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### RESEARCH



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# In vivo imaging reveals sigmoidal growth kinetic of $\beta$ -amyloid plaques

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### Abstract

A major neuropathological hallmark of Alzheimer's disease is the deposition of amyloid plaques in the brains of affected individuals. Amyloid plaques mainly consist of fibrillar  $\beta$ -amyloid, which is a cleavage product of the amyloid precursor protein. The amyloid-cascade-hypothesis postulates A $\beta$  accumulation as the central event in initiating a toxic cascade leading to Alzheimer's disease pathology and, ultimately, loss of cognitive function. We studied the kinetics of  $\beta$ -amyloid deposition in Tg2576 mice, which overexpress human amyloid precursor protein with the Swedish mutation. Utilizing long-term two-photon imaging we were able to observe the entire kinetics of plaque growth *in vivo*. Essentially, we observed that plaque growth follows a sigmoid-shaped curve comprising a cubic growth phase, followed by saturation. In contrast, plaque density kinetics exhibited an asymptotic progression. Taking into account the fact that a critical concentration of A $\beta$  is required to seed new plaques, we can propose the following kinetic model of  $\beta$ -amyloid deposition *in vivo*. In the early cubic phase, plaque growth is not limited by A $\beta$  concentration and plaque density increases very fast. During the transition phase, plaque density stabilizes whereas plaque volume increases strongly reflecting a robust growth of the plaques. In the late asymptotic phase, A $\beta$  peptide production becomes rate-limiting for plaque growth. In conclusion, the present study offers a direct link between *in vitro* and *in vivo* studies facilitating the translation of A $\beta$ -lowering strategies from laboratory models to patients.

Keywords: Alzheimer's disease, β-amyloid, Aggregation kinetics, Plaque growth, Two-photon imaging, In vivo

### Introduction

Alzheimer's disease (AD) is the most common form of dementia and is characterized by the accumulation of amyloid plaques as one major neuropathological hallmark [1]. Amyloid plaques consist mainly of A $\beta$  peptide (A $\beta$ ) that is produced by sequential cleavage of amyloid precursor protein (APP) by  $\beta$ -secretase and  $\gamma$ -secretase [2]. The amyloidcascade-hypothesis postulates that A $\beta$  accumulation is the central event in the etiology of AD, initiating a toxic cascade that culminates in widespread neurodegeneration [3-6]. Indeed, recent biomarker studies in humans have shown that changes in A $\beta$  metabolism and A $\beta$  accumulation initiate about 20 years before the onset of clinical symptoms [7-10].



In recent years several studies investigated the kinetics of A $\beta$  aggregation into amyloid plaques in AD mouse models [15]. Collectively, these studies revealed that newly formed plaques were initially very small and enlarged slowly over long periods of time during the early stages of amyloid pathology [16-20], while late-stage amyloid pathology lacked further plaque formation [17,19,21] and growth [16,17,20,22].

In the present study we aimed at extending this basic knowledge of *in vivo* plaque growth kinetics by investigating the close relationship between plaque density and plaque growth. Furthermore, we were able to monitor the complete plaque growth kinetics over an observation



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period of 15 months that reflect both the early and late stage amyloid pathology. In this manner, we found that plaque growth follows an initial cubic and a late asymptotic phase.

### Materials and methods

### Transgenic mice

Heterozygous Tg2576 [9] mice (B6;SJL-Tg(APPSWE) 2576Kha from Taconic, Cologne, Germany) were kindly provided by neuroanatomy group of Abbott (Ludwigshafen, Germany). Tg2576 mice express a human APP with the Swedish mutation (K670M/N671L) under a hamster prion protein promoter. Heterozygous APPPS1 mice coexpress a human APP with the Swedish mutation and a mutated PS1 (L166P) under the neuron-specific Thy1-promoter [23]. Mice were of both sexes (four females in the young cohort and one male/three females in the old cohort) and group-housed under pathogen-free conditions until surgery, after which they were single-housed. All procedures were performed in accordance with an animal protocol approved by the University of Munich and the Government of Upper Bavaria [Az. 55.2-1.54-2531-110-06].

### Cranial window surgery

A cranial window was implanted over the right cortical hemisphere as previously reported [24-27]. In short, the mice were anesthetized with an intraperitoneal injection of ketamine/xylazine (0.13/0.01 mg/g body weight; WDT/ Bayer Health Care, Garbsen/Leverkusen, Germany). Additionally, dexamethasone (0.02 ml at 4 mg/ml; Sigma) was intraperitoneally administered immediately before surgery [28]. A circular piece of the skull (about 5 mm in diameter) over the right hemisphere (centered over the parietal bone, approx. 5.5 mm caudal from the bregma and 5.5 mm lateral from midline) was removed using a dental drill (Schick-Technikmaster C1; Pluradent; Offenbach, Germany). The craniotomy was closed immediately with a round coverslip (5 mm in diameter), held with dental acrylic. A small metal bar, containing a hole for a screw, was glued next to the coverslip to allow repositioning of the mouse during subsequent imaging sessions. After surgery, mice received subcutaneous analgesic treatment with carprophen (7.5 mg/kg body weight; Rimadyl; Pfizer, New York, USA) and antibiotic treatment with cefotaxim (0.25 mg/g body weight; Pharmore, Ibbenbüren, Germany).

### Long-term two-photon in vivo imaging

Imaging started 3 to 4 weeks after the cranial window preparation to allow the animals to recover from surgery. For amyloid staining methoxy-X04 [29] was intraperitoneally injected 24 h before imaging. Initially, we administered a loading dose of 2 mg/kg body weight and in subsequent weekly imaging sessions a maintenance dose of 0.4 mg/kg body weight [17]. Two-photon imaging was performed on a LSM 7 MP (Zeiss, Jena, Germany) equipped with standard photomultiplier detectors and a 20x water-immersion objective (W Plan-Apochromat 20x/1.0 DIC, 1.0 NA, Zeiss, Jena, Germany). Methoxy-X04 was excited at 750 nm by a Ti:Sa laser (MaiTai DeepSee, Spectra-Physics, Darmstadt, Germany) and emission was collected from 440 to 500 nm. Image stacks of  $850 \times 850 \times 250-400 \ \mu m^3$ were acquired using the "tilescan" mode of the microscope control software (Zen2009/Zen2010 64bit) that performs automatic stitching of several fields of view (2×2 were used) with a lateral resolution of 0.83 µm and 3 µm separation distance between consecutive images. Mice were anesthetized with isoflurane (Forene®, Abbott, Wiesbaden, Germany) for imaging and fixed to a custom-made holder using the glued metal plate. In subsequent imaging sessions, previously imaged volumes were identified by eye using the unique blood vessel pattern and fine adjusted by the positions of preexisting plaques. This allowed a precise alignment of the same imaging volume over a period of up to 15 months. The laser intensity was adjusted to keep the emitted fluorescence stable at different depths using the zcorrection tool in the microscope control software and also at subsequent imaging sessions.

The following volumes were imaged: young cohort 2.888 mm<sup>3</sup> (12 positions), old cohort 3.111 mm<sup>3</sup> (16 positions), 12 to 18 months 1.076 mm<sup>3</sup> (4 positions) and 12 to 27.5 months 0.559 mm<sup>3</sup> (2 positions). The following numbers of plaques were analyzed for each imaging cohort: 50 newly formed and 101 preexisting plaques (12 to 14 months), 786 plaques (18 to 20 months) and 90 plaques (12 to 27.5 months).

### Image processing

The images with data depth of 12 bit were analyzed as time series of three-dimensional (3D) images in Imaris (Versions 6.2.1/7.4.2, Bitplane, Zurich, Switzerland). First, images were contrast-normalized (i.e., based on the average and standard deviation of intensities of 3D stacks). Plaque volumes were extracted by 3D-surface-rendering with background subtraction and a threshold of 500. All detected plaques were tracked by utilizing a custom-written Matlab plugin for Imaris or the surface tracking module of Imaris. Newly formed plaques were tracked back to the first time point when they appeared and were only assessed when present for at least 3 time points. Kinetic volume data were exported as Excel files. 3D stacks are either displayed as 3D-volume-rendered images ("normal shading" algorithm, Imaris) or as 3D-surface-rendered images in figures. Only the image data of detected plaques are shown in figures to focus the viewer's eye on the essential information within the images. Background signals from autofluorescence were removed whereas contrast settings within time series were kept constant.

### Data analysis and statistics

For each plaque the radii from each time point were calculated from the kinetic volume data assuming a spherical shape of plaques [19]. Plaque densities were calculated for each imaged position. All other calculations, curve fitting and graphs were done in Prism (Version 5.04, GraphPad, La Jolla, USA). Data were tested for normality using D'Agostino-Pearson omnibus K2 test. Plaque densities and sizes had log-normal distributions and were therefore logarithmized which allowed for parametric statistical testing. These data were displayed with an "antilog" scale to represent the real measurement dimensions. For each plaque a linear growth rate was determined as the slope of a linear regression from radii calculations over time. Linear plaque growth rates were not normally distributed. All statistical tests are specified in the figure legends.

### Results

## Long-term in vivo imaging over 15 months reveals sigmoid-shaped plaque growth kinetics

Tg2576 mice express human amyloid precursor protein with the Swedish mutation under the control of a hamster prion protein promoter [9], leading to the accumulation of A $\beta$ -peptide and deposition of amyloid plaques starting at 8 to 10 months of age [9,30]. The fluorescent marker methoxy-X04 was injected [29] to visualize fibrillar amyloid through open-skull cranial windows over long time periods. We imaged two different cohorts starting at 12 and 18 months of age over at least 2 months, in weekly intervals (Figure 1a). These time-frames were chosen since plaque development is very dynamic in the young cohort whilst being more static in the older mice [17,22]. The mice from the young cohort were imaged for as long as technically feasible, limited by the clarity of the cranial window, in order to reconcile the measurements within both age cohorts in a single time series (Figure 1a). For precise measurement of the plaque density and in order to maximize the number of analyzable plaques, big volumes that typically sized 850  $\times$  850  $\times$  250–400  $\mu$ m<sup>3</sup> (x, y, z dimensions) were scanned to 27.5 months (Figure 1b; see Additional file 1 for the complete time series and the Material and Methods section for all integrated volumes of each cohort). All images were of optimal quality and did not suffer from motion artifacts due to breathing or heart beating of the animal. The volume of plaques and cerebral amyloid angiopathy (CAA) were assessed by 3D surface-rendering and tracked over time. This analysis revealed a nonlinear sigmoid-shaped curve for the mean ( $\mathbb{R}^2$  0.983, Figure 1c) and also integrated volume ( $\mathbb{R}^2$  0.988, Figure 1d) of plaques. The integrated volume of the CAA followed a sigmoid function, too ( $\mathbb{R}^2$  0.952, Figure 1e). In contrast, the plaque density kinetics exhibited an asymptotic progression and could be fitted to a one-phase association function  $(\mathbb{R}^2)$ 0.975, Figure 1d).

Next, we analyzed the growth kinetics for each individual plaque separately, rather than the mean of all plaques as previously (Figure 2). We found that single plaque kinetics could also be fitted into a sigmoid function (Figure 2a,b). A comparison of all fitted curves for each individual plaque (originating from the example in Figure 1b) highlighted the fact that only a few plaques became very large whereas the majority of plaques showed much flatter curves with slower A $\beta$  accumulation (Figure 2c).

### Quantification of plaque densities, sizes and growth rates in young and aged mice

The differences in plaque densities, sizes and growth rates between the young and old age cohorts are summarized in Figure 3. Mean plaque density started very low in the young cohort at 26.2 mm<sup>-3</sup> (13.8-49.8 mm<sup>-3</sup> CI), but increased to 42.0 mm<sup>-3</sup> (23.2-75.7 mm<sup>-3</sup> CI) during the 2-month observation period (Figure 3a,b). This robust increase in plaque density was attributed to 50 newly formed plaques that were added to the 101 already-existing plagues at the beginning of the study. In contrast, within the old cohort only 3 new plaques were observed alongside 786 preexisting plaques. As expected, mean plaque density was significantly higher at 18 months of age  $(191 \text{ mm}^{-3})$ ; 126-292 mm<sup>-3</sup> CI) compared to 12 and 14 months of age (Figure 3b). Long-term imaging revealed no further significant increase in mean plaque density, which peaked at a plateau of 160 mm<sup>-3</sup> (100-255 mm<sup>-3</sup> CI) at 24 months of age (Figure 3c, solid line).

The size of a plaque can be represented by its radius, calculated from the measured volume assuming a spherical plaque shape to facilitate comparisons [19]. At 12 months, newly-formed plaques had a mean radius of 2.27  $\mu$ m (2.12-2.43  $\mu$ m CI; Figure 3d), while the mean radius of preexisting plaques was 5.08  $\mu$ m (4.45-5.80  $\mu$ m CI). With ageing, mean plaque radii increased further and reached 7.98  $\mu$ m at 14 months (7.31-8.72  $\mu$ m CI), 10.8  $\mu$ m at 18 months (10.4-11.2  $\mu$ m CI) and 16.7  $\mu$ m at 27 months (Figure 3d, 15.0-18.6  $\mu$ m CI). Radii from newly formed plaques showed the narrowest spectrum and were exclusively normally distributed, while sizes from preexisting plaques covered a wide range and had a log-normal distribution (Figure 3e).

A linear regression of plaque radii over time revealed linear growth rates reflected by the slope of the lines [19]. Nascent plaques exhibited the fastest median growth rate at 0.475  $\mu$ m/week (Figure 3f, 0.326-0.589  $\mu$ m/week interquartile range, IQR). Preexisting plaques from the young cohort grew at almost the same rate (0.326  $\mu$ m/week, 0.217-0.479  $\mu$ m/week IQR; Figure 3f), while the old cohort had the slowest growth rate (0.099  $\mu$ m/week, -0.070-0.262  $\mu$ m/week IQR; Figure 3f). Nevertheless, an overall net growth could be detected even in the old cohort (Figure 3g).



were investigated from 12 to 14 and 18 to 20 months of age, respectively, at a weekly imaging intervals. (b) Time series of 3D volume-rendered images acquired with two-photon fluorescence microscopy showing amyloid plaques and cerebral amyloid angiopathy (CAA). The length of one side of the squares equals 100  $\mu$ m. (c) Mean volumes of all plaques over time (open circles). The black line indicates a fitted sigmoid function (R<sup>2</sup> 0.983). Error bars show the 95% confidence intervals (CI). (d) Integrated volume of all plaques over time (open circles). The black line indicates a fitted sigmoid function (R<sup>2</sup> 0.988). In addition, the temporal development of the plaque density is depicted (gray line) which can be fitted to an exponential function of one phase association (R<sup>2</sup> 0.975). (e) Integrated volume of the cerebral amyloid angiopathy (open circles) with the corresponding fit of a sigmoid function (R<sup>2</sup> 0.952).

As the long-term data over 15.5 months displayed both phases (cubic and asymptotic) of growth a bifid analysis of the growth rates was performed to compare them to the young and old cohorts (Figure 4a). The cubic and asymptotic growth phases are divided by the inflection point of the fitted sigmoid function that lies at the age of 23.5 months. Therefore, plaque growth rates were determined from 12 to 23.3 and from 23.5 to 27.5 months of



age for each single plaque (Figure 4b). The linear plaque growth rate was fastest in the cubic phase (median 0.365  $\mu$ m/week, 0.258-0.453  $\mu$ m/week IQR), whereas a strong decline was observed for the asymptotic phase (Figure 4c, median 0.159, 0.049-0.297  $\mu$ m/week IQR).

## Characteristic relationships between quantitative plaque parameters

Relationships between plaque densities, growth rates, size and age at plaque formation were analyzed to enable a detailed characterization of plaque formation and growth. In the early stage of plaque pathology (young cohort) plaque density at the start of imaging correlated with plaque formation rate, the latter being an indicator of the number of plaques formed per volume and time unit (Figure 5a). The average plaque formation rate in the young cohort amounted to 3.40 week<sup>-1</sup> mm<sup>-3</sup> (1.63-5.16 week<sup>-1</sup> mm<sup>-3</sup> CI). A global comparison of all data collected in the young and old cohort revealed an inverse correlation between averaged plaque growth rates and plaque densities (Figure 5b). In addition, analysis of a single plaque over a period of 15.5 months covering the dynamic cubic and the asymptotic phases of plaque development provided further insight into plaque growth characteristics. The largest plaques at the end of imaging had formed early during plaque development, which leads to a strong inverse correlation between plaque radius and age of the mouse at plaque formation (Figure 5c). Moreover, plaque radius at the end of imaging correlated strongly with the corresponding growth rates of these plaques (Figure 5d).

There was a wide distribution of individual plaque growth rates which lead to the question whether there is a specific spatial relationship between growth rates of neighboring plaques. This analysis was performed using data from long-term imaging, the main reason being that a longer observation period facilitates more precise measurements of plaque growth rates. For a visual impression, plaques from overview image of Figure 1b were categorized in five classes according their growth rates. These plaques were displayed as 3D surfacerendered objects with their corresponding growth rate class coded in different colors (Figure 6a). Analysis involved calculating the shortest Euclidean distance of a plaque to its nearest neighbor, which averaged 65.8 µm (55.3-78.4 µm CI; Figure 6b). In the case of a spatial relationship, growth rates of neighboring plaques would be expected to be similar. The analysis of two imaged positions and 46 pairs of plaques (90 plaques total), however, revealed no such relationship (Figure 6c). Moreover, the differences between growth rates of neighboring plaques



Figure 3 Summary or plaque densities, radii and growth rates from all imaging conorts. (a) Representative examples of imaged volumes from the young and old cohort displayed as 3D volume-rendered images. The length of one side of the squares represents 50  $\mu$ m. (b) Mean plaque densities from the young (12 and 14 months) and old cohorts (18 months). (c) The kinetic of plaque density from long-term imaging (mean from 2 positions) is shown as black line. In addition, plaque densities for each imaged position with their respective means from different cohorts are displayed at the corresponding age. (d) The mean plaque radius and their corresponding cumulative frequency distributions (e) are depicted for each imaging cohort including newly formed plaques from young cohort (two month imaging). For the long-term imaging cohort plaque sizes at 27 months are shown. (f) Box plot for linear plaque growth rates and (g) their corresponding cumulative frequency distributions. Plaque growth from 18 to 20 months of age was significant different from zero (P < 0.0001). Error bars show 95% CI (b-d), Whiskers represent 10th and 90th percentile and outliers are not shown (f). Plaque densities and radii were logarithmized before analysis (b-e). Statistical tests: one-way ANOVA with Tukey-Kramer Post-hoc test (b, d), Kruskal-Wallis test with Dunn's Post-hoc test (f), Wilcoxon signed-rank test (f, 18 to 20 months), paired *t*-test (b, 12 vs 14 months). \*\*\* P < 0.0001

showed a wide distribution and reflect the visual impression from Figure 6a.

### Discussion

Novel insights into plaque growth kinetics arose from several *in vivo* studies in recent years. Assessment of basic plaque growth parameters in the present study gave results consistent with the majority of findings from the literature. The most prominent feature is the increase of plaque size during ageing [16-20,31]. Plaques in their nascent stage were smallest and constituted a distinct size class that showed a normal distribution. Preexisting plaques were much larger, depending on the age of the mice, and displayed a log-normal distribution as also found in humans [32]. In contrast, the rate of plaque growth decreased with ageing [16,17,20,22]. Several reasons could

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Figure 5 Correlations between quantitative parameters of plaque growth dynamics. (a) Plaque formation rate as a function of plaque density. Black line, linear regression (slope is significantly different from zero, p = 0.016). Each circle represents one imaged position. (b) Plaque growth rate as a function of plaque density. Error bars show the 95% Cl. Black line shows a linear regression of the data. The slope is statistically significant different from zero (p < 0.001). (c) Radius of newly formed plaques at the end of long-term imaging over 15.5 months as a function of the age of the mouse at plaque formation (N = 81 newly formed plaques from 2 positions). (d) Plaque radius at the end of long-term imaging over 15.5 months as a function of the data. The slope is statistically significant different from zero. Statistical tests: F-test (a, b, d), Spearman correlation (c).



be responsible for the decline in plaque growth: (i)  $A\beta$ production becomes rate limiting, because an everincreasing total surface area of β-amyloid plaques requires an increase in the amount of  $A\beta$  to maintain a constant linear growth rate. (ii) The dissociation of  $A\beta$ aggregates from  $\beta$ -amyloid plaques increases. (iii) The number and volume of microglia around plaques increases and changes plaque maintenance [33]. In addition, the arguments (ii) and (iii) may explain negative plaque growth rates in aged animals, whereas we can not completely exclude that at least in part data noise is responsible for negative values. A recent study proposed clustering of plaques as different mechanism of plaque growth [34]. In some rare instances we also observed fusion of plaques which we found as a consequence of their volume growth in combination with close proximity to each other (see image series Figures 1 and 3, Additional file 1). Furthermore, two studies found no steady plaque growth [17,19,21] or plaque growth and shrinkage [33] in the early phase of plaque development. Possible reasons for the inconsistent findings are discussed within the literature and beyond the scope of this publication [16-20].

For the first time we were able to chart the entire kinetics of growth of a single plaque, and demonstrate a sigmoidal growth curve comprising both a cubic and an asymptotic phase (Figures 1 and 2, Additional file 2: Figure S1a-b). Recent studies in humans using positron emission tomography (PET) applying an amyloid tracer (Pittsburgh compound B, PiB) [10,35,36] also found a plateau and/or a sigmoid-shaped progression of  $\beta$ -amyloid accumulation [7,13]. Long-term imaging over at least 6 months revealed an asymptotic increase in plaque densities that could be fitted to a one-phase association function. This finding is congruent with a previously described decline in the rate of plaque formation with ageing [16-20,31]. Interestingly, plaque growth and plaque density are inter-related, as shown by correlation analysis of both parameters.

During the dynamic phase of plaque development (young cohort) a distinct correlation between plaque density and plaque formation rate was observed. In humans, a similar relationship was found using PiB-PET which revealed a higher increase in PiB retention in the second PET scan after 18 months if those subjects had a positive PiB signal in the first PET scan [37]. Fibrillization of  $A\beta$ has been convincingly shown by in vitro studies to consist of a two-step process that requires first the nucleation of a seed, before polymerization into amyloid fibrils [14,38]. Furthermore, a direct relationship was measured in vitro between  $A\beta$  concentration and the ability to form nucleation seeds [8,39]. For a mouse model of AD, a similar correlation between A<sup>β</sup> concentration in the interstitial/ cerebrospinal fluid and plaque load (a relative measure of the area of a brain slice that is covered by plaques) was reported [40,41]. Taking together the knowledge from these studies with our results, a link between plaque density and AB production can be established since there is a strong correlation between plaque density and plaque load (Additional file 2: Figure S1c). Interestingly, plaque densities varied over a wide range which suggests that  $A\beta$  production varies between individual Tg2576 mice a fact that was already reported

[42]. A possible reason could be the mixed genetic background of Tg2576 mice which may be lead to differences in epigenetic regulation and/or promotor activities [9]. Nevertheless, all 4 animals of the young cohort covered the complete range of plaque densities measured in 30 animals of a further study (Additional file 2: Figure S1d). In addition, the predictive value of plaque density regarding  $A\beta$  production could be limited in mouse models without a constant  $A\beta$  production over time [23]. Another limitation worth noting is that methoxy-X04 in mice, and also PiB in humans, detect insoluble fibrillar but not soluble Aß species [29,36,43,44]. Furthermore, in vivo two-photon imaging using methoxy-X04 allows for the resolution of single amyloid plaques in brain regions accessible by this technique while PET imaging allows to image the whole brain, but with a resolution limit of about 1 mm<sup>3</sup>.

The observation of a steady linear increase in plaque radius over several months during the cubic growth phase presumes that  $A\beta$  concentration is not rate limiting. Otherwise a decline in growth would be expected due to the ever-increasing surface area of  $\beta$ -amyloid plaques which requires a proportional increase in the amount of  $A\beta$  to maintain a constant linear growth rate. A comparison of plaque growth of the Tg2576 with the APPPS1 mouse model that was investigated by two other studies, using exactly the same methods and data analysis, supports this hypothetical assumption [16-20,31]. Although APPPS1 mice produce much more A $\beta$  and accumulation starts much earlier than in Tg2576 mice [9,19,23,30] median plaque growth rates were about 0.3 µm/week in both mouse models [16-20,31]. In conclusion, there appears to be an upper limit for median plaque growth in vivo that was also identified in vitro for the growth of A $\beta$  fibrils [45]. The higher A $\beta$  production in APPPS1 mice causes a much higher plaque density compared to Tg2576 mice (631 mm<sup>-3</sup>, 521-762 mm<sup>-3</sup> CI vs. 26.2 mm<sup>-3</sup>, 13.8-49.8 mm<sup>-3</sup> CI, Additional file 2: Figure S1e, Figure 3b). Accordingly, a 10-fold higher rate of plaque formation was reported for APPPS1 mice [19] compared to Tg2576 mice (35 vs.  $3.40 \text{ week}^{-1} \text{ mm}^{-3}$ ). Although median plaque growth rates were similar in both mouse models, individual plaque growth rates showed a big variance. In addition, plaque growth rates of neighboring plaques did not correlate. Combining both results, it is most likely that different conditions within the microenvironment of plaques are causal for the different growth rates. Such conditions could include pH, ionic strength and hydrophobicity of the A $\beta$  peptide, as well as interactions with membranes [46-49]. Even though the analysis of a single animal is statistically not sufficient, the unique dataset containing a complete plaque growth kinetic further supports this idea. It highlights that large plaques were formed early during A $\beta$  accumulation and have the highest growth rates. This in turn suggests that plaques are first formed at sites where the best conditions exist.

In conclusion, the comprehensive plaque density and growth kinetics point to a three-stage model of  $\beta$ -amyloid accumulation (Figure 7). According to the kinetics of plaque volume three phases can be distinguished: (1) a cubic (2) a transition and (3) a saturation phase. Initially, a high concentration of  $A\beta$  above the critical concentration leads to the formation of many new plaques (cubic phase). Later, the large number of plaques may serve as "dumps" for free A $\beta$ , so that its concentration gradually drops, causing the formation of fewer plaques during the transition phase, while existing plaques continue growing. In the saturation phase, single plaques show an overall slower growth due to the large number of plaques, while no new plaques are deposited, thus AB production becomes rate-limiting. Furthermore, due to the same sigmoid-shaped characteristic of plaque accumulation in humans, albeit over one to two decades, such a model may help extrapolate observations from AD mouse models to the situation in humans, particularly with regards to pre-clinical testing of  $A\beta$ -lowering therapeutics [7,13]. In particular, the determination of plaque density kinetics may be more sensitive than analyzing plaque volume kinetics when evaluating anti-aggregation agents, since  $A\beta$  concentration is not rate-limiting to plaque growth in the early stage of amyloid deposition.



Figure 7 Model for the relationship between plaque density kinetics and amyloid plaque growth *in vivo*. The term  $c_{critical}$  refers to the minimum critical concentration of A $\beta$  that is necessary to form plaque seeds (a prerequisite discovered by in vitro studies). This initial step of plaque formation was observed by recording the plaque density kinetics *in vivo*. According to the plaque density kinetics and applying the aforementioned requirement different relations of the A $\beta$  concentrations comparative to the critical minimum concentration can be assigned to the different growth phases.

### **Additional files**

**Additional file 1:** This video shows the complete plaque growth **kinetic from 12 to 27.5 months of age (image data from Figure 1b).** Data is displayed as 3D-volume rendered images. Big squares represent 100 μm.

Additional file 2: Figure S1. (a) Plaque volume data of two additional mice over 6 months. Mean plaque volumes were best fit with either a sigmoid function (mouse1: open triangles) or a cubic function (mouse2: open circles). Error bars show 95% CI. (b) Kinetics of the plaque densities (dotted lines) and mean volumes (colored lines showing nonlinear fitted functions) over time for all 3 mice imaged at least over a period of 6 months. Vertical dotted lines indicate time points when plaque densities did not further increase. (c) Relationship of amyloid plaque load (integrated plague volume/imaging volume) and plague density for each imaged position. Linear regression was performed for each age cohort. The slopes were statistically significant different from zero (F-test). (d) Comparison of the plaque densities between the young cohort of the current study and another group of mice of a further study. Line and error bars display mean with 95% CI. The means were not significant different (t-test). (e) Plaque density measured in APPPS1 mice at 4 months of age. Line and error bars display mean with 95% Cl. The equations were used for curve fitting.

### **Competing interests**

The authors declare that they have no competing interests.

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