Tailoring the Antibody Response to Aggregated Aß Using Novel Alzheimer-Vaccines

Markus Mandler1,*, Radmila Santic1, Petra Gruber1, Yeliz Cinar2, Dagmar Pichler1, Susanne Aileen Funke2, Dieter Willbold2, Achim Schneeberger1, Walter Schmidt1, Frank Mattner1

1 AFFiRiS AG, Karl-Farkas-Gasse 22, A-1030, Vienna, Austria, 2 Institute for Structural Biochemistry (Institute of Complex Systems 6), Forschungszentrum Jülich, 52425, Jülich, Germany

These authors contributed equally to this work.

✉ Current address: Abbott GmbH & Co. KG, Max Planck Ring 2, 65205, Wiesbaden, Germany
¤ Current address: University for Applied Sciences and Arts, Faculty of Science, Bioanalytics, Friedrich-Streib-Straße 2, 96450, Coburg, Germany
* markus.mandler@affiris.com (MM); achim.schneeberger@affiris.com (AS)

Abstract

Recent evidence suggests Alzheimer-Disease (AD) to be driven by aggregated Aß. Capitalizing on the mechanism of molecular mimicry and applying several selection layers, we screened peptide libraries for moieties inducing antibodies selectively reacting with Aß-aggregates. The technology identified a pool of peptide candidates; two, AFFITOPES AD01 and AD02, were assessed as vaccination antigens and compared to Aβ1-6, the targeted epitope. When conjugated to Keyhole Limpet Hemocyanin (KLH) and adjuvanted with aluminum, all three peptides induced Aß-targeting antibodies (Abs). In contrast to Aß1-6, AD01- or AD02-induced Abs were characterized by selectivity for aggregated forms of Aß and absence of reactivity with related molecules such as Amyloid Precursor Protein (APP)/secreted APP-alpha (sAPPa). Administration of AFFITOPES-vaccines to APP-transgenic mice was found to reduce their cerebral amyloid burden, the associated neuropathological alterations and to improve their cognitive functions. Thus, the AFFITOME-technology delivers vaccines capable of inducing a distinct Ab response. Their features may be beneficial to AD-patients, a hypothesis currently tested within a phase-II-study.

Introduction

Alzheimer’s disease (AD) is the most prevalent neurodegenerative disorder currently affecting 28 million people worldwide [1]. It typically presents with a characteristic amnestic dysfunction associated with other cognitive-, behavioral- and neuropsychiatric changes impairing a given individual’s (social) function and ultimately resulting in its death [2]. Available treatments include three acetylcholinesterase inhibitors (AChEI) and one N-Methyl-D-aspartate
(NMDA) antagonist. Their effects are small and only symptomatic in nature [3]. Thus, there is a high medical need for a disease-modifying drug.

Accumulation of Amyloid Beta (Aβ) appears to be an early event and central to the disease process. Aβ is a proteolytic fragment of the amyloid precursor protein (APP) [4, 5, 6]. APP-cleavage results in several peptides including Aβ1-40 and Aβ1-42, which are subject to further processing. Recent studies suggest Aβ-variants and aggregates drive the disease process [7, 8].

Immunotherapy offers the possibility to specifically address Aβ-variants and aggregates. However, targeting self-proteins by immunological means bears the risk of autoimmunity [9]. This is exemplified by autoimmune reactions following the administration of cancer vaccines [10]. While regarded as immune privileged, the brain is not excluded from such reactions but represents a relevant target organ as experienced with AN1792 [11] or deduced from the existence of paraneoplastic autoimmune Central Nervous System (CNS) syndromes [12].

With regard to pathological autoimmunity, both cellular- and humoral effector mechanisms need to be considered. Avoidance of T-cell responses against CNS-targets is crucial as demonstrated by AN1792-triggered cases of meningoencephalitis. All second generation AD-vaccines in clinical development, are designed to avoid activation of target-specific T-cells by restricting antigen length to <8 amino acids (aa) or by excluding bona-fide T-cells epitopes (CAD106, ACC001, UB-311, ACI-24 [13, 14, 15]).

The risk of pathological humoral autoimmunity is primarily related to the antigenic epitopes addressed. Efficient control of this risk requires selective targeting of structures exclusively expressed in disease, so called neo-epitopes. The free N-terminus of native, aggregated Aβ is an excellent example of a neo-epitope. Exclusive reactivity to this structure would preclude antibodies (Abs) induced to cross-react with APP and related molecules such as secreted APP-alpha (sAPPa).

Conventional Aβ-vaccines [13, 14, 15, 16] are conjugates of an N-terminal Aβ-fragment and a carrier. The N-terminus of Aβ is accessible in monomers, aggregates and amyloid plaques. Abs elicited by conventional conjugate-vaccines typically fail to discriminate between the various Aβ-aggregation states. Given the fact that Aβ-monomers possess physiological functions [17, 18, 19, 20] while aggregates exert neurotoxic and synaptotoxic effects [21, 22, 23, 24], a potential benefit of vaccines may require them to elicit Abs selectively addressing Aβ-aggregates.

To generate a vaccine that integrates both, targeting the Aβ-N-terminus and selective recognition of Aβ-aggregates, we devised a technology based on mechanisms of molecular mimicry. Peptide libraries were screened for peptides exhibiting both features. This yielded several hits. Two of them, AD01 and AD02, were characterized in more detail. Both did exhibit the intended specificity, and were found to reduce pathological alterations and to ameliorate behavioral deficits of APP-transgenic Tg2576-mice. Results obtained not only suggested them to be disease-modifying but to have a safety profile superior to conventional Aβ1-6-based vaccines.

**Material and Methods**

**AFFITOPE identification and vaccine formulation**

AFFITOPE-peptides were identified by screening of peptide libraries (phage display: New England BioLabs, USA; randomized synthetic hexa- and hepta-peptide libraries: Mimotopes Pty., France or MULTIPIN peptide technology), with monoclonal antibodies (mAbs, AFFiRiS, Austria) specific for the free N-terminus of Aβ1-40/42. Identified peptides (EMC microcollections, Germany) were conjugated to KLH (Biosyn, Germany) using N-gamma-Maleimidobutyryloxysuccinimide ester (GMBS, Thermo Scientific, USA) and adsorbed to Aluminum-hydroxide (ALUM, Brenntag, Denmark). 30μg peptide/vaccine-dose containing 0.1% ALUM were applied to animals.
Animal experiments

All animal experiments were performed in accordance with the Austrian Animal Experiments Act (TVG2012) using 8–12 week old female C57Bl/6 mice (Charles River, Germany), or Tg2576-mice (Taconic Farms, USA; 129S6/SvEvTac). Experiments were performed under approval numbers: LF1-TVG-22/004-2007; M58/007052/2011/7 and LF1-TVG-22/0102011. General health was checked by modified Smith Kline Beecham, Harwell, Imperial College, Royal London Hospital, phenotype assessment (SHIRPA) primary observational screen [25]. Mice were injected s.c. 3–6 times in monthly or biweekly intervals. Blood was taken in regular intervals, plasma prepared and stored until further use. At study end mice were sacrificed, cerebrospinal fluid (CSF), brains were collected and hemispheres separated. One hemisphere was fixed in 4% Paraformaldehyde (PFA, Sigma Aldrich, USA), dehydrated and paraffin-embedded. Brain tissue was sectioned at 7µM using a sliding microtome (Leitz, Germany) and sections were mounted on Superfrost Plus Slides (Menzel, Germany). The second hemisphere was quick-frozen at -80°C until further extraction.

Titer determination by ELISA

Standard enzyme-linked immunosorbent assay (ELISA) technology was used to measure levels of vaccine-induced antibodies in plasma and CSF [26]. Substrates used included murine (Anaspec, USA) and human (BACHEM, CH) Aβ1-40/42 (each at 5µg/ml), KLH (1µg/ml), recombinant sAPPa (1µg/ml, Sigma-Aldrich, USA), peptide-Bovine serum albumin (BSA) conjugates (1µM), or Aβ-aggregates (5µg/ml, immobilized via Streptavidin). Optical density (OD) was measured at 405nm using a micro-well reader (Tecan, CH). ODmax/2 was calculated. For determination of antibody selectivity for different Aβ species (monomers, oligomers and fibrils), relative units were calculated as the ratio of OD values for individual measurements: e.g. OD405nm of Oligomer-specific ELISA signals and OD405nm of Monomer-specific ELISA signals. Abs 3A5 (AFFiRiS, Austria), mAbP2-1 (Life-Technologies, USA) and 6E10 (Covance, USA) served as positive controls.

Preparation and characterisation of Aβ-monomers, -oligomers and —fibrils

Preparation of Aβ mono and oligomers (<100kd) was performed according to Johansson et al. with slight modifications [27]. Pure C-terminally biotinylated Aβ1-42 was used to prepare seedless Aβ monomers. A 1/10 mix of biotinylated and unmodified Aβ1-42 (Anaspec, USA) was used for oligomer- and fibril-production. For preparation of Aβ-mono and oligomers Aβ was first resolved in Hexafluoro-2-propanol (HFIP) over night and subsequently removed by vacuum centrifugation. Aβ peptides were then resuspended and separated using a Superdex 75-10/300 column (GE Healthcare, UK). Elution of monomers and oligomers was determined by detection at 214nm with oligomers eluting at 8 ml and monomers at 14,5 ml, respectively. Column calibration was done according to manufacturers protocol (LMW Gel Filtration Calibration Kit; GE Healthcare, UK). For fibril preparation Aβ peptides were resuspended in 1xPBS and fibrils were assembled by constant rotation of peptide solutions for 24h at 350rpm (37°C). Fibril-preparations were then centrifuged and the pellet was resuspended in elution buffer used for gelfiltration. Aggregation of Aβ-species was confirmed by Thioflavin-T, Western- and Dot blot analysis (see Appendix).

APP-FACS analysis

To test for APP-specific antibodies a Fluorescence-activated cell sorting (FACS) assay based on Chinese hamster ovary (CHO)-cells stably expressing a fusion protein of human APP and enhanced green fluorescent protein (eGFP) (APP-751-EGFP in pCMV-Sport 6, APP: NP_958816, pCMV-Sport 6 eGFP-FLAG-tagged (Gift from J.M.Peters, IMP, Austria)) was
used. A mixture of transfected and un-transfected CHO-cells (50% each) was exposed to diluted plasma and analysed for double positive cells (eGFP and APP) with a FACScan (BD Biosciences, USA). mAbP2-1 served as positive control. For each sample 10,000 events were acquired and analysed using CellQuest software (BD Biosciences).

**Behavioral tests**

To analyse cognitive dysfunction immunised Tg2576 animals were subjected to Modified Morris water maze task (MWM, with changes) and contextual fear conditioning (CFC, with changes), both analyzed using AnyMaze software (Stoelting Co, USA). MWM was subdivided into cued-, hidden task, and probe-trial. Animals were trained in a tap-water filled 110-cm pool, allowed to swim for 60s with platform occupancy for 10s prior to the next trial. 24h after the hidden training, memory retention was determined in a single 60s probe-trial without a platform. The percent of distance swam and time spent in each quadrant was determined.

For CFC, on day 1 mice were placed in the conditioning chamber (AFFiRiS), allowed to habituate for 2 min. and received three 0.8mA foot-shocks in 2 min intervals plus 30s rest. To assess contextual learning on day 2, animals were readmitted to the chamber and monitored for 5 min. with s120-240 chosen as time frame for analysis (time freezing = lack of movement except for respiration). The first two minutes of day 1 were considered as baseline-freezing which was subtracted from day 2 values. Cognitive testing was initiated 4 weeks prior to sacrifice which was required to complete both cognitive tests for the individual animals including habituation phases at the site of testing.

**Immunohistochemistry (IHC), immunofluorescence (IF) and analysis of cerebral Aβ**

IHC/IF analysis was done as described previously [26]. Reactivity of vaccine-induced antibodies to Aβ and APP was determined using an adapted Tissue Amyloid Plaque Immunoreactivity (TAPIR) analysis on untreated Tg2576- and human AD-brain sections (n = 4, obtained from Novagen, USA (n = 1) or the UCSD ADRC Brain Bank (n = 3); patients analysed were n = 3 female and n = 1 male; females: Braak stage VI and male patient Braak stage V) using plasma samples and an APP-specific mAb (22C11, EMD Millipore, USA) as control. Competition experiments of AD01- and AD02 induced antibodies were performed using specific AFFITOPE-peptides at a final concentration of 10 μM. Control antibody used for amyloid staining on human brain sections was the monoclonal antibody BAM10 (Sigma, USA). For murine sections monoclonal antibody 6E10 (Signet, USA) was used as control antibody.

For Aβ-specific IF-staining, brain sections of immunized Tg2576 were processed for analysis of amyloid load and incidence of amyloid bearing vessels using mAb 3A5 (AFFiRiS AG, Austria) [26]. All secondary reagents were obtained from Vector Labs (USA). For TAPIR analysis, color reactions were performed using DAB Substrate Kit. For IF, sections were mounted and counterstained using DAPI-containing VECTASHIELD-HardSet Mounting Medium. Sections were examined using MIRAX-SCAN (Carl Zeiss AG, Germany). AD-like pathology in animals was assessed by determining the total tissue area of coronal cross sections of the total brain as well as the 3A5 positive area on the respective brain sections were determined and the relative cerebral area occupied by amyloid deposits was calculated using a semi-automated area recognition program (eDefiniens Architect XD; www.definiens.com). For analysis three slides/animal and ≤ five individual sections/slide were assessed. Sections carrying tissue artifacts or aberrant staining were excluded. To assess the number of Aβ-positive vessels, 3A5 stained sections (n = 3 slides/animal covering cortex and hippocampus and up to five individual sections per slide) have been analysed. Aβ-positive vessels were manually counted in sub-
regions of the cortex as well as in the hippocampus. Number of positive vessels per mm² was determined.

**Analysis of micro-hemorrhaging**

To assess the number of micro-hemorrhages, sections were stained using the Iron Stain Kit (Sigma Aldrich, USA) according to manufacturer’s protocol. 3 slides/animal covering cortex and hippocampus and up to five individual sections per slide have been analysed. Prussian blue-positive spots were manually counted in sub-regions of the cortex as well as in the hippocampus. Number of positive spots per animal was determined.

**Analysis of cerebral levels of Aβ by ELISA**

The frozen brain hemispheres were thawed and homogenized in homogenisation buffer (50mM HEPES (pH 7.3), 5mM EDTA, with proteinase inhibitor cocktail: Complete Mini, Roche, CH) and centrifugated at 4°C for 30 minutes at 40.000 rpm. The supernatant was aliquoted and stored at -80°C as soluble fraction. The pellet was re-homogenized in Guanidine-HCl buffer (5M Guanidine-HCl, 50mM HEPES (pH 7,3), 5mM EDTA with proteinase inhibitor cocktail: Complete Mini, Roche, CH) and centrifuged at 1600g. The supernatant was dialysed against PBS, aliquoted and stored at -80°C as insoluble fraction. Fractions were analysed for protein content using the Quick Start Bradford Protein Assay according to manufacturer’s protocol (BioRad, USA).

For quantification of Aβ40 and Aβ42 peptides in soluble and insoluble fractions, an ELISA analysis was used (Human Amyloid Beta 40 and Human Amyloid Beta 42 ELISA kits, EMD-Milipore, USA). The concentration of amyloid peptides in ng/mg of total protein was calculated for Aβ40 and Aβ42 in both fractions (soluble and insoluble).

**ELISPOT analysis**

Animals (C57Bl/6 mice) were immunized three times in biweekly intervals with AD01-conjugate (30µg net peptide content/mouse/immunization), AD02-conjugate (30µg net peptide content/mouse/immunization) or Ovalbumin (100µg/mouse/immunisation), adjuvanted with CpG/polyR as adjuvant for T-cell stimulation (CpG (ODN1668: 5’ TCC ATG AGC TTC CTG ATG CT 3’, Invivogen, San Diego, USA) 32 µg/mouse; polyR 100 µg/mouse; Sigma-Aldrich). 1 week after the final immunization animals were sacrificed, splenocytes isolated and analysed for the presence of target specific T-cells by ELISPOT analysis. ELISPOT analysis was performed using Ready-SET-Go kits obtained from eBioscience (San Diego, USA) according to the manufacturer’s protocol. Full length Aβ1-42 (50µg/ml), carrier (KLH, 50µg/ml) or short MHC-I or MHC-II restricted Ovalbumin-derived peptides Ova 244 (TEWTSSNVMEERKIKV; MHC class II restricted; 10µg/ml) and Ova 245 (SIINFEKL; MHC class I restricted; 10µg/ml) as positive control for T-cell induction were used for splenocyte restimulation. Stimulated cells were assayed for the secretion of either Interleukin 4 (IL4) or Interferon gamma (IFNg). The stimulation was controlled by application of two positive control stimulators, for IL4 secretion, Phorbol-12-Myristate-13-Acetate (PMA, working conc.: 20nM) and ionomycin (working conc.:750nM) and for IFNγ secretion Concanavalin A (ConA); working conc.:10 µg/ml) were used, respectively.

**Statistical analysis**

All experiments were done blind-coded. To determine statistical significance, values were compared using (i) one-way analysis of variance for unpaired samples with Tukey’s Multiple
Comparison Tests, (ii) unpaired T-tests with Welch correction or (iii) Kruskal-Wallis-H-Test with Dunn’s Multiple Comparison Tests. For correlation analysis a Spearman-Rank-Correlation has been calculated.

**Results**

To generate an Aß-vaccine not activating Aß-specific T-cells but inducing Abs selectively recognizing aggregated Aß and at the same time being specific for the Aß-N-terminus, we screened peptide libraries with Abs applying various selection filters. Specifically, mAbs directed against the N-terminus of intact, full length Aß1-40/42; (aa1-6: DAEFHR) were used to screen $10^9$ peptides from different hexa- and hepta-peptide libraries for 6–7 mer peptides for binding. Specificity of peptide hits was assessed by competition with Aß1-6 (DAEFRH). Several rounds of selection yielded 68 candidates fulfilling both of the above criteria. Comparing the sequence of the n = 68 peptides to the one of native Aß revealed no candidate with only 1aa exchange and a difference of n = 2aa in 16%, n = 3aa in 31%, n = 4aa in 23.5% or n = 5aa in 9%. The remaining 20.5% of the peptides differed at all positions. As a next step, out of the 68 candidates, 17 were randomly picked and tested for their ability to elicit antibodies reacting to the peptides used for immunization (=immunizing peptide) and, at the same time, Aß. To this end, they were coupled to KLH, which served as carrier, adsorbed to aluminum (ALUM), the adjuvant used, and subcutaneously injected into C57BL/6 and Tg2576-mice. While all 17 induced Ig-Abs reactive with the respective peptide, only 14 elicited Abs recognizing Aß1-10 coupled to BSA. Those conjugates were used to mimic binding to Aß-aggregates, as peptide-BSA conjugates should show a local enrichment of Aß-N-termini probably similar to the situation present in full length, native Aß-aggregates. Two examples, AD01 and AD02, characterized by a difference of 50% in their amino acid sequence compared to the targeted Aß epitope, were characterized in more detail and compared to Aß1-6-KLH-vaccine.

**Immunogenicity of AD-AFFITOPEs**

To test the immunogenicity of AD01 and AD02 in comparison to Aß1-6, Tg2576-mice were injected 6x, s.c., at 4-week intervals with either conjugate-vaccine containing 30μg net peptide. Vaccination induced Abs were measured in plasma samples at defined time points after immunization (AD01 (n = 9), AD02 (n = 8) and Aß1-6 conjugate (n = 9)). All 3 elicited strong and comparable IgG titers towards the peptide used for immunization (Fig 1A). Both AFFITOPEs, AD01 and AD02, elicited Abs to the N-terminus of human Aß at levels comparable to the Aß1-6-KLH conjugate-vaccine (Fig 1B). Of note, the IgG responses triggered by the 3 conjugate-vaccines followed the same kinetics (Fig. 1C). Titers reached a plateau after 2 immunizations, which was stable during the treatment period. Analyzing the CSF of AD02-immunized Tg2576-mice demonstrated the presence of peptide-/Aß-specific Abs at a level of 0.1–0.7% (0.31% +/- 0.05%) of the respective plasma levels (Fig. 1D).

**Specificity of the AFFITOPE induced antibody response**

We next assessed the specificity of the Abs induced in more detail. Neither AD01-, AD02- nor Aß1-6-induced plasma samples reacted with irrelevant control peptides such as Aß11-20 offered as BSA conjugates (ELISA, not shown). The reactivity of AD01- and AD02-induced Abs towards murine Aß was limited and comparable, whereas the signal obtained with Aß1-6-induced sera was 4 times higher (Fig. 1E). Interestingly, while AD01-elicited plasma samples reacted strongly with AD01 offered as BSA conjugate in an ELISA setting they barely did so with AD02-BSA; the opposite was true for AD02-induced samples (approx. 13-fold difference,
This lacking reactivity towards the respectively other AFFITOPE while retaining reactivity towards the Aß-N-terminus is most probably explained by the fact that the amino acid sequences of the two AFFITOPEs tested in this experiment differ from each other by 67% (n = 4/6aa) but show a similar difference to the native Aß sequence of 50% (n = 3/6aa).
Exclusion of APP reactivity

Abs used to identify the AD01/AD02 AFFITOPE-family were characterized by recognition of the Aβ-N-terminus and a lack of reactivity with full length APP, the precursor of Aβ. To check whether, as intended, AD01- and AD02-induced Abs would mirror this characteristic of the paternal mAbs, plasma of AFFITOPE-vaccinated animals were analyzed for APP-binding employing a FACS assay based on CHO-cells expressing human APP on their surface. Plasma of immunized animals were analyzed in comparison to APP-specific mAbP2-1, which showed a dose dependent signal (Fig. 2A-C). Of note, such a signal was not seen in plasma from AD01 (n = 8) or AD02 immunized animals (n = 30; representative example in Fig. 2E and data not shown). By contrast, a substantial portion of plasma from Aβ1-6-immunized animals, n = 6/30, was found to contain APP-specific Abs (Fig. 2F).

In addition, plasma samples were also tested for Abs directed against sAPPα, an important mediator of APP-function. None of the AFFITOPE samples tested (n = 20; Fig. 2G) contained sAPPα specific Abs detectable by ELISA. This differentiated them from Aβ1-6-based vaccines, which induced sAPPα-reactive Abs in all animals tested (n = 10/10; Fig. 2G). Interestingly, this APP cross-reactivity was not directly correlated with the absolute anti-Aβ titer in these samples (Spearman-Rank-Correlation r = 0.4316, p = 0.2129; Fig. 2H), implying that this reactivity against cleaved forms of APP is a unique feature of a fraction of Abs present within the oligoclonal response elicited by Aβ1-6-based vaccines.

As a third method to assess the potential cross reactivity of AD01- and AD02-induced antibodies to human APP/sAPPα and an APP-eGFP fusion protein, a Western blot analysis was performed (see Appendix, S2A Fig.). In this assay, both AD01- and AD02 induced plasma samples failed to detect full length human APP/sAPPα in brain extracts from 12 month old Tg2576 animals or in cell extracts from CHO cells stably expressing a fusion protein of human APP and eGFP (also used in the FACS based analysis mentioned above). As expected the APP specific positive control antibody 22C11 was able to detect APP/sAPPα and APP-eGFP using this method.

Differential reactivity towards Aβ-aggregation states

To characterize AD01- and AD02-induced plasma samples with regard to their reactivity towards defined Aβ-aggregation states we devised ELISA systems covering monomers, oligomers and fibrils. 6E10, known to bind Aβ in all its aggregation states (Fig 3), was used as standard. Of note, the patterns of reactivity seen with AD01, AD02 and Aβ1-6 were found to differ substantially. Aβ1-6-induced antibodies behaved like 6E10 reacting equally well with all Aβ-aggregation states tested (Fig. 3). By contrast, AFFITOPE-elicited Abs exhibited a differential recognition pattern of the various Aβ-aggregation states. AD01-induced plasma reacted with aggregated (both oligomers and fibrils) but not with monomeric Aβ. AD02-induced sera were found to recognize fibrillar Aβ only (Fig. 3) and showed only limited reactivity towards oligomeric Aβ preparations and no reactivity with monomeric Aβ using ELISA based analyses.

In a second set of experiments, Aβ aggregate specificity of AD01 and AD02 induced antibodies was analysed employing peptide ELISAs with aggregated Aβ both as bait coated on the ELISA plate and as peptide used as competitor for antibody binding to the immobilized Aβ aggregates (see Appendix, S2C Fig.). Indeed this competition experiment revealed a concentration dependent, specific reduction of the binding to aggregates and hence further substantiates the claim of selective aggregate recognition by AFFITOPE induced antibodies.

In addition, AD01 and AD02 induced antibodies were also tested for binding to monomeric, dimeric and aggregated Aβ by Western blot analyses. As suggested by ELISA results (see
Fig. 3), AD02 induced antibodies showed a lack of reactivity to monomeric and dimeric Aβ and against low molecular weight (MW) Aβ aggregates (<100kD). Reactivity could only be detected to high MW aggregates (>100kd, see Appendix, S2B Fig.). In contrast to ELISA based results, AD01 induced antibodies displayed reactivity to monomeric and dimeric Aβ probably due to different sensitivity of the assays used. In line with previous results (see Fig. 3 and
they also reacted against all aggregated Aβ forms present on Western blots similar to the non-conformer specific control antibody 4G8 (see Appendix; S2B Fig.).

Furthermore, we assessed the reactivity of AD01- and AD02-induced plasma on brain tissue of Tg2576-mice and of AD-patients (n = 4) applying a standard DAB immuno-histochemical protocol [26] and using 22C11 as comparator reacting with full length APP. AD01- or AD02-elicited plasma were found to exclusively stain amyloid deposits and to spare neuronal surfaces with a comparable staining pattern as the Aβ specific control antibody 6E10 in brain sections of Tg2576-mice (Fig. 4 A, B, D, E, J and K). In addition, a loss of immunoreactivity could be detected when AD01- or AD02-elicited plasma was pre-adsorbed with the respective AFFITOPE-peptides to inhibit antibody binding to amyloid plaques, indicating specificity of the AFFITOPE-induced antibody staining observed in these animals (Fig. 4G, H). Immunohistochemical analysis of APP reactivity showed an opposite staining pattern with specific reactivity on neuronal cell walls and plaque-surrounding neuritic alterations both in the hippocampus and the cortex of Tg2576 mice (Fig. 4M and data not shown)

A comparable staining pattern was seen on human AD-brain sections. The analysis of human sections of n = 4 patients revealed a specific amyloid deposit staining of AD01-(Fig. 4F)
and AD02-induced Abs (Fig. 4I) present in murine plasma similar to staining obtained by using control antibody BAM10 (Fig. 4C). No staining was detectable with plasma derived from a naïve, untreated animal (Fig. 4L). These findings corroborated the Aβ-specificity described above characterized by the lack of APP cross-reactivity (Fig. 2).

Figure 4. AFFITOPE-induced antibodies detect amyloid deposits but spare neuronal APP on murine and human brain sections. Sections prepared from the hippocampus (A, D, G, K, M) and the cortex (B, E, H), of a 12 month old Tg2576 mouse were incubated with plasma of AD01- (D, J) or AD02-treated mice (E, K) and, for control purposes, with the antibodies 6E10 (A+B) and mAb 22C11 (J, K, M), recognizing Aβ and full length APP. G) and H) show a loss of immunoreactivity on Tg2576 brain sections incubated with AD01- and AD02-induced samples which were mixed with the respective AFFITOPE peptide to inhibit AFFITOPE specific staining (i.e. G: AD01-induced plasma + 10µM AD01 peptide, H: AD02-induced plasma + 10µM AD02 peptide). Sections prepared from the cortex of a female AD patient (C, F, I, L Braak stage VI,) were incubated with the Aβ specific control antibody BAM10 (C), AD01- (F) and AD02-induced plasma (I) or plasma from naïve control mice as negative control (L). Binding of the Abs was detected using a standard DAB immunohistochemistry protocol. The analysis of human sections reveals a specific amyloid deposit staining of AD01-(F) and AD02-induced Abs (I) present in murine plasma similar to staining obtained by using control antibody BAM10 (C), whereas no staining was detectable with plasma derived from a naïve, untreated animal (L). Pictures were taken at a magnification of 20x. Scale bars: 200µm; circles in G and H indicate amyloid deposits devoid of amyloid specific staining by AD01- or AD02- induced antibodies following peptide-specific competition.

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AFFITOPE vaccination does not activate Aβ-specific T-cells

AD01 and AD02 are 7 amino acids long (6 containing the mimicry information +1 residue used for conjugation). Therefore, these AFFITOPE peptides should be too short to bind to MHC molecules and activate peptide specific T-cells. In addition, their amino acid sequences differ from the one of the N-terminus of Aβ. To formally test whether conjugate vaccines containing AFFITOPEs AD01 and AD02 would activate AFFITOPE-peptide or Aβ-specific T-cells, splenocytes of immunized, non-transgenic animals were analyzed by ELISPOT. To this end, groups of n = 6 C56BL/6 mice were immunized 3 times at 2 week-intervals with AD01-KLH, AD02-KLH or ovalbumin (OVA). One week after the last immunization, splenocytes were isolated and stimulated in vitro with the carrier (KLH), Aβ or ovalbumin-derived MHC class I- (IFNg assay) and MHC class II (IL-4 assay) binding peptides. Cultures were assessed for IL-4- and IFNg-producing cells, which, given the stimulation conditions, reflect T-lymphocytes that had been primed during vaccination. Assay controls included stimulation with PMA/ionomycin (IL-4 assay) and concanavalin A (IFNg assay) and confirmed cell viability/functionality (Fig. 5A and B). Restimulation with the carrier protein demonstrated that both AFFITOPE vaccines had led to the induction of a KLH-specific T-cell response. Such a response was not evident in OVA-immunized animals. However, in vitro stimulation of splenocytes derived from AD01 and AD02-immunized animals with either AD01- or AD02-peptides as well as with recombinant Aβ did not yield a signal over background confirming the expected inability of the two AFFITOPEs of activating either AFFITOPE peptide- as well as Aβ-specific T-cells. This view is supported by experimental evidence from transgenic animals undergoing active immunotherapy using AD01 and AD02: To test whether AD01 and AD02 immunotherapy would lead to brain infiltration of T-cells, brain sections of n = 20 AD02-immunized-, n = 9 AD01-immunized-, and n = 10 carrier-treated Tg2576 mice were subjected to immunohistochemical examination using CD3 specific antibodies to detect potential CD3+ T cells (see Appendix, S3 Fig.). Despite that fact that in most of the AD01- and AD02- treated animals immunisation had resulted in a reduction of amyloid deposition (see Fig. 6 and 7), none of the brains was found to be infiltrated by CD3+ T cells.

Both vaccine candidates lower cerebral Aβ without triggering cerebral amyloid angiopathy (CAA) or micro-hemorrhages (MH)

To test whether AD01 and AD02 would lower cerebral amyloid load, groups of 6-months old Tg2576-mice (n = 10/group) were vaccinated 6x at monthly intervals with either vaccine (independent experiments), and sacrificed at 14 months of age. Their brains were assessed for diffuse and dense-cored plaques by IF-staining using monoclonal antibody 3A5. Cortical as well as hippocampal sections of KLH/ALUM-injected controls were covered by numerous amyloid plaques. They covered on average 2.00% (AD01 experiment) and 0.69% (AD02 experiment) of the area analyzed. By contrast, respective brain areas of AD01- and AD02-immunized Tg2576-mice contained significantly less deposits (Fig. 6A-D) covering 0.21% (Fig. 6E) and 0.77% (Fig. 6F), respectively. Thus, AD01 reduced the area covered by amyloid by 62% (p<0.05) and AD02 by 70% (p<0.05).

In addition to the analysis of amyloid deposition in situ we also assessed the effect of AFFITOPE vaccination on the cerebral level of Aβ1-40 and Aβ1-42 by peptide ELISA. Therefore, brain samples of AD01 and AD02 treated Tg2576 animals were extracted and soluble and insoluble brain fractions were subjected to Human Aβ40 and Human Aβ42 ELISA (EMD-Millipore, USA) analysis. Neither AD01 nor AD02 treated animals showed a significant change of soluble Aβ1-40 and Aβ1-42 following immunotherapy (see Fig. 7A and B). In contrast for both vaccines, insoluble Aβ was reduced significantly following immunotherapy (see Fig. 7C).
For AD02 a 60% reduction of Aβ1-40 (p < 0.05) and a 62% (p = 0.056) reduction of Aβ1-42 could be detected. AD01 showed a reduction of 69% (Aβ1-40, p < 0.05) and 78% (Aβ1-42, p < 0.01), respectively. These differences are most probably reflecting a selective removal of aggregated and deposited Aβ while soluble forms were only reduced to a low amount.

Figure 5. AD01 and AD02 immunization does not induce self-reactive T-Cells. Neither AD01 nor AD02 treated mice showed any sign of Aβ-specific T-cell activation in two ELISPOT assays (A+B). Re-stimulation using the carrier (KLH) was resulting in a stimulation of IL4 and Interferon gamma (INFγ) secretion, indicative of the presence of carrier specific T-cells following immunization with AD01 and AD02. The positive control Ovalbumin was able to induce a slightly higher Interferon gamma secretion than the carrier used in the AFFITOPE vaccines (B). A+B depict two representative ELISPOT analyses following vaccination of Ovalbumin, AD01 and AD02. A) IL4 secretion following splenocyte restimulation using carrier (KLH) and Aβ compared to the controls OVA244 (TEWTSSNVMEERKIKV; MHC class II restricted to demonstrate Ovalbumin induced T-cells) and PMA/ionomycin (PMA/Ion); B) INFγ secretion following splenocyte restimulation compared to the positive controls OVA245 (SIINFEKL; MHC class I restricted to demonstrate Ovalbumin induced T-cells) and Concanavalin A (ConA). Bg describes the background of secretion in non-stimulated cells in this assay. Numbers are the total number of spots per million of cells seeded on the ELISPOT plates.

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As amyloid removal appears to partially occur via the vasculature and peripheral sink mechanisms [31, 32, 33] and can be associated with enhanced micro-bleedings following active and passive immunotherapy [34], blood-vessels of relevant brain regions (cortex and hippocampus) were analyzed for their amyloid content by 3A5 staining and the same regions were assessed for microbleedings by Prussian Blue staining, respectively. At the time point assessed, the number of 3A5-positive vessels per mm² is comparable for control- and AFFITOPE-treated
Figure 7. AFFITOPE immunization reduces cerebral amyloid levels in Tg2576 mice (ELISA). Groups of Tg2576 mice (n = 10/group) received 6 monthly injections of KLH/ALUM or AD01-, AD02-conjugate vaccines. Brains were isolated, 8 weeks after the 6th immunization, extracted and soluble and insoluble brain fractions were subjected to Human Aβ40 and Human Aβ42 ELISA (EMD-Milipore, USA) analysis. Neither AD01- (A) nor AD02 treated animals (B) showed a significant change of soluble Aβ1-40 and Aβ1-42 following immunotherapy as compared to control immunized animals. Insoluble Aβ was reduced significantly following immunotherapy. C) AD01 treated animals showed a 69% reduction of Aβ1-40 levels (p = 0.005) and a 78% reduction of Aβ1-42 (p = 0.015), respectively. D) For AD02 a 60% reduction of Aβ1-40 (p = 0.033) and a 62% (p = 0.056) reduction of Aβ1-42 could be detected. Results are expressed as average ± SEM and are given as ng/mg total protein. Black bars represent Aβ1-40 and white bars represent Aβ1-42 values. Asterisks in C+D indicate statistical significant difference (*p<0.05, **p<0.01); doi:10.1371/journal.pone.0115237.g007

Figure 8. Cerebral amyloid angiopathy and microhemorrhaging are unchanged following AFFITOPE-immunization. The analysis of the incidence of amyloid bearing vessel in AFFITOPE- and control treated animals by assessing 3A5 staining in cerebral vessels reveals no significant differences (A-C). A) Example from a cortical section of a control animal. B) Example from a cortical section of an AD02-treated animal. C) Quantitative analysis demonstrating the average number of 3A5 positive vessels/mm² (avg. ± SEM) in control (n = 9), and AD02-treated animals (n = 8). The analysis of the incidence of cerebral micro-hemorrhages in these animals by assessing Hemosiderin staining (= Prussian Blue) did not show significant differences at 14 months of age (D-F). D) Example from a cortical section of a control animal. E) Example from a cortical section of an AD02-treated animal. F) Quantitative analysis demonstrating the average number of Hemosiderin positive vessels/animal in control (n = 9), and AD02 treated animals (n = 8, respectively); Arrows indicate 3A5 positive vessels, arrowheads indicate amyloid deposits. Scale bar: 50µm; pictures taken at 20x magnification doi:10.1371/journal.pone.0115237.g008
animals (see Fig. 8) compatible with the view that AFFITOPE vaccines do not enhance CAA after repeated immunization. Moreover, the MH-number was low and comparable for both immunized and control mice (Fig. 8) indicating no effect on the occurrence of micro-hemorrhages following AFFITOPE immunization.

Both vaccine candidates improve functional deficits of APP-transgenic mice

To evaluate the effect of AFFITOPE-vaccination on cognitive functions, we applied, MWM (AD01 and AD02) and CFC (AD02 only) analyzing spatial and contextual memory and learning in Tg2576-mice. In the MWM learning phase, both AD02- and control-treated mice (receiving KLH/ALUM) showed a similar learning capability (Fig. 9A). During probe-trials for assessing memory retention, AD02-treated mice performed significantly better than control mice (Fig. 9B, p < 0.05). No differences in swim speed between the two groups were detectable during the probe trial (data not shown). Analysing the percentage of mice per group searching in the target quadrant for >25% of the time, showed that 82% of the AD02-treated animals were able to correctly remember the former platform position compared to 40% in the control group (data not shown). A similar MWM-analysis using AD01-immunized animals revealed a similar improvement in spatial memory. However, while AD01-immunized animals performed better than controls, the effect did not reach statistical significance (Fig. 9C, p = 0.1).
CFC demonstrated that AD02-treated mice were superior to control animals (Fig. 9D+E). Although showing slightly lower freezing levels in response to day 1 foot-shocks (Fig. 9D), animals froze significantly more on day 2 during the representative 2 min period of the retention phase of the test (p<0.01, Fig. 9D+E). The significant improvement of AD02-treated animals was also detectable by averaging the performance in CFC during the whole 5 minutes on day 2 (p<0.05, data not shown). Taken together, these findings demonstrate AFFITOPE-vaccination to effectively reduce memory defects in Tg2576-mice in two learning paradigms.

**Discussion**

The work presented aimed at generating novel Aβ-targeting AD-vaccines with specific features. Specifically, they were designed to (i) trigger Abs specific for the Aβ-N-terminus (ii) being selective for aggregated Aβ and (iii) preclude the activation of AFFITOPE peptide- or Aβ-specific T-cells. This was accomplished via the mechanism of molecular mimicry and by applying sequential selection filters. It led to the identification of two candidates, AD01 and AD02, which fulfill the predefined criteria and exhibit disease-modifying activity in the models tested.

Molecular mimicry, in terms of humoral immunity, denotes the phenomenon of Abs not only recognizing a single epitope but more than one resembling each other and, thus, being indistinguishable for the Ab. This is not uncommon. Most examples we are aware of are related to negative effects. They include autoimmune reactions as a result of bacterial- or viral infections, neoplasias (e.g., paraneoplastic CNS disorders) or vaccination (e.g., AN1792-triggered cases of meningoencephalitis) [12, 35, 36]. We explored the possibility of exploiting molecular mimicry for the development of AD-vaccines with optimized safety- and efficacy features. To this end, Abs known to bind the Aβ-N-terminus (DAEFRH), were exposed to a pool of 10^9 6- or 7-mer peptides. In addition to DAEFRH, a total of 68 peptides were found to bind the Abs employed (frequency of cross-reactive peptides: 4.0x10^-8). 20.5% of the hits differed at every aa-position from the original epitope, the remaining had 2 or more aa-exchanges. For all 68 peptides, binding could be competed with DAEFRH arguing for their interaction with the antigen binding sites of the Abs. Out of the 68 peptides, 17 were picked and tested for their ability to elicit Abs when administered as peptide-KLH conjugates adjuvanted with aluminum. All 17 elicited an Ab-response to the immunizing peptide, 14 of them induced Abs reacting with Aβ1-10-BSA conjugates, which resemble to some extent Aβ-aggregates given the high density of binding sites on BSA following conjugation. These data confirm and quantify the phenomenon of molecular mimicry for mAbs primarily known to bind to the Aβ-N-terminus. They also demonstrate, at least for mice, that it is possible to reverse and hence exploit the process. That is, cross-reacting peptides, foreign to the human proteome, can trigger Abs that recognize the „original”epitope and have imprinted „additional”features, e.g. selective Aβ-aggregate recognition and selective recognition of peptides derived from the same screen.

Beyond pathophysiology, design of AD-vaccines has to consider physiological functions and the dynamics of the ensuing Ab-response. While AD-pathophysiology is complex, it appears that toxicity resides within the aggregated Aβ fraction affecting neurons and synapses [21, 22, 23, 24]. By contrast, monomeric Aβ as well as sAPPa and APP possess physiological functions. Monomeric Aβ regulates the proliferation of neural progenitors and contributes to synaptic function [17, 18, 19, 20, 37]. APP and sAPPa are involved in the development and plasticity of the nervous system, regulation of neurite outgrowth, neuronal proliferation and contribute to cognitive performance and memory [38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50]. Abs recognizing physiological elements of the Aβ-pathway, such as the ones induced by Aβ1-6-KLH, could have negative effects via various, mutually non-exclusive mechanisms:
interference with the above molecules and their functions, Ab-triggered cytotoxicity. Moreover, APP/sAPPa and serum Aβ would sequester such Abs thereby lowering their levels. Of note, the advantages of Abs with the above specificity may not be discernible in AD-models used, which are characterized by a strong over-expression of Aβ and sAPPa. So far, AFFITOPE-vaccines are the first second-generation vaccines which report to spare binding to the above molecules. Other examples either do not provide any analyses on cross-reactivity [14, 15, 16] or report a lack of APP/sAPP reduction without providing data or an analysis of antibody binding other than on fixed tissue, therefore probably underestimating potential cross-reactivity in vivo [13].

Beyond differential targeting of Aβ-variants, AD01 and AD02 reduced cerebral amyloid load by 62- and 70%, respectively. This compares favorably to conventional vaccines [13, 14, 15, 16, 51]. In addition to the IF analysis, assessing the amount of insoluble Aβ showed a selective and significant reduction following AD01 and AD02 immunotherapy, whereas soluble forms of Aβ1-40 and Aβ1-42 were not significantly changed. This selective removal of insoluble and deposited Aβ could further support the selectivity of AFFITOPE-vaccines for Aβ-aggregates.

Amyloid reduction was not associated with an increase in detectable MH or CAA, as seen with other vaccines [52, 53], but with improvement of cognitive dysfunction as assessed by MWM and CFC. Furthermore, we could also demonstrate the inability of the two AFFITOPE vaccines of activating either AFFITOPE peptide- as well as Aβ-specific T-cells. This is in line with similar experimental results obtained using AFFITOPE vaccines targeting alpha Synuclein (aSyn) in animal models of synucleinopathies [54]. In these experiments, no aSyn AFFITOPE peptide- or target specific T-cells (i.e. alpha Synuclein) could be detected by ELISPOT- or immunohistochemical analyses following active immunotherapy in mice using peptide conjugate vaccines [54].

In conclusion, data presented support the feasibility of the proposed technology based on molecular mimicry. Vaccine candidates identified, AFFITOPEs AD01 and AD02, exhibit high specificity (Aβ-aggregates but no monomers) defining their high safety (e.g., sparing of APP/ sAPPa-recognition) and efficacy profiles. Given their disease-modifying potential both have been introduced to clinical testing in mild to moderate AD with AD02 being currently assessed in a multicentre phase II study in early AD-patients.

Supporting Information

S1 Fig. Aggregation analysis of Aβ-monomers, -oligomers and — fibrils. To assess aggregation status of Aβ-monomers, -oligomers and — fibrils, ThT Fluorescence analysis (A) as well as Dot blot (B) and Western blot (C) were performed. (A) Monomer preparations show relative fluorescence units (RFU) close to background indicating the absence of fibrillar Aβ. Oligomeric and fibrillar preparations contained ThT positive aggregates with fibrillar preparations containing significantly more positive aggregates (RFUs >5000) than oligomeric preparations (RFUs of ca. ≤2000). (B) Dot Blot analysis using NAB 228 showed equal signals for Aβ-monomers, -oligomers and — fibrils whereas analysis using A11 did show only oligomer specific signals and failed to detect Aβ-monomer and — fibril preparations indicating that only the oligomer preparations were also containing oligomeric species, not detectable in the other two preparations. (C) Western Blot analysis using NAB 228 showed equal signals for Aβ-monomers and — oligomers. Oligomeric preparations contained Aβ-dimers, -trimers and -tetramers as well as oligomers with a size of approx. 35–40kd (weak signal in C) in this analysis. No fibril specific signals could be detected.

(TIF)
S2 Fig. Reactivity of AD01- and AD02-induced antibodies to Aβ and sAPPa. The reactivity of AD01- and AD02-induced Abs towards full length APP/sAPPa/APP-eGFP as well as different forms of Aβ was assessed by Western blot analysis (A+B). Specificity of AFFITOPE-induced antibodies for aggregated Aβ was assessed by competition ELISA (C). A) Western blot analysis using brain extracts form a 12 month old female Tg2576 mouse and from CHO cells stably expressing a human APP-eGFP fusion protein showed a lack of reactivity of AD01- and AD02- induced antibodies against full length APP/sAPPa and APP-eGFP fusion protein whereas the positive control antibody 22C11 (APP-specific) was able to detect APP/sAPPa and APP-eGFP, respectively. B) Western Blot analysis of aggregated recombinant Aβ revealed a lack of reactivity of AD02-induced Abs against monomeric and dimeric Aβ as well as low molecular weight (MW) Aβ aggregates (<100kD). AD02-induced Abs react predominantly against high MW Aβ aggregates (>100kD). AD01 induced antibodies, as the non-conformer specific antibody 4G8 showed binding to Aβ-monomers,—dimers, as well as low and MW Aβ aggregates. C) ELISA experiment demonstrating the selectivity of AD01- and AD02- induced antibodies for aggregated Aβ by concentration dependent, specific competition using aggregated Aβ. Bars represent the means of OD values (at 405nm) of individual samples derived from single animals immunized with AD01 or AD02. Reactivity of sera was tested against aggregated Aβ1-42 immobilised on ELISA plates (1μM). Competition was done using plasma samples (dilution of 1/100) and aggregate concentrations of 0.5µg/ml and 1µg/ml, respectively. A: 1. . . Tg2576 brain extract; 2. . . CHO APP-eGFP cell extract; B: m+d. . . Aβ monomer and dimer, l+h. . . low and high MW Aβ aggregates; C: sec. only. . . secondary antibody used as background control for the ELISA; grey and black bars indicate OD values for AD01 (grey) and AD02 (black) induced antibodies (+/- aggregated Aβ)

(TIF)

S3 Fig. T-cell response to immunization with AD01 and AD02. Immunostaining of T-cells present in the perivascular space with an anti-CD3 antibody. No CD3-positive cells were observed in brains of Control (A), AD01 (B) or AD02 (C) immunized animals. CD3 positive cells could be detected in murine splenic tissue sections used as positive controls for staining (D). Pictures in A-C show CA1 region of brains from 14 month old Tg2576 animals undergoing immunotherapy. Per mouse, a total of ≤20 individual brain sections were assessed. Scale bar = 50 µm, pictures taken at a 20x magnification.

(TIF)

S1 Appendix. Materials And Methods.

(DOCX)

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Author Contributions

 Responsible for study design and study conduct: MM. Contributed equally to analysis of immune reactions and pathologic assessment: MM RS. Responsible for behavior experiments: PG. Responsible for assessment of aggregate reactivity: YC SAF DW. Wrote the paper: MM AS FM WS. Responsible for study design and study conduct: MM. Contributed equally to analysis
of immune reactions and pathologic assessment: MM RS. Responsible for behavior experiments: PG. Responsible for assessment of aggregate reactivity: YC SAF DW. Critical review of the manuscript and agreement on final version: RS PG YC DP SAF DW.

References

1. Wimo A, Prince M (2010) World Alzheimer Report 2010: The glocal economic impact of dementia.
2. Dubois B, Feldman HH, Jacova C, Cummings JL, Dekosky ST, et al. (2010) Revising the definition of Alzheimer’s disease: a new lexicon. Lancet Neurol 9: 1118–1127. doi: 10.1016/S1474-4422(10)7023-4 PMID: 20934914
3. Lyketsos CG, Colenda CC, Beck C, Blank K, Doraiswamy MP, et al. (2006) Position statement of the American Association for Geriatric Psychiatry regarding principles of care for patients with dementia resulting from Alzheimer disease. Am J Geriatr Psychiatry 14: 561–572. doi: 10.1097/01.JGP.0000221334.65330.55 PMID: 16816009
4. Haass C, Koo EH, Mellon A, Hung AY, Selkoe DJ (1992) Targeting of cell-surface beta-amyloid precursor protein to lysosomes: alternative processing into amyloid-bearing fragments. Nature 357: 500–503. doi: 10.1038/357500a0 PMID: 1406936
5. Seubert P, Vigo-Pelfrey C, Esch F, Lee M, Dovey H, et al. (1992) Isolation and quantification of soluble Alzheimer’s beta-peptide from biological fluids. Nature 359: 325–327. doi: 10.1038/359325a0 PMID: 1439760
6. Shoji M, Golde TE, Ghiso J, Cheung TT, Estus S, et al. (1992) Production of the Alzheimer amyloid beta protein by normal proteolytic processing. Science 258: 126–129. doi: 10.1126/science.1439760 PMID: 1439760
7. Tomic JL, Pensalfini A, Head E, Glabe CG (2009) Soluble fibrillar oligomer levels are elevated in Alzheimer’s disease brain and correlate with cognitive dysfunction. Hum Brain Dis 35: 352–358. doi: 10.1016/j.nbd.2009.05.024 PMID: 19523517
8. Benilova I, Karran E, De Strooper B (2012) The toxic Abeta oligomer and Alzheimer’s disease: an emperor in need of clothes. Nat Neurosci 15: 349–357. doi: 10.1038/nn.3028 PMID: 22286176
9. Schneeberger A, Mandler M, Mattner F, Schmidt W. (2010) AFFITOME(R) technology in neurodegenerative diseases: the doubling advantage. Hum Vaccin 6: 948–952. doi: 10.4161/hv.6.11.13217 PMID: 20980801
10. Le Poole IC, Luiten RM (2008) Autoimmune etiology of generalized vitiligo. Curr Dir Autoimmun 10: 227–243. doi: 10.1159/000131485 PMID: 18460889
11. Orregozo JM, Gilman S, Dartigues JF, Laurent B, Puel M, et al. (2003) Subacute meningoencephalitis in a subset of patients with AD after Abeta42 immunization. Neurology 61: 46–54. doi: 10.1212/01.WNL.0000073623.84147.A8 PMID: 12947195
12. Rosenfeld MR, Dalmau J. (2010) Update on paraneoplastic and autoimmune disorders of the central nervous system. Semin Neurol 30: 320–331. doi: 10.1055/s-0030-1255223 PMID: 20577938
13. Wiessner C, Wiederhold KH, Tissot AC, Frey P, Danner S, et al. (2011) The second-generation active Abeta immunotherapy CAD106 reduces amyloid accumulation in APP transgenic mice while minimizing potential side effects. J Neurosci 31: 9323–9331. doi: 10.1523/JNEUROSCI.0293-11.2011 PMID: 21697382
14. Muhls A, Hickman DT, Pihlgren M, Chuard N, Girienon V, et al. (2007) Liposomal vaccines with conformation-specific amyloid peptide antigens define immune response and efficacy in APP transgenic mice. Proc Natl Acad Sci U S A 104: 9810–9815. doi: 10.1073/pnas.0703137104 PMID: 17517995
15. Wang CY, Finsrud CL, Wallfeld AM, Sia C, Sokoll KK, et al. (2007) Site-specific UBTh amyloid-beta vaccine for immunotherapy of Alzheimer’s disease. Vaccine 25: 3041–3052. doi: 10.1016/j.vaccine.2007.01.032 PMID: 17287052
16. Liu B, Frost JL, Sun J, Fu H, Grimes S, et al. (2013) MER5101, a novel Abeta1-15:DT conjugate vaccine, generates a robust anti-Abeta antibody response and attenuates Abeta pathology and cognitive deficits in APPswe/PS1DeltaE9 transgenic mice. J Neurosci 33: 7027–7037. doi: 10.1523/JNEUROSCI.5924-12.2013 PMID: 23595760
17. Giuffrida ML, Caraci F, Pignataro B, Cataldo S, De bona P, et al. (2009) Beta-amyloid monomers are neuroprotective. J Neurosci 29: 10582–10587. doi: 10.1523/JNEUROSCI.1763-09.2009 PMID: 19710311
18. Chen Y, Dong C (2009) Abeta40 promotes neuronal cell fate in neural progenitor cells. Cell Death Differ 16: 386–394. doi: 10.1038/cdd.2008.94 PMID: 18566800
19. Kamenetz F, Tomita T, Hisieh H, Seabrook G, Borchelt D, et al. (2003) APP processing and synaptic function. Neuron 37: 925–937. doi: 10.1016/S0896-6273(03)00124-7 PMID: 12670422
20. Sothibundhu A, Li QX, Thangnipon W, Coulson EJ (2009) Abeta(1–42) stimulates adult SVZ neurogenesis through the p75 neurotrophin receptor. Neurobiology Aging 30: 1975–1985. doi: 10.1016/j.neurobiolaging.2008.02.004 PMID: 18374455

21. Freir DB, Fedrani R, Scully D, Smith IM, Selkoe DJ, et al. (2011) Abeta oligomers inhibit synapse remodelling necessary for memory consolidation. Neurobiology Aging 32: 2211–2218. doi: 10.1016/j.neurobiolaging.2010.01.001 PMID: 2097446

22. Hartley DM, Walsh DM, Ye CP, Diehl T, Vasquez S, et al. (1999) Protodiffusible intermediates of amyloid beta-protein induce acute electrophysiological changes and progressive neurotoxicity in cortical neurons. J Neurosci 19: 8876–8884. PMID: 10516307

23. Mucke L, Selkoe DJ (2012) Neurotoxicity of amyloid beta-protein: synaptic and network dysfunction. Cold Spring Harb Perspect Med 2: a006338. PMID: 22762015

24. Reed MN, Hofmeister JJ, Jungbauer L, Welzel AT, Yu C, et al. (2011) Cognitive effects of cell-derived and synthetically derived Abeta oligomers. Neurobiology Aging 32: 1784–1794. doi: 10.1016/j.neurobiolaging.2009.11.007 PMID: 20031278

25. Rogers DC, Jones DN, Nelson PR, Jones CM, Quilter CA, et al. (1999) Use of SHIRPA and discriminant analysis to characterise marked differences in the behavioural phenotype of six inbred mouse strains. Behav Brain Res 105: 207–217. doi: 10.1016/S0166-4328(99)00072-8 PMID: 10563494

26. Mandler M, Rockenstein E, Ubhi K, Hansen L, Adame A, et al. (2012) Detection of peri-synaptic amyloid-beta pyroglutamate aggregates in early stages of Alzheimer’s disease and in AbetaPP transgenic mice using a novel monoclonal antibody. J Alzheimers Dis 28: 783–794. PMID: 22064070

27. Johansson AS, Berglindh-Delin F, Karlsson G, Edwards K, Gellerfors P, et al. (2006) Physiochemical characterization of the Alzheimer’s disease-related peptides A beta 1–42Arctic and A beta 1–42wt. FEBs J 273: 2618–2630. doi: 10.1111/j.1742-4658.2006.05263.x PMID: 16817891

28. Morris R (1984) Developments of a water-maze procedure for studying spatial learning in the rat. J Neurosci Methods 11: 47–60. doi: 10.1016/0165-0270(84)90007-4 PMID: 6471907

29. Comery TA, Martone RL, Aschmies S, Atchison KP, Diamantidis G, et al. (2005) Acute gamma-secretase inhibition improves contextual fear conditioning in the Tg2576 mouse model of Alzheimer’s disease. J Neurosci 25: 8898–8902. doi: 10.1523/JNEUROSCI.2693-05.2005 PMID: 16192379

30. Hock C, Konietzko U, Streffer JR, Tracy J, Signorell A, et al. (2003) Antibodies against beta-amyloid slow cognitive decline in Alzheimer’s disease. Neuron 38: 547–554. doi: 10.1016/S0896-6273(03)00294-0 PMID: 12765607

31. DeMattos RB, Bales KR, Cummins DJ, Dodart JC, Paul SM, et al. (2001) Peripheral anti-A beta antibody alters CNS and plasma A beta clearance and decreases brain A beta burden in a mouse model of Alzheimer’s disease. Proc Natl Acad Sci U S A 98: 8850–8855. doi: 10.1073/pnas.151261398 PMID: 11438712

32. DeMattos RB, Bales KR, Cummins DJ, Paul SM, Holtzman DM (2002) Brain to plasma amyloid-beta efflux: a measure of brain amyloid burden in a mouse model of Alzheimer’s disease. Science 295: 2264–2267. doi: 10.1126/science.1067568 PMID: 11910111

33. Lemere CA, Spooner ET, LaFrancois J, Malester B, Mori C, et al. (2003) Evidence for peripheral clearance of cerebral Abeta protein following chronic, active Abeta immunization in PSAPP mice. Neurobiol Dis 14: 10–18. doi: 10.1016/S0969-9961(03)00044-5 PMID: 13678662

34. Wilcock DM, Colton CA (2009) Immunotherapy, vascular pathology, and microhemorrhages in transgenic mice. CNS Neurol Drug Disc Targets 8: 50–64. doi: 10.2174/187152709787601858 PMID: 19275636

35. Oyarbide-Valencia K, van den Boorn JG, Denman CJ, Li M, Carlson JM, et al. (2006) Therapeutic implications of autoimmune vitiligo T cells. Autoimmun Rev 5: 486–492. doi: 10.1016/j.autrev.2006.03.012 PMID: 16920575

36. Fujinami RS, von Herrath MG, Christen U, Whilton JL (2006) Molecular mimicry, bystander activation, or viral persistence: infections and autoimmune disease. Clin Microbiol Rev 19: 84–90. doi: 10.1128/CMR.19.1.80-94.2006 PMID: 16418524

37. Yankner BA, Caceres A, Duffy LK (1990) Nerve growth factor potentiates the neurotoxicity of beta amyloid. Proc Natl Acad Sci U S A 87: 9020–9023. doi: 10.1073/pnas.87.22.9020 PMID: 2174172

38. Heber S, Herm J, Gajic V, Hainfellner J, Aguzzi A, et al. (2000) Mice with combined gene knock-outs reveal essential and partially redundant functions of amyloid precursor protein family members. J Neurosci 20: 7951–7963. PMID: 11050115

39. Herm J, Anliker B, Heber S, Ring S, Fuhrmann M, et al. (2004) Cortical dysplasia resembling human type 2 lissencephaly in mice lacking all three APP family members. EMBO J 23: 4106–4115. doi: 10.1038/sj.emboj.7600390 PMID: 15385965
40. Leyssen M, Ayaz D, Hebert SS, Reeve S, De Strooper B, et al. (2005) Amyloid precursor protein promotes post-developmental neurite arborization in the Drosophila brain. EMBO J 24: 2944–2955. doi: 10.1038/sj.emboj.7600757 PMID: 16052209

41. Wang P, Yang G, Mosier DR, Chang P, Zaidi T, et al. (2005) Defective neuromuscular synapses in mice lacking amyloid precursor protein (APP) and APP-Like protein 2. J Neurosci 25: 1219–1225. doi: 10.1523/JNEUROSCI.4660-04.2005 PMID: 15689559

42. Priller C, Bauer T, Mitteregger G, Krebs B, Kretzschmar HA, et al. (2006) Synapse formation and function is modulated by the amyloid precursor protein. J Neurosci 26: 7212–7221. doi: 10.1523/JNEUROSCI.1450-06.2006 PMID: 16822978

43. Hayashi Y, Kashiwagi K, Ohta J, Nakajima M, Kawashima T, et al. (1994) Alzheimer amyloid protein precursor enhances proliferation of neural stem cells from fetal rat brain. Biochem Biophys Res Commun 205: 936–943. doi: 10.1006/bbrc.1994.2755 PMID: 7999135

44. Small DH, Nurcombe V, Chang P, Zaidi T, et al. (2005) Defective neuromuscular synapses in mice lacking amyloid precursor protein (APP) and APP-Like protein 2. J Neurosci 25: 1219–1225. doi: 10.1523/JNEUROSCI.4660-04.2005 PMID: 15689559

45. Priller C, Bauer T, Mitteregger G, Krebs B, Kretzschmar HA, et al. (2006) Synapse formation and function is modulated by the amyloid precursor protein. J Neurosci 26: 7212–7221. doi: 10.1523/JNEUROSCI.1450-06.2006 PMID: 16822978

46. Taylor CJ, Ireland DR, Ballagh I, Bourne K, Marechal NM, et al. (2008) Endogenous secreted amyloid precursor protein-alpha regulates proliferation of neural stem cells and enhances LTP in hippocampal slices. Neurobiol Dis 31: 250–260. doi: 10.1016/j.nbd.2008.04.011 PMID: 18585048

47. Ohsawa I, Takamura C, Morimoto T, Ishiguro M, Kohsaka S (1999) Amino-terminal region of secreted form of amyloid precursor protein stimulates proliferation of neural stem cells. Eur J Neurosci 11: 1907–1913. doi: 10.1046/j.1460-9568.1999.00601.x PMID: 10336659

48. Meziane H, Dodart JC, Mathis C, Little S, Clemens J, et al. (1998) Memory-enhancing effects of secreted forms of the beta-amyloid precursor protein in normal and amnestic mice. Proc Natl Acad Sci U S A 95: 12683–12688. doi: 10.1073/pnas.95.21.12683 PMID: 9770546

49. Pride M, Seubert P, Grundman M, Hagen M, Eldridge J, et al. (2008) Progress in the active immunotherapeutic approach to Alzheimer’s disease: clinical investigations into AN1792-associated meningoencephalitis. Neurodegener Dis 5: 194–196. doi: 10.1159/000113700 PMID: 18322388

50. Wilcock DM, Gharkholonarehe N, Van Nostrand WE, Davis J, Vitek MP, et al. (2009) Amyloid reduction by amyloid-beta vaccination also reduces mouse tau pathology and protects from neuron loss in two mouse models of Alzheimer’s disease. J Neurosci 29: 7957–7965. doi: 10.1523/JNEUROSCI.1339-09.2009 PMID: 19553436

51. Mandler M, Valera E, Rockenstein E, Weninger H, Patrick C, et al. (2014) Next-generation active immunization approach for synucleinopathies: implications for Parkinson’s disease clinical trials. Acta Neuropathol 127(6): 861–79. doi: 10.1007/s00401-014-1256-4 PMID: 24525765