Amyloid plaque formation precedes dendritic spine loss

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Abstract Amyloid-beta plaque deposition represents a major neuropathological hallmark of Alzheimer’s disease. While numerous studies have described dendritic spine loss in proximity to plaques, much less is known about the kinetics of these processes. In particular, the question as to whether synapse loss precedes or follows plaque formation remains unanswered. To address this question, and to learn more about the underlying kinetics, we simultaneously imaged amyloid plaque deposition and dendritic spine loss by applying two-photon in vivo microscopy through a cranial window in double transgenic APPPS1 mice. As a result, we first observed that the rate of dendritic spine loss in proximity to plaques is the same in both young and aged animals. However, plaque size only increased significantly in the young cohort, indicating that spine loss persists even many months after initial plaque appearance. Tracking the fate of individual spines revealed that net spine loss is caused by increased spine elimination, with the rate of spine formation remaining constant. Imaging of dendritic spines before and during plaque formation demonstrated that spine loss around plaques commences at least 4 weeks after initial plaque formation. In conclusion, spine loss occurs, shortly but with a significant time delay, after the birth of new plaques, and persists in the vicinity of amyloid plaques over many months. These findings hence give further hope to the possibility that there is a therapeutic window between initial amyloid plaque deposition and the onset of structural damage at spines.

Keywords Alzheimer’s disease · Two-photon in vivo imaging · Dendritic spines · Amyloid plaque · Kinetics · Structural plasticity

Introduction

Alzheimer’s disease (AD) is the most common form of dementia, which is characterized by two neuropathological hallmarks, namely β-amyloid plaque deposition and intracellular neurofibrillary tangles. The progression of disease pathology is further accompanied by a substantial loss of neurons and synapses. Controversial findings have, however,
been reported regarding the relationship of synapse loss and plaque load [3, 33]. For instance, synapse loss has been reported as an early event in the brains of AD patients and it represents the pathological feature that best correlates with cognitive impairment [11, 23, 26, 39, 47, 48, 56]. Indeed, AD pathology includes a substantial decrease in the number of dendritic spines [16, 33, 44], which are thought to be a primary sign of cognitive impairment in AD patients and amyloid plaque load has also been extensively investigated in the absence of plaques [1, 6, 28, 35, 45] as well as in their vicinity, in various AD mouse models [6, 19, 32, 34, 40, 53, 54, 59]. However, little is known about the underlying kinetics of these processes. For instance, Tsai et al. [59] reported an increased spine elimination and formation over 4–5 weeks, whereas Spires-Jones et al. only found a significant increase in spine elimination over 1 h [37, 53]. To shed more light on the chronology of these events, we investigated dendritic spines over several weeks in a well-characterized AD mouse model [7, 22, 46]. We focused on two age cohorts: (1) 3–4 months of age, when amyloid pathology is still dynamically unfolding, and (2) 18–19 months, which represents the phase of late-stage pathology. The kinetics of dendritic spine formation and elimination were analyzed in the vicinity and further away from plaques in AD transgenic mice as well as in control animals within somatosensory cortex. At the same time, we monitored the plaque growth kinetics. This combined approach enabled, for the first time to the best of our knowledge, to image individual dendritic spines before and during the appearance of amyloid plaques and hence correlate their respective kinetics. A similar approach has recently been applied to investigate the sequence of neuritic and glial changes [41, 52].

Materials and methods

Transgenic mice

APPs1 mice are double transgenic for APPKM_670/671NL and PS1L_166P mutations [46]. We want to thank Matthias Jucker from the University of Tübingen and German Center for Neurodegenerative Diseases (DZNE), Tübingen, Germany, who kindly provided these mice. Heterozygous mice of this line were crossed with mice heterozygous for YFP-H [15] (B6.Cg-Tg(Thy1-YFP)2Jrs/J from The Jackson Laboratory, Bar Harbor, USA). Heterozygous triple transgenic offspring of mixed gender were used in the experiments. Single transgenic YFP-H littermates were used as controls. Mice were group-housed under pathogen-free conditions until surgery, after which they were singly housed. All procedures were 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 over the right cortical hemisphere was surgically implanted as previously described [5, 17]. The mice were anesthetized with an intraperitoneal injection of ketamine/xylazine (0.13/0.01 mg g⁻¹ body weight). Additionally, dexamethasone (0.02 ml at 4 mg ml⁻¹) was intra-peritoneally administered immediately before surgery [25]. A circular piece of the skull over the somatosensory cortex (4 mm in diameter) was removed using a dental drill (Schick-Technikmaster C1; Pluradent; Offenbach, Germany). This was immediately covered with a circular glass coverslip (5 mm in diameter), which was glued to the skull using dental acrylic (Cyano-Veneer fast; Heinrich Schein Dental Depot, Munich, Germany) to close the craniotomy. 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 a subcutaneous analgesic dose of carprofen (Rimadyl; Pfizer, New York, NY, USA) for 3 days (5 mg kg⁻¹). Imaging began following a 21-day recovery period after surgery.

Long-term two-photon in vivo imaging

Long-term two-photon in vivo imaging was performed as previously described [5, 18, 29]. Less than 50 mW of laser power was delivered to the tissue to avoid laser-induced phototoxicity. For amyloid plaque staining, methoxy-X04 [31] (1 mg/kg) was intraperitoneally injected 24 h before imaging. YFP and methoxy-X04 were excited by a Ti:Sa laser (MaiTai, Spectra-Physics, Darmstadt, Germany) at 880 and 750 nm and the emission was collected from 527 to 582 nm and 460 to 500 nm, respectively (LSM 7 MP, Zeiss, Jena, Germany). For overview images, z-stacks of 230 × 230 × 150 μm³ with a 2 μm z-resolution and 1024 × 1024 pixels per image frame (0.22 μm/pixel) were taken with a 40× IR-Achroplan water immersion objective (0.8 NA, Zeiss, Jena, Germany) to analyze amyloid plaques. For higher-resolution images to count dendritic spines, the...
same objective was used with 1 μm z-resolution and 512 × 512 pixels per image frame (0.11 μm/pixel). The high-resolution (2048 × 2048 pixels with 0.14 μm/pixel) and large volume (283 × 283 × 300 μm³) images with 2 μm z-resolution to monitor dendritic spines before and after plaque deposition were taken with a 20× W Plan-Apochromat water immersion objective (1.0 NA, Zeiss, Germany).

Image processing and data analysis

All images were deconvolved using the adaptive blind 3D deconvolution algorithm of AutoDeblur with ten iterations (Version x2.0.1, Media Cybernetics Inc., Bethesda, MD, USA). The images were maximum intensity projected (Imaris 6.1, Bitplane, Zurich, Switzerland). In some figures, distracting neighboring dendritic elements were removed. Spines were counted in z-stacks by manually scrolling through the images of subsequent time points of the same position. The spine scoring method has previously been described [5, 18, 25]. Spine densities refer to the amount of spines per dendrite length in μm from which they protrude. Spine densities investigated on dendrites closer than 50 μm from a plaque were only analyzed on the segment that is located within the 50 μm radius from the plaque border. Spine morphology classification was performed by using the filament tracer module for 3D reconstruction in Imaris software (Version 7.4.2, Bitplane, Zurich, Switzerland). The volume measurements were done with the following parameters: small diameter 0.112 μm, large diameter 2 μm, and contrast threshold 0.3. Dendritic spines were classified due to their morphology into thin, stubby, and mushroom spines [21]. Filopodia spines were not found in the investigated brain region at 18–19 months of age. The geometrical classification rules from Harris et al. were interpreted by the following hierarchical expressions in the Imaris XT spine classification module: mushroom spines = “max_width (head)/min_width(neck) > 1.4 and max_width(head) > 0.5 and min_width(neck) > 0”; stubby spines = “length(spine)/mean_width (neck) ≤ 3 or min_width (neck) = 0 or min_width (neck) > 0.5”; thin spines = “length(spine)/mean_width (neck) > 3” [29]. The volume of amyloid plaques was automatically calculated using Imaris software (Version 6.2, Bitplane, Zurich, Switzerland) as previously described [8]. The size of new born plaques was directly presented as volume (Fig. 5), whereas for pre-existing plaques that were already present at the first imaging time point the radius was calculated assuming a spherical plaque shape [22]. All data are presented as mean ± SD or ±SEM or 95 % confidence intervals (95 % CI). Error types are stated where appropriate. Statistical differences in measurements over time were determined using repeated-measures ANOVA while statistical comparison between two groups was performed with Student’s t test. Multiple group comparison was done by one-way ANOVA followed by Tukey–Kramer post hoc test. The slope from a linear regression was tested for statistical difference from zero by F test. Distributions of morphological subtypes of dendritic spines were tested for differences by Chi-square test. All statistical analysis and graphs were done using Prism 5.04 (GraphPad Software Inc., La Jolla, CA, USA). Figures were arranged using Adobe Illustrator CS4.

Results

In the present study, we used the APPPS1 mouse model, which co-expresses human amyloid precursor protein with the Swedish mutation (KM 670/671 NL) and L166P mutated presenilin 1 [46]. In these mice, the first amyloid plaques become visible before the age of 2 months and amyloid plaque load reaches almost 10 % at 8 months within the cerebral cortex [46]. These features make APPPS1 mice an excellent model to study amyloid plaque-related AD pathology. In order to investigate synapse pathology in the vicinity of plaques, these mice were intercrossed with a mouse line that expresses yellow fluorescent protein in a subset of layer III and V pyramidal neurons within the cortex (YFP-H line) [15]. Amyloid plaques were stained by the fluorescent dye methoxy-X04 [31], which crosses the blood–brain barrier and allows to study plaque growth kinetics by repeated injection before every imaging session. The implantation of an open-skull cranial window above the somatosensory cortex provides direct optical access to the brain and deep tissue imaging using two-photon microscopy can be performed over several weeks [12, 24, 55]. This experimental setup set the stage to co-investigate and correlate dendritic spine kinetics with plaque growth kinetics in two separate age groups (Fig. 1). The first group ranged from 3 to 4 months, an age when plaque growth is still very dynamic in this mouse model [22]. The second age group (18–19 months) represents the end-stage of amyloid pathology. In both groups, a single time-point analysis revealed a significant lower spine density in the vicinity (<50 μm) of amyloid plaques compared to areas distant to plaques (>50 μm) with respect to wild-type control animals (Fig. 2a, 3 months: 0.470 ± 0.084 μm⁻¹ control, 0.511 ± 0.075 μm⁻¹ > 50 μm from plaque, 0.264 ± 0.092 μm⁻¹ < 50 μm from plaque, p < 0.05; Fig. 2c, 18 months: 0.467 ± 0.081 μm⁻¹ control, 0.466 ± 0.034 μm⁻¹ > 50 μm from plaque, 0.238 ± 0.086 μm⁻¹ < 50 μm from plaque, p < 0.01, one-way ANOVA with Tukey–Kramer post hoc test, errors represent SD). The dendritic spine density in the vicinity of amyloid plaques did not significantly differ between age groups (Supplementary Fig. a). It is important to mention that we were actively looking for spine-bearing dendrites to
analyze in the surrounding volume of amyloid plaques, which were subjectively more difficult to find, yet still present, in the older cohort. During the same time interval, plaque size increased by almost threefold, from a mean radius of 5.413 μm at 3 months to 13.410 μm at 18 months (Fig. 2b, 95 % CI 4.980–5.846 μm and 11.750–15.060 μm, p < 0.0001, t test).

After this single time-point analysis, we focused on the kinetics of amyloid plaque growth and spine loss over 4 weeks. Therefore, the spine density at mature plaques and at nascent plaques was analyzed in young and old cohorts. Within the peri-plaque region (<50 μm around plaques) we found a significant reduction in spine density in both young and aged mice (Fig. 3a, b 57.9 ± 28.3 % at 3 months (n = 80) compared to 18 months (n = 41) of age (p < 0.0001, t test).
4 months and Fig. 4a, b 67.8 ± 20.8 % at 19 months, \( p < 0.001 \), repeated measures ANOVA with Tukey–Kramer post hoc test). Over the same time period, no spine loss was detected >50 μm distant from plaques or in wild-type animals having no plaques at all. Notably, spine loss kinetics were not significantly different between both age groups (Supplementary Fig. c). Long-term in vivo imaging makes it possible to follow individual spines over time and to determine the fraction of lost and gained spines. Consequently, we could identify the factor responsible for a net decrease in spine density. Such an analysis revealed that the overall decline in spine density could be attributed to a relative increase in the fraction of lost spines over 4 weeks. 

The fraction of gained spines, on the other hand, remained the same when compared to areas distant to plaques or in control animals in both age groups (Figs. 3c, 4c, \( p < 0.001 \), one-way ANOVA with Tukey–Kramer post hoc test). In other words, the observed loss of spines at pre-existing plaques is caused by an increase in spine elimination rather than by a malfunction in the process to form new spines. 

We further investigated whether a specific morphological subtype of dendritic spine is lost in the vicinity of amyloid plaques. With this aim, we classified dendritic spines into “thin”, “stubby”, and “mushroom” spines based on their morphology [21] (Fig. 5a). For every dendrite we analyzed the first and last imaging time-point, 4 weeks later.
We expected to see differences, if any, in the old cohort at 18–19 months of age, since pathology here is in an advanced stage. However, we found no significant differences in each morphological category over time (Fig. 5b, c, Chi-square test). There were also no differences in the distribution of the three dendritic spine subtypes between the control and in the vicinity of plaques (Fig. 5b, c, Chi-square test). Thus, we conclude that no specific morphological subtype is preferentially eliminated.

Plaque growth was quantified by converting the cubic root from volumes in 3D images to yield a linear measure, analogous to the radius [22]. In the young cohort, we could confirm a significant growth of $0.320 \pm 0.066 \, \mu m \, week^{-1}$, which had previously been reported by Hefendehl et al. [22] (Fig. 3b, $p < 0.01$, $F$ test, $F = 23.811, DFn = 1, DFd = 7$). In contrast, plaque radius in aged mice remained unchanged (Fig. 4b, $-0.189 \pm 0.084 \, \mu m \, week^{-1}$, $p \geq 0.05$, $F$ test, $F = 4.992, DFn = 1, DFd = 7$). Lack of plaque growth is supported by a similar finding in aged Tg2576 mice, another mouse model of AD [8, 9].

Finally, we analyzed the kinetics of individual dendritic spines before and during the de novo formation of new plaques. The observation of plaque birth is a very rare event [8, 22, 41, 61] and for spine analysis to be feasible in proximity to an appearing plaque, two prerequisites have to be fulfilled. Firstly, plaque birth has to occur in the vicinity of a dendrite lying mainly in the imaging plane (due to...
resolution limits in the axial direction) [12, 55]. Secondly, the resolution of the images has to be high enough to resolve individual dendritic spines over long periods of time. To meet both criteria, 12 volumes of the dimensions $283 \times 283 \times 300 \, \mu m^3$ (totaling $0.288 \, mm^3$) were imaged over a period of 3 months. In these large volumes of high resolution, we were able to detect the exceptional event of plaque birth in close proximity to dendrites seven times over. Imaging was started at 2–3 months of age when few amyloid plaques were present, with new plaques expected to form during the following months [22]. To the best of our knowledge, this is the first time that individual dendritic spines were monitored before and during the formation of an amyloid plaque in their proximity (Fig. 6a, b). Interestingly, the spine density of these dendrites remained unchanged prior to plaque formation and did not decline immediately after the plaques first appeared (Fig. 6c; total length of dendritic segments $444 \, \mu m$, $n = 247$ dendritic spines). A significant reduction in spine density did however occur 4.5 weeks after initial amyloid plaque formation ($p < 0.01$, repeated measures one-way ANOVA with Tukey–Kramer post hoc test). Over the same time period, plaque volume increased significantly immediately after the plaque’s appearance (Fig. 6c, $p < 0.05$, Wilcoxon signed-rank test against theoretical value $0 \, \mu m^3$). Thus, there seems to be a latency of about 4 weeks between plaque formation and the onset of dendritic spine elimination in the vicinity of plaques.

Discussion

We co-investigated and correlated amyloid plaque growth with dendritic spine kinetics in APPPS1 mice by long-term two-photon in vivo imaging. Several studies previously reported dendritic spine loss around amyloid plaques in a range of different mouse models and brain regions by applying both ex vivo and in vivo approaches [6, 19, 32, 34, 40, 53, 54, 59]. Here, we were able to confirm spine loss in proximity of plaques in the APPPS1 mouse model. However, the primary focus of this study was to analyze the underlying kinetics of spine loss around plaques. We were especially interested in the chronology: Which event comes first, dendritic spine loss or plaque formation? While a recent study investigated changes of neuritic curvature after plaque appearance [41], we were able to observe the fate of individual dendritic spines not only after the appearance of fibrillar plaques, but also before. We investigated spine loss kinetics in two age groups: a young cohort, 3–4 months old, when plaque growth is highly dynamic and an older cohort, 18–19 months old, when the plaque pathology is in the terminal stages.

As expected, we noted significant plaque growth in the young cohort, in line with several other studies [8, 10, 22, 61]. Moreover, our analysis showed a similar rate of plaque growth ($0.320 \pm 0.066 \, \mu m \, week^{-1}$, Fig. 3b) as reported by Hefendehl et al. [22] in the same mouse model and at a similar age ($\sim 0.3 \, \mu m \, week^{-1}$). Hefendehl et al. imaged mice up to an age of 10 months and observed no decline in plaque growth kinetics by that age. However, when we extended the observation period to an age of 18–19 months, we found that plaque growth declined essentially to zero (Fig. 4b). This finding corroborates the results from two other studies which also found no plaque growth in aged Tg2576 mice [8, 9]. Further support for a slow and gradual growth of plaques over extended periods of time is given by the large difference in the mean plaque sizes between the two age cohorts (Fig. 2b).

In the second part of our study, we identified a significant loss of spines in close proximity to amyloid plaques...
50 l m) compared to areas farther away from plaques and in age-matched control animals. Interestingly, similar results were found for both age cohorts. Thus, when monitoring the relative spine density over 4 weeks at already-existing plaques, we found a reduction to 57.9 ± 28.3 % in young mice and 67.8 ± 20.8 % in aged mice (Figs. 3a, b, 4a, b). Linear regression analysis showed no statistically significant difference between these results (Suppl. Fig. c). This finding suggests that spine loss around plaques persists for more than 1 year in this mouse model.

Taking also into consideration the results from plaque growth kinetics, we can envision a situation in which plaques are steadily growing and, of course, the volume around the plaques, where spine density is affected, increases as well. Nevertheless, even when plaque growth is not detectable anymore in aged mice, synapses are still being lost. In this context, it is worth noting that the spine density of dendrites in the surrounding volume of amyloid

Fig. 6 Spine density kinetics before and during the formation of amyloid plaques. a Maximum intensity projections of two-photon in vivo images of YFP-labeled dendrites (grey) are shown in a weekly imaging interval. At week 0, a new plaque (blue) appeared in direct vicinity to the dendrite in the center. The black rectangle marks the dendritic segment, which is shown in greater magnification in b. Important to note, the dendrite does not take course directly through the plaque, but the plaque is located above the dendrite (for a 3D view see supplementary figure). Scale bar indicates 10 μm. b Time series of the maximum intensity projected YFP labeled dendrite from a. The grey highlighted time scale indicates the time period when the amyloid plaque is already present. Blue arrows indicate maintained spines, red arrows lost spines, and green arrows gained spines (only some spines are exemplarily marked). Scale bar represents 2 μm. c Quantification of the dendritic spine kinetics are shown in black and plaque growth kinetics in blue. Spine densities were normalized to the spine densities at the first time point. The time point when amyloid appeared was set to 0 and is marked by a dashed line. Individual traces from seven dendrites (n = 2 mice, 444 μm dendrite length, n = 247 dendritic spines) are indicated by dashed lines, whereas the solid line shows mean with 95% CI. Dendritic spine loss became first significant 4 weeks after plaque appearance (p < 0.01, repeated measures ANOVA with Tukey–Kramer post hoc test). In contrast, the increase in amyloid plaque volume became significant directly with appearance (p < 0.05, Wilcoxon signed-rank test against theoretical value 0 μm³).
plaques in both age groups were quite similar (Suppl. Fig. a), while the number of healthy-looking dendrites was subjectively decreased in older mice. A previous study made a similar observation in aged (18–24 months) Tg2576 mice. Over 40 % aspiny dendrites were found in the surrounding area of plaques compared to about 15 % in control animals [53].

In the adult rodent brain, a small fraction of dendritic spines undergoes a certain turnover with spines being newly gained or lost, while the majority of spines is stable over long periods of time [20, 25, 58, 63]. In Tg2576 mice, increased spine elimination was found in proximity to plaques over 1 h [53]. In another AD mouse model (PSAPP), an increase in both spine formation as well as elimination was reported over 4–5 weeks [59].

Thus, besides confirming the fact that spines are lost around amyloid plaques, we also determined whether the spine loss was attributable to a decrease in gained spines or an increase in lost spines. In fact, we did not observe any changes in the fractions of gained spines, but a strong and significant increase in the fraction of lost spines measured in the vicinity of plaques over 4 weeks (Figs. 3c, 4c). In parallel to the reduced spine density, the fractions of lost spines were equally elevated in both age groups (Suppl. Fig. b). Moreover, the increased number of lost spines was due to the elimination of stable spines (data not shown). A further analysis revealed no preferential decline of a specific morphological subtype of dendritic spines over time within the old cohort (Fig. 5a–c). There was also no difference when comparing the morphological classification of dendritic spines from control animals with dendrites from the region around amyloid plaques (Fig. 5b, c). In conclusion, we detected a disturbed spine turnover within the periplaque region that was caused by a loss of stable spines. These stable spines are considered to represent the structural basis for long-term information storage [2, 4]. Increased spine elimination might be related to a higher concentration of toxic amyloid-β species in proximity to amyloid plaques as previously proposed [27, 30, 34, 49].

However, intriguingly, spine density did not decline immediately after the first appearance of a plaque, but rather following a delay of at least 4 weeks. This finding answers the question of what comes first, spine loss or amyloid plaque formation, [52] in favor of the latter. There even seems to be a significant delay between amyloid plaque formation and commencement of dendritic spine loss. Combining this finding with the result that spine loss around plaques is a protracted process taking place over a long period of time, the present study may explain the delay of 5–10 years between the accumulation of amyloid-β in human brains and onset of cognitive decline [50, 51]. However, we would like to emphasize that dendritic spine loss is only one part of the underlying multifactorial pathology of AD that contributes to cognitive decline besides neuron death, tauopathy, and hyperexcitability, to mention just some. Notwithstanding, the findings presented here may give hope to the possibility that even after initial plaque formation there might be a therapeutic window where potentially irreversible structural damage to synapses could still be prevented.

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