Effects of Early IL-17A Neutralization on Disease Induction in a Primate Model of Experimental Autoimmune Encephalomyelitis

Yolanda S. Kap · S. Anwar Jagessar · Nikki van Driel · Erwin Blezer · Jan Bauer · Marjan van Meurs · Paul Smith · Jon D. Laman · Bert A. ‘t Hart

Guarantor of the work: Dr. Bert A. ‘t Hart (Chairman of the Department of Immunobiology, BPRC)

Abstract We report on the effect of antibody-mediated neutralization of interleukin (IL)-17A in a non-human primate experimental autoimmune encephalomyelitis (EAE) model induced with recombinant human myelin oligodendrocyte glycoprotein (rhMOG). We tested a human-anti-human IL-17A-antibody in two doses (3 and 30 mg/kg) against placebo (PBS). The treatment was started 1 day before EAE induction and continued throughout the experiment. Although all monkeys developed clinically evident EAE, the onset of neurological signs was delayed in some monkeys from both treatment groups. Total CNS lesion volumes, demyelination, or inflammation did not differ between the different groups. Immune profiling revealed an altered distribution of IL-17A producing cells in the lymphoid organs of antibody-treated monkeys. Comparable numbers of IL-17A producing cells were observed in the brain. RhMOG-induced T cell proliferation in the lymph nodes was slightly reduced after anti-IL-17A antibody treatment. To summarize, we found that anti-IL-17A antibody as a single treatment from disease induction effects a trend towards delayed neurological disease progression in the marmoset EAE model, although the effect did not reach statistical significance. This suggests a role of IL-17A in late stage disease in the marmoset EAE model, but IL-17A may not be the dominant pathogenic cytokine.

Keywords Th17 · IL-23 · Antibody therapy · Non-human primate · Neuroimmunology

Introduction

Multiple sclerosis (MS) is a progressive neurological disease of unknown origin characterized by the progressive accumulation of inflammation and degeneration in the grey and white matter of the central nervous system (CNS). The accumulation of injury involves a combined cellular and humoral autoimmune attack on the protective myelin sheaths around axons leading to disturbed saltatory pulse conduction (Compston and Coles 2008; Sospedra and Martin 2005). This immunopathogenic process is modeled in the experimental autoimmune encephalomyelitis (EAE) model, an experimental disease induced in genetically susceptible laboratory animals including mice, rats, guinea pigs, and monkeys. The EAE model in common marmosets, a small-sized neotropical primate, combines a remarkable
clinical and neuropathological similarity with MS and a high
genetic and immunological proximity to humans (‘t Hart and
Massacesi 2009). The model is therefore exquisitely suitable
for translational research into immunopathogenic mechan-
isms and the efficacy evaluation of therapeutic antibodies,
which because of their high target specificity do not cross-
react with lower species. The marmoset EAE model thus
forms a bridge narrowing the gap between EAE models and
MS (‘t Hart et al. 2004; Kap et al. 2010).

Research conducted in the past years has shown that the
100% disease incidence and variable progression rate of
neurological deficit is associated with the mobilization of
highly reactive T cells present in the normal repertoire with
variable specificities for myelin oligodendrocyte glycopro-
tein (MOG) (Jagessar et al. 2010; Kap et al. 2008). Whereas
disease induction could be mapped to the Caja-
DRB1*W1201 restricted activation of T helper (Th1) cells
specific for MOG34–56 (Brok et al. 2000), the rate of disease
progression appeared to be regulated by natural killer-like T
cells specific for MOG34–56 (Jagessar et al. 2010; Kap et al.
2008). Further research confirmed that this T cell subset is
capable to induce clinical signs and pathology in the CNS
white and grey matter without the support of innate antigen
receptor ligands in the inoculum or autoantibodies against
rhMOG or the inducing peptides (Jagessar et al. 2010).
These data led to the hypothesis that MOG34–56-specific T
cells have a central pathogenic role in the regulation of
disease progression in the marmoset EAE model.

The most obvious immunological characteristics of
MOG34–56-specific T cell lines generated from EAE-
affected monkeys were specific cytolytic activity towards
MOG34–56-pulsed target cells and production of IL-17A
(Jagessar et al. 2010; Kap et al. 2008). However, typical
Th1 cell-derived cytokines, such as interferon (IFN)-γ or
tumor necrosis factor alpha (TNF-α), that are abundantly
produced by the MOG34–56-specific Th1 cells involved in
EAE initiation (Brok et al. 2000) were not produced by
MOG34–56-specific T cells (Jagessar et al. 2010). This may
well be due to the fact that for the activation of
encephalitogenic Th1 cells in mice as well as in marmosets
strong bacterial adjuvants are needed, i.e. complete
Freud's adjuvant (CFA) (Billiau and Matthys 2001). By
contrast, our data in the marmoset EAE model demonstrates
that encephalitogenic T cells specific for MOG34–56 can
also be activated by immunization with MOG34–56 in
incomplete Freud's adjuvant (IFA) (Jagessar et al. 2010).

These observations prompted us to test whether
antibody-mediated neutralization of IL-17A modulates
disease progression in the EAE model in marmosets,
elicited by a single immunization with recombinant protein
of the extracellular domain of human MOG (MOG1–125) in
CFA. The most prevalent disease course in this model is
clinically characterized by the sudden appearance of
progressively accumulating motor defects after an episode
of variable length, during which MRI-detectable lesions do
not lead to evident neurological symptoms (Brok et al.
2000; Kap et al. 2008). A typical relapsing-remitting
disease course is only occasionally observed. Our previous
studies have revealed that intervention with an antibody
against the shared p40 subunit of IL-12 and IL-23 has a
clear beneficial effect in the rhMOG-induced EAE model (‘t
Hart et al. 2005b; Brok et al. 2002). This supports an
important pathogenic role of IL-12 and IL-23, which may
be exerted via the inhibition of IL-17A production, as IL-23
promotes IL-17A production (Langrish et al. 2005).

Here, we report that prophylactic treatment with anti-IL-17A
monoclonal antibody (mAb) at doses achieving plasma through
levels above 1 μg/ml induced a moderate delay of EAE onset.
These data indicate that IL-17A is only one of several cytokines
driving progression of the marmoset EAE model to clinically
evident neuroinflammatory disease; absence of IL-17A being
possibly compensated by Th1 cytokines that are abundantly
produced in the rhMOG/CFA model.

Material and methods

Animals

The common marmoset monkeys used in this study were
randomly selected from the outbred colony kept at
the Biomedical Primate Research Centre (Rijswijk, The
Netherlands) under conventional and not SPF conditions.
Animals were only included after a complete physical,
hematological, and biochemical check-up had been per-
formed. During the study, monkeys were pair-housed in
spacious cages and remained under intensive veterinary care.
The daily diet consisted of commercial food pellets for New
World monkeys (Special Diet Services, Witham, Essex,
UK), supplemented with rice, raisins, peanuts, marshmal-
lows, biscuits, fresh fruit, grasshoppers, and maggots.
Drinking water was provided ad libitum. In accordance
with the Dutch law on animal experimentation, all study protocols
and experimental procedures have been reviewed and
approved by the Institute's Ethics Committee.

The study comprised 22 male and two female monkeys,
M05011 and M04049. The average age of the monkeys was
33±16 months, which is adult age for marmosets. The
average body weight was 377±35 g.

RhMOG-induced EAE

EAE was induced by immunization with a recombinant
protein encompassing the extracellular domain of human
MOG residues 1–125, which was produced in Escherichia
coli and purified as previously described (Kerlero de Rosbo
et al. 1997). The inoculum, containing 100 μg rhMOG in 300 μl phosphate buffered saline (PBS) emulsified with 300 μl CFA containing mycobacterium butyricum (Difco Laboratories, Detroit, MI), was injected at four locations into the dorsal skin under ketamin anesthesia (40 mg/kg; AST Pharma, Oudewater, The Netherlands).

Clinical signs were scored daily by two independent observers using a previously documented semi-quantitative scale (’t Hart et al. 2008): 0=no evident clinical signs; 0.5=apathy, loss of appetite, altered walking pattern without ataxia; 1=lethargy, anorexia, tail paralysis, tremor; 2=ataxia, optic disease; 2.25=monoparesis; 2.5=paraparesis, sensory loss; 3=para- or hemiplegia. For ethical reasons monkeys were sacrificed before or once complete paralysis of hind limbs (score ≥3.0) was observed, or at the pre-determined endpoint of the study (post-sensitization day (psd) 113). Body weight measurements of conscious monkeys, which is used as a surrogate disease marker, were performed three times per week.

Monkeys selected for necropsy were first deeply sedated by intramuscular injection of ketamin (50 mg/kg) and subsequently euthanised by infusion of pento-barbital sodium (Euthesate®; Apharmo, Duiven, The Netherlands).

Reactivity and dosing regimen of anti-IL-17A mAb

The test substance was produced by UCB Celltech (UK) as a humanized IgG4κ mAb specific for human IL-17A, coded as 497.g2. The antibody has been extensively characterized in vitro in terms of bioassay, affinity for IL-17A, and cross-reactivity against marmoset IL-17A. The affinity of the antibody with marmoset IL-17A is twofold lower than with human IL-17A when assessed by Biacore and fourfold less potent in a bioassay compared with humans.

The animals were subcutaneously injected once a week starting 1 day before immunization until the predetermined end of the study at day 113. Animals were randomly assigned to three experimental groups. Eight animals received 3 mg/ml/kg anti-IL-17A mAb diluted in PBS, eight animals were injected with 30 mg/ml/kg anti-IL-17A mAb diluted in PBS, and eight control animals received sterile PBS (1 ml/kg) as placebo treatment. All animals received the same volume per kg body weight. One monkey (M04063) in the 30 mg/kg antibody dose group succumbed at psd 69 unexpectedly without prior signs of EAE and was therefore excluded from further analyses. Autopsy revealed that the cause of death was not related to the test substance or EAE, but to perforation of the gastro-intestinal tract by plant material, possibly originating from the branches used for cage enrichment.

Blood sampling and plasma levels of anti-IL-17A mAb

Venous blood was collected into heparinized vacutainers (Greiner, Sölingen, Germany) under ketamin anesthesia (40 mg/kg; AST Pharma, Oudewater, The Netherlands) at psd 0, 6, 34, 62, and at necropsy.

After centrifugation plasma was collected and stored frozen at −20°C until analysis of test substance levels was performed. Test substance plasma levels were determined by ELISA. Microtitre plates were coated with human IL-17A (R&D Systems, Minneapolis, MN) at 0.5 μg/ml in PBS overnight, blocked with PBS/1% BSA, glazed with PBS/5% lactose/0.1% BSA, dried, sealed in foil pouches, and stored at 2–8°C. The standard curve was prepared by making serial doubling dilutions of the 497.g2 top standard (starting at 200 ng/mL) in PBS/1% BSA/1% BGG/1% human plasma. 50 μL of each standard, interassay control (IAC), and sample (diluted at least 1/100) were added to the appropriate wells containing 50 μL PBS/1% BSA/1% BGG. The IAC concentrations were nominally 80, 20, and 8 ng/mL. Standards, IAC, and samples were tested in duplicate. The plate was covered and incubated with agitation at RT for 2 h. The plate was washed with PBS/0.1% Tween-20 four times and incubated with goat anti-human Kappa-HRP conjugate (1/10,000) in PBS/1% BSA/1% BGG at RT for 30 min. The plate was washed again with PBS/0.1% Tween-20 four times and incubated with 100 μL Tetramethyl benzidine substrate for 10 min. The reaction was stopped with 50 μL/well of 2.5 M H2SO4 and measured at 450 nm (and 630 nm as a reference).

Magnetic resonance images

Post-mortem magnetic resonance images (MRI) of one brain hemisphere were recorded to assess differences in the CNS lesion load between treated and control monkeys (Blezer et al. 2007). Half of the brain collected at necropsy was fixed in 4% buffered formalin and transferred into buffered saline containing sodium azide after two weeks.

MRI experiments were performed on a 9.4 T horizontal bore MRI scanner (Varian, Palo Alto, CA). The formalin-fixed brains were submerged in a non-magnetic oil (Fomblin; perfluorinated polyether, Solvay Solexis, Weesp, The Netherlands) to prevent unwanted susceptibility artifacts.

The following quantitative images were obtained (field of view=2.5×2.5 cm², matrix=256×256, slice thickness=0.75 mm, number of slices=41, number of experiments=2);

- T2 maps, which were calculated from the images obtained of a multi echo sequence using the following parameters: repetition time=4,000 ms, echo spacing=...
14.75 ms, echo train length=4. T2 maps were the result of a mono-exponential fit of the MRI signal intensities as a function of TE. The T2-weighted images with TE of 14.75 ms were used for region of interest (ROI) analyses.

- Magnetization transfer ratio (MTR) maps were calculated from two T1-weighted spin echo images with and without a magnetization transfer-saturation pulse, with \( MTR = 100\left[\frac{M_{\text{unsaturated}} - M_{\text{saturated}}}{M_{\text{unsaturated}}}\right] \). Repetition time=1675 ms; echo time=23 ms; MT-pulse: 8.19 ms Gaussian-shaped pulse, nominal flip angle 1,000, offset –9.4 kHz.

ROI were defined using the free available medical image processing, analysis and visualization (MIPAV version 4.3.0, National Institutes of Health, Bethesda, MD) package. ROI of lesions, defined as areas with abnormal increased signal intensities, were automatically outlined in all T2-weighted images containing white matter structures using the level-set method of MIPAV. Volumes, T2, and MTR values were calculated from these ROI.

Histology and immunohistochemistry

Frozen and fixed tissues were processed for histological and immunohistochemical techniques as previously described (t’Hart et al. 1998; Laman et al. 1998). Demyelination was visualized by staining myelin and myelin degradation products with Klüver Barrera method (Luxol Fast Blue with Periodic Acid-Schiff). Inflammation was visualized by staining infiltrating cells with hematoxylin and eosin. Of each animal eight sections, approximately 6 cm² in total, were analyzed.

Snap-frozen sections of the brain, spleen, and axillary (ALN), inguinal (ILN), and cervical lymph nodes (CLN) were used to determine the number of IL-17A-producing cells. Sections of 6 µm were thaw mounted on gelatin/chrome alum coated glass slides and stored overnight in humidified atmosphere. Next day, sections were air-dried for 1 h at room temperature. Within 2 weeks sections were fixed at room temperature in fresh acetone containing 0.02% H2O2. After air-drying, sections were washed with PBS and incubated for 7 min at room temperature with 1% blocking reagent (TSA HRP kit, Invitrogen). Sections were incubated for 10 min with tyramide-biotin dissolved in amplification buffer (TSA HRP kit, Invitrogen) containing 0.15% H2O2. Next, sections were again incubated for 30 min with streptavidin-HRP. To reveal peroxidase activity, sections were incubated with AEC (3-amino-9-ethylcarbazole, Sigma) substrate for 10 min resulting in a bright red precipitate. Sections were counterstained with hematoxylin and embedded with glycerol-gelatin. In control stainings in which the first antibody was omitted or using an isotype control, no immunopositive cells could be detected. For the lymph nodes, the number of IL-17A producing cells is the mean of the number of IL-17A-positive cells of three non-adjacent sections.

Analysis of T cell reactivities

Ex vivo T cell responses against different antigens were analyzed as described previously (Brok et al. 2000). Briefly, PBMC were isolated by density gradient centrifugation over lymphocyte separation medium (Axis-Shield PoC AS, Oslo, Norway). Mononuclear cell (MNC) suspensions were prepared from aseptically removed spleen and ALN. Cells were cultured in triplicate for detection of proliferative responses towards rhMOG and a panel of MOG peptides. All antigens were tested at 5 µg/ml. After 48 h of culture, 0.5 µCi/well of tritiated thymidine (³H-Thy) was added and incorporation of radiolabel was determined after 18 h using a matrix 9600 β-counter (Packard 9600; Packard Instrument Company, Meriden, CT). Results are expressed as the mean stimulation index (SI), which is defined as the counts per minute (cpm) of stimulated cells divided by the cpm of unstimulated cells. SI values above 2.0 were considered to be relevant.

The phenotype of proliferating cells was determined as described previously (Kap et al. 2008). Briefly, 4x10⁶ viable MNC from ALN or spleen were suspended in 1 ml PBS and incubated for 7 min at room temperature with CFSE (final concentration 1.5 µM; Fluka, Deisenhofen, Germany). Labeled MNC were cultured for 7 days with rhMOG or MOG peptides. For flow cytometric analysis we used the following commercially available labeled monoclonal antibodies directed against human CD markers: Alexa Fluor 700-labeled anti-CD3 (BD Biosciences), APC-labeled anti-CD4 (DAKO, Glostrup, Denmark), biotinylated anti-CD8 (Serotec, Düsseldorf, Germany), and streptavidin PerCP (BD Biosciences). Flow cytometric analysis was performed on a FACSSort using FACSDiva software (BD Biosciences).
Statistics

Log-rank test, one-way ANOVA, and student’s t test were performed to statistically analyze the results. Results were considered statistically different at $p<0.05$.

Results

Plasma levels of anti-IL-17A mAb in rhMOG-immunized monkeys

The marmosets included in the study were randomly assigned to three treatment groups. The control group ($n=8$) received 1 ml/kg PBS and the two treatment groups received anti-IL-17A mAb at 3 mg/ml/kg ($n=8$) or 30 mg/ml/kg ($n=7$). The test substances were subcutaneously administered once per week starting 1 day before immunization. Plasma levels of the anti-IL-17A mAb were determined 7 days after each administration (Fig. 1). As expected, plasma concentrations of the antibody in the 3 mg/kg antibody dose group were about tenfold lower compared with the 30 mg/kg antibody dose group. In two animals (M04099 and M03144) of the 3 mg/kg antibody dose group plasma antibody trough levels, measured 1 week after administration, were remarkably low, i.e., below 1 μg/ml at psd 34, while these were above 1 μg/ml in all the other monkeys of this group. In the 30 mg/kg antibody dose animals, the plasma levels of the test substance were above 10 μg/ml from psd 6 throughout the experiment.

Clinical signs after anti-IL-17A mAb treatment

EAE was induced by a single immunization with rhMOG emulsified in CFA. The course of clinical signs in individual animals is depicted in Fig. 2. Solid lines represent clinical scores and dotted lines represent percentage body weight loss, serving as a surrogate disease marker in this model. The figure shows a similar disease course in the monkeys that developed full-blown clinical EAE. Three monkeys from the 3 mg/kg antibody dose group and one monkey in the 30 mg/kg antibody dose group did not reach the stage of hind-limb paralysis (score 3) within the pre-determined observation period of 113 days psd. In the 3 mg/kg antibody dose group one monkey (M05020) did not develop any signs of EAE within the pre-defined observation period. A second monkey of this group (M9913) reached EAE score 2 around psd 85, remitted and developed a second episode of score 2 EAE which remained present until the end of the observation period (psd 106). Two monkeys (M04008 and M05004) had to be sacrificed for ethical reasons with score 2 EAE due to serious body weight loss. It is noteworthy that the two monkeys with the lowest anti-IL-17A antibody plasma levels at day 34 (M03144 and M04099) were amongst the first monkeys to be sacrificed.

The EAE progression rate to overt neurological deficit was defined as the time interval between the day of rhMOG/CFA inoculation (psd 0) and the first day that clinical score 2 was observed. At the level of this parameter we observed a delayed EAE progression in the 3 mg/kg antibody dose group (mean psd, 69.3; range, 48–113) compared with the placebo group (mean psd, 57.1; range, 39–91) and the 30 mg/kg antibody dose group (mean psd, 60.0; range, 51–69). It is noteworthy that the variation between animals of this parameter was much lower in the 30 mg/kg antibody dose group (18 days) than in the placebo group (52 days) and the 3 mg/kg antibody dose group (>65 days).

The disease progression from EAE score 2 (ataxia) to the ethical end-points at score 3.0 (paraplegia) or excessive body weight loss (>20%) was more rapid in placebo group
monkeys (mean, 10.8 days; range, 3–20) than in monkeys from the 3 mg/kg antibody (mean, 12.9; range, 5–28) and 30 mg/kg antibody (mean, 13.7; range, 6–31) dose group. Statistical evaluation of both disease phases (score, 0–2 and score, 2 to >2) was done by survival analysis, but no significant differences were found (Fig. 3).

In summary, these data show a moderate inhibitory effect of the antibody treatment on the EAE course, although differences between the groups did not reach statistical significance.

Effect of anti-IL-17A mAb on white matter lesion load

Prior to sectioning for histological examination, postmortem T2-weighted MR images were made of the formalin-fixed cerebral hemispheres to visualize and quantitate the total lesion load. The analyzed parameters were lesion volume, T2, and MTR values of white matter lesions. In fixed brains, an increased T2 signal reflects demyelination, while a reduction of MTR values correlates well with the intensity of macrophage infiltration (Blezer et al. 2007).

Significant differences were not observed between the three groups (Table 1). In the analyzed hemispheres of two animals of the control group cerebral white matter lesions were absent. White matter lesion volumes were slightly increased in the 30 mg/kg antibody dose group compared with the placebo group. Especially monkey M05004 had clinical score 2 for a long time, which might explain the high lesion load. The lesions in the 3 and 30 mg/kg antibody dose group had a slightly higher T2 and lower MTR value compared with placebo (Table 1).

These results were confirmed by the quantitative immunohistochemistry data (Table 1). Both inflammation and demyelination were moderately increased, although not significant, in the two antibody-treated groups compared with the placebo group.

For the correct interpretation of these data it is pertinent to emphasize that lesion formation in this model starts well before neurological deficits can be observed. Moreover, most animals were killed at the stage of clinically evident EAE, which occurred at different time intervals after immunization. It may therefore not be surprising that major effects of the antibody treatment on the CNS pathology were not observed, neither in positive nor in negative direction.

IL-17A expression in the brain and lymphoid organs

To assess the effect of anti-IL-17A mAb treatment on in situ expression of IL-17A, cryosectioned brains and lymphoid organs were examined with immunohistochemistry. In the brains of control and anti-IL-17A mAb-treated animals IL-17A-positive cells were found in the perivascular space or in the vicinity of blood vessels (Fig. 4a). No differences were observed between placebo and anti-IL-17A mAb-treated animals.

IL-17A-producing cells in secondary lymphoid organs were mainly found in the red pulp of the spleen and in the medulla of the lymph nodes. We observed a trend towards higher numbers of IL-17A producing cells in the spleen of animals dosed with 3 mg/kg anti-IL-17A mAb compared with animals that had received placebo or 30 mg/kg antibody, but differences were not significant (Fig. 4a–b). By contrast, we detected lower numbers of IL-17A producing cells in the ALN and ILN, which drain the immunization sites, and the brain-draining CLN, in both treatment groups compared with the placebo group (Fig. 4a–b). Taken together, these
data suggest that the antibody treatment caused altered distribution of IL-17A producing cells over the different lymphoid compartments.

Effect of anti-IL-17A mAb on T cell proliferation

Cellular immune parameters of the three groups were examined to underpin the observed in vivo effects of the IL-17A mAb. To this end, PBMC and MNC of spleen and ALN were tested for proliferation against rhMOG and a panel of 23-mer overlapping MOG peptides.

Similar to previous studies (Kap et al. 2008), proliferation of PBMC during the disease course and at necropsy was low and remained confined to the immunizing rhMOG protein and a few peptides, i.e. MOG_{14-36}, MOG_{24-46}, and MOG_{34-56} (Fig. 5). No clear differences were observed between the three groups.

More robust proliferative responses were observed in the lymphoid organs, especially in the ALN and spleen. Figure 5 shows that in most animals the highest proliferative response was against rhMOG followed by peptides MOG_{14-36} and MOG_{24-46}. MNC from the ALN of the anti-IL-17A mAb-treated groups displayed a lower proliferative response against rhMOG and MOG peptides than MNC from ALN of the placebo group (Fig. 5). No differences between the three groups were observed in the spleen (Fig. 5).

Next, we analyzed the phenotype of the rhMOG responsive cells using the CFSE fluorescent dye dilution
Fig. 4 Differential expression of IL-17A in the brain and lymphoid organs. IL-17A expression was detected by immunohistochemistry. a (200×) Representative examples of each group for brain, spleen, and axillary (ALN), inguinal (ILN), and cervical (CLN) lymph nodes. b IL-17A positive cells per mm² (mean ± SEM). Only one CLN of the control group was analyzed. No IL-17A producing cells were detected in the ILN of the 30 mg/kg treated group. Statistical differences between the numbers of IL-17A positive cells were analyzed by One-way ANOVA. Results were considered statistically different at \( p < 0.05 \). No statistical differences were found between the three groups.
technique in combination with cell phenotyping for surface markers. RhMOG responsive CD3+CD4+ and CD3+CD8+ cells with CFSE dye dilution were visualized by flow cytometry. The percentages of CD3+CD4+ and CD3+CD8+ cells determined by fluorescent dye dilution did not differ significantly between placebo and anti-IL-17A mAb-treated groups (data not shown), suggesting that the treatment with anti-IL-17A mAb did not cause a skewing of the CD4/CD8 ratio of rhMOG-specific T cells.

In conclusion, treatment with the anti-IL-17A antibody did not exert an obvious effect on the quality or quantity of the cellular autoimmune reactions. However, we like to emphasize again that the analyzed lymphoid organs were collected at different time intervals after the EAE induction, creating a possible bias in the data.

**Discussion**

Key pathogenic elements for EAE development in rhMOG-immunized marmosets comprise CD40, IL-12, and IL-23 ('t Hart et al. 2005a; Boon et al. 2001; Brok et al. 2002; Laman et al. 2002). We have shown that an antibody against the shared p40 subunit of IL-12 and IL-23 caused profound suppression of disease symptoms when dosed from disease induction. One possible mechanism of action could have been a reduction in IL-17A that is one of the downstream cytokines from IL-23. However, the modest effect of an anti-IL-17A antibody in the current study indicates that a reduction in IL-17A does not account for the activity of IL-12 p40 antibody and that other cytokines, perhaps IL-12 directly or other cytokines downstream of IL-12 and/or IL-23, are more important in the induction phase of this non-relapsing model.

The results of the current study demonstrate that prophylactic treatment with an anti-IL-17A mAb induced only a moderate extension of the disease-free time interval after EAE induction. Furthermore, immune parameters of the disease, such as T cell proliferation against rhMOG or MOG peptides was only somewhat reduced, whereas also a marginal aggravation of CNS pathology was observed. However, the variations were all within the normal range of variation in this outbred model and may also be attributable to the longer survival. Using immunohistochemistry only a few IL-17A-positive cells could be detected in the brains of placebo and treated monkeys, but no significant differences

![Fig. 5](image.png)

Fig. 5 T cell responses after anti-IL-17A mAb treatment. PBMC and mononuclear cells isolated from the axillary lymph node and spleen at necropsy were tested for proliferation against rhMOG and MOG peptides. The stimulation index (SI) was calculated by dividing the counts per minute (cpm) of stimulated wells by cpm of unstimulated wells. The stimulation index is divided into groups with different shading as indicated in the figure.
between groups were observed. The data show a trend towards increased numbers of IL-17A-positive cells in the spleen in the 3 mg/kg group and a decreased number in the axillary and inguinal lymph nodes of both groups compared with placebo animals (not significant). The higher number of Th17 cells in the spleen of monkeys that have been treated with anti-IL-17A antibody compared with placebo monkeys may be explained by feedback regulation. The observation that IL-17A receptor deficient mice produce more IL-17A (Smith et al. 2008) indicates that IL-17A production may be regulated by a negative feedback loop. In conclusion, the current results fail to show a robust effect of the anti-IL-17A mAb treatment on the EAE course in rhMOG/CFA-immunized marmosets.

The moderate protection against EAE induction is in agreement with data reported by some groups in mouse EAE models. Also in IL-17A−/− mice (Komiyama et al. 2006), in mice treated with anti-IL-17A mAb (Chen et al. 2006; Hofstetter et al. 2005; Langrish et al. 2005), and in mice vaccinated against IL-17A (Rohn et al. 2006) only moderately delayed onset of clinical signs has been observed. However, other groups have demonstrated more robust effects of IL-17A on the EAE course in mice. Examples are by vaccination with IL-17A-OVA complexes that induce IL-17A blocking antibodies (Uyttenhove and Van Snick 2006) or adoptive transfer of IL-17+ and IL-17− T cells (Jager et al. 2009; Komiyama et al. 2006; Langrish et al. 2005). Furthermore, in the chronic-relapsing EAE model in Biozzi AB mice, prophylactic dosing with an anti-IL-17A antibody produced a small delay in onset of the acute phase of disease, whereas treatment with the antibody after the acute phase or during the secondary-progressive phase was very efficacious, and could even reverse the chronic motor disability (Smith, unpublished observations). Other groups suggested that Th17 are not pathogenic at all (Haak et al. 2009) or that besides Th17 cells, Th1 cells are also essential in EAE (Kroenke et al. 2008; Lees et al. 2008; O’Connor et al. 2008).

How can it be explained that IL-17A neutralization does not fully protect against EAE in the rhMOG-induced marmoset model? One possible explanation is the high complexity of the model. Unraveling of the pathogenic mechanism showed that the disease is initiated by MOG24–36-specific Th1 cells, which induce small inflammatory lesions (Brok et al. 2000). The moderate inflammatory injury caused by Th1 cells can be dramatically amplified by antibodies against conformational epitopes of MOG (Genaia et al. 1995), which are clearly produced in the rhMOG model. Another subset of highly reactive T cells with natural killer-cytotoxic T-lymphocyte characteristics and specific for MOG34–56 becomes involved at a later stage in the disease (Kap et al. 2008). Our previously reported studies in marmoset EAE models induced with MOG34–56 in CFA (Kap et al. 2008) or IFA (Jagessar et al. 2010) demonstrated that the same clinical and pathological endpoints can be reached via different immunopathogenic routes. It can thus be envisaged that after neutralization of IL-17A in the complex rhMOG/CFA-induced model other pathogenic mechanisms, such as Th1 cells, may come into play. It could also be possible that Th1 and Th17 cells act in different phases of the disease. In the previously reported study in the rhMOG marmoset model it was observed that Th1 cells specific for MOG24–36 are engaged in EAE induction, whereas MOG34–56-specific T cells, which produce mainly IL-17A, mediate disease progression (Jagessar et al. 2010; Kap et al. 2008). This suggests that, consistent with the results in the Biozzi ABH mouse model, neutralizing IL-17A in a later phase of the disease is more effective.

Yet, another possible explanation is that IL-17A, although being the signature cytokine of Th17 cells, is not the dominant pathogenic factor produced by Th17 cells. This explanation is supported by data in mice. IL-12/IL-23−/− mice are completely resistant to EAE, whereas only clinical scores and not the disease incidence is reduced in IL-17A−/− mice (Becher et al. 2002; Cua et al. 2003; Gran et al. 2002; Komiyama et al. 2006). Moreover, treatment of mice with anti-IL-23 antibody prevented either EAE induction or relapse, in contrast to the anti-IL-17A antibody treatment that blocked relapse, but had a smaller effect on the induction phase (Chen et al. 2006). This may suggest that other Th17 cytokines induced by IL-23 may be pathogenetically more relevant than IL-17A in the induction phase, such as IL-21 or IL-22 (reviewed in (Fouser et al. 2008)). In addition, Chen et al. suggested that IL-23 may also directly activate macrophages to produce inflammatory cytokines, such as IL-6, IL-1, and TNF-α, which may cause inflammation and demyelination even when IL-17A is neutralized (Chen et al. 2006).

Is there a place for an IL-17A neutralizing antibody in the treatment of MS? Axtell et al. reported that on the basis of clinical response to IFN-β two groups may be distinguished in the MS patient population. Responsiveness to IFN-β treatment requires a Th1 cytokine profile, whereas cases with a Th17 dominated cytokine profile are non-responders to IFN-β treatment (Axtell et al. 2010). It is tempting to speculate on the basis of these and our data that non-responder cases to IFN-β treatment in the MS population may benefit from treatment with IL-17A antibody. The ideal study design to obtain proof-of-principle may be a parallel effectivity analysis in the marmoset EAE model, where in one sibling of a chimeric twin EAE is induced with MOG34–56 in CFA, which is a Th1 prone model (Jagessar et al. 2010; Kap et al. 2008) and in the other twin sibling with MOG34–56 in IFA, which is a more Th17 prone disease (Jagessar et al. 2010; Kap et al. 2008).
In summary, we found that treatment with anti-IL-17A antibody induces a moderate delay of clinical EAE in marmosets, but that EAE is not completely abrogated. This suggests a pathogenic role for IL-17A in the marmoset EAE model and maybe in MS, but IL-17A may not be the only key pathogenic cytokine.

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Conflicts of Interest The authors of this manuscript do not report conflict of interest.

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