Research Paper

Dynamic assessment of tau immunotherapies in the brains of live animals by two-photon imaging

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ABSTRACT

Our original findings, showing the effectiveness of active and passive tau immunizations in mouse models, have now been confirmed and extended by many groups, with several clinical trials underway in Alzheimer’s disease and progressive supranuclear palsy. Here, we report on a unique and sensitive two-photon imaging approach to concurrently study the dynamics of brain and neuronal uptake and clearance of tau antibodies as well as the acute removal of their pathological target in live animals. This in vivo technique is more sensitive to detect clearance of pathological tau protein than western blot tau analysis of brain tissue. In addition to providing an insight into the mechanisms involved, it allows for an efficient in vivo assessment of the therapeutic potential of tau antibodies, and may be applied to related protein misfolding diseases.

1. Introduction

Tau immunotherapies are a promising approach for Alzheimer’s disease (AD) and related tauopathies. Our group published the original reports showing the effectiveness of active [1,2] and passive tau immunizations in mouse models [3,4]. These findings have now been confirmed and extended by multiple groups and several clinical trials have already been initiated [5,6]. However, the mechanisms of antibody-mediated clearance of tau aggregates are relatively unclear, although work by us and others has clarified to some extent various pathways that may be involved [5,6].

Two monoclonal antibodies (mAbs) that we have generated against the P-Ser396, 404 tau region, 4E6 and 6B2, have markedly different properties. 4E6 is overall more effective in various culture-, ex vivo- and in vivo models in preventing/reducing tau pathology and associated cognitive impairments, whereas 6B2 or rather its smaller derivatives, may be better suited as a diagnostic imaging marker [7–10]. We showed that the two mAbs are taken up by neurons and microglia and that their neuronal uptake is primarily via clathrin-dependent FcγII/III endocytosis [7–10]. Within tauopathy neurons, these antibodies bind to pathological tau protein primarily within the endosomal/lysosomal system, whereas their uptake/retention is limited in wild-type neurons [7–10].

Advances in imaging techniques now allow direct visualization of brain tissue in live animals by two-photon imaging. Compared to other imaging strategies, two-photon microscopy allows us to monitor the dynamics of mAbs in layers of the cortex (layerII/III) with minimal photo bleaching as reported for amyloid-β (Aβ) antibodies [11]. This permits the study of the time course of the entrance of antibody into the brain following an intravenous injection, its subsequent uptake into neurons as well as its binding to and clearance of tau aggregates in vivo. Monitoring tau clearance can be achieved by co-administration of a pathological tau marker, 1-fluoro-2,5-bis (3-carboxy-4-hydroxystyryl) benzene (FSB). It has been shown to cross the blood-brain-barrier and bind to tau aggregates in the brain as assessed postmortem [12].

In addition to providing such mechanistic insight, our findings show that this technique is a more sensitive way to acutely determine antibody efficacy in clearing pathological tau than by other traditional means such as by western blot analysis. In this regard, it complements...
2. Materials and methods

2.1. Animals

The htau model (Jackson Laboratories, stock #004808) [13] was crossed with a model that expresses the PS1 M146L human mutation [14]. The htau mice express unmutated human tau protein on a mouse tau knockout background (mtau KO) and develop tau pathology and tangles with age. The htau X PS1 model on mtau KO background, referred to as htau/PS1, has an earlier onset and more aggressive progression of tau pathology than the htau model. However, since our initial report on this cross [15], its severity has shifted to older age. For all the experiments, female htau/PS1 mice and wild-type (WT) controls of the same gender were at 18–20 months of age. For the study of antibody binding to and clearance of pathological tau (Figs. 2, 4–5), 6 htau/PS1 mice were used per group (IgG1, 4E6, or 6B2). These numbers were based on our past experience with this model, taking into account the data that was acquired sequentially. Analyses of the animal data was randomized and blinded.

All animals were housed at NYU School of Medicine animal facilities and cared for by the veterinary staff in AAALAC-approved facilities. All the procedures were approved by the Institutional Animal Care and Use Committee (IACUC) of the university, and are in accordance with NIH Guidelines, which meet or exceed the ARRIVE guidelines.

2.2. Injected antibodies

Monoclonal antibodies (mAbs) 4E6G7 (hereafter referred to as 4E6) and 6B2G12 (hereafter referred to as 6B2) were purified from our hybridomas by Genscript (Paramus, NJ). IgG1 (hereafter referred to as IgG1) of the same isotype as the tau mAbs served as control (eBioscience, sc-359,845) was injected into the femoral vein, with or without an immediate follow up injection of 100 μg of one of the Alexa Fluor 568 labeled antibodies (IgG1, 4E6, or 6B2) to label brain tau aggregates in vivo. The cranial window and the titanium bar in these mice were prepared in the same manner as described above. The mice were individually housed for 2 weeks after surgery and then FSB and/or antibody were injected followed by serial two-photon imaging at 1 h, 1 day, 4 days, 7 days, and 14 days after injection.

2.3. Animal preparation for two-photon imaging

Surgery was performed using aseptic techniques under isoflurane anesthesia (3% for induction, 1.5–2% for surgery). Briefly, the mouse was placed on a heated pad to maintain a body temperature of ~37 °C. The depth of anesthesia was monitored frequently during the surgery. The animal was placed in a stereotaxic frame, incision was made and the skull exposed to reveal the landmarks (bregma, lambda and midline) needed for cranial window. Subsequently, a section of the skull, about 3 mm in diameter, over the right somatosensory cortex was removed.

2.4. Intravenous injections

2.4.1. Antibodies in neurons

Alexa Fluor 568 labeled IgG1, 4E6, or 6B2 were injected (100 μg) into the femoral vein of the htau/PS1 or age-matched wild-type mice with the GFP labeled cortical neurons. Serial two-photon images were obtained at 4 h, 1 day, 4 days, 7 days, and 14 days after injection.

2.4.2. Antibodies bound to pathological tau

In a separate group of mice, 1-fluoro-2,5-bis(3-carboxy-4-hydroxystyryl) benzene (FSB; 0.1%, 300 μL, one injection; Santa Cruz, sc-350,845) was injected into the femoral vein, with or without an immediate follow up injection of 100 μg/200 μL of one of the Alexa Fluor 568 labeled antibodies (IgG1, 4E6, or 6B2) to label brain tau aggregates in vivo. The cranial window and the titanium bar in these mice were prepared in the same manner as described above. The mice were individually housed for 2 weeks after surgery and then FSB and/or antibody were injected followed by serial two-photon imaging at 1 h, 1 day, 4 days, 7 days, and 14 days after injection.

2.5. In vivo two-photon imaging

In vivo imaging was performed on anaesthetized mice (1.0% isoflurane) using Olympus two-photon microscope (FV1000MPE) equipped with a LUMPLFLN 60× water immersion lens (NA1.00) or Olympus two-photon microscope (FVMP-E) equipped with a 20× water immersion lens (NA 1.05). The mice were secured under a custom-built head fixation plate. FSB and Alexa 568 labeled antibodies were imaged using 800 nm excitation wavelength and AAV infected neurons were imaged using 920 nm excitation wavelength. Nondescan detector filter cube MRV/G-RXD1 (420–460 nm, EmDM-485) was used for FSB, MRG/R-RXD2 (575–630 nm, EmDM-570) was used for Alexa 568 labeled mAbs and MRG/R-RXD1 (495–540 nm, EmDM-570) was used for neuron-GFP.

Image stacks were analyzed using NIH ImageJ software. The fluorescence intensity of each time point (F) was normalized to Day 1 (F0) when FSB and mAbs had already entered into brain. Neuron/mAbs and FSB/mAbs colocalization coefficient (R) was analyzed using NIH ImageJ software with Colocalization Threshold and Colocalization Test plugins. For neuron and mAbs colocalization, R means Pearson’s correlation coefficient for pixels where both 6B2 channel and neuron channel are above their respective threshold within 6B2-positive neurons. For FSB and mAbs colocalization, R means Pearson’s correlation coefficient for pixels where both mAb channel and FSB channel are above their respective threshold.
2.6. Tau extraction and western blots

The left hemisphere of the brain was homogenized in (5 × vol/w) modified RIPA buffer (50 mM Tris-HCl, 150 mM NaCl, 1 mM EDTA, 1% Nonidet P-40, pH 7.4) with protease and phosphatase inhibitors (1 μg/mL of protease inhibitor mixture (Complete, Roche), 1 mM NaF, 1 mM Na3VO4, 1 mM PMSF, 0.25% sodium deoxycholate). The brain homogenate was centrifuged (20,000 × g) for 20 min at 20 °C and supernatants were collected as soluble tau fraction LSS (Low Speed Supernatant). For the sarkosyl insoluble fraction, equal amounts of protein from LSS were mixed with 10% sarkosyl solution to final 1% sarkosyl concentration, and the sample mixed for 30 min at room temperature, then centrifuged at 100,000 × g for 1 h at 20 °C. The sarkosyl pellet (SP) fraction was then air dried for 30 min, mixed with 50 μL of modified O+ buffer (1:5). LSS and SP in O+ buffer were boiled, and electrophoresed on 12% SDS-PAGE gels and transferred to nitrocellulose membranes. The blots were blocked in Superblock® (Thermo Fisher Scientific, incubated with PHF1 (1:1000, gift from Peter Davies), CP27 (1:500, gift from Peter Davies), GAPDH (1:3000, Cell Signaling Technology, D16H11), AT8 (1:500, Thermofisher Scientific) AT100 (1:500, Thermofisher Scientific) and HRP-conjugated anti-Fab secondary antibody (Thermofisher Scientific), and detected with ECL substrates (ThermoFisher Scientific). Images of immunoreactive bands were acquired by Fuji LAS-4000 imaging system and the bands quantified by MultiGauge software.

2.7. ELISA

To verify that the antibodies were not binding to FSB, the tau labeling dye, IgG1, 4E6 or 6B2 (0.5 μg per well) were coated onto an ELISA plate (Immulon, 2HB, Thermofisher Scientific) overnight at 4 °C in coating buffer (carbonate-bicarbonate buffer 50 mM, pH = 9.6). Then, FSB (1:10 to 1:100 K) was incubated with the coated plates at room temperature for 3 h. Following three washes in 0.1% TBST, fluorescence
endpoint signal was detected by BioTek Synergy™ 2 (Biotek Instrument, Inc) with the excitation wavelength 395 nm – 445 nm and emission wavelength 475 nm – 495 nm. For positive control, wells were loaded with the FSB dilutions immediately before signal detection.

2.8. Immunohistochemistry

After imaging studies, brains were collected for immunohistochemistry assays. Briefly, the mice were perfused transaortically with PBS and the extracted brains were fixed in 4% paraformaldehyde at 4 °C overnight, followed by at least 24 h immersion in 2% DMSO and 20% glycerol in phosphate buffer at 4 °C, and then cryo-sectioned coronally at 40 μm as previously described in detail [16]. FSB and Alexa-568 labeled monoclonal antibodies were directly visualized using a Nikon C-1 confocal microscope. For immunostaining, the brain sections were permeabilized with 0.3% Tween-20 and incubated with respective primary antibodies overnight and secondary antibodies for 1 h at room temperature. Brain FSB signal after its intravenous injection is mostly cleared during standard defatting and rehydration histological processing of mounted tissue sections. Hence, sections were restained with FSB (0.1%) in the same manner as standard Thioflavin S staining [16], to examine its postmortem co-localization with injected 4E6 or PHF1-stained sections.
2.9. Statistical analyses

GraphPad Prism 7 was used for statistical analyses. The intensity changes of mAbs and FSB over time were analyzed using two-way ANOVA, repeated measures, with Tukey's post-hoc test. Colocalization and Western blot data was analyzed using one-way ANOVA with Dunnett's post-hoc test. Significance levels were set at $P < .05$. All data are presented as mean ± SEM.

3. Results

3.1. Tau mAbs and control IgG cross the blood-brain barrier and are primarily detected within neurons. Neuronal 4E6 is cleared faster than 6B2 in htau/PS1 mice

To determine blood-brain-barrier (BBB) permeability, neuronal uptake and clearance of tau antibodies in live htau/PS1 tauopathy mice, neurons were GFP tagged by AAV virus infection, and Alexa 568-IgG1 control, Alexa 568-4E6 tau mAb or Alexa 568-6B2 tau mAb (100 μg) were injected into the femoral vein of the tauopathy mice (18–20 months). Two-photon live images were collected at Day 0, 1, 4, 7 and 14 from the same animal. Most of the 6B2 mAb (≥ 95%) was still in the blood vessels 4 h after injection (Fig. 1a). At 24 h (Day 1), 6B2 was primarily found within neurons ($R = 0.78$) and gradually decreased up to 14 days post-injection (Fig. 1b-c). Consistently, about 80% of the antibody was found within neurons, indicating comparable clearance from different cell types / binding sites (Fig. 1b). The distribution of the tau 4E6 mAb and control IgG1 from blood to neurons within the brain followed a similar pattern as for 6B2, although the kinetics differed (Fig. 1c). IgG1, 4E6 and 6B2 were decreased within neurons to 24%, 22% and 47% on Day 14 compared to Day 1. Two-way ANOVA analysis showed significant time and group effects ($F (3, 35) = 117.8, p < .0001$ and $F (2, 35) = 25.68, p < .0001$, respectively). 6B2 was cleared more slowly on Day 4, Day 7 and Day 14 ($p = .0047, .0004$ and .0129, respectively; Tukey's post-hoc test), compared to IgG1 on the same days [8].

3.2. Neuronal 6B2 signal is cleared more slowly in htau/PS1 mice than in wild-type mice

To explore whether pathological tau levels affect the clearance of mAbs in live mice, Alexa 568-4E6 or Alexa 568-6B2 (100 μg) were separately injected into htau/PS1 mice or wild-type (WT) mice. Changes in...
the fluorescence signal were detected by two-photon microscope at Day 1, Day 4, and Day 7. Similar to our findings in htau/PS1 mice (Fig. 1a), most of mAb signal in WT mice was in blood vessels 4 h after mAbs injection, leading to extensive neuronal signal within the brain 24 h after injection (data not shown). In all the groups, mAbs signal decreased over the 7 day imaging period (Fig. 1d). Antibody clearance was comparable in 4E6 injected WT and htau/PS1 mice and 6B2 injected WT mice but was much less in 6B2 injected htau/PS1 mice. Two-way ANOVA analysis showed significant time (F [2, 24] = 217.2, p < .0001) and group (F [3, 12] = 15.11, p = .0002) effects. At 7 days post-injection, neuronal 6B2 signal had decreased to 23% in WT vs. 71% in htau/PS1 mice, compared to Day 1. The neuronal 6B2 signal differed significantly in these two groups on Day 4 (p = .0342) and Day 7 (p = .0196; Tukey’s post-hoc test).

3.3. 4E6 and 6B2 colocalize with a tau aggregate dye probe in htau/PS1 mice

To clarify if the antibodies are binding to pathological tau in htau/PS1 mice, 1-fluoro-2,5-bis (3-carboxy-4-hydroxystyryl) benzene (FSB) was used to label tau tangles [12]. FSB (0.1%, 300 μL) with Alexa 568-IgG1/4E6/6B2 (100 μg) were injected into the femoral vein of the htau/PS1 mice. Serial two-photon images were then collected of the same brain region over 14 days and FSB intensity was analyzed. Representative two-photon in vivo live images showing excellent co-localization of FSB and tau mAbs 4E6 and 6B2 4 days after their injection. Quantification of FSB colocalization with different mAbs. R is Pearson’s correlation coefficient for pixels where both mAb and FSB channels are above their respective threshold. F [2, 9] = 83.16, p < .0001 one-way ANOVA. The tau mAbs 4E6 and 6B2 co-localized more significantly with FSB (R = 0.55, R = 0.64, respectively, p < .0001; Dunnett’s post-hoc test), compared to IgG1 (R = 0.25). FSB intensity in the brain decreased over time, and more with 4E6 than 6B2 or IgG1. F [3, 35] = 20.56, p < .0001 for time factor and F [2, 35] = 13.17, p < .0001 for group factor. FSB signal decreased faster when it was co-injected with 4E6 on Day 4, Day 7 and Day 14 (p = .0246, .0056 and .0427, respectively), compared to IgG1 on the same day (two-way ANOVA, repeated measures, Tukey’s multiple comparisons test). F/F0: Ratio of average fluorescence intensity on Day 1 vs. other days. Scale bar = 10 μm. All data are presented as mean ± SEM. *: p < .05; **: p < .01; ****, p < .0001.

![Fig. 4](https://example.com/figure4.png)

**Fig. 4.** Two-photon in vivo live imaging of FSB colocalization with tau mAbs and 4E6 is more effective than 6B2 in clearing tau aggregates. FSB (0.1%, 300 μL) with Alexa 568-IgG1/4E6/6B2 (100 μg) were injected into the femoral vein of the htau/PS1 mice. Serial two-photon images were then collected of the same brain region over 14 days and FSB intensity was analyzed. a. Representative two-photon in vivo live images showing excellent co-localization of FSB and tau mAbs 4E6 and 6B2 4 days after their injection. b. Quantification of FSB colocalization with different mAbs. R is Pearson’s correlation coefficient for pixels where both mAb and FSB channels are above their respective threshold. F [2, 9] = 83.16, p < .0001 one-way ANOVA. The tau mAbs 4E6 and 6B2 co-localized more significantly with FSB (R = 0.55, R = 0.64, respectively, p < .0001; Dunnett’s post-hoc test), compared to IgG1 (R = 0.25). c. FSB intensity in the brain decreased over time, and more with 4E6 than 6B2 or IgG1. F [3, 35] = 20.56, p < .0001 for time factor and F [2, 35] = 13.17, p < .0001 for group factor. FSB signal decreased faster when it was co-injected with 4E6 on Day 4, Day 7 and Day 14 (p = .0246, .0056 and .0427, respectively), compared to IgG1 on the same day (two-way ANOVA, repeated measures, Tukey’s multiple comparisons test). F/F0: Ratio of average fluorescence intensity on Day 1 vs. other days. Scale bar = 10 μm. All data are presented as mean ± SEM. *: p < .05; **: p < .01; ****, p < .0001.
their partial colocalization (Fig. 3c). One-way ANOVA showed significant group effect on antibody colocalization with FBS in the cortex of the mice when analyzed 4 days post-injection ($F \{2, 9\} = 83.16$, $p < .0001$). The tau mAbs 4E6 and 6B2 significantly co-localized with FSB ($p < .0001$, Dunnett’s post-hoc test; $R = 0.55$, $R = 0.64$, respectively), compared to IgG1 ($R = 0.25$, Fig. 4a-b).
3.4. 4E6 is more effective than 6B2 in clearing tau aggregates

To clarify if the antibodies are promoting clearance of pathological tau, two-photon images were collected after FSB co-injection with IgG1, 4E6 or 6B2 (100 μg) on Day 1, Day 4, Day 7 and Day 14. FSB intensity in the brain decreases over time, and more with 4E6 than 6B2 or control IgG1 (Fig. 4c). Two-way ANOVA analysis showed significant time and group effects (F (3, 35) = 20.56, p < .0001 and F (2, 35) = 13.17, p < .0001). FSB signal decreased faster, when it was co-injected with 4E6, on Day 4, Day 7 and Day 14 (p = .0246, .0056 and .0427, respectively, compared to IgG1 on the same day, Tukey's post-hoc test). However, FSB injected with 6B2 was cleared at a similar rate as with IgG1. Following imaging on Day 14, the brains were collected for biochemical analysis of tau to determine if these differences in FSB clearance rate were reflected in soluble (Fig. 5a-h) and insoluble tau fractions (Fig. 5i-k). One-way ANOVA analysis showed significant group effect on soluble phospho-tau AT100 and PHF1/CP27 (Fig. 5d, f, AT100, F [2, 14] = 6.349, p = .0109; PHF1/CP27, F [2, 14] = 4.299, p = .035). Both 4E6 and 6B2 decreased soluble phospho-tau AT100 levels (Fig. 5d, p = .0115 and .0165; Dunnett’s post-hoc test). 4E6, but not 6B2, decreased soluble phospho-tau PHF1/CP27 levels (Fig. 5f, p = .0215; Dunnett’s post-hoc test). There was no significant group effect on insoluble tau level (Fig. 5i-k).

4. Discussion

In this study, we measured and analyzed dynamic changes in fluorescently labeled tau mAbs and in a tau β-sheet dye in live mice by two-photon imaging. We found that the tau mAbs crossed the BBB and were primarily detected within neurons (Fig. 1a-b). One of the tau mAbs, 6B2, was cleared more slowly and to a lesser extent in tauopathy mice than the other tau mAb, 4E6, which was cleared at a similar rate as a control IgG (Fig. 1c). In WT mice, both tau antibodies were cleared at a similar pace and to a comparable degree (Fig. 1d). These clearance differences between the two tau mAbs can be explained by their different binding properties. Although 4E6 and 6B2 were generated to target the same tau peptide P-Ser396, 404 region, 4E6 binds better to soluble tau and 6B2 binds more strongly with insoluble tau [10].

Previously, we identified 6B2 and its scFv derivate as potential imaging ligands to detect tau pathology using an In Vivo Imaging System (IVIS) [9]. The 4E6 tau mAb gives overall less IVIS signal in the same mice than 6B2, based on our preliminary results [17], which fits well with the two-photon findings.

Apart from providing detailed information on the vascular vs cellular distribution of the ligand, which is not possible with the IVIS approach, the two-photon method is much more sensitive. That is as expected because the skull and skin obviously diminish the IVIS signal. With the two-photon technique, we found that most of the mAb signal was in blood vessels 4 h after injection, and approximately 25–50% of the antibody remained in the brain, predominantly within neurons (about 80%) 14 days following the intravenous injection (Fig. 1a-b). In our prior IVIS study, the signal remained relatively stable for a few hours and then gradually subsided over a week back to background values [9]. The key advantage of the IVIS method is its non-invasiveness as a signal can be obtained from an intact animal. However, unlike the two-photon approach, it cannot provide information on the vascular vs cellular distribution of the ligand(s) under study. The large percentage of the antibodies detected within neurons is analogous to our prior report on tau antibody uptake in brain slice culture derived from mouse tauopathy brain [8]. There, about 10% of the tau antibodies added to the brain slice culture were detected within microglia. One would perhaps expect a larger percentage of antibodies within those phagocytic cells, but their clearance of antibody-tau complexes is likely more efficient than in neurons, resulting in an underestimate of antibody uptake into microglia. In addition, most of pathological tau resides within neurons. That fact, coupled with their slower clearance of antibody-tau complexes explains the high percentage of neuronal tau antibodies.

Co-injection of the tau β-sheet dye FSB and tau antibodies clearly showed that these markers associate within neurons, where most of pathological tau is found in these animals (Fig. 4a-b). Such intraneuronal association of tau antibody with pathological tau in vivo fits prior reports from us and others, showing with different methodologies in postmortem tissue that tau antibodies enter the brain and neurons, in which they bind to pathological tau [2,7–10,18–20]. Similar intraneuronal binding of antibodies to other protein aggregates has been reported by various laboratories [21–24]. FSB has previously been shown to label brain tau aggregates upon postmortem examination after peripheral injection [12], and not to give a two-photon signal in wild-type mice [25]. Another β-sheet dye that labels various amyloids, including tau and Aβ, has previously been shown to label tau tangles in mice as detected by two-photon imaging following topical application [26], as unlike FSB, it does not readily cross the blood brain barrier. Not all tau antibodies are rapidly taken up into neurons [27–29] as such uptake is likely charge-dependent [6].

FSB co-localizes well with the tau antibodies within brain neurons in the two-photon images when both are injected (Fig. 4a). However, only a partial co-localization is seen when brain sections are stained with FSB following 4E6 antibody injection (Fig. 3b). Such direct staining showed more binding than can be expected with two-photon imaging. Furthermore, 4E6 binding to tau may diminish further FSB binding to that particular site. For these two reasons, overlap between FSB and tau antibody is only partial in the histological sections (Fig. 3b). Gradient binding to tau of FSB and tau antibody following an intravenous injection can be expected to result in a more mosaic pattern of binding and therefore better colocalization as observed. In addition, the 4E6 signal in the histological sections is more condensed-vesicle-like than following two-photon imaging. This may be explained by the different time period from injection to imaging in these images (7 days in Fig. 3b vs 4 days in Fig. 4a). With time, 4E6 becomes more condensed within neurons, presumably as it goes down the endosomal/lysosomal pathway as we have previously shown it is mainly confined within after neuronal uptake [8].

It is of note as well that one of the tau antibodies, 4E6, was associated with faster and more extensive clearance of FSB than 6B2 or control IgG1 (Fig. 4c). This acute efficacy difference is analogous to our recent report showing acute cognitive benefits of 4E6 whereas 6B2 was ineffective [10]. Here, in as that prior report, these benefits were not associated with extensive clearance of pathological tau as detected on Western blots. However, significant decrease was detected in soluble hyperphosphorylated tau in the 4E6 group compared to the 6B2 or IgG control groups following a single intravenous injection of 100 μg of antibody (about 3 mg/kg; Fig. 5d, f). That therapeutic effect is comparable to our prior study, in which a different tauopathy mouse model received three intraperitoneal injections of 4E6 or 6B2 (10 mg/kg) [10]. Hence, this type of two-photon approach analyzing antibody-mediated clearance of FSB in real time, allows for an efficient and sensitive in vivo screen of the therapeutic potential of tau antibodies. It is not surprising that it is more sensitive than western blot measurements because it entails direct analysis of live animals without the potentially confounding variables associated with the numerous steps involved in postmortem tissue processing for western blots. In addition, although we show here and previously that antibody-mediated tau clearance can be detected by staining with antibodies against various epitopes [2,4,15], some epitopes may be cleared better than others as shown here. This adds another variable to such analysis. In future studies, measurement of CSF tau levels in conjunction with two-photon imaging should further clarify target engagement of tau antibodies. Likewise, since FSB gradually fades with time (about 50% in two weeks), its injection prior to each imaging session may allow better detection of antibody efficacy. In addition, ELISA measurements of tau levels may be more sensitive than Western blot analyses and provide further insight into antibody-mediated tau clearance.
With the advantages of penetration into brain tissue and low phototoxicity, in vivo two-photon imaging is widely used in AD mouse models [30]. This includes to image neurites, dendritic spines, microglia, astrocytes, Aβ plaques, tau tangles and pre-tangles or neuronal and astrocytic activity. In particular, antibody therapies have been shown to clear Aβ plaques based on reduced signal from the fluorescent dye that labels Aβ [11]. Furthermore, camelid single-domain antibodies have been shown to cross the BBB and label Aβ deposits and neurofibrillary tau tangles in vivo using two-photon microscopy [31]. However, we are not aware of other two-photon studies showing antibody-mediated clearance of tau aggregates as we are reporting.

In summary, two-photon imaging provides a unique insight into the dynamics of uptake and clearance of mAbs, as well as the removal of their pathological targets in live animals. In addition to elucidating the mechanisms involved, this approach allows for an efficient in vivo assessment of the therapeutic potential of tau antibodies, and may be applied to related protein misfolding diseases, assuming that appropriate β-sheet dyes exist for those aggregates.

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Declaration of interests

EMS is an inventor on patents on tau immunotherapy and related diagnostics that are assigned to New York University. Some of this technology is licensed to and is being co-developed with H. Lundbeck A/S.

Author contributions

Designing experiments: EMS, QW, JG. Performing experiments: QW, YL, JG. Analyzing data and writing manuscript: EMS, QW. All authors read and approved the final manuscript.

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