thought to involve the differential activation of TCR signaling pathways (Sloan-
752 Lancaster and Allen, 1996). Although the signaling pathways that underlie
753 differential TCR-mediated effects remain to be fully elucidated, our findings add that
754 effector T cell cytokine production, in the absence of adequate proliferative
755 response, is not sufficient for the induction of EAE.

756

757 The mechanism by which mannan-conjugated myelin peptides reduce the
758 encephalitogenic function of effector T cells therefore appears to be different from
759 those previously described for immune tolerance induced by APC targeting which
760 include association with CD5 expression (Hawiger *et al.*, 2004), reduction of IL-17 cell
761 and increase of regulatory T cell differentiation (Stern *et al.*, 2010) and immune
762 deviation towards an immunoregulatory profile (Apostolopoulos *et al.*, 2000),
763 (Apostolopoulos *et al.*, 1995). It most closely resembles tolerance induced by
764 mannosylated antigens, as previously described Nagelkerken and his group. Like
765 mannan-MOG, mannosylated PLP₁₃₉₋₁₅₁ protected SJL mice against the development
766 of PLP-EAE through an active mechanism, because it inhibited disease when
767 administered after EAE induction by active immunization with PLP or adoptive
768 transfer of PLP₁₃₉₋₁₅₁-reactive T cell blasts (Luca *et al.*, 2005), (Kel *et al.*, 2007). Also
769 lymph node cells isolated from mice that had been immunized with mannosylated
770 PLP₁₃₉₋₁₅₁ showed equal cytokine and chemokine production but reduced
771 proliferation responses compared to cells primed with non-mannosylated PLP₁₃₉₋₁₅₁
772 (Kel *et al.*, 2008). This defect was associated with poor Th1 effector functions as
773 shown by reduced IgG2a antibody levels, reduced DTH responses and EAE symptoms
774 (Luca *et al.*, 2005), (Kel *et al.*, 2007). However, unlike tolerance induced by OM-MOG,
775 which was resistant to PTx, that induced by mannosylated peptides was abrogated
776 by PTx administration (Kel *et al.*, 2008). PTx is widely used to increase antigen-
777 specific T cell responses and disease susceptibility in EAE models through multiple
778 effects including prevention of antigen-induced peripheral T cell anergy (Kamradt,
779 Soloway, Perkins, and Geftter, 1991), induction of IL-17 production (Hofstetter, Grau,
780 Buttman, Forsthuber, Gaupp, Toyka, and Gold, 2007) and reduction of regulatory T
781 cells (Chen, Winkler-Pickett, Carbonetti, Ortaldo, Oppenheim, and Howard, 2006). It
782 is possible therefore, that mannosylated and mannan-conjugated peptides induce T
783 cell tolerance through a similar mechanism and that differences in sensitivity of the
784 two approaches to immune adjuvants reflect differences in the strength of tolerance
785 induced.

786

787 The MR is expressed by most tissue macrophages, other cell types including hepatic
788 and lymphatic endothelia, a subpopulation of DC in lymphoid organs that drain the
789 periphery and in brain meningeal macrophages and perivascular microglia in the
790 mouse (McKenzie, Taylor, Stillion, Lucas, Harris, Gordon, and Martinez-Pomares,

2007), (Linehan *et al.*, 1999) as well as additional skin and gut DC in humans (Engering, Geijtenbeek, van Vliet, Wijers, van, Demaurex, Lanzavecchia, Fransen, Figdor, Piguet, and van, 2002). It provides an efficient internalization system for the recognition, transport and clearance of host-derived glycoproteins and microbe-derived ligands (Taylor, Martinez-Pomares, Stacey, Lin, Brown, and Gordon, 2005). Previous studies showed that engagement of the MR by mannosylated lipoarabinomannans, mannan or an anti-MR antibody inhibited LPS-induced IL-12 production by human DC (Nigou, Zelle-Rieser, Gilleron, Thurnher, and Puzo, 2001) and Toll-like receptor-dependent IL-12 production in mouse macrophage cells (Pathak, Basu, Bhattacharyya, Pathak, Kundu, and Basu, 2005). Also some, not all, natural MR ligands activated an anti-inflammatory program in human monocyte-derived DC, which included inhibition of IL-12 production and potential to polarize Th1 effector cells (Chieppa, Bianchi, Doni, Del, Sironi, Laskarin, Monti, Piemonti, Biondi, Mantovani, Introna, and Allavena, 2003). These latter studies show that appropriate engagement of the MR on DC can elicit immunosuppressive effects. To investigate the cellular basis of MOG-specific T cell tolerance in OM-MOG-vaccinated mice, we used DC cultures derived from mouse bone marrow and unexpectedly found that OM-MOG-loaded DC displayed phenotypic and functional characteristics of immune-promoting mature DC. First, DC loaded with OM-MOG or MOG efficiently presented antigen to 2D2 MOG-specific T cells and equally induced the differentiation of IFN γ -producing and IL-17-producing T cells. Second, both OM and OM-MOG increased the surface expression of CD40, CD80 and CD86 in DC to levels at least as high as LPS-treated DC, showing that mannan induces the phenotypic maturation of bone marrow-derived DC, a property consistent with its functional characteristics as a PAMP, and in agreement with our previous results on the effects of OM-MUC1 in bone marrow-derived DC as well as *in vivo* splenic and lymph node DC (Sheng *et al.*, 2006). Third, OM-MOG DC showed CD40-inducible production of the Th1 and Th17 polarizing cytokines, IL-12 and IL-23 respectively, properties that are essential for the immune-promoting functions of mature DC (Caux, Massacrier, Vanbervliet, Dubois, Van, Durand, and Banchereau, 1994), (Cella, Scheidegger, Palmer-Lehmann, Lane, Lanzavecchia, and Alber, 1996), (Koch, Stanzl, Jennewein, Janke, Heufler, Kampgen, Romani, and Schuler, 1996), (Iezzi *et al.*, 2009). These results first of all suggest that T cell tolerance induced by the administration of OM-MOG in mice is not due to an intrinsic defect in APC function. Our finding that OM-MOG induces T cell tolerance, and that *in vitro* grown OM-MOG-loaded DC are sufficient to transfer tolerance into mice with ongoing EAE therefore appears paradoxical. A number of previous studies have already challenged the concept that only immature and semi-mature DC mediate T cell tolerance by showing that phenotypically mature DC can also induce tolerance (Albert *et al.*, 2001), (Menges, Rossner, Voigtlander, Schindler, Kukutsch, Bogdan, Erb, Schuler, and Lutz, 2002), (Fujii *et al.*, 2004), (Sporri *et al.*, 2005). In some of these studies (Albert *et al.*, 2001), (Fujii *et al.*, 2004), but not all (Menges *et al.*, 2002), stimulation of CD40 changed the outcome of tolerance to immunity. Since various cell types express the MR in mice it is possible that tolerance induced by OM-MOG is mediated by APC other than DC, for example by meningeal macrophages or perivascular microglia (Linehan *et al.*, 1999) which are known to have APC properties (Greter, Heppner, Lemos, Odermatt, Goebels, Laufer, Noelle, and Becher, 2005). Nevertheless, when we administered

838 FGK45 *in vivo* it did not change the outcome of protection against EAE in OM-MOG-
839 vaccinated mice while, as expected, it exacerbated disease in PBS-vaccinated mice.
840 Taken together, our findings show that mannan-peptide-targeted APC do not show
841 overt functional defects; on the contrary they show characteristics of mature
842 immunogenic APC; and suggest they play a nonautonomous role in mediating the
843 tolerogenic effects of OM-MOG to T cells.

844

845 Lymphocytes derived from OM-MOG-tolerized mice and MOG-specific T cells
846 exposed to OM-MOG DC exhibited a selective defect in proliferation, but not
847 cytokine responses to antigen stimulation and this unresponsiveness was overcome
848 by exogenous IL-2. Clonal T cell anergy, characterized by reduction of antigen-
849 specific proliferation responses but not necessarily changes in cytokine production,
850 has been described as a mechanism of T cell tolerance induced by oral
851 administration of MBP (Whitacre *et al.*, 1991), (Chen *et al.*, 1996) and PLP₁₃₉₋₁₅₁
852 (Karpus *et al.*, 1996), by myelin peptide-coupled splenocytes (Vandenbark, Celnik,
853 Vainiene, Miller, and Offner, 1995) and by prostaglandin 2 (Mannie, Prevost, and
854 Marinakis, 1995). In oral tolerance, low doses of antigen generate regulatory T cells
855 whereas high doses induce T cell anergy or deletion. Our data suggest that OM-MOG
856 administration in mice results in strong stimulation of self-reactive T cells by APC,
857 through combined MOG-TCR-specific and MR-induced innate immune signals, that
858 results in partial T cell anergy. This would be in line with our previous findings, where
859 OM-MUC1 greatly enhanced T cell responses to MUC1 but, in that case, promoted
860 anti-tumor immunity (Apostolopoulos *et al.*, 1995). Our finding that T cell tolerance
861 in OM-MOG vaccinated EAE mice and OM-MOG DC-T cell cultures was not altered by
862 CD40 co-stimulation is a strong indication of clonal T cell anergy because mice that
863 are deficient in CD40 do not develop pathogenic Th17 T cells and are completely
864 resistant to EAE (Becher *et al.*, 2001), (Iezzi *et al.*, 2009). It is established that *de novo*
865 processing and presentation of CNS antigens in the context of MHC class II is
866 absolutely required for the development of EAE (Tompkins *et al.*, 2002), (Becher,
867 Durell, and Noelle, 2003), (Kawakami, Lassmann, Li, Odoardi, Ritter, Ziemssen,
868 Klinkert, Ellwart, Bradl, Krivacic, Lassmann, Ransohoff, Volk, Wekerle, Lington, and
869 Flugel, 2004) probably by APC associated with the meninges and CNS vasculature
870 (Greter *et al.*, 2005). Previous studies showed that mannosylated PLP could
871 ameliorate EAE when administered after the induction of EAE by adoptive transfer of
872 activated myelin-reactive T cells (Kel *et al.*, 2007), showing that tolerance is actively
873 induced in already activated T cells, possibly preventing their reactivation in the
874 periphery or CNS. Similarly in this study administration of OM-MOG, either as i.d.
875 injection or transferred with DC, actively induced tolerance in resting or already
876 activated T cells, thereby reducing their proliferation and encephalitogenic potential
877 and protecting mice against EAE. In line with the hypothesis that OM-MOG acts by
878 reducing the capacity of MOG-specific T cells to be restimulated by endogenous
879 antigen in the target tissue, is the finding that CD3⁺ T cells accumulated in the
880 leptomeninges and vasculature of the spinal cord and showed markedly reduced
881 infiltration of the parenchyma after the induction of active EAE. Alternatively, it is
882 possible that OM-MOG exerts a decoy effect, diverting the migration of MOG-
883 specific T cells away from targets in the CNS to the site of injection in the skin, and
884 further *in vivo* experiments will be needed to investigate this possibility.

885

886 Overall, mannan-conjugated peptides administered as prophylactic vaccines or
887 therapeutically induce potent and reproducible peptide-specific protection of mice
888 from the clinical signs of EAE with all protocols tested even in the presence of strong
889 APC maturation stimuli. The protective effect of mannan-conjugated peptides is
890 especially important in a setting where EAE is induced in the presence of adjuvants,
891 like PTx and CD40, that have the ultimate role of breaking tolerance indicating that it
892 might be a powerful strategy to silence autoimmunity in a human setting where
893 immune challenge by microbial antigens is thought to play a key role in the
894 breakdown of immune tolerance to self-antigens. Further studies are needed to
895 characterize the mechanism of anergy triggered by TCR stimulation by mannan-
896 peptide conjugates, especially since they constitute interesting targets of clinical
897 application for therapeutic intervention in MS. Treatment with mannan-conjugated
898 autoantigens could differ from current therapeutic regimens that are
899 immunosuppressive and could be a step towards the design of personalized
900 therapies for the different immunodominant peptide epitopes found in autoantigen
901 specific immune reactions in MS and relevant demyelinating disorders.

902

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914

915 **Abbreviations:** Ag, antigen; APC, antigen-presenting cells; APL, altered peptide
916 ligands; CFA, complete Freund’s adjuvant; CNS, central nervous system; cpm, counts
917 per minute; DC, dendritic cells; DLN, draining lymph nodes; DTH, delayed-type
918 hypersensitivity; EAE, experimental autoimmune encephalomyelitis; FCS, foetal calf
919 serum; GA, glatiramer acetate; H&E, haematoxylin & eosin; LPS, lipopolysaccharide;
920 MAb, monoclonal antibody; MOG, myelin oligodendrocyte glycoprotein; MR,
921 mannose receptor; MS, multiple sclerosis; MUC1, mucin 1; OM, oxidized mannan;
922 PAMP, pathogen-associated molecular pattern; PLP, proteolipid lipoprotein; POL,
923 GA-type polypeptide; PTx, *Bordetella pertussis* toxin; RM, reduced mannan; RP-HPLC,
924 reverse phase-high performance liquid chromatography; SI, stimulation index; TCR, T
925 cell receptor; TFA, trifluoroacetic acid

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927

928 **Figure legends**

929 **FIGURE 1.** Administration of OM-MOG or RM-MOG in prophylactic (vaccination) or
930 therapeutic protocols attenuates the development of MOG-induced EAE in C57BL/6
931 mice. **(a)** Mean clinical scores of MOG-EAE in groups of mice that were vaccinated i.d.
932 with dilute soluble OM-MOG, RM-MOG, OM, RM or PBS at indicated time points
933 (arrows) prior to the induction of EAE by immunization with MOG/CFA/PTx (n = 6 for
934 each group). **(b)** Mean clinical scores of MOG-EAE in groups of mice that received i.d.
935 administration of dilute soluble peptides on day 0 and 7 relative to immunization for
936 EAE induction (n=5 for each group). **(c)** Mean clinical scores of MOG-EAE in groups of
937 mice injected i.d. with dilute soluble peptides at indicated time points after
938 immunization for EAE induction (n = 5 for each group). The results shown are from one
939 representative of two **(a, c)** or three **(b)** independent experiments. **(d, e)** Vaccination
940 with OM-MOG protects C57BL/6 mice from spinal cord inflammation and demyelination
941 during MOG-EAE. **(d)** Inflammatory cell infiltration was visualized by H&E (left column)
942 and demyelination by Luxol fast blue (right column) staining of spinal cord sections
943 taken from representative mice in the different vaccination groups on day 24
944 following immunization for the induction of MOG-EAE. Representative sections from
945 1 of 5 animals per group are shown. **(e)** Quantification of spinal cord inflammation
946 (black bars) and demyelination (grey bars) as well as brain demyelination (white bars) in
947 all experimental groups. Representative data from 5 animals per group are shown.
948 Statistical significance after pair-wise comparisons of each experimental group with the
949 non-vaccinated control (PBS) group is shown (*, $p < 0.05$). Triangles (a) indicate time
950 points where pairwise comparison between OM-MOG and RM-MOG groups also show
951 significant differences.

952
953 **FIGURE 2.** Protection of mice against EAE by mannan-peptides is peptide-specific. **(a)**
954 Vaccination of SJL/J mice with OM-PLP₁₃₉₋₁₅₁, but not OM-MBP₈₃₋₉₉, attenuated the
955 development of PLP-EAE. Mean clinical scores of PLP-EAE in groups of mice that were
956 vaccinated with OM- PLP₁₃₉₋₁₅₁, OM- MBP₈₃₋₉₉ or PBS on the days indicated (arrows)
957 prior to immunization for the induction of PLP-EAE (n = 5 for all groups). **(b)** Vaccination
958 of C57BL/6 mice with OM-MOG₃₅₋₅₅, but not OM-PLP₁₇₈₋₁₉₁, attenuated the
959 development of MOG-EAE. Mean clinical scores of MOG-EAE in groups of mice that
960 were vaccinated with OM-MOG₃₅₋₅₅, OM-, PLP₁₇₈₋₁₉₁ or PBS on the days indicated
961 (arrows) prior to the induction of PLP-EAE by immunization with PLP/CFA (n = 5 for all
962 groups). Data are from one representative of two independent experiments. Statistical
963 significance after pair-wise comparisons of each experimental group with the non-
964 vaccinated control (PBS) group is shown (*, $p < 0.05$).

965
966 **FIGURE 3.** Antigen-specific responses of T cells exposed to OM-MOG show reduced
967 proliferation but normal expansion of Th1, Th17 and regulatory T cell populations.
968 **(a)** Proliferation of splenocytes isolated from vaccinated mice 25 days post-
969 immunization for MOG-EAE. Splenocytes were stimulated *ex vivo* with MOG (mice
970 vaccinated with PBS n = 4; OM-MOG n = 6; RM-MOG n = 6; OM n = 5; RM n = 4). **(b)**
971 Proliferation of CFSE-labelled MOG-specific 2D2 CD45.1⁺ cells *in vivo* after i.v. transfer
972 into recipient CD45.2⁺ mice that had received a complete vaccination protocol with
973 mannan-conjugated peptides, on day 2 after MOG-EAE induction. DLN were isolated
974 from mice 7 days after immunization with MOG/CFA/PTx, and on the CD45.1⁺ gate the

975 percentages of proliferating cells (showing low and intermediate levels of CFSE staining)
976 were measured (mice vaccinated with MOG n = 5; OM-MOG n=5; OM n=5; PBS n=5).
977 Right panels show representative histogram plots showing CFSE dilution in MOG-
978 specific 2D2 CD45.1⁺ cells in DLN isolated from OM- and OM-MOG-vaccinated mice. **(c-
979 d)** Antigen priming of Th1 and Th17 T cells is normal in mice vaccinated with OM-MOG.
980 Mice were vaccinated with OM, OM-MOG, MOG or a mix of unconjugated OM and
981 MOG and immunized for the induction of MOG-EAE. **(c)** The production of IFN- γ and IL-
982 17 by CD4⁺ T cells from DLN isolated 7 days after MOG immunization was measured by
983 intracellular cytokine staining (n=5 for the MOG and OM-MOG groups and n=4 for the
984 OM and OM/MOG groups). **(d)** The production of IL-17 by CD4⁺ T cells from DLN (left
985 panel) and spleen (right panel) isolated on days 10 (pre-onset) and 15 (peak) of EAE was
986 also measured by intracellular cytokine staining (n = 5 for all groups). **(e)** The
987 production of IL-10 by CD4⁺ T cells from DLN isolated on day 13 of EAE was also
988 measured by intracellular cytokine staining (n = 4 for all groups). **(f)** Proportions of
989 CD4⁺FoxP3⁺ cells in DLN cells from vaccinated mice 7 days post-immunization for MOG-
990 EAE, and stained for surface CD4 and intracellular FoxP3 (n=5 mice in each group). Right
991 panel, representative dot plots are shown from each experimental group. Data are
992 from one (e) or one representative of two independent experiments. Statistical
993 significance after pair-wise comparisons (using Student's t test) of each experimental
994 group with the non-vaccinated control (PBS) group is shown.
995

996 **FIGURE 4.** OM-MOG-loaded DC up-regulate cell surface maturation markers, Th1 and
997 Th17 polarising cytokines and down-regulate PD-L1 and are sufficient to transfer
998 tolerance into EAE mice. **(a)** Flow cytometry of DC loaded with PBS or peptide
999 conjugates or incubated with LPS (1 μ g /ml), showing frequency of CD11c⁺ cells
1000 expressing DC surface maturation markers. **(b)** Enzyme-linked immunosorbant assay of
1001 cytokines in culture supernatants of DC loaded with OM-MOG or MOG in the absence
1002 or presence of FGK 45 agonistic anti-CD40 antibody (10 μ g/ml) for 72 hr. **(c)** Flow
1003 cytometry of DC loaded with PBS or peptide conjugates showing frequency of CD11c⁺
1004 cells expressing PD-L1 surface marker. **(d)** Mean clinical scores of mice that were
1005 immunized for the induction of MOG-EAE and 8 days later received i.v. transfer of bone
1006 marrow-derived DC loaded *ex vivo* with OM-MOG (n=9), unconjugated MOG (n = 8),
1007 OM (n = 7) or PBS (n = 8) at the indicated time point (arrow). Data are from two **(b, d)**
1008 or three **(a)** independent experiments. Statistical significance after pair-wise
1009 comparisons between groups or **(d)** or each experimental group with the non-
1010 vaccinated control (PBS) group (*, $p < 0.05$) is shown.
1011

1012 **FIGURE 5.** OM-MOG DC induce normal maturation of MOG-specific Th1 and Th17
1013 responses but reduced T cell proliferation and CD40 costimulation responses. **(a, b)**
1014 Antigen presentation assays between MOG DC and OM-MOG DC and 2D2 MOG-
1015 specific lymphocytes at a 1:5 DC:T cell ratio in the absence or presence of FGK45
1016 agonistic anti-CD40 antibody (10 μ g/ml) for 72 hr. **(a)** Proliferation of 2D2 lymphocytes
1017 by peptide-loaded DC as measured by [³H] thymidine incorporation. **(b)** Production of
1018 IFN- γ and IL-17 by CD4⁺ 2D2 T cells in response to peptide-loaded DC as measured by
1019 intracellular cytokine staining. **(c)** Annexin V⁺ surface expression by CD4⁺ 2D2 MOG-
1020 specific T cells stimulated by OM-MOG, MOG, OM and PBS DC. Annexin V translocation
1021 was measured at day 7 of culture. Statistical significance after pair-wise comparisons

1022 (using Student's t test) between 2D2 cells stimulated with PBS DC versus Ag DC is
1023 shown. **(d)** Mean clinical scores of MOG-EAE in groups of mice that were vaccinated i.d.
1024 with dilute soluble OM-MOG or PBS and were further left untreated or treated by
1025 twice weekly i.p. injections of FGK45 agonistic anti-CD40 antibody (90
1026 µg/mouse/injection) from day 2 post-immunization (n = 8 for the OM-MOG and OM-
1027 MOG + aCD40 groups; n = 4 for the PBS and the PBS + aCD40 groups). **(e)** Proliferation
1028 of 2D2 lymphocytes by peptide-loaded as measured by [³H] thymidine incorporation in
1029 the absence or presence of recombinant IL-2 (10 ng/ml). Representative results from
1030 one of two independent experiments are shown. Statistical significance after pair-wise
1031 comparisons between cells or mice treated in the absence or presence of agonistic anti-
1032 CD40 antibody (a-d) (*, *p*<0.05) or IL-2 (e) is shown.

1033

1034

1035 **FIGURE 6.** CD3⁺ T cell and immune cell infiltration of the spinal cord in OM-MOG-
1036 vaccinated mice is sparse and limited to leptomeninges or exceptional compact
1037 white matter lesions. **(a)** Double immunofluorescence staining with anti-CD3 and anti-
1038 Ki67 antibodies identifies proliferating CD3⁺ T cells in spinal cord sections taken at the
1039 peak of disease from chimeric C57BL/6 mice reconstituted with bone marrow from
1040 TgEGFP mice, vaccinated with OM-MOG or OM control and immunized for MOG-EAE.
1041 Numerous CD3-immunoreactive infiltrating cells are seen in the parenchyma of PBS-
1042 vaccinated mice, many of which also show Ki67-immunoreactivity (clinical score 3.5)
1043 (upper panel, arrowheads). CD3-immunoreactive infiltrates are markedly reduced in
1044 OM-MOG vaccinated mice (clinical score 0) where they are mainly restricted to
1045 leptomeninges and the associated perivascular spaces (lower panel). **(b)** Oxidative burst
1046 in infiltrating cells shown by immunostaining of sections from the same mice as in a)
1047 with anti-p22phox (p22) and iNOS antibodies in areas of CD3⁺ T cell infiltration.
1048 Representative results from 5 mice per group are shown. Scale bars 500 µM (a).

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