Amyloid beta 4-42 in Alzheimer’s disease: Target, Therapy, Mechanism

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Gregory Antonios

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Abstract

Targeting Aβ has recently been the main objective in Alzheimer’s disease therapeutic approaches. Passive immunization trials have encountered undesirable side effects but the therapy remains a promising option. Aβ_{4-42} has not been previously considered as a target for AD immunotherapy. Tg4-42 mouse model has been recently established and validated as an advantageous research tool in AD. The current work deals with Aβ_{4-42} and explores its potential as a target, while elucidating therapeutic mechanism and crystallizing the NT4X Fab in complex with Aβ_{4-19} aiming to further reveal the structural basis of antibody: target affinity.

In the present work, the novel monoclonal antibody NT4X specifically reacts with N-truncated Aβ at position 4 of Aβ. It binds N-truncated Aβ under native and denaturing conditions and rescues in vitro toxicity of Aβ_{4-42} and that of pyroglutamate Aβ_{pE3-42}. The Fab fragment of the antibody was also able to prevent the in vitro toxicity caused by Aβ_{4-42} in rat primary cortical neuron cultures.

Aβ_{4-42} intracerebroventricular injection into wildtype mice induced a behavioral deficit, shown as a reduction in alteration rate in a Y-Maze, which was prevented using the NT4X. The Fab fragment of the antibody, at a higher dosage, was also able to prevent the in vivo behavioral deficit in a replicate experiment.

The Tg4-42 homozygous mouse model, expressed Aβ_{4-42} and allows for is intraneuronal accumulation. At 6 months of age, the model already exhibits 50% neuronal loss in the CA1 region of the hippocampus and severe reference memory deficits in a Morris water maze. Preventative passive immunotherapy with the NT4X antibody and its Fab fragment was able to mitigate neuron loss significantly and rescue spatial memory deficits as compared to an isotype control group.

Crystallization of the NT4X Fab in complex with Aβ_{4-19} has been successful. Diffraction data has been collected at 2.8 Å. Efforts to
resolve the crystal structure of the complex are ongoing. Issues with antibody-target engagement in terms of affinity, species and conformation of Aβ bound may be dealt with before going to a clinical setting, with the help of information arising from the crystal structure of the NT4X Fab: Aβ_{4-19} complex.
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List of Abbreviations

Ach: acetylcholinesterase
AD: Alzheimer’s disease
AICD: APP intracellular domain
APOE: Apolipoprotein E
APP: Amyloid precursor protein
Aβ: amyloid beta
BACE: Beta secretase
BBB: blood-brain barrier
C99: C-terminal 99-residue fragment
CA1: Region I of hippocampus proper
CAA: Cerebral amyloid angiopathy
CTF: ABP Carboxy-terminal fragment
ELISA: Enzyme-linked Immunosorbent Assay
EOAD: early onset Alzheimer’s disease
FW: flow-through
GWAS: genome-wide association studies
HRP: horseradish peroxidase
ICV: intracerebroventricular
LOAD: late onset Alzheimer’s disease
MRI: magnetic resonance imaging
MWCO: molecular weight cut-off
NaOAc: Sodium Acetate
NFT: Neurofibrillary tangles
NMDA: N-methyl-D-aspartate receptor
PBS: Phosphate buffered saline
PCBs: polychlorinated biphenyls
PEG: polyethylene glycol
PFA: paraformaldehyde
PHF: paired helical filaments
PS-1: Presenilin 1
PS-2: Presenilin 2
RT: room temperature
SDS-PAGE: sodium dodecyl sulfate polyacrylamide gel electrophoresis
TBS-T: Tris- buffered saline supplemented with Tween-20
TRH: thyrotropin releasing hormone
WHO: World Health Organization
1. Introduction

1.1. Dementia

The clinical syndrome referred to as “Dementia” encompasses a wide range of underlying pathologies. Most commonly these include Alzheimer’s disease (AD), Lewy body dementia, vascular dementia and frontotemporal dementia. The common grounds underlying the aforementioned diseases is a progressive deterioration in cognitive ability and capacity for independent living, brought on via a slow process of neurodegeneration (Sousa et al., 2009, 2010). Attention, problem solving, language, memory and visual-spatial are the most common areas affected by the resulting neurodegeneration (Lee, 2011). In the initial stages of dementia, manifestations of the symptoms are very subtle and often go unnoticed.

Accounting for around 60 – 80% of dementia cases, Alzheimer’s disease is the most common form of dementia (Alzheimer’s Association, 2012).

1.2. Alzheimer’s disease

Senile dementia, before the term Alzheimer’s disease was coined at the turn of the previous century following the description of Auguste Deter’s case, was well known, dating back to the ancient Greek and Roman philosophers and physicians. However, the histological findings and concepts presented by early researchers of AD, most notably Dr. Alois Alzheimer, were a turning point for the understanding of dementia (Berchtold & Cotman, 1998). Estimates in France, Sweden, the United States and the Netherlands, consider Alzheimer’s disease the most costly disease for society after the age of 65. The epidemiology of AD is closely linked with the increase in human life span (Bonin-Guillaume et al., 2005).
1.3. Epidemiology

The World Health Organization (WHO) projected that in 2005, 0.379% of people worldwide had dementia, and that the prevalence would increase to 0.441% in 2015 and to 0.556% in 2030. A more recent WHO study estimates that in 2010, 47.5 million people had dementia, 35 million people of whom had Alzheimer’s disease (WHO, 2010, 2015). In line with the previous, another study estimates that by 2050, the prevalence of AD will quadruple, by which time 1 in 85 persons worldwide will be living with the disease (Brookmeyer et al., 2007).

The death attributable to Alzheimer’s disease and other dementias, seems to be correlated to both yearly income (World Bank income categories) and living region (WHO region). Where the low-income category contributes to 0.41% of deaths as compared to 2.84% for the high-income category. Correspondingly, the Eurozone contributes to 2.04% of AD and other dementias related deaths as compared to 0.1% for Africa, 1.47 for the Americas and 0.42% for Eastern Mediterranean countries (WHO, 2006).

1.4. Risk Factors

1.4.1. Health

Data from cross-sectional and longitudinal studies relating health risk factors to the risk of developing Alzheimer’s disease have mostly remained inconsistent.

Hypertension has been suggested to increase the risk of AD through protein extravasation into brain tissue, caused by a decrease of the vascular integrity of the blood-brain barrier (BBB) (Kalaria, 2010). Likewise, Types 2 diabetes, in observational studies, has been linked to almost doubling the risk of AD (Leibson et al., 1997; Luchsinger, 2001).
On the other hand, a lower risk of AD has been associated with case-control studies examining smoking (Ferini-Strambi et al., 1990; Tyas, 1996). Other health risk factors studied have been cerebrovascular disease (Weishaupt, 2003), body weight (Gustafson et al., 2009; Razay, 2005), plasma lipid levels (Kuo et al., 1998; Michikawa, 2003; Muckle & Roy, 1985) and traumatic brain injury (Mayeux et al., 1995; Rasmussen et al., 2009).

1.4.2. Environmental

Some studies have examined the influence of toxic chemical exposures on the risk of dementia, Alzheimer’s disease and eventually, cognitive decline. Cognitive function including, verbal memory/learning, language, visual memory and more importantly visual spatial/motor functions, have been recently shown to be adversely affected by low-level additive exposure to lead outside of the work setting (Shih et al., 2006).

The potential role of aluminum in Alzheimer’s disease has been controversial since 1965. The relevance of a recent laboratory observation to actual real-life conditions has yet to be established since the prediction of aluminum absorption is not mainly extrapolated from the quantity of aluminum ingested. The study had found that a nano-molar amount of aluminum was able to induce gene expression, which promotes inflammation and cell death, similar to observed in AD. The study however, was conducted ex vivo on human neural cells (EFSA, 2008; Lukiw et al., 2005).

Moreover, evidence has begun to develop regarding studies implicating pesticides, polychlorinated biphenyls (PCBs) and particulate air pollution to AD.
1.4.3. **Genetic factors**

Alzheimer’s disease may be classified into early onset AD (EOAD, onset <65 years), and late-onset AD (LOAD, onset ≥65 years) that accounts for more than 95% of cases. Less than 1% of AD cases are those of EOAD, with the majority of cases belonging to the category of LOAD (Blennow et al., 2006).

While clinically indistinguishable from LOAD, EOAD is generally associated with a more rapid rate of progression and a Mendelian pattern of inheritance.

1.4.3.1. **Early onset Alzheimer’s disease**

The Amyloid precursor protein (APP) as the name entails, is involved in the generation of amyloid beta (Aβ). Mutations in presenilin 1 and 2 (PS-1, PS-2) proteins results in an increase in production of full-length Aβ40/42. They have been firmly implicated in the pathology of EOAD through mutations in the gene encoding their proteins. Considered diagnostic biomarkers of EOAD, these mutations exhibit autosomal dominant inheritance and eventually lead with certainty to Aβ aggregation and an early onset of the disease (Reitz & Mayeux, 2014; Waring & Rosenberg, 2008).

1.4.3.2. **Late onset Alzheimer’s disease**

Apolipoprotein E (APOE) is a lipid-binding protein and is expressed in humans as three common isoforms coded for by three alleles, APOEε2, ε3, and ε4. Located on chromosome 19q13, the APOEε4 allele is the only clearly recognized susceptibility gene in Alzheimer’s disease. While inhering a single copy of the APOEε4 allele increases the risk of developing AD by 2-3 fold, and lowering the age of onset by 6-7 years, it does not guarantee it. Having two copies of the APOEε4 allele is associated with an even higher risk of more than 5 fold increase (Corder et al., 1993; Gomez-Isla et al., 1996b; Kuusisto et al., 1994).
1.4.4. Genome wide association studies

More recent genome-wide association studies (GWAS) have found 19 areas in genes that appear to affect the risk of developing AD. These genes include: CASS4, FERMT2, INPP5D, MS4A, PICALM, NME8, CD2AP, CELF1, HLA-DRB5, EPHA1, ZCWPW1, ABCA7, MEF2C, CLU, SORL1, BIN1, SLC24A4, PTK2B and CR1 (Lambert et al., 2013).

1.5. Neuropathological Hallmarks

1.5.1. Amyloid deposits

Consisting of folded $\beta$-sheet structures, amyloid depositions are complex mosaic accumulations of protein. Referred to as amyloid plaques, they may be classified into either neuritic or diffuse plaques depending on size, shape and composition (Small, 1998; Yamaguchi et al., 1988).

Neuritic plaques, as the name suggests, are extracellular depositions of fibrillar amyloid beta enclosed by dystrophic neurites. Dystrophic neurites are accumulations of enlarged axons and dendrites amongst others enlarged lysosomes and numerous mitochondria. Ranging in size between 10 to 120 $\mu$m, in AD patients neuritic plaques contain many other proteins such as Tau, ubiquitin and the amyloid precursor protein (Norfray & Provenzale, 2004; Selkoe, 1989; Su et al., 1998).

Diffuse plaques, on the other hand, are not enclosed by dystrophic neurites and may vary in size from 50 $\mu$m to a couple of hundred $\mu$m. They consists of amorphous and non-fibrillar $A\beta$ depositions and were thought to represent the precursors of neuritic plaque generation (Dickson, 1997).

Cerebral amyloid angiopathy (CAA) is another form of amyloid beta deposition, where $A\beta$ accumulates inside the walls of blood vessels, and is often observed in AD patients (Alafuzoff et al., 2008).
1.5.2. **Neurofibrillary tangles**

Neurofibrillary tangles (NFT) are constituted of filament pairs, roughly 10 nm in width, wound into paired helical filaments (PHF) whose composition consists of hyperphosphorylated tau protein (Grundke-Iqbal et al., 1986; Kidd, 1963; Lee et al., 1991). Tau is primarily expressed in axons of the central nervous system. The hyperphosphorylation of tau leads to its dissociation from microtubules, where its primary function is their promotion and stabilization, thus aggregating into PHFs and destabilizing axons (Alonso et al., 2001; Hirokawa et al., 1996). NFTs are used for postmortem staging of AD, although they are acknowledged to be downstream amyloid pathology, yet they correlate better with clinical stages as compared to neuritic plaques (Braak & Braak, 1991).

1.5.3. **Brain atrophy and neuron loss**

Being a neurodegenerative disease, another characteristic hallmark of AD pathology is brain atrophy. At the macroscopic level, this is exhibited via the enlargement of the ventricles, widening of the sulci and shrinkage of the gyri (Blennow et al., 2006). Marked by reduction in both volume and weight, magnetic resonance imaging (MRI) may be used to detect this alteration that is noticeable predominantly in the entorhinal cortex, amygdala and hippocampus (Bottino et al., 2002). Studies have shown that hippocampal volume and brain volume are correlated with neuron number, thus pointing towards neuron loss as the driving factor behind brain atrophy (Kril et al., 2004).

The underlying cause driving the progression and severity of neuron loss in AD is a highly debated matter. While correlations have been made between neuron loss and NFTs (Gomez-Isla et al., 1997), more recently other groups have proposed that the key role is rather neurotoxicity induced by intraneuronal Aβ (Bayer & Wirths, 2010; Haass & Selkoe, 2007). Regardless of the reason, analysis of AD cases...
using unbiased stereology have shown significant neuron loss in the entorhinal cortex (Gomez-Isla et al., 1996a), while later studies have also shown neuron loss in the superior temporal gyrus, amygdala, substantia nigra, supramarginal gyrus and region I of hippocampus proper (CA1) (Fig. 1.1) (Duyckaerts et al., 2009).

![Figure 1.1. Brain atrophy in Alzheimer's disease.](image)

**Figure 1.1. Brain atrophy in Alzheimer's disease.** Progressive brain atrophy in AD as compared to a preclinical stage. (Image modified from the National Institute of Health, Open Access source).

### 1.6. The Amyloid Precursor Protein

The gene for the amyloid precursor protein (APP), in humans, is located on chromosome 21 (Yoshikai et al., 1990). It encodes an integral membrane protein whose primary function is yet unknown, yet has been proposed to be involved in iron export (Duce et al., 2010), neural plasticity (Turner et al., 2003) and synapse formation (Priller et al., 2006). As a precursor molecule, its subsequent pathological cleavage results in the generation of beta amyloid, a key player in Alzheimer’s disease.

#### 1.6.1. Amyloidogenic and non-amyloidogenic processing

In the normal physiological “non-amyloidogenic” pathway, α-secretases that are membrane-bound and predominantly active at the cell surface (Sisodia, 1992), cleave APP around the middle region of the Aβ sequence (between amino acids 16 and 17, both Leucine). This leads to the release of the soluble sAPPα fragment into the extracellular space (Anderson et al., 1991; Sisodia et al., 1990).
Thereafter, γ-secretase cleaves the remaining C-terminal fragment (C83), thus producing the P3 fragment, which in contrast to Aβ is rapidly degraded, simultaneously producing the APP intracellular domain (AICD) that is thought to be a transcriptional regulator (Haass et al., 1993; Hartmann, 1999).

However, in the pathological “amyloidogenic” pathway, APP is rather cleaved upstream of the α-secretase cleavage site by β-secretase. This consequently releases a soluble N-terminal fragment (sAPPβ) and a longer C-terminal 99-residue fragment (C99), which remains membrane associated. γ-secretase then proceeds to cleave C99 at various sites within the transmembrane domain, generating Aβ peptides ranging from 37 to 43 amino acids in length. This also results in the production of the AICD fragment (Citron et al., 1995; De-Paula et al., 2012) (Fig. 1.2).

It is of worth to note that the main constituent of amyloid plaques is Aβ containing 42 amino acids, while under normal biological conditions mainly Aβ containing 40 amino acids is generated and represents the major portion of the total Aβ (Zhang et al., 2011).
Figure 1.2. APP processing and imbalance in age-related neurodegeneration. The amyloid precursor protein is processed either by an amyloidogenic pathway (left) or a canonical pathway (right). (Left) Beta secretase (BACE) cleaves APP resulting in increased amounts of ABP Carboxy-terminal fragment (CTFβ) and Aβ peptide (light blue) being secreted outside the cell. The increased extracellular accumulation of toxic Aβ species, particularly Aβ42, results in the formation of Aβ oligomers. (Right) Normally, most APP is cleaved by the α-secretase, secreting sAPPα. CTFα is endocytosed and then processed by γ-secretase, resulting in formation of the p3 peptide, which is secreted, and releasing the AICD into the cytosol. BACE processing of APP does occur to generate Aβ (blue), but these are degraded and cleared. While few small plaques may accumulate with aging, they are much smaller and fewer in number than those associated with disease (Figure and legend modified from (Webb & Murphy, 2012), Open Access source).
1.7. Amyloid Beta

As previously mentioned, amyloid beta (Aβ) is generated by the sequential cleavage of beta and gamma secretase resulting in the production of peptides varying in length. Apart from so called “full length” Aβ consisting of Aβ1-40 and Aβ1-42 several truncated versions have been reported such as C-truncated Aβ1-17, Aβ1-38 and Aβ1-39 (Portelius et al., 2012; Wiltfang et al., 2002). New data has even described peptides similar to Aβ, which span amino acids 663-711, before the beta and gamma secretase cleavage sites (Kaneko et al., 2014). Different Aβ isoforms, whether C- or N-truncated, possess a carrying degree of aggregation, stability and structure and neurotoxic activity (Bouter et al., 2013; Jarrett et al., 1993; Pike et al., 1995; Wirths et al., 2010). Of particular interest lately in AD therapy have been N-truncated Aβ peptides.

1.7.1. N-truncated amyloid beta

The enzymatic activities that lead to N-terminal truncations are not fully understood. In some cases the question arises if the truncation has occurred after Aβ1-42 release or if it is an independent event. Meprin-β has recently been proposed to be responsible for generation of Aβ2-x peptides through cleavage of APP (Bien et al., 2012). Neprilysin (NEP) cleaves between Arg-2 and Glu-3 or between Glu-3 and Phe-4, to generate N-truncated Aβ peptides (Howell et al., 1995; Leissring et al., 2003). Furthermore, β-secretase, the enzyme responsible for the generation of Aβ1-x peptides, can also generate Aβ11-x peptides, through between Tyr-10 and Glu-11 (Vassar et al., 1999). An overview of further proteases thought to be involved in N-truncated Aβ generation are seen in Figure 1.3.
**Figure 1.3. Cleavage sites of enzymes potentially involved in generation of N-truncated Aβ peptides.** Sequence of the first 17 amino acids of the N-terminus of human Aβ is shown in three-letter and one-letter-code. Amino acids (AA) with charged polar side-chains are shown in red, AA with uncharged polar side-chains in green and hydrophobic non-polar AA in blue. (Figure adapted from (Bayer & Wirths, 2014), Open Access source)

### 1.7.2. N-truncated Aβ peptides in murine AD models

The relative amount of N-truncated Aβ in human AD is much higher than levels reported in AD mouse model (Rüfenacht et al., 2005). Nonetheless, several N-truncated forms of Aβ have been reported in murine models. The APP/PS1 KI model has measureable levels of Aβ4/5-x and AβpE3-x (Casas et al., 2004). Another study assessed the levels of AβpE3-x in the 5XFAD, Tg2576, APP/PS1ΔE9 and 8 other models (Frost et al., 2013). Aβ5-42 has also recently been reported in both the 3xTg and 5XFAD models (Guzman et al., 2014).

To further elucidate the toxicity of N-truncated Aβ peptides, 2 models expressing only the AβpE3-x (TBA42) (Wittnam et al., 2012) and Aβ4-42 (Tg4-42) (Bouter et al., 2013), led to the intracellular accumulation of the respective peptides, accompanied by neuron loss and behavioral deficits.
1.7.3. **Amyloid hypothesis**

Initial explanations of the pathogenesis of Alzheimer’s disease arose from the amyloid cascade hypothesis that pointed the finger towards Aβ deposition as the causative factor in the etiology of AD. This thereby linked Aβ deposition to the eventual pathology in AD, claiming it lead to neuron and synapse loss, brain atrophy and eventually leading to dementia (Hardy & Higgins, 1992). Plaque formation caused by the accumulation of Aβ_{40} and Aβ_{42}, caused by increased Aβ production or lack of its clearance, triggered the cascade (Pimplikar, 2009).

Although the theory was heavily supported by a stream of evidence, mostly from familial AD cases studies (Bertram et al., 2010), murine AD models based on familial AD mutations (Duyckaerts et al., 2008), down syndrome and triplication of the APP gene (Rumble et al., 1989; Schupf & Sergievsky, 2002), the hypothesis had major flaws. The most important indication that the hypothesis was faulty came from evidence showing that cognitively normal individuals, showing no signs of dementia, had shown strong plaque deposition (Pimplikar, 2009). Furthermore, both in human studies (Snowdon, 2003) and in AD mouse models (Walsh & Selkoe, 2007), plaque load does not correlate consistently with cognitive deficits/dementia.

1.7.4. **Revised amyloid hypothesis**

In contrast to the classical amyloid hypothesis, a revised version dependent on the accumulation of intracellular Aβ peptides has been proposed. It proposes that intracellular Aβ could be secreted and re-up taken from cells. Thus elevating levels of intracellular accumulation, is causative in downstream events such as neuronal and synaptosomal dysfunction leading to AD (Wirths et al., 2004). This is supported by several observations, such as, in Down Syndrome intracellular Aβ is detectable at a young age and accumulates less with age as plaques begin to develop (Gyure et al.,
Furthermore, classical AD mouse models, most of which carry a familial AD mutation and harbour strong plaque pathology, reflect human AD inflammatory pathology, but lack neuron loss (Wirths et al., 2004). To this end, recent mouse models such as the 5XFAD (Oakley et al., 2006), the APP/PS1KI (Casas et al., 2004) and the Tg4-42 mouse model (Bouter et al., 2013) exhibit an early accumulation of intracellular Aβ and a correlation of behavioral deficits with neuron loss. This implies that Aβ oligomers, soluble intracellularly and extracellularly, play a pivotal role in the development and progression of early AD.

1.8. Treatment

Current drugs designed to alleviate the clinical symptoms of AD mostly rely on neurotransmitter regulation. Since reduction in the activity of cholinergic neurons in AD has been extensively studied, 4 drugs are acetylcholinesterase (Ach) inhibitors employed to slow down the rate of degradation of Ach (Birks & Harvey, 2006; Bullock et al., 2005). These are galantamine, rivastigmine, donepezil and tacrine, which are marketed under different commercial names. The 5th drug is an N-methyl-D-aspartate receptor (NMDA) antagonist that blocks the glutamatergic system overstimulation by glutamate (Lipton, 2006). The benefit from their use is very limited and none of the drugs delays the progression of the disease or halts the clinical symptoms (Birks & Harvey, 2006). Observational studies suggest that combination treatment may increase the time before patients require nursing home care (Lopez et al., 2009).
1.8.1. Targeting Aβ

Targeting Aβ has recently been the main target in therapeutic approaches. Drugs have been aimed at either decrease the production of Aβ or enhance its clearance. The primary research focus has been on enzymes involved in Aβ production and their inhibition. β- or γ-secretase inhibition has been attempted with limited success and side effects, as it has been difficult to develop drugs that penetrate the BBB, and precisely inhibit β- or γ-secretase. Unwanted side effects have been the cleavage of alternative substrates such as sodium channel subunits and Notch (Citron, 2010; De Strooper et al., 2010). Research focus has been on immunization approaches to reduce Aβ levels and enhance clearance (Lannfelt et al., 2014).

1.8.2. Active immunization

Initial pre-clinical trials in active immunization were a breakthrough, using PDAPP transgenic mice, immunization with preparations of full length Aβ_{42} resulted in the clearance of amyloid plaques (Schenk et al., 1999). Other studies using different mouse models were similar in results and even lead to the amelioration of behavioral symptoms in the mice (Dodart et al., 2002; Janus et al., 2000). Given the previous results, clinical trials in humans were approved hurriedly. Using Aβ preparations in AD patients initially reported no adverse effects (Schenk, 2002). However, the clinical phase 2A trials were subsequently halted due to the development of symptoms of central nervous system inflammation, in some patients, and some deaths due to pulmonary embolism (Ferrer et al., 2004; Gilman et al., 2005; Nicoll et al., 2003). The field afterwards turned to passive immunization as a means to overcome the negative side effects caused by active immunization.
1.8.3. Passive immunization

As with any form of therapy that is taken from the bench to bedside application, passive immunization offers both advantages and disadvantages. The major disadvantage of passive immunization lies in the fact that repeated infusions of antibody over the lifetime of a patient would be necessary to maintain a desired therapeutic effect. The therapy however, offers a precise delivery of target specific antibodies to patients and the opportunity to rapidly decrease antibody titer if side effect should develop (Lannfelt et al., 2014).

The first evidence of success in passive immunotherapy in an AD mouse model was demonstrated by Bard and colleagues in 2000. Wherein the systemic administration of the N-terminus 3D6 monoclonal antibody eventually resulted in a decreased plaque burden (Bard et al., 2000). The humanized version of 3D6, Bapineuzumab, proceeded in phase I, II and III clinical trials but terminated after phase III trials did not achieve significant clinical efficiency (Lannfelt et al., 2014). A recent version of the antibody, AAB-003, engineered to reduce risk of vasogenic edema and micro-haemorrhage, is now in phase I study.

Other humanized antibodies currently still in various stages of clinical trials are Crenezumab (Genentech), BAN2401 (Biogen), LY3002813 (Eli Lilly & Co), SAR228810 (Sanofi) and Solanezumab (Eli Lilly & Co) (Clinicaltrials.gov).

Although passive immunization trials have encountered undesirable side effects, and their long-term cost to the healthcare system is highly debated, the therapy remains a promising option.
1.9. Crystallization

Crystallization is a feature of precipitation, wherein the process of formation of solid crystals is driven through precipitation from a solution or gas. The two major events of crystallization are nucleation and crystal growth. The clustering of solute molecules, dispersed in the solvent, is a nanometer scale event referred to as nucleation. Conditions such as temperature, supersaturation, pH and more, will determine whether a cluster the critical size in order to become a stable nucleus (Fig. 1.4). Supersaturation is the driving force of crystal growth, with either nucleation or growth being predominant thus affecting the different sizes and shapes of crystals obtained. Crystallization is complete once supersaturation is drained and the solid-liquid system reaches equilibrium.

Figure 1.4. Adjustable parameters for crystallization include precipitant or additive concentration, pH and temperature. The four major crystallization methods are represented: (i) microbatch, (ii) vapor diffusion, (iii) dialysis and (iv) FID. Each involves a different route to reach the nucleation and metastable zones, assuming the adjustable parameter is precipitant concentration. The filled black circles represent the starting conditions. Two alternative starting points are shown for FID and dialysis because the undersaturated protein solution can contain either protein alone or protein mixed with a low concentration of the precipitating agents. The solubility is defined as the concentration of protein in the solute that is in equilibrium with crystals. The supersolubility curve is defined as the line separating conditions under which spontaneous nucleation (or phase
separation or precipitation) occurs from those under which the crystallization solution remains clear if left undisturbed. Figure and legend taken from (Chayen & Saridakis, 2008), used with permission from Nature Publishing Group, license number 3770641097789).

1.9.1. **X-ray crystallography**

Is an experimental technique that is used to determine the three dimensional structure of a substance (e.g. protein, RNA, DNA, small molecules). It exploits the fact that periodically arranged atoms in the substance crystal diffract X-rays. Based on the resulting diffraction patterns, the electron density can be reconstructed and the structural model describing position of every atom of the substance can be built. Macromolecular crystallography is a technique used to study biological molecules such as nucleic acids and proteins to a resolution higher than 5 Å. It allows for the elucidation of protein structure and function and may also demonstrate interaction between 2 or more proteins. The crystallographic analysis of antigen-antibody complexes has been demonstrated (Edmundson et al., 1996).

1.9.2. **Crystallization of anti-amyloid beta antibodies**

Alzheimer’s disease researchers have come to believe that the success of Aβ antibodies, currently in clinical development, in terms of eventual success in therapeutic trials, may be correlated with the precise spatial details of how the antibodies engage their antigen. A growing collection of crystal structures are building a knowledge base, in hopes of improving therapeutic outcome and minimize collateral activity such as vasogenic edema. Each of the clinical anti-Aβ antibodies targets specific parts of the notorious peptide, though some of the binding profiles overlap. Some of the published crystal structures include that of Bapineuzumab (Crespi et al., 2014; Feinberg et al., 2014b; Miles et al., 2013), Solanezumab (Crespi et al., 2015), Ponezumab (La Porte et al., 2012) and Gantamerumab (Bohrmann et al., 2012).
1.10. Objectives

Targeting Aβ has recently been the main objective in Alzheimer’s disease therapeutic approaches. Passive immunization trials have encountered undesirable side effects but the therapy remains a promising option. Amyloid beta 4-x has not been previously considered as a target for AD immunotherapy.

The current work's objectives are the following:

- The characterization of a novel antibody (NT4X-167) which recognizes N-truncated amyloid beta
- The large scale purification of the NT4X antibody and its respective Fab fragment
- The characterization of NT4X binding specificity and sensitivity
- The investigation of NT4X in vitro and in vivo rescue of amyloid beta
- The validation of Tg4-42 as a valid AD model for therapeutic intervention
- The passive immunization of Tg4-42 with NT4X and determination of the mechanism of target engagement
CHAPTER 2

2. Materials and Methods

2.1. Animals

2.1.1. General considerations

Murine lines were maintained in the central animal facility of the University Medicine Göttingen under standard housing conditions. For behavioral analysis, mice were kept under constant 12 h/12 h dark/light conditions and supplied with food and water ad libitum.

Animal experiments were approved by the local animal protection authorities (Niedersächsisches landesamt für Verbraucherschutz und Lebensmittelsicherheit) under the approval number 14/1450. All experiments were conducted in accordance with the approved protocols.

2.1.2. Transgenic mice

Tg4-42 mice express human Aβ4-42 fused to the murine thyrotropin releasing hormone (TRH) signal peptide under the control of the neuronal Thy-1 promoter on a C57Bl/6J genetic background (Bouter et al., 2013).

For this study only homozygous Tg4-42 mice were used (Tg4-42_hom).

2.2. Animal Experiments

2.2.1. Passive immunization

The potential therapeutic effects of NT4X-167 were studied using a passive immunization approach in Tg4-42_hom. Passive
immunization was performed by intraperitoneal injections of the Aβ-specific monoclonal antibody NT4X (against the N-terminus of Aβ_{4-42}) or its Fab fragment and compared to control groups using antibody of the same immunoglobulin class as NT4X (IgG2b).

Tg4-42_{hom} mice were immunized by injections of the NT4X antibody, 10mg/kg body weight, diluted in sterile Phosphate buffered saline (PBS) (pH 7.4). Mice received weekly injections beginning at 3 months of age. Each mouse received a total of 12 injections. Behavior testing started between the 10th and the 11th injection. Animals were sacrificed after the last injection.

Control groups comprised of Tg4-42_{hom} mice received intraperitoneal IgG2b antibody (10mg/kg body weight) diluted in PBS. Control mice were treated exactly the same as the NT4X group receiving 12 injection of the respective control solution starting at 3 months of age. Furthermore, a non-treated Tg4-42_{hom} group served as an additional control.

2.2.2. **Morris water maze**

Spatial reference memory in Tg4-42_{hom} mice was evaluated using a modified version of the Morris water maze (Morris, 1984).

Briefly, mice learn to use visual spatial cues to locate a hidden platform in a circular pool filled with opaque water. The pool was divided into four virtual quadrants that were defined based on their spatial relationship to the platform: left, right, opposite and target quadrant, which contains the submerged goal platform.

ANY-Maze video tracking software (Stoelting Co., Wood Dale, USA) was used to record escape latency, swimming speed and quadrant preference.

The experiment began with three days of cued training during which the submerged platform was made visible with a triangular flag that extended upwards. Both the location of the platform and the
position where mice were introduced into the pool changed between trials. Each mouse received four training trials per day.

Forty-eight hours after the last day of cued training, mice performed five days of acquisition training. For this part of testing, the flag was removed from the submerged platform. Both distal cues, existing in the room, and proximal visual cues attached to the outside of the pool, were used as learning cues. The platform location remained stationary for each mouse throughout training. Each mouse again received four training trials per day.

Twenty-four hours after the last acquisition trial, a probe test was performed to assess long-term spatial reference memory. The platform was removed from the pool, and mice were introduced into the water from a novel entry point. Mice were then allowed to swim freely for one minute while their swimming path was recorded.

Mice were sacrificed 24-48 hours after the probe trial.

2.2.3. **Intracerebroventricular injection of soluble Aβ**

12-week-old male C57BL/6J mice (Janvier, Le Genest-St-Isle, France) were intracerebroventricularily (icv) injected under anesthetization. Freshly prepared Aβ42 peptides (50 pmol in 1 µL; 0.1M PBS (pH 7.4)) or freshly prepared 50 pmol Aβ42 in combination with the NT4X antibody [1 or 10 pmol respectively in 1 µL; 0.1M phosphate-buffered saline (pH 7.4)] was injected into the right ventricle, with stereotaxic coordinates from the bregma (AP –0.22, L – 1.0 and D 2.5 in mm). Vehicle (0.1M phosphate-buffered saline) or vehicle in combination with 1 or 10 pmol NT4X respectively were injected into the right ventricle as a control. ICV injections were made using a 10-µl Hamilton microsyringe fitted with a 26-gauge needle. Four days following ICV infusion of Aβ peptides, working memory was assessed using the Y-maze test (Experiments performed by SynAging, Nancy, France).
2.2.4. **Y-Maze task**

Immediate spatial working memory performance in icv injected 12-week-old male C57BL/6J wildtype mice was assessed by recording spontaneous alternation behavior in a Y-maze as described previously (Miller et al., 2008; Youssef et al., 2008). The Y-maze task was carried out on day four after Aβ_{4-42} and NT4X application. The maze was made of opaque plexiglas and each arm was 40 cm long, 16 cm high, 9 cm wide and positioned at equal angles. Mice were placed at the end of one arm and allowed to move freely through the maze during a 5 min session. The series of arm entries were recorded visually and arm entry was considered to be completed when the hind paws of the mouse were completely placed in the arm. Alternation was defined as successive entries into the three arms on overlapping triplet sets. The percentage alternation was calculated as the ratio of actual (total alternations) to possible alternations (defined as the number of arm entries minus two), multiplied by 100 (Experiments performed by SynAging, Nancy, France).

2.3. **Electrophoresis of Peptides**

2.3.1. **Abeta peptide preparation**

Stock solutions of synthetic peptides for blotting (1mg/ml in 10 mM NaOH; PSL, Heidelberg) were prepared, sonicated for 5 min in water bath (Sonorex RK 100H, Bandelin electronic), snap frozen in liquid nitrogen and stored at −80°C.

2.3.2. **Western blot of synthetic peptides**

For Western blot analysis under reducing conditions 7 μg peptide per well (mixed with an equal volume of sample buffer) were loaded on 4-12% Tris-Tricin VarioGels (Anamed) and run at a constant current of 120 V. After electrophoresis, the peptides were
transferred onto 0.45 μm nitrocellulose membranes for 30 minutes per membrane at constant 25 mA in a semi-dry transfer chamber. Free binding sites were blocked with 10 % (w/v) non-fat dry milk dissolved in Tris-buffered saline supplemented with Tween-20 (TBS-T) (50 mM Tris(hydroxymethyl)- aminomethane (Tris) pH 8.0 supplemented with 0.05 % (v/v) Tween-20) for one hour at room temperature (RT).

For detection, the primary antibodies IC16 (1 μg/ml), 1–57 (1 μg/ml) and NT4X (1 μg/ml) were dissolved in TBS-T and incubated on a shaker over night at 4 °C. After three, 5 min, washing steps with TBS-T, the secondary rabbit-anti-mouse horseradish peroxidase (HRP)-conjugated antibody was diluted 10000-fold in TBS-T and incubated with the membrane for 2 hours at RT. Blots were developed using Luminata Crescendo Western HRP Substrate (Millipore) and exposed with the ODYSSEY Fc (LI-COR).

2.4. NT4X-167 Antibody

The novel oligomeric Aβ specific antibody NT4X (IgG2b; official name of cell line Aβ4-40 NT4X-167; DSM ACC3162) was generated by immunizing three Balb/c mice with unconjugated Aβ4-40. After preparation of the lymph nodes they were fused with the myeloma cell line P3-X63-Ag8 for generation of the hybridoma cells. The hybridoma supernatants of mixed clones were screened by Ezyme-linked Immunosorbent Assay (ELISA) and immunohistochemistry and subcloned. The idea behind the generation of novel oligomeric antibodies was that in solution Aβ4-40 peptides are forming stable aggregates that can be used as an epitope for antibodies that specifically bind at the N-terminus of Aβ4-40. Therefore Aβ4-40 was used for immunizing mice and positive clones were screened in four steps. (Antonios et al., 2013).
2.4.1. NT4X hybridoma cell culture

The NT4X hybridoma cell line was cultured in a serum-free media for hybridoma culture ISF-1 (Biochrom) and maintained in fed batch/continuous perfusion mode at 25% Oxygen, 5% CO2 and 37°C in a 3L glass vessel bioreactor (Applikon) controlled using ADI 1030 (Applikon). The cell-culture supernatants were harvested by centrifugation at 500 g and the antibody containing media was collected.

2.4.2. NT4X purification

Media was further centrifuged at 10,000 g for 30 min prior to Protein A affinity chromatography and pH was stabilized by addition of 20% (v/v) of PBS buffer. Media was loaded at 5 ml/min onto HiTrap™ Protein A HP (GE Healthcare) connected to an AKTAxpress (GE Healthcare). Further column washing and elution was done according to Protein A column manufacturer instructions. The eluted sample was further purified by size-exclusion chromatography using a HiLoad 26/60 Superdex 200 pg run in PBS on an AKTAxpress (GE Healthcare).

2.4.3. NT4X Fab generation and purification

Fab fragments were produced utilizing papain digestion of the intact NT4X monoclonal antibody. Papain agarose (Sigma) was pre-activated, shaking at 1400 rpm, with 10 mM cysteine (Sigma) in PBS for 30 min at room temperature. Previously purified NT4X antibody (in PBS) was incubated for 4 hours at 37°C with pre-activated papain agarose (1 U agarose/5 mg antibody). Papain agarose was subsequently removed by filtration through a 0.45 µm filter. Filtrate was then dialyzed into Buffer A (20 mM Sodium Citrate, 20 mM NaCl, pH= 5.5) and afterwards loaded onto a MonoS 10/100 GL (GE Healthcare). Column was eluted using 30 column volumes, reaching a
10% gradient of Buffer B (20 mM Sodium Citrate, 1 M NaCl, pH= 5.5). Column eluate was immediately neutralized using Buffer C (1 M Tris, pH= 9.0).

Fab Eluate was further purified and buffer exchanged by size-exclusion chromatography using a HiLoad 26/60 Superdex 75 pg run in PBS or Tris 50 mM buffer on an AKTAxpress (GE Healthcare).

2.4.4. Antibody storage

Both full length antibody and Fab fragment were concentrated to 2 mg/ml with a 10 kDa molecular weight cut-off (MWCO) centrifugal concentrator (Amicon Ultra, Millipore), snap frozen in liquid nitrogen and stored at -80°C until further use.

2.5. In vitro neuroprotection assays

2.5.1. Full-length NT4X antibody neuroprotection assay

Neurons were prepared from the cortex of 18 day old embryonic CD rat brains using a papain digestion kit (Worthington, Biochemical Corporation). Cells were plated at 3 × 10^5 cells/ml in 96 well Cell Coat, Poly-D-Lysine coated plates (Greiner) in Neurobasal media with Pen/Strep, L-glutamine and B27 supplement with antioxidants (Gibco, ThermoFisher). The plates were incubated in a humidified incubator at 37°C, 5% CO2 for two days prior to adding the antibodies and amyloid peptides. Media was removed from the plates and fresh Neurobasal media containing B27 supplement without antioxidants (Invitrogen) added to perform the assay. Purified antibody stocks (mouse NT4X IgG2b and an isotype control antibody) were diluted in sterile PBS (Ca/Mg free) and 10 µl added to a total volume of 100 µl per well to give a series of dilutions at the required concentration. HFIP treated and dried amyloid peptide aliquots (stored in a desiccator at -80°C) were dissolved in 100 mM NaOH and further diluted in Neurobasal media without antioxidants to achieve final peptide concentration.
concentrations of 5 µM for both Aβ_{pE3-42} and Aβ_{1-42} and 10 µM for Aβ_{4-42} peptide in the cell assay (Anaspec, Fremont, CA (Eurogentec) and California Peptide Research, San Francisco). The plates were incubated for seven days prior to LDH measurement with the CytoTox96 Non-Radioactive cytotoxicity assay kit (Promega). The LDH assay was performed on 50 µl/well culture supernatant in fresh 96 well assay plates (Costar). The plates were read at 490 nm on a Tecan Safire II plate reader. The inhibition assay was repeated n = 2 to 3 for each peptide with the two antibodies detailed. (Experiment performed by MRC Technology, London, United Kingdom)

### 2.5.2. NT4X Fab fragment neuroprotection assay

Cortical neurons from embryonic day 16–17 Wistar rat fetuses were prepared as previously described (Pillot et al., 1999). In brief, dissociated cortical cells were plated at 50,000 cells/well in 48-well plates precoated with 1.5 mg/mL polyornithine (Sigma). Cells were cultured in a chemically defined Dulbecco’s Modified Eagle’s/F12 medium free of serum (Gibco) and supplemented with hormones, proteins and salts. Cultures were kept at 35°C in a humidified 5% CO2 atmosphere, and at 6–7 DIV, cortical population was determined to be at least 97% neurons by immunostaining as previously described (Youssef et al., 2008). At 6 DIV, the medium was removed and cortical neurons were incubated for 24 h with vehicle (cell culture medium) or Aβ peptides (dissolved in cell culture medium) at the indicated concentrations. Following a 24 h incubation of primary cortical neurons with Aβ peptides, cell viability was determined using a calcein-AM assay (Invitrogen, Molecular Probes).

Briefly, cells were washed twice with PBS and incubated protected from light for 30 min at room temperature in the presence of 2 µM calcein-AM solution prepared in PBS. Cells were then washed twice with PBS and incubated for 15 min at room temperature in PBS containing 1% Triton X-100 (v/v). The level of calcein fluorescence was
monitored by fluorescence emission at 530 nm after exciting at 485 nm, using a Fluostar microplate reader (BMG-Lab technologies, France). (Experiment performed by SynAging, Nancy, France)

2.6. Tissue Experiments

2.6.1. Collection and preservation

For all intended purposes, mice tissue was collected and preserved after transcardial perfusion of anaesthetised mice.

First mouse is anaesthetised using 10 mL/kg intraperitoneal injection of 1% Ketamin (Medistar, Ascheberg, Germany) and 0.1% Xylazin (Ecuphar, Belgium) in injectable grade water. Following deep anaesthesia, transcardial perfusion is carried out using ice-cold PBS buffer, followed by 4% paraformaldehyde (PFA) in PBS. Brain hemispheres were carefully divided at the midline. Left hemispheres were post fixed in 4% PFA overnight followed by 30% sucrose in PBS overnight. Following day, the tissue is frozen on dry ice and stored at -80°C until further processing.

Right brain hemispheres were post fixed in 4% buffered formalin (Roth, Karlsruhe, Germany) at 4°C before the tissue was embedded in paraffin by dehydration in a series of ethanol of 50%, 60%, 70%, 80%, 90%, and 2x 100% ethanol followed by Xylol treatment. Finally, tissue was immersed in liquid paraffin for 2 hours before being embedded in solid paraffin blocks.

2.6.2. Neuronal quantification

2.6.2.1. Cresyl violet staining

For neuronal stereology, frozen left hemispheres brains were cut in 10 series of 30 μm coronal sections with collection of every 10th section throughout the brain and stored frozen at -80°C until further processing.
One series comprising every 10\textsuperscript{th} section from each animal was carefully mounted in PBS onto super frost glass slides and left to dry overnight at room temperature.

Sections were washed 2 x 10 min in 0.04 M Sodium acetate, 0.1\% acetic acid solution (Solution 1), delipidated 20 min in 0.025\% Triton X-100; 75\% ethanol, washed 2 x 10 min in solution 1 and stained for 2 x 8 min in 0.01\% cresyl violet dissolved in solution 1.

The sections were then washed 3 x 1 min in solution 1, 3 min in 100\% ethanol, 10 min in isopropanol, and 2 x 5 min in xylol before being embedded in Roti-Histokitt (Roth) mounting medium.

2.6.2.2. Unbiased stereology

Stereological analysis was performed as previously described (Bouter et al., 2013). Briefly, mice were anaesthetized and transcardially perfused with 4\% paraformaldehyde. Brains were carefully removed from the skull, dissected and post-fixed. The left brain hemispheres were cryoprotected in 30\% sucrose, quickly frozen and cut frontally into entire series of 30 \mu m thick sections on a cryostat (Microm HM550, Germany). Every tenth section was systematically sampled, stained with cresyl violet and used for stereological analysis of the neuron number in the CA1. The hippocampal cell layer CA1 (Bregma -1.22 to -3.52 mm) was delineated on cresyl violet stained sections. Using a stereology workstation (Olympus BX51 with a motorized specimen stage for automatic sampling), StereoInvestigator 7 (MicroBrightField, Williston, USA) and a 100x oil lens (NA = 1.35), neuronal nuclei were sampled systematically using optical dissector probes, and the total number of neurons was subsequently estimated by the fractionator method using a 2 \mu m top guard zone.

Age and sex-matched mice were analyzed for all groups. Samples were blinded to avoid biased counting.
2.7. Crystallization

2.7.1. General crystallography setup

Crystallization of the Fab portion of the NT4X antibody or the Fab bound to respective Aβ peptides was carried out at 20 °C using the sitting drop vapor diffusion method. Different commercial crystallization reagents were screened to identify initial conditions. Initial crystals were obtained by mixing 60 nl protein solution with 60 nl reservoir solution in 96-well MRC plates using a Cartesian liquid dispensing robot with 8 channels. For grid screening with seeding, crystals were obtained by mixing 100 nl protein solution, 100 nl reservoir solution, and 20 nl seed stocks/dilutions in 96-well INTELLI-PLATEs using the Gryphon liquid handling system (Art Robbins Instruments). For growing larger crystals, manual drops of 1 µl protein, 1 µl reservoir solution and 200 nl seed stock/dilution were set up in 24-well plates. Crystal growth was monitored by imaging with RockImager (Formulatrix, Bedford, United States).

2.7.2. Crystallization and diffraction data collection

Fab-Aβ complex preparation was done as follows. Peptide corresponding to residues 4-19 of the human amyloid beta sequence (FRHDSGYEVHHQKLVF) was purchased from Peptide Specialty Laboratories (Germany) at 95% purity. Peptide was re-suspended in 50 mM Tris, 150 mM NaCl (pH= 7.5) and added to Fab in a Fab:Aβ ratio of 1:5 and left on ice for 4 hours to allow for antibody antigen binding. Final Fab concentration used for successful crystallization trials was 5 mg/ml.

Crystal seed stocks were prepared with Seed Bead™ kit and done according to manufacturer’s instructions (Hampton Research, United States). Seeding was attempted by the use of horse’s tail hair but was unsuccessful. Direct seeding resulted in diffraction quality crystals,
where seed stocks or dilutions were directly added to protein + reservoir combination.

X-ray diffraction data were collected at beamline PXII of SLS (Paul Scherrer Institute, Villigen, Switzerland) and beamline 14.2 of BESSY II (HZB, Berlin, Germany). An example of data collection parameters (example) is summarized in Table 2.1 below.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
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</thead>
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<td>Exposure time (s)</td>
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<tr>
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</tr>
<tr>
<td>Number of frames</td>
<td>720</td>
</tr>
</tbody>
</table>
CHAPTER 3

3. Results

3.1. Full-length NT4X Purification

Initial NT4X antibody purification was attempted using Ammonium Sulfate precipitation (50-70% w/v). However, due to the use of serum-free culture media, antibody-containing hybridoma supernatant could be loaded onto a Protein A column (GE Healthcare) after pelleting cell debris through centrifugation. Elution with low pH solution resulted in antibody disengagement from column. Antibody buffer was subsequently exchanged into appropriate storage/usage buffer (Fig 3.1).
Figure 3.1. **NT4X antibody purification from hybridoma cell line DSM ACC3162 using immobilized protein A.** Left peak, antibody purification from culture media using protein A. Right peak, antibody buffer exchange using HiPrep 26/10 desalting column (GE Healthcare).

NT4X antibody purification was tracked at various stages of purification through sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) gels stained either by coomassie or silver staining. Hybridoma media contained antibody and various other proteins that were required for hybridoma cell line maintenance. Following Protein A column, only full antibody and some antibody fragments were collected in eluate. Column flow-through (FW) still contained antibody and was re-loaded on protein A column (2-3 times) until no more antibody was detectable in FW (Fig 3.2). An antibody yield of approximately 50-60 mg/L of hybridoma media was achievable.
To further purify the NT4X from smaller antibody fragments that eluted along with full-length antibody, the resulting eluate from the Protein A column was concentrated and injected into a HiLoad 26/60 Superdex 200 pg size-exclusion column in PBS. Fractions from different peaks were analyzed. Peak 1 contained full-length antibody (Fig 3.3).
Figure 3.3. Coomassie stained SDS-PAGE gel (non-reducing) tracking NT4X antibody purification following size-exclusion chromatography. Using HiLoad 16/600 Superdex 200 pg. Fractions from Peak 1 are represented on gel demonstrating a high purity antibody preparation.

In order to validate antibody presence in hybridoma media and throughout purification steps; a reducing PAGE-SDS gel was run and either stained by coomassie (Fig 3.4A) or transferred to a nitrocellulose membrane and probed using a mouse secondary antibody (Fig 3.4B). Coomassie gel revealed the presence of heavy and light chains (50 kDa and 25 kDa respectively) and probed membrane identified that antibody chains are murine as is expected.
Figure 3.4. Validation of antibody presence during purification process

(A) Coomassie stained SDS-PAGE gel (reducing), verifying presence of both heavy-chain and light chain fragments of antibody and (B) Western blot of similar gel using a mouse secondary HRP conjugated antibody, revealed that antibody being purified is murine.
3.2. NT4X Fab Generation and Purification

Antibody was incubated with papain for the duration of the reaction. Papain was initially not pre-activated. Reaction at room temperature (a) resulted in partial digestion of antibody while reaction at 37° C resulted in the creation of the fab fragment (at 50 kDa) within the time frame. Smaller digestion products at 15 kDa and 25 kDa were also noticeable. The constant region of the antibody (Fc) was subsequently removed by binding digest on a Protein A column where the Fc fragment would bind but the Fab would pass in the FW (Fig. 3.5).

Figure 3.5. NT4X antibody digestion using papain (a) antibody incubated at room temperature for duration of digestion (b) antibody incubated in a 37°C water bath for duration of digestion. Time scale: 0-4 hours.
Numerous purification strategies were tried for the purpose of further purifying the Fab fragment from smaller unwanted digestion products. Size exclusion chromatography was not successful in separating components. Ion exchange chromatography using a strong cation exchanger (Mono S prepacked with MonoBeads) was able to bind products resulting from papain digestion and successfully separated one component (peak 4) using a very shallow gradient. Peaks 1, 2, and 3 overlapped in elution profiles and were not fully separated, this was due to the fact that they eluted at 3.3 mS/cm, 3.45 mS/cm and 3.6 mS/cm respectively (Fig 3.6).

![Figure 3.6. Mono S purification chromatogram showing 2 main eluted components, where peaks (1, 2 and 3) overlapped and peak 4 was separate.](image-url)
Fractions from Mono S purification were analysed using SDS-PAGE and visualized using coomassie. Peak 1 contained two proteins (50 kDa), peak 2 early fractions contained same as peak 1 while fractions that eluted later contained 3 proteins (50 kDa, 25 kDa and 15 kDa), peak 3 contained same proteins as late fractions from peak 2 and finally peak 4 contained a double band of protein (15 kDa) (Fig 3.7). As was apparent from the Mono S purification chromatogram, peaks 1, 2, and 3 were not properly separated while peak 4 was successfully separated from the rest.

![Figure 3.7. Mono S purification fractions SDS-PAGE](image)

**Figure 3.7. Mono S purification fractions SDS-PAGE.** Numbers represent respective peaks in Mono S purification chromatogram.

A modified digestion protocol using pre-activated papain was eventually used and proved successful in reducing unwanted digestion products. Papain was incubated with cysteine for 30 minutes in order to activate the enzyme. Buffer was then exchanged to one not containing cysteine for the duration of digestion. Digestion products were loaded unto protein A column several times to bind undigested full antibody and Fc fragments. After 3rd pass on protein A column, Fab fragment (50 kDa, flow-through 3 and 4) was obtained in higher purity and buffer was exchanged depending on later usage into
either PBS or Tris buffer. Eluate from protein A column contained undigested (150 kDa), partially digested (100 kDa), and F_c fragment of antibody (Fig 3.8).

**Figure 3.8. Modified antibody digestion protocol.** Using pre-activated papain and subsequent purification with protein A column. Left to right. Papain digestion, FWs from successive loading on protein A column (FW 3 and 4 contain pure Fab) and eluate from protein A column showing undigested antibody and F_c fragment.
3.3. Crystallization

Intended workflow for the purification and crystallization the NT4X Fab fragment and the Fab bound to a respective amyloid beta peptide may be found below. As the previous parts have already described the purification part, this part will deal with the crystallization portion of the work.

![Flowchart describing intended process of crystallization](image)

**Figure 3.9. Flowchart describing intended process of crystallization.** The large scale purification has been described in previous section. Crystallization screening and optimization is described in following section.
3.3.1. Initial screening

Screening for crystallization conditions for the Fab fragment of the NT4X antibody was performed to expose the protein to a variety of agents in order to find “hits” that point to conditions that may result in crystallization. The most common way to adjust crystallization conditions is by varying the concentration of protein, type and concentration of precipitant, and pH or temperature. In the initial screening, conditions were tested out in various concentrations of the Fab in order to first ascertain the appropriate concentration while simultaneously screening precipitants and buffers.

The primary objective of the screening process was to find conditions in which the Fab either formed crystals, crystalline precipitate or phase separation from the reservoir buffer used. Screening was done in 96 well format plates using the NeXtal tube suits (Qiagen) (Fig 10).

Initial screening included the following suites:

- AmSO4
- Anions
- Cations
- Classics
- Classics II
- Classics Lite
- MbClass
- MbClass II
- MPD
- PEGs
- PEGs II
- pHClear
- pHClear II

Figure 3.10. Initial screening for crystallization conditions using NeXtal tube suits.
The vapor diffusion method was used where the protein solution was set up in a sitting drop that equilibrated against crystallizing agents at either higher or lower concentrations than in the drop. This provided a way to sample the crystallization parameter space, as the conditions vary continuously as equilibration proceeds. Screening was performed using 3 concentrations: 5, 13 and 20 mg/ml. Heavy precipitation was noticeable in both the 13 and 20 mg/ml (Fig 3.11b) screens while light precipitation was noticeable in the 5 mg/ml (Fig 3.11a) screens. Moreover, screens which contained polyethylene glycol (PEG), of various molecular weights and concentrations, at 5 mg/ml were the ones that contained initial hits.
Figure 3.11. Different concentrations used in screening for crystallization conditions of the NT4X Fab fragment. Representative plates from Qiagen PEG screen. Images taken at time zero immediately after setting plates. (a) some protein precipitation (b) immediate heavy protein precipitation.
Crystals were first obtained in various conditions in which PEG was the main precipitant. Crystal growth was monitored and observed 0-10 days after drops were set. Depending on the crystallization condition in the respective drops, crystals took between 1-3 days to appear. Crystals for the Fab grew in 25% PEG 3000 and 4000, 0.1 M Tris HCl reservoir solutions (Fig 3.12).

![Figure 3.12. Representative images from initial crystals of Fab fragment obtained through screening (a) Crystals may be seen below the surface of the drop (b) several superimposed crystals, amorphous in shape.](image)

To verify that crystals obtained were not salt crystals, best crystals obtained from screening were fished from corresponding drops and soaked/washed in reservoir buffer before being loaded on an SDS-PAGE along with loading buffer (Fig 3.13). Lane (1) demonstrates protein band of crystal visualized using silver staining as compared to lane (2) which contained protein sample used to set up crystallization screens, both of which ran at around 50 kDa. This demonstrated that crystals grown were indeed formed by the Fab fragment of the NT4X antibody.
Figure 3.13. Verification that crystals grown were consistent with Fab fragment where crystal from (b) was loaded on gel (a) and visualized using silver staining.

3.3.2. Fab + Aβ4-19

In order to better study the structural basis for Aβ recognition of the NT4X antibody, the NT4X Fab was co-crystallized together with an N-truncated Aβ peptide. The Aβ4-19 peptide was used as the antibody has the highest affinity to the N-terminal part of Aβ starting at position 4. Initial crystals of the Fab bound to the Aβ4-19 peptide grew in similar conditions all containing PEG as precipitant. Initial hits were in following reservoir solutions (Fab+Aβ4-19: 5.5 mg/mL):

- 25% PEG 2000 MME, 0.1 M NaOAc trihydrate pH 4.6 (Fig 3.14a)
- 25% PEG 3000, 0.1 M NaOAc trihydrate pH 4.6 (Fig 3.14b)
- 25% PEG 3350, 0.1 M NaOAc trihydrate pH 4.6
- 25% PEG 6000, 0.1 M NaOAc trihydrate pH 4.6 (Fig 3.14c)
- 25% PEG 8000, 0.1 M NaOAc trihydrate pH 4.6
Figure 3.14. Crystals from initial hits of Fab+Aβ_{4-19} complex. Crystals grown in reservoir solution containing 0.1 M NaOAc trihydrate pH 4.6 buffer and (a) 25% PEG 2000 MME (b) 25% PEG 3000 (c) 25% PEG 6000.

3.3.2.1. Refinement

Refinement grids were consequently designed around the conditions where initial hits were observed. Grids varied the concentration of PEG using used (between 20-30% in small 1% increments) and the pH of the NaOAc trihydrate buffer used (between pH 4 and 5, in 0.1 pH increments). Best crystals obtained were in 25% PEG 2000 MME and 24% PEG 8000, both in 0.1 M NaOAc trihydrate pH 4.6. Replicates of the conditions were then setup in both 96 and 24 well formats. Best diffraction quality crystals obtained were from 96 well plates. Crystals from PEG 2000 MME reservoirs took 3 days to appear and a total of 10 days to grow and diffracted up to 9 Å (Fig 3.15a). Crystals from PEG 8000 reservoirs took 2 days to appear and a total of 6 days to grow and diffracted up to 5 Å (Fig 3.15a).
3.3.2.2. Seeding

Best crystals grown in PEG 8000 were used to prepare seed stock solutions. Seeding was performed in 96 well format and in 24 well plates. Seeding was done using gradients of different PEGs (3000, 6000, and 8000) and even using commercial PEG I and II screens from Qiagen. Best crystals resulting from seeding were from 23-24% PEG 3000, and were again used to prepare fresh seed stocks from which replicate plates of 23-24% PEG 3000, 0.1 M NaOAc trihydrate pH 4.6 were set in both 96 and 24 well format. Seeding was done in this sequential manner several times. Best diffraction quality crystals resulting from seeding, grew in 23.27% PEG 3000 in both 200 nl (Fig 3.16a) drops and 2 µl drops (Fig 3.16b), both of which diffracted well. Crystals grown in 200 nl drops grew between 50-100 µm, while crystals grown in 2 µl drops grew up to 200 µm.
3.3.3. Crystal diffraction

Several complete datasets from crystals diffraction were collected, the best of which was at 2.8 Å (Fig 3.17) and collected from a crystal grown in 2 µl drops and cryoprotected using 30% glycerine in reservoir solution of 0.1 M NaOAc trihydrate pH 4.6 buffer and 23.27% PEG 3000.

Figure 3.17. Crystal diffraction. Collected at 2.8 Å from crystal grown in 23.27% PEG 3000, 0.1 M NaOAc trihydrate pH 4.6. Cryo: 30% glycerine.
3.4. Characterization of NT4X

3.4.1. Aβ binding properties of NT4X under reducing conditions

Freshly dissolved Aβ peptides were subjected to SDS-PAGE to dissect the binding specificity of the three tested antibodies. Under denaturing conditions NT4X reacted with both N-terminally truncated AβpE3-X and Aβ4-X variants, but not with Aβ1-X. In addition to monomers and dimers, trimers and tetramers of AβpE3-42 and Aβ4-42 were recognized. AβpE3-40 and Aβ4-40 produced primarily monomers and dimers. Antibody 1–57 stained only AβpE3-40 and AβpE3-42, but no other bands as previously demonstrated. IC16 recognized the N-terminus of full-length Aβ1-40 and Aβ1-42, but not any of the N-truncated peptides (Fig 3.18).

Figure 3.18. SDS-PAGE Western blot analysis of IC16, 1–57 and NT4X antibodies. Freshly dissolved synthetic Aβ variants were probed to a membrane. (a) IC16 detects Aβ1-40 and Aβ1-42 monomers and low molecular weight aggregates. (b) 1–57 recognizes AβpE3-40 and AβpE3-42 monomers, low molecular weight and larger aggregates of AβpE3-42. (c) NT4X recognizes monomers and low molecular weight aggregates derived from AβpE3-40/42, Aβ4-40/42.
3.4.2. **Sensitivity of NT4X to Aβ4-42**

In order to analyze the sensitivity of NT4X, a dilution series was performed with freshly dissolved synthetic Aβ4-42 and the staining was visualized using a Western blot under reducing conditions. NT4X detected monomers and dimers between 1 and 0.03 μg corresponding to a minimum of approximately 7 picomoles of Aβ4-42 (Fig 3.19).

![Figure 3.19. SDS-PAGE Western blot analysis of Aβ4-42 for sensitivity testing of NT4X using freshly dissolved peptides. NT4X detects monomers and dimers of Aβ4-42 up to 0.03 μg peptide.](image)

3.5. **NT4X and Fab Fragment in vitro Rescue of Aβ**

**Toxicity**

*In vitro* assays were set up using primary rat cortical cultures to investigate the protective properties of the NT4X antibody against the various amyloid peptides. Concomitant with the binding properties of the antibody, NT4X rescues AβpE3-42 and Aβ4-42 oligomer induced toxicity but not of full-length Aβ1-42 induced toxicity (Fig 3.20). Experiment was performed by MRC Technology (London, United Kingdom).
Figure 3.20. NT4X protection of Aβ_{pE3-42} and Aβ_{4-42} induced toxicity in primary rat neuronal cultures. NT4X and an isotype IgG2b control antibody were assayed to assess potential protective properties in an amyloid peptide induced cellular toxicity assay using primary rat cortical cultures. Cellular toxicity was measured using an LDH release assay and results converted to percentage inhibition of cell death compared to control wells (n=3 repeats on cell preparations from 3 separate rats). Values plotted are means +/- SEM. The NT4X antibody shows inhibition of Aβ_{pE3-42} and Aβ_{4-42} induced toxicity but has no effect on the toxicity induced by full length Aβ_{1-42}.

In vitro toxicity rescue of the Fab fragment of the NT4X was also studied in primary neurons using a calcein assay. Treating the cells with freshly prepared Aβ_{4-42} resulted in a dose-dependent reduction in cell viability. NT4X significantly rescued toxicity of Aβ_{4-42} (Fig 3.21). Experiment was performed by SynAging (Nancy, France).
3.6. NT4X and Fab Fragment in vivo Rescue of Aβ Toxicity

In order to study the therapeutic effects of NT4X and its Fab fragment in wild-type mice, the animals were given an intracerebroventricular (ICV) injection of freshly prepared 50pmol Aβ4-42 and Aβ4-42 in combination with the NT4X antibody or the Fab fragment, respectively. Working memory was assessed using the Y-maze (Fig 3.22). The alternation rate was significantly reduced after injection of Aβ4-42 reaching chance level (dashed line). In contrast, mice injected with Aβ4-42 and 1 pmol NT4X antibody behaved like vehicle controls and learned well (compared to Aβ4-42 injection; one-way ANOVA: p<0.05). The alternation rate in mice treated with 10 pmol NT4X further increased (compared to Aβ4-42 injection; one-way ANOVA: p<0.01). Furthermore, mice treated with Aβ4-42 and 10pmol Fab NT4X also showed no deficits in working memory and behaved like the vehicle control (compared to Aβ4-42 injection; one-way ANOVA: p<0.01).
Mice treated with only the NT4X antibody or its Fab fragment showed a similar alternation rate than mice treated with the vehicle control alone. Experiment was performed by SynAging (Nancy, France).

Therefore, the NT4X antibody and its Fab fragment were able to rescue the learning deficits induced through Aβ4-42 injection in a dose-dependent manner.

![Graph showing results of NT4X and Fab fragment rescue](image)

**Figure 3.22.** Full-length and Fab fragment of NT4X rescue Aβ4-42 induced working memory deficits in wildtype mice. (a,b) Working memory deficits were induced by intraventricular injection of Aβ4-42 as the mice performing at chance level (dashed line). Both treatment with (a) full-length antibody and (b) Fab fragment rescued memory deficits in a dose-dependent manner. Mice treated with the vehicle control and the vehicle in combination with full-length or Fab fragment of NT4X respectively demonstrated normal working memory performance. One-way analysis of variance (ANOVA) followed by Bonferroni multiple comparisons; n=6–8 per group; **p<0.01; *p<0.05; data presented as mean±S.E.M.
3.7. Spatial Reference Memory Deficits in Homozygous Tg4-42 Mice

Memory deficits have previously been described in homozygous Tg4-42 mice at 8 months of age (Bouter et al., 2013). In order to identify the onset of spatial reference memory deficits Tg4-42 hom mice were analysed at six months (m) of age and compared to age-matched wild type controls using the Morris water maze.

All mice showed progressively decreased escape latencies over three days of cued training (Fig 3.23a; two-way repeated measures ANOVA, 6 m Non-treated (NT) and 6m Wild-type (WT): main effect of days: $p < 0.00001$). All mice showed comparable swimming speeds (Fig 3.23b).

**Figure 3.23. Cued training of the 6 months homozygous Tg4-42 water maze.** Reveals that Tg4-42 exhibit intact eyesight and motor abilities. Homozygous Tg4-42 mice and WT littermate controls were examined at 6 months of age. All mice showed progressively shorter escape latencies in response to training (main effect of days: $p < 0.00001$). Swimming speed did not differ between genotypes. Escape latencies and swimming speed: two-way repeated measures ANOVA. n = 8-9 per group; data presented as mean ± S.E.M; m = months.
Throughout the five days of acquisition training, both Tg4-42hom mice and aged-matched WT mice, showed a significant decrease in the escape latencies (Fig 3.24). A significant main effect of days could be found for escape latencies (two-way repeated measures ANOVA, days: $p=0.0291$). Also, a significant difference was detected between the genotypes (two-way repeated measures ANOVA, genotype: $p=0.0038$). Since a significant difference in average swimming speed was also identified between genotypes (two-way repeated measures ANOVA, genotype: $p=0.0073$), distance travelled before reaching goal platform was also calculated to ascertain that genotype difference did not arise from swimming speed. Again, a significant effect of days could be found (two-way repeated measures ANOVA, days: $p=0.0123$) as well as a significant genotype difference (two-way repeated measures ANOVA, days: $p=0.0206$).
Figure 3.24. Acquisition training of the Morris water maze reveals that 6 months Tg4-42hom mice perform inferior to WT controls. Homozygous Tg4-42 mice and WT littermate controls were examined at 6 months of age. Across the 5 days of acquisition both groups showed a significant decrease in the escape latencies and in distance travelled before reaching platform. (a,c) At 6 months Tg4-42hom performed significantly inferior to WT mice both in escape latency and distance travelled (b) A significant difference in swimming speed was detected between genotypes. (a)-(c). Escape latencies, distance travelled and swimming speed: two-way repeated measures ANOVA, (**p < 0.01, *p < 0.05. n = 8-9 per group; data presented as mean ± S.E.M; m = months.)

48 hours after the last acquisition trial, a probe trial was conducted to assess spatial reference memory. Six-month-old Tg4-42hom mice displayed no significant preference for the target quadrant as compared to WT controls that displayed a significant preference to the target quadrant (Fig 3.25a, one-way ANOVA followed by Bonferroni’s multiple comparisons test for WT group, T vs. L, R and
Since a significant difference in swimming speed was also identified between genotypes (Fig 3.25b, two-way repeated measures ANOVA, *genotype:* *p*=0.023), distance travelled in all quadrants was also assessed to confirm results. Again 6 months old Tg4-42hom displayed no preference for distance travelled in target quadrant as compared to WT group that preferentially covered more distance in target quadrant (Fig 3.25c, one-way ANOVA followed by Bonferroni’s multiple comparisons test for WT group, T vs. L, R and O: *p* < 0.001). Results are also evident in the group occupancy plots reflecting search strategy during the probe trial (Fig 3.25d) where the WT group shows preference for the target quadrant while 6 months old Tg4-42hom display a random search strategy.

Taken together results indicate severe spatial memory deficits of Tg4-42hom at 6 months of age.
Figure 3.25. Spatial reference memory deficits in 6 months old Tg4-42hom mice shown in the probe trial of the Morris water maze. Homozygous Tg4-42 mice and WT littermate controls were tested at 6 months of age. (a) Homozygous Tg4-42 mice did not show a preference for time spent in target quadrant. However, same-aged WT mice had a significant preference for the target quadrant. (b) Swimming speed was significantly different between genotypes. (c) Homozygous Tg4-42 mice did not show a preference for distance travelled in target quadrant as compared to WT group that showed preferentially covered more distance in the target quadrant. (d) Group occupancy plots reflecting search strategy during the probe trial. (Abbreviations: T = target quadrant; L = left quadrant; R = right quadrant; O = opposite quadrant. Quadrant preference: ANOVA followed by Bonferroni multiple comparisons, ***p < 0.001. Swimming speed: two-way ANOVA, *p < 0.05. n = 8-9 per group; data presented as mean ± S.E.M; m = months.)
3.8. Preventative Passive Immunization of Tg4-42hom Mice with NT4X and Fab

3.8.1. Rescue of spatial reference memory deficits

Severe spatial reference memory deficits in Tg4-42hom were detected at 6 months of age. Therefore, 6-month-old Tg4-42hom mice that had received weekly injections with the full-length or Fab of NT4X antibody, and those injected with an IgG antibody control for a period of 12 weeks were assessed using the Morris water maze. All Tg4-42hom treatment groups showed progressively decreased escape latencies in the cued training. A significant main effect of days could be found for escape latencies (Fig 3.26a, two-way repeated measures ANOVA, days: p < 0.0001). All mice showed comparable swimming speeds (Fig 3.26b).

The cued training revealed that all mice had an intact vision and appropriate motor abilities to swim.

Figure 3.26. Cued training of the water maze reveals that Tg4-42 treatment groups exhibit intact eyesight and motor abilities. Homozygous Tg4-42 treatment groups were examined at 6 months of age. All mice showed progressively shorter escape latencies in response to training. Swimming speed did not differ between treatment groups. Escape latencies and swimming speed: two-way repeated measures ANOVA. n = 10-13 per group; data presented as mean ± S.E.M.
All groups showed a significant decrease in escape latencies over the five days of acquisition training (Fig 3.27a, two-way repeated measures ANOVA, days: p < 0.0001). Different treatments did not have a significant effect on escape latency. Swimming speed between groups was comparable (Fig 3.27b).

**Figure 3.27. Spatial learning was assessed in the acquisition training of the Morris water maze.** Tg4-42hom treatment groups were examined at 6 months of age. (a) Across the 5 days of acquisition all groups showed a significant decrease in escape latency. (b) Swimming speed did not differ between treatment groups. Escape latencies: two-way repeated measures ANOVA, effect of days: p < 0.0001. n = 10-13 per group; data presented as mean ± S.E.M.

48 hours after the last acquisition trial, a probe trial was conducted to assess spatial reference memory. No quadrant preference was found for IgG control treatment group. Both full-length and Fab NT4X immunized Tg4-42hom mice displayed a significant preference for the target quadrant, as indicated by the percentage time spent in different quadrants of the pool (Fig 3.28a, one-way ANOVA, full-length NT4X: p < 0.01 T vs. L and O quadrant; Fab: p<0.001 T vs. L and O, T vs R p <0.01 quadrants). Swimming speed did not differ between treatment groups (Fig 3.28b).
This indicates that both the full-length NT4X antibody and the Fab fragment of NT4X were able to rescue spatial reference memory deficits in 6 months old Tg4-42hom mice.

No significant difference in Cued training, Acquisition training or the Probe trial test were found between the Non-treated 6 months old Tg4-42hom (Fig 3.28) and the IgG control treatment group (Fig 3.28).

**Figure 3.28.** Passive immunization rescues spatial reference memory deficits in NT4X and Fab treated Tg4-42hom mice shown in the probe trial of the Morris water maze. Homozygous Tg4-42 mice were tested at 6 months of age. (a) Tg4-42hom IgG treated control mice did not show a preference for the target quadrant. However, NT4X and Fab treated mice showed significant preference for the target quadrant. (b) Swimming speed did not differ between the groups. (Abbreviations: T = target quadrant; L = left quadrant; R = right quadrant; O = opposite quadrant. Quadrant preference and swimming speed: One-way analysis of variance (ANOVA) followed by Bonferroni multiple comparisons, ***p < 0.001; **p < 0.01. n = 10-13 per group; data presented as mean ± S.E.M.)

### 3.8.2. Deceleration of neuron loss

In order to study the potential therapeutic effect of NT4X on the observed neuron loss in 6 months old Tg4-42hom the neuron numbers in the CA1 were quantified using design-based stereology.
The number of neurons was compared between 6-months-old Tg4-42hom immunized with full-length NT4X or Fab NT4X and a control group immunized with an IgG2b antibody control. Stereology revealed a significant treatment effect (Fig 3.29, one-way ANOVA, treatment: p = 0.0007). Following ANOVA, the individual groups were then analysed using Dunnett’s multiple comparisons. Compared to IgG control injected mice (mean=136,170; SEM±5455), the neuron numbers in Tg4-42hom were significantly increased after immunization with full length NT4X (mean=160,156; SEM±5096) and Fab (mean=165,844; SEM±4717) (Fig 3.29; one-way ANOVA: NT4X vs IgG injected: p<0.01; Fab vs IgG injected: p<0.001).

NT4X immunized Tg4-42 mice displayed 14.97% more neurons in the CA1 layer of the hippocampus than control IgG treated mice. In comparison, Fab immunized mice displayed 17.89% more neurons as compared to control IgG treated mice.

Figure 3.29. NT4X and Fab mitigate neuron loss in Tg4-42hom mice. Quantification of neurons in the CA1 of 6-months-old Tg4-42hom using unbiased stereology. Tg4-42hom mice that received weekly injections with the NT4X antibody or its respective Fab fragment and an IgG control group for a period of 12 weeks were analysed at 6 months of age. Tg4-42hom mice immunized with NT4X full length or Fab fragment both displayed significantly more neurons than age-matched IgG control injected mice. One-way analysis of variance (ANOVA) followed by Dunnett’s multiple comparisons; n=10–14. (**p<0.01; ***p<0.001; data presented as mean±S.E.M.)
CHAPTER 4

4. Discussion

4.1. N-Truncated Amyloid Beta 4-x: The Target

4.1.1. Properties of Aβ4-x

Part of this work has been dedicated to confirm N-truncated amyloid beta starting at position 4 (phenylalanine) as a viable target in AD research and therapy. To this end, the aggregation properties of Aβ4-x, under reducing conditions, was demonstrated using the N-terminal specific antibody NT4X (Fig 3.18). As previously reported by Jan and colleagues for full length Aβ (Jan et al., 2008), we also mark that the aggregation properties of N-terminally truncated AβpE3-x and Aβ4-x is affected by the carboxyl-terminal length of the peptide. Wherein, under reducing conditions, AβpE3-40 and Aβ4-40 migrate as monomers (4 kDa) and dimers (8 kDa), in contrast to AβpE3-42 and Aβ4-42, both of which exhibited higher molecular weight aggregations of trimer (12 kDa) and tetramers (16 kDa) in addition to monomers and dimers. We had previously shown that in contrast to full length Aβ1-42, which tends to remain in equilibrium with its monomeric state for longer, AβpE3-42 and Aβ4-42 tend to form stable oligomers faster (Bouter et al., 2013). It has previously also been reported that the aggregation propensities of Aβ may be directly affected by varying either terminal length of the peptide (Jan et al., 2008; Jarrett et al., 1993; Pike et al., 1995).

4.1.2. Aβ4-x as a target

The neurotoxicity of N-terminal truncated variants of Aβ has been known for over two decades (Pike et al., 1995). First demonstrated by Pike and colleagues, it was later suggested that they may initiate or nucleate the pathological deposition of Aβ into plaques.
(Russo et al., 2002). These observations were recently extended by our lab, showing that Aβₚₑ₃₋₄₂ and Aβ₄₋₄₂ have specific structural features that render their soluble aggregates neurotoxic. The stable aggregates formed by Aβₚₑ₃₋₄₂ and Aβ₄₋₄₂ were different in size and morphology than those formed by full length Aβ₁₋₄₂, as shown by far UV CD spectroscopy, dynamic light scattering and NMR spectroscopy (Bouter et al., 2013).

The exact levels of N-truncated Aβ peptides that comprise a part of the total Aβ pool of AD patient’s has been hard to determine. This is mostly due to a variation of methodology and tools used to extract this information. It is in agreement nonetheless, that N-truncated Aβ peptides appear during Alzheimer disease progression (Bayer & Wirths, 2014). It is of interest that intraneuronal Aβ has also been reported to be N-truncated and that Aβ₄₋ₓ, starting with phenylalanine is the variant that is mostly reported in pre-symptomatic AD cases (Gouras et al., 2000). In good agreement, N-terminal truncated variants (beginning with position 4 “Phenylalanine”, position 5 “Arginine”, position 8 “Serine”, position 9 “Glycine” and position 3 with a pyroglutamyl residue) comprised the first stage of amyloid deposition in non-demented individuals. At this stage all Aβ variants detected were not truncated at their C-terminus (Aβₓ₋₄₂) (Sergeant et al., 2003).

Masters and colleagues had first reported that Aβ₄₋ₓ was a component of amyloid deposits analyzed from human AD brains (Masters et al., 1985). In agreement with these findings, cotton wool plaques in patients with familial AD, were found to harbor high amounts of Aβ₄₋₄₂ (Miravalle et al., 2005). Along those lines, we have also shown that Aβ₄₋ₓ intraneuronal accumulation precedes that of Aβₚₑ₃₋ₓ in the established 5XFAD mouse model (Antonios et al., 2013).

To that end, the Tg₄₋₄₂ mouse model has been recently established (Bouter et al., 2013) and validated as an advantageous research tool in AD research as it exclusively expresses Aβ₄₋₄₂, allows
for its intraneuronal accumulation, and permits the study of a previously neglected therapeutic option in AD; the Aβ_{4-42} peptide.

4.1.3. The Tg4-42 model: Aβ_{4-42} in vivo

N-truncated Aβ peptides have been reported in several AD mouse models. A 2D electrophoresis analysis combined with mass spectrometry, has enabled the detection of a wide variety of N-truncated Aβ variants in APP/PS1KI mice. At 2.5 months Aβ_{2/3-42}, Aβ_{4-5-42} and Aβ_{8/9/10-11-42} were found, and the appearance of Aβ_{pE3-x} was noticeable from 6 months of age (Casas et al., 2004). Another typical AD mouse model to exhibit N-truncated Aβ peptides is the 5XFAD model. Confirmed variants in the 5XFAD model have been Aβ_{pE3-40/42}, Aβ_{4-40/42} and Aβ_{5-40} (Antonios et al., 2013; Guzman et al., 2014; Jawhar et al., 2012; Oakley et al., 2006; Richard et al., 2015). Other models reported to have N-truncated Aβ peptides are the PDAPP, Tg2576 and APP23 mouse models, all of which also exhibit intracellular Aβ accumulation (Chapman et al., 1999; Games et al., 1995; Hsiao et al., 1996; Kelly et al., 2003; Masliah et al., 1996; Rijal Upadhaya et al., 2013; Rockenstein et al., 1995; Sturchler-Pierrat et al., 1997).

The disadvantage that had previously accompanied transgenic mouse models of AD was the fact that they relied overly on the over-expression of APP or the expression of mutations that affected the processing of APP. This has led to models that mostly reflect the familial form of AD from which these models were inspired. They also neglected contributions from N-truncated Aβ forms, which were found in small amounts relative to full length Aβ_{1-40/42} in these models. Although highly beneficial for basic research into the etiology of AD disease, the models might not provide perfect tools for therapeutic intervention that may be translated into the clinic.

To this end, our lab has recently developed the Tg4-42 mouse model. Expressing only Aβ_{4-42} under the THY1 promoter, the model
exhibits some aspects of AD pathology without the reliance on APP overexpression or processing. $\text{A}\beta_{42}$ intraneuronal accumulation eventually leads to neuron loss and spatial memory deficits, without extracellular $\text{A}\beta$ accumulation in the form of plaques (Bouter et al., 2013). This further boosts our previous hypothesis on the importance of intraneuronal $\text{A}\beta$ being the trigger for respective downstream pathological events (Wirths et al., 2004).

4.1.4. **Neuron loss and spatial reference memory: Age-dependent decline**

Efforts to model AD-relevant cognitive deficits in mouse models have met with partial success (Kobayashi & Chen, 2005). The hippocampus is one of the first and most severely touched brain regions in AD patients. Various mouse models have tried to assess reference memory as this particular memory function is highly specific for the hippocampal region (West, 1993). We have previously shown that at 8 months of age, homozygous Tg4-42 mice have developed severe hippocampus neuron loss and spatial reference memory deficits (Bouter et al., 2013). It was of interest to the current work, to determine the onset of neuron loss and subsequent spatial reference memory deficit in the Tg4-42 homozygous mice.

We show that the Tg4-42hom mice exhibit an age-dependent neuron loss in the CA1 region of the hippocampus where neuron loss starts at 4 months of age and gradually decreases with age to reach 50% at 6 months of age and 64% at 8 months of age (Antonios et al., 2015). More importantly, this coincides with age-dependent spatial memory deficits where the mice at 6 months of age have developed a severe spatial reference memory deficit as compared to aged matched WT control mice both in acquisition training and in the probe trial of the Morris water maze (Fig 3.24 and Fig 3.25). In rats, spatial memory impairment has been reportedly linked to bilateral dorsal hippocampal lesions, where the onset of impairment coincided with
lesions that encompassed 30-50% of total volume and worsened dramatically as lesion size increased above 50% of total hippocampal volume (Broadbent et al., 2004). At 5 months of age, the Tg4-42hom mice suffer from a 43% neuron loss in the CA1 and signs of mild spatial reference memory deficits as compared to 6 months where neuron loss has increased to 50% and spatial reference memory deficits have become severe (Antonios et al., 2015). This is in agreement with other findings which have shown that a significant neuron loss in the hippocampus may still be compensated as only 20-40% of the total hippocampus is required for effective spatial learning (Moser et al., 1995).

Other studies have also shown an age dependent reference memory decline in transgenic mouse models of AD. Westerman and colleagues have shown that spatial reference memory deteriorates progressively after 6 months of age, as shown in a Morris water maze, in Tg2576 mice. Severe deficits, however are only apparent after 12 months of age (Westerman et al., 2002). Another study utilized the radial arm water maze to show an age dependent reference memory decline in APP/PS1 KI mice. Again, memory deficits are only observable starting at 11 months of age and become severe at 15 months (Webster et al., 2013).

This goes to show the importance of the Tg4-42hom mice as a tool for AD therapeutic intervention. Since the mice already display tremendous neuron loss and reference memory deficits at 6 months of age, the timeline for therapeutic intervention is drastically cut down as compared to other mouse models.

4.2. **NT4X: The Therapeutic Tool**

Since Aβ4-40 peptides were found to form stable aggregates in solution (Bouter et al., 2013), we have used them in order to generate a monoclonal antibody that binds at the N-terminus of Aβ.
The novel oligomeric Aβ specific antibody NT4X was generated by immunizing three Balb/c mice with unconjugated Aβ4-40. After preparation of the lymph nodes they were fused with the myeloma cell line P3-X63-Ag8 for generation of the hybridoma cells. After fusion, the hybridoma cells were screened by an enzyme-linked immune-absorbent assay (ELISA) for an antibody that recognizes Aβ4-10/40 peptides but not the C-terminal end of amyloid beta, thus not recognizing the Aβ36-40 peptide. Positive antibody clones were further screened by IHC staining of human brain sections. The intention of the screening procedure was to find a clone that would not preferentially bind to amyloid plaques thereby identifying NT4X (Antonios et al., 2013).

4.2.1. Antibody purification

The hybridoma clone B4C11 was subsequently chosen as it fit all previously mentioned selection criteria for the desired antibody. Although small-scale cultures of the hybridoma cell line were maintained in media containing serum, the need for large-scale antibody purification (Fig 3.1) required the conditioning of the cell line with a serum-free culture media. This was done to achieve higher purity of antibody purified. It is well known that serum containing media would contain high titers of antibody from the host organism thus interfering with purification (Howard & Kaser, 2013).

For the purpose of this study, large amounts of purified antibody were necessary. Since the small-batch production of milligram amounts of antibody was not sufficient, we turned to culture using bioreactors to improve the amount and efficiency of purification. It has been previously reported that a perfusion culture of hybridoma cells drastically increases the number of viable cells (cells/mL) and thus leads to higher overall amounts of antibody produced (mg/L) per liter of culture media (Tang et al., 2007). Our results are in good agreement with this as we achieved antibody yields...
upwards of 50-60 mg/mL as compared to 30 mg/mL from previous purifications using traditional methods.

Tracking of antibody purification reveals a high degree of purity was achievable through the designed purification protocol, while the presence of both heavy and light chains confirm antibody presence (Fig 3.3 and Fig 3.4).

4.2.2. Antibody characterization

In order to determine the specificity of the purified NT4X antibody against Aβ species, it was utilized as a primary antibody in western blot analysis of various N-truncated and full length Aβ species. Since the antibody itself was raised against unconjugated Aβ4-40, it was no surprise that it detected Aβ4-40 and Aβ4-42 as shown under reducing conditions (Fig 3.18). The antibody however also recognized the pyroglutamate modified N-truncated Aβ species as well, recognizing both AβpE3-40 and AβpE3-42 peptides under reducing conditions (Fig 3.18). This is further corroborated by results obtained from a similar experiment performed under native conditions where the antibody also recognized N-truncated Aβ species (Antonios et al., 2013).

It is worth noting that, as compared to the IC16 antibody, the NT4X antibody does not recognize full length Aβ1-40/42 as we have shown in both reducing and native conditions. This is due to the fact that the antibody binds most strongly to the phenylalanine as position 4 of Aβ, as demonstrated by a pepscan analysis, and the additional peptides contained in full length Aβ might be blocking it from binding to its preferred site (Antonios et al., 2013).

Previously characterized antibodies have been raised against pyroglutamate modified N-truncated Aβ species (Frost et al., 2012; Wirths et al., 2010), most have been raised against full length Aβ (Bard et al., 2000; Dodart et al., 2002; Wilcock et al., 2004c) and the NT4X antibody is unique in its ability to recognize both AβpE3-x and Aβ4-x species. Moreover, it is highly sensitive and may bind as little as
30 ng of Aβ4-42, corresponding to a sensitivity of approximately 7 picomoles under reducing conditions (Fig 3.19).

We further show that the NT4X antibody, as previously selected to be, reacts to a very low extent with extracellular plaques in human AD tissue (Antonios et al., 2013). Our working group has further expanded and recently confirmed the previous result in comparison to the biosimilar antibodies of Bapineuzumab, Crenezumab, Solanezumab in sporadic AD cases and mouse models (Bouter et al., 2015). Similarly, this description has been valid for the confirmation specific AβpE3-x antibody 9D5 (Wirths et al., 2010). In murine tissue the NT4X reacts both with intracellular and extracellular Aβ and we were able to show that in the homozygous 5XFAD model (Richard et al., 2015), that the intraneuronal accumulation of Aβ4-x precedes that of AβpE3-x, underlying a fact which was not previously presented (Antonios et al., 2013).

4.2.3. **In vitro and in vivo target engagement and rescue**

To further out work in characterizing the NT4X antibody and its therapeutic potential, collaborators at SynAging (Nancy, France) and MRC Technology (London, United Kingdom) have performed in vitro and in vivo target engagement and rescue assays with the NT4X antibody.

To go in parallel with our results showing that the NT4X recognized both N-truncated AβpE3-x and Aβ4-x species, but that full length Aβ1-x (Antonios et al., 2013), NT4X was also found to rescue the toxicity resulting from both AβpE3-42 and Aβ4-42 in primary rat neuronal cultures (Antonios et al., 2015). Even when antibody concentration is increased well above 1:1 molar ratio (antibody:peptide) the NT4X does not rescue toxicity induced by full length Aβ1-42 as measured by an LDH release assay. Not surprisingly, the antibody seems also to rescue Aβ4-42 preferably as it requires only 133.7 µg/ml of NT4X to reach 50% of cell death inhibition caused by 10 µM of Aβ4-42, as compared to 109.5 µg/ml antibody to reach the same level of cell
death inhibition (50%) caused by a lesser amount of 5 µM Aβ_pE3-42 (Fig 3.20). The molecular mechanism of Aβ-induced neuronal cell death is not well understood, though studies have reported the involvement of reactive oxygen species (ROS) in Aβ induced neuronal cell death (Behl et al., 1995). It was more recently determined that Aβ induces neuronal cell death through ROS mediated ASK1 activation, and the authors have proposed ASK1 as a therapeutic target for prevention and treatment of AD (Kadowaki et al., 2005).

It was previously demonstrated that Aβ peptide ICV injection induced short term memory impairment in rats (Stepanichev et al., 2003). Recently, we have shown that N-truncated Aβ variants (including Aβ_4-42) induce working memory deficits in wild type mice (Bouter et al., 2013). In order to demonstrate NT4X in vivo target engagement, a similar experimental setup was reproduced. ICV injection of Aβ_4-42 caused a working memory deficit as demonstrated by a Y-maze, the effect of which is completely rescued by the NT4X antibody in a dose dependent manner (Fig 3.22a). The work is essential in showing that the NT4X is capable of target engagement and rescue of Aβ_4-42 in vivo, under physiological conditions where the peptide would otherwise be toxic.
4.3. Passive Immunization of the Tg4-42hom AD Model Using NT4X and its Fab

4.3.1. Timeline and dosage

Pre-clinical passive immunization studies usually differ in at least one of the following parameters: Dosage/time course, animal model employed and antibody used for immunization. Previous therapeutic and preventative passive immunization trials against N-truncated Aβ peptides have attempted both subcutaneous and intraperitoneal administration and opted for a dosage of 12.5 mg/kg body weight (DeMattos et al., 2012; Frost et al., 2012). Moreover, mouse models frequently used in immunization studies such as the Tg2576 and PDAPP exhibit a difference in the morphology and stoichiometry of Aβ deposits (Fryer et al., 2003; Hsiao et al., 1996; Sasaki et al., 2002). This goes to reflect the difficulty in comparing different preclinical trials. It also reflects as to why the outcome of clinical studies poorly reflected the results from pre-clinical immunizations (Lannfelt et al., 2014).

At 4 months of age, neuron loss in the CA1 region of the hippocampus in the homozygous Tg4-42 is marked with a 17% loss, and precedes the appearance of behavioral deficits. As the mice are still pre-symptomatic at that stage, still capable of learning and have an intact reference memory till the age of 4 months in a MWM, we opted for the purpose of this study to perform a preventative passive immunization. For early intervention in AD by passive immunization, it would be necessary to identify patients inclined to more likely develop cognitive impairment. Though this is currently not possible, it has driven research into the defining of preclinical biomarkers in AD (Fiandaca et al., 2014).

The dosage, administered intraperitoneally, chosen for this study was 10 mg/kg body weight to be injected weekly. In this treatment window, mice would be immunized with the NT4X antibody starting 3 months (12 weeks) of age where the animals are pre-
symptomatic, for the duration of 12 weeks reaching an age of 6 months. This has allowed us to examine the preventative therapeutic effect of the NT4X at an endpoint where the mice usually exhibit reference memory deficits (Fig 4.1).

![Passive Immunization timeline of Tg4-42](image_url)

**Figure 4.1. Passive Immunization timeline of Tg4-42.** Onset and progression of neuron loss and of spatial reference memory decline are displayed.
4.3.2. **Rescue of neuron loss**

Almost all pre-clinical immunization studies, have relied on Aβ burden as a measurable outcome. While some approaches were therapeutic, with immunization starting after the onset of plaque pathology, other attempted a preventative approach. In therapeutic trials, some studies have reported no clearance of existing plaques (Dodart et al., 2002; Levites et al., 2006), while other report limited clearance (Tucker et al., 2008). Not all trials were unsuccessful however, with several studies suggesting a significant reduction in pre-existing plaques (Frost et al., 2012; Wilcock et al., 2004b). On the other hand, preventative passive immunization has been more successful, with numerous reports of significant inhibition of plaque pathology (Bard et al., 2000, 2003; Bussiere et al., 2004; Frost et al., 2012; Lord et al., 2009).

In spite of the previous, it is well known that Aβ levels do not necessarily correlate with the therapeutic outcome. Reversal of memory deficits in mice, for example, has been shown in an AD mouse model, even though the preclinical passive immunization study did not reduce brain Aβ burden (Dodart et al., 2002). On the other hand, passive immunization of APPswe/PS1ΔE9 transgenic mice with an antibody against the N-terminus of pyroglutamate Aβ_{3-42} significantly reduced total plaque deposition in the hippocampus and cerebellum, without however, affecting Aβ levels as measured by ELISA (Frost et al., 2012).

In this study, we propose a new measurable outcome for the assessment of passive immunization in the homozygous Tg4-42 mice. As previously discussed, the Tg4-42 mice exhibit an early loss of neuron in the CA1 region of the hippocampus. Preventative passive immunization using the NT4X antibody or its Fab fragment, was able to mitigate neuron loss as compared to an IgG control group (Fig 3.29).
4.3.3. **Rescue of reference memory deficits**

While most passive immunization studies do not tackle the issue of behavioral deficits, most likely due to the fact that results were negative, others have reported findings. The rescue of memory deficits has been reported for a variety of mouse models such as the 5XFAD (Wirths et al., 2010), PDAPP (Dodart et al., 2002) and the Tg2576 (Wilcock et al., 2006; Wilcock, et al., 2004a). Is it important to mention that some have even reported the reversal of memory deficits with no reduction in plaque load, further asserting the fact that plaque burden and behavioral deficits are not correlated (Dodart et al., 2002). Moreover, most mouse models in which an amelioration of behavioral deficits is reported, are APP overexpressing or enzymatic mutation of APP related models, reflecting more cases of familial AD than those of sporadic AD.

We were able to achieve a reversal of spatial reference memory deficits in the Tg4-42 homozygous mice after passive immunization with the NT4X antibody and its Fab fragment. The preventative treatment, allowed for mice to maintain an intact spatial reference memory, as one exhibited in the pre-symptomatic phase (Fig 3.28). This goes in line with the direction of on-going clinical trials where most passive immunization trials are now focused on prevention and very early treatment of AD (Lemere, 2013).

4.3.4. **Mechanism of NT4X passive immunotherapy**

The cornerstone theory of Aβ engagement, suggested that antibodies may act catalytically to prevent Aβ aggregation or dissolve its preformed aggregates (Solomon et al., 1997). This was later on supported by the first active immunization trials which demonstrated plaque removal in response to immunization with Aβ_{1-42} aggregates, allowing the immune system to develop anti- Aβ antibodies (Schenk et al., 1999). Though active immunization per-say was abandoned due to life-threatening side effects, the field still moved forward in research.
concerning removal of deposited amyloid peptides. It is estimated that less than 1% of antibody injected in passive immunization crosses the BBB to reach its target, treatment effects have nonetheless been positive (Lemere, 2013). Bapineuzumab was the first humanized antibody in clinical trials, its murine predecessor 3D6 target full length Aβ_{1-42}. It failed however to improve clinical outcomes in a phase III placebo-controlled study of patients with mild to moderate AD (Salloway et al., 2014). Studies have even suggested that the solubilization of fibrillar Aβ may have a noxious effect (Benilova et al., 2012; Haass et al., 1994). In this study, we circumvent this issue by the use of the Tg4-42 model, which does not exhibit Aβ deposits in the form of plaques, and allows for target engagement of the NT4X antibody without the repercussions faced by other passive immunotherapy studies which employed models harboring plaque pathology.

A different hypothesis for antibody Aβ clearance, the peripheral sink hypothesis, does not require the penetration of the BBB. It has been shown that Aβ can be rapidly transported to the periphery (Ghersi-Egea et al., 1996; Shibata et al., 2000). In this light, it has been proposed that a shift in Aβ equilibrium, driving a higher efflux from the central nervous system would allow peripherally circulating antibodies to facilitate Aβ clearance (Bacskai et al., 2002; Das et al., 2003). In reality however, reports have been contradictory. While some studies utilizing the murine predecessor of Solanezumab, the m266 antibody, have reported an Aβ/m266 complex in the plasma and cerebrospinal fluid of treated PDAPP mice, thus supporting the peripheral sink hypothesis (Dodart et al., 2002b). Other studies have reported that the m266 neutralizes intracerebral rather than peripheral Aβ (Yamada et al., 2009).

Microglial activation in immunotherapy has been suggested as a key player in the clearance of Aβ (Wang et al., 2011). The mechanism proposed that the clearance of amyloid deposits occur via an interaction of the Fc receptor on microglia cells with the Fc art of the
therapeutic antibody bound to Aβ (Bard et al., 2000). Results obtained in this study, do not support this hypothesis, as the passive immunization with the Fab fragment of the NT4X antibody, resulted in the same therapeutic effect of the full length antibody itself in mitigating neuron loss and reversing spatial reference memory deficits (Fig 3.29 and Fig 3.28). Along those lines, another study using Tg2576 APP mice that were further engineering as Fc receptor-gamma chain knock-outs, still demonstrated an effect of anti-Aβ antibodies on Aβ deposition (Das et al., 2003), disproving the need for Fc receptor mediated phagocytosis.

This study has been able to discuss several aspects of the treatment mechanism in AD passive immunotherapy. Primarily, we show that mouse models of AD needn’t harbor plaque pathology and that the Tg4-42 mouse model presents a chance to tackle soluble Aβ42 aggregates of as possible therapeutic targets. We have introduced another paradigm in terms of measurable pre-clinical outcome for passive immunotherapy; neuron loss, and have demonstrated that the Fab fragments of antibodies are viable options for therapy.

4.4. Outlook in Immunotherapy

4.4.1. Autoantibodies against Aβ

The presence of natural antibodies against Aβ has been reported in plasma samples and the cerebrospinal fluid of both AD patients and healthy controls (Britschgi et al., 2009). One such antibody presently in clinical trials is Aducanumab (Biogen). The most recent developments at the AD/PD conference (March 2015) reported interim analysis of first data sets from Aducanumab (Biogen). Developed from aged, cognitively healthy, donors using reverse translational medicine, it is a fully human IgG1 monoclonal antibody against a conformational epitope found on Aβ. Results have so far been promising with a dose dependent reduction in amyloid deposition in six cortical regions. Results communicated in the Alzheimer’s
Association International Conference (AAIC) in August 2015, the 6 mg/kg immunotherapy group was numerically reported to be closer to the 1 mg/kg than the 3 or 10 mg/kg group, in the mini-mental state examination (MMSE), but overall dose dependence remained significant. Phase III trials began in August 2015 and are set to run till 2022 (Alzforum.org).

4.4.2. Passive immunotherapy and vascular pathology

A diet deficient in folate and vitamins B6 and B12, has been reported to cause vascular cognitive impairment in WT mice (Sudduth et al., 2013). APP/PS1 mice, on a similar diet as previously mentioned, had a remarkable redistribution of Aβ to the vasculature from the parenchyma, although total Aβ levels remained unchanged. The mice also worsened as compared to control groups, in a radial arm maze (Sudduth et al., 2014). This has raised questions regarding the co-morbidity effect of passive immunotherapy and vascular pathology. Aβ-related imaging abnormalities (ARIA), which have been linked to vasogenic edema, were part of the demise of the Bapineuzumab program, and are being investigated in the ongoing trials with Aducanumab as well (Alzforum.org).

4.4.3. Intrabodies in immunotherapy

Intracellular antibodies, or Intrabodies, are small, usually single-chain antibodies that are able to penetrate cells through the use of viral or other types of vectors. They are designed to bind and inactivate target molecules inside cells (Chen et al., 1994).

The use of protein engineering, has further allowed the targeting of intrabodies to specific subcellular compartments. By adding a KDEL amino acid motif, targeting the antibody to the endoplasmic reticulum, the scFvA13 intrabody, which only recognizes Aβ in a sequence and confirmation selective manner, a study has been able to decrease the production of Aβ oligomers in 7PA2 cells expressing the
antibody (Meli et al., 2014). It was previously shown that 7PA2 cells produce large quantities of \( \text{A}\beta_{42} \) and exports various forms of the peptide out of the cell (Walsh et al., 2005). Meli and colleagues are now investigating, using in vivo experiments, the targeting of the scFvA13 intrabody to the hippocampus of AD mouse models (Alzforum.org).

4.5. Crystallization of NT4X Fab and Target Peptide

4.5.1. Generation of Fab fragment

In order to study the antibody: antigen binding properties of the NT4X antibody, the Fab fragment of the antibody had to be generated and ensured to be functional. Antibody subclasses exhibit diverse physical properties and show a substantial difference in their susceptibility to cleavage sites. Experimental parameters must be examined and adjusted for each individual subclass to ensure a complete digestion and more importantly a homogenous one. Antibody fragments have been routinely obtained by papain digestion. It has been suggested that the monitoring of antibody papain digestion by electrospray ionization mass spectrometry is of benefit (Bennett et al., 1997). Initial results from antibody digestion with papain resulted in a heterogeneous mix of Fab subtypes and that of over digested products (Fig 3.5). The separation of resulting fragments was not possible via size exclusion chromatography nor through ion exchange chromatography (Fig 3.6).

Although it was believed that the digestion of antibody fragments using pre-activated papain will result in the generation of \( F(\text{ab}')2 \) fragment (100 kDa), it was later suggested that a closer monitoring of digestion conditions with regard to antibody subtype can result in a homogenous digestion resulting in an Fab fragment using pre-activated papain (Adamiczyk et al., 2000). Our results are in agreement with this, as digestion with pre-activated papain resulted in
a homogenous digestion of the NT4X antibody, resulting in a more pure Fab fragment (Fig 3.8).

**4.5.2. Fab functional *in vitro* and *in vivo***

In order to ensure the Fab fragment of the NT4X antibody has the same binding and functional properties as its full-length counterpart, our collaborators in Synaging have performed *in vitro* and *in vivo* toxicity rescue assays. The *in vitro* cellular toxicity assay using the Fab fragment, demonstrates the capability of the Fab fragment to bind and rescue the toxicity caused by Aβ_{4-42} in a dose dependent manner (Fig 3.21). On the other hand, an *in vivo* assay has demonstrated the Fab’s ability to rescue an Aβ_{4-42} induced working memory deficit in WT mice, as tested by a Y-maze alteration assay (Fig 3.21b). The experiments demonstrate that the NT4X Fab is functional and exhibits similar binding capabilities as the full-length antibody. We have also used the Fab in passive immunization of the Tg4-42 mice where it has demonstrated an ability to bind its target in an AD mouse model and rescue both neuronal loss and spatial reference memory deficits (Antonios et al., 2015).

**4.5.3. Crystallization conditions***

To examine the structural basis of NT4X binding to Aβ, we endeavored to isolate the NT4X as a Fab fragment in complex to the N-truncated Aβ_{4-19} peptide and went further to crystallize the complex. Initial screening for crystallization conditions tackled both matters of appropriate conditions and that of protein concentration (Chayen & Saridakis, 2008). Initial crystals obtained were of the unconjugated Fab fragment of the NT4X (Fig 3.12) and SDS-PAGE followed by silver staining confirmed that crystals observed were of respective size corresponding to a Fab fragment (Fig 3.13).
We next pursued the crystallization of the Fab: Aβ₄₋ₓ peptides complex. Several Aβ C-terminal lengths were tested, including: Aβ₄₋₁₀, Aβ₄₋₁₉ and Aβ₄₋₂₂. Crystallization was successful using the Fab: Aβ₄₋₁₉ complex. The corresponding peptide sequence of Aβ₄₋₁₉ is the following:

\[
\text{Phe}^4 - \text{Arg} - \text{His} - \text{Asp} - \text{Ser} - \text{Gly} - \text{Tyr} - \text{Glu} - \text{Val} - \text{His} - \text{His} - \text{Gln} - \text{Lys} - \text{Leu} - \text{Val} - \text{Phe}^{19}
\]

Initial crystals of the Fab: Aβ₄₋₁₉ complex grew in PEG 2000 MME. Though some were amorphous in shape (Fig 3.14b,c), others were overlapping crystals (Fig 3.14a) of a distinct nature that were used to develop screening grids in conjunction with seeding. Seeding in crystallization, is used to bypass excess nucleation. By introducing ready-made nuclei directly into the metastable zone, the presence of a limited number of nuclei may ensure growth into larger single crystals (Bergfors, 2003; Chayen, 2003). Our results demonstrate the previous by (Fig 3.15 and Fig 3.16). We also report an improvement in the diffraction potential of crystals, where successive rounds of seeding have improved the diffraction resolution limit from 9Å (Fig 3.15a) and 5Å (Fig 3.15b) to 2.8Å (Fig 3.16b). It is worth noting that all crystals from respective seeding rounds grew in very similar conditions consisting of PEG (3000, 6000 or 8000) as precipitant and (0.1M, pH = 4.6) sodium acetate trihydrate.

The crystallization conditions of the NT4X Fab: Aβ₄₋₁₉ complex, is very comparable to previously published data. Crystals of the WO2 antibody, the first anti-Aβ antibody to be crystallized complexed with Aβ, was crystallized in 25% PEG 400, 100 mM MES pH 6.5 (Wun et al., 2008). Later on, Bapineuzumab, the first anti-Aβ antibody to go into AD passive immunotherapy, was crystallized in complex with Aβ₁₋₂₈ in 20% PEG 3350, 0.2 M sodium formate (Crespi et al., 2014).
4.5.4. Crystallization of clinically relevant antibodies

It is the future objective of this work, to determine the molecular basis of the Aβ recognition of NT4X. In context of clinical relevance, this will allow the development and humanization of the NT4X in a way which enhances its binding to its target. A report has previously demonstrated the conservation of antigen contact residues and conformation of antigen recognized between the 3D6 antibody and its humanized counterpart Bapineuzumab (Feinberg et al., 2014). It has been previously demonstrated that the introduction of mutations to an antibody, following information gathered from its crystallographic analysis, induced a 500 fold improvement in affinity for a helical peptide ligand. The mutated residues did not actually interact with the ligand directly, it was rather through subtle changes in the framework surrounding the binding sites (Zahnd et al., 2004). Moreover, the crystal structure of a Bapineuzumab Fab-Aβ peptide complex revealed that it captured Aβ in a monomeric helical conformation at the N-terminus (Miles et al., 2013). Furthermore, the mobilization of toxic Aβ species from inert plaque material, might partially explain the failure of Bapineuzumab in clinical trials (Salloway et al., 2014).

A recent study suggested that amino acids that interacted with Aβ in both the Solanezumab and Crenezumab antibodies had nearly the same sequence, suggesting they bind Aβ in the same way (Crespi et al., 2015), although reports had other suggested that they recognize different forms of Aβ (Table 4.1).
Table 4.1. Conformational binding properties of some clinically relevant Aβ antibodies.

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Epitope Structure</th>
<th>Conformations Recognized</th>
<th>Aria-E</th>
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<tbody>
<tr>
<td>Solanezumab*</td>
<td>AA 16-26 Helix - β coil</td>
<td>Yes Unknown Unknown Yes Low</td>
<td></td>
</tr>
<tr>
<td>Crenezumab*</td>
<td>AA 16-26 Helix - β coil</td>
<td>Yes Yes Yes Low</td>
<td></td>
</tr>
<tr>
<td>Bapineuzumab*</td>
<td>AA 1-7 Helix</td>
<td>Yes Unknown Yes High</td>
<td></td>
</tr>
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<td>Gantenerumab*</td>
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<td>Aducanumab*</td>
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*Information retrieved from Alzforum.org

Adolfsson and colleagues have reported that, unlike Solanezumab, Crenezumab binds aggregated forms of Aβ (Adolfsson et al., 2012). On the other hand, we have recently shown that, both antibodies showed a very similar staining profile in formalin-fixed and paraffin-embedded tissue and fresh frozen tissue and were able to detect plaques and CAA in humans in addition to plaques and intraneuronal Aβ in mice (Bouter et al., 2015). This goes to show the importance of structural studies of antibody-Aβ recognition, as IHC, though an important basis for diagnostic pathology, might not reflect antibody efficacy or action *in vivo*.

Issues with antibody-target engagement in terms of affinity, species and conformation of Aβ bound may be dealt with before going to a clinical setting, with the help of information arising from the crystal structure of the NT4X Fab: Aβ*4-x* complex.
5. Summary & Conclusions

The NT4X antibody has been raised against the human Aβ_{4-40} peptide. We present here its characterization data as an antibody which recognizes both Aβ_{4-x} and Aβ_{pE3-x} N-truncated peptides in reducing and native conditions in PAGE where the NT4X is used as a primary antibody. Further analysis demonstrates that the phenylalanine at position 4 of Aβ is essential for binding its target. Having previously shown that the Aβ_{4-42} peptide is toxic in vitro and in vivo, we now demonstrate that the NT4X antibody is able to rescue the peptide’s toxicity.

In this study, we have endeavored to present a new target for Aβ passive immunotherapy. In an AD mouse model, the Tg4-42, Aβ_{4-42} expression leads to a progressive neuronal loss in the CA1 region. This is accompanied by a gradual loss of spatial reference memory from moderate (5 months) to severe at 6 months of age, in the homozygous line. This short timeframe allowed us to perform a preventative therapeutic passive immunization trials do commence before neuronal loss and reference memory deficits begin and to stop when they are severe. NT4X and its Fab fragment demonstrated a significant mitigation of neuron loss and rescue of spatial reference memory deficits in this treatment paradigm. In this regards, we also present neuronal loss in the Tg4-42 model as an alternative, more relevant, measurable clinical outcome.

In light of failures in translation from pre-clinical studies to the clinic, our work has strived to present a wholesome picture of the therapeutic tool being used in complex to its target. We have been successful in co-crystallizing the NT4X Fab and the Aβ_{4-19} peptide. Our collaborators at the Macromolecular crystallography group (Max Planck for Biophysical Chemistry) are currently working on solving the crystal structure of the NT4X Fab:Aβ_{4-19} complex. Subsequent information may
be used to support the humanization process of the NT4X currently underway at MRC Technology.
CHAPTER 6

6. Bibliography


Biological Sciences, 90(3), 104–17.


affect cultured neuron and astrocyte survival. *Journal of Neurochemistry, 82*(6), 1480–9.


Turner, P. R., O’Connor, K., Tate, W. P., & Abraham, W. C. (2003). Roles of amyloid precursor protein and its fragments in regulating neural


Yamada, K., Yabuki, C., Seubert, P., Schenk, D., Hori, Y., Ohtsuki, S., ... Iwatsubo, T. (2009). Abeta immunotherapy: intracerebral sequestration...


7. **Curriculum Vitae**

**Gregory Antonios**

**Personal information**

Address
Föhrenweg 1, 37077 Göttingen, Germany

Telephone(s)
+49 0551 40153408

Mobile
+49 01525 5691510

E-mail
gregory.antonios@gmail.com

**Occupational field**

**Molecular Medicine**

**Work experience**

**October 2012 – March 2016**

**Position held**
PhD candidate in Molecular Medicine

**Employer**
Principal Investigator: Prof. Dr. Thomas Bayer
Department of Molecular Psychiatry, University Medical Center, Göttingen, (Germany)

**Main activities and responsibilities**

Lab interests: Alzheimer’s Disease molecular mechanisms & mouse models

Project: Generation and characterization of NT4X-167 monoclonal antibody
- Characterized NT4X-167 antibody in vitro binding properties
- Performed in vivo and in vitro toxicity assays analysis

Project: Passive immunization of Tg4-42 murine line with NT4X-167 antibody
- Established and performed preventative immunization of Alzheimer’s disease mouse model
- Performed Behavioral & Biochemical analysis
- Designed and implemented Unbiased Stereology for Neuronal counting

**Dates**
March 2013 – March 2016

**Position held**
Research Fellow

**Collaborator**
Principal Investigator: Dr. Vladimir Pena
Macromolecular Crystallography, Max Planck Institute for Biophysical Chemistry (Germany)

**Main activities and responsibilities**

Lab interests: Structural basis of spliceosomal assembly, remodelling and catalysis at atomic level
Projects: NT4X Fab fragment generation and crystallization with target amyloid beta peptide
- Established large-scale culture and production of NT4X-167 antibody
- Designed purification protocol for antibody Fab fragment
- Implemented and designed crystallization of NT4X Fab bound to target Aβ4-19 peptide
- Performed antibody stability analysis
- Collected crystal diffraction data at the Swiss Light Source, Switzerland
- Obtained European radiation passport
Curriculum vitae of Gregory Antonios

Dates: July 2011 – August 2012

Position held: Research Assistant in Molecular Diagnostics Lab

Employer: Principal Investigator: Dr. Rami Mahfouz; Co-Investigator: Dr. Soha Yazbek
American University of Beirut - Medical Center (AUBMC), Beirut (Lebanon)

Main activities and responsibilities:
Lab interests: Genetically Inherited Diseases

- Microarray expression profiling of samples from patients clinically and genetically identified with FMF
- Collection of patient samples and consent forms

Dates: July 2011 - September 2012

Position held: Research Assistant in Proteomics Lab

Employer: Director: Dr. Brigitte Wex
Lebanese American University (LAU), Byblos (Lebanon)

Main activities and responsibilities:
Lab interests: Protein Identification and Sequencing through Matrix Assisted Laser Desorption Ionization (MALDI) Mass Spectrometry

Project: “Effect of Solute Receptor SLC35b4 on cellular O-Glycosylation and its association with glucose production” through protein knock-down (HepG2 cells):
- Purification of glycosylated protein
- 2D gel electrophoresis and protein extraction
- Quantification and MALDI analysis

Teaching

Dates: August 2013 – Present

Position Held: Teaching and supervision

Employer: Universitätsmedizin Göttingen

Responsibilities:
Clinical Neurosciences and Higher Brain Functions
- Neuroscience Graduate program course
- Presented and elucidated molecular aspects of neurodegenerative dementias
- Prepared and corrected student exams

Behavioral and neuropathological analysis of transgenic mouse models:
- GGNB Graduate program course
- Prepared and gave lab module in immunohistochemistry

Masters and rotation students:
- Planned long-term projects
- Supervised day-to-day learning and progress

Dates: October 2009 - June 2011

Position held: Teaching Assistant

Employer: Lebanese American University, Byblos (Lebanon)

Activities and Responsibilities:
Genomics Research Methods Graduate course, Spring 2011:
- Preparation and Assistance in the three Lab Modules
- Molecular Biology, Microbiology & Cell Biology

Introduction to Biology Undergraduate lab, Fall 2010:
- Preparation of Lab Experiments & Exams, and Teaching
Education

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Skills and competences

Organisational skills and competences
- Establishment and follow-up on numerous national and international collaborations
- Organization of and follow-up on stock availability of lab kits, chemicals, reagents & consumables
- Scheduling and follow-up on technical staff timetable and experimental workload
- Organization of Conferences and Workshops

Technical skills and competences
- **Molecular Biology**: PCR, Real-time PCR, Sequencing, PFGE, Transformation, Restriction digestion, DNA & plasmid extraction
- **Cell Biology**: Cell culture & maintenance, Overexpression and knock-down of proteins, DNA/RNA/Protein extraction from fresh and FFPE tissues, Western blot and protein quantification, Immunohistochemistry and imaging of cells & tissues, Live cell imaging, Wound-healing assays, Protein preps
- **Microbiology**: API, Antimicrobial testing, Bacterial isolation, enrichment, culturing & cryobanking
- **Proteomics**: 2D electrophoresis, protein purification and MALDI data acquisition,
- **Crystallography**: Antibody purification, crystallization, and diffraction data collection

Computer skills and competences
- Bioinformatics: Proficient in NCBI, UniProt, Mascot server
- Software: Image J, Zeiss Imaging, GraphPad Prism 5, and Microsoft Word, Excel, PowerPoint

Languages
- **Lebanese Arabic** (mother tongue)

Self-assessment

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<td><strong>Writing</strong></td>
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Artistic skills and hobbies
- Photography, Cooking, Hiking
Publications:  

Conferences & Workshops:  
“Arbeitsgemeinschaft für Neuropsychopharmakologie und Pharmakopsychiatrie”, Symposium, München, September 2015  
“12th International Conference On Alzheimer’s & Parkinson’s Diseases” Nice, France March, 2015  
“Ageing Brain” Workshop, Göttingen, September 2014  
“Alzheimer’s Disease Research from Different Perspectives”, Workshop: Society for Neuroscience, San Diego, November 2013  
“LAAS 18th International Science Meeting: New Discoveries in Science” Conference, NDU, March 2012  
“Green Economy in a Changing Arab World” AFED Annual Conference, Beirut, October 2011  
“Enhancing Graduate Research in the Biomedical Sciences” Conference, AUST, June 2011  
“Advanced Microscopy in Cell Biology” Workshop, LAU - Byblos & AUB, May 2011  
“4th Poster Conference in Biology & Biomedical Sciences”, LAU - Byblos, May 2011  
“5th Environmental Symposium of German-Arab-Scientific Forum for Environmental Studies”, LAU - Byblos, Sep. 2010  

References:  
• Thomas A. Bayer, Prof. Dr. rer. nat. (PhD Advisor)  
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Tel: 0551-201-1046  
Email: Vlad.Pena@mpibpc.mpg.de  

• Oliver Wirths, PhD (Molekulare Psychiatrie, Lab Manager)  
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Tel: 0551-201-10290  
Email: Oliver.Wirths@medizin.uni-goettingen.de