Intraneuronal aggregation of the β-CTF fragment of APP (C99) induces Aβ-independent lysosomal-autophagic pathology

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Abstract Endosomal-autophagic-lysosomal (EAL) dysfunction is an early and prominent neuropathological feature of Alzheimer’s disease, yet the exact molecular mechanisms contributing to this pathology remain undefined. By combined biochemical, immunohistochemical and ultrastructural approaches, we demonstrate a link between EAL pathology and the intraneuronal accumulation of the β-secretase-derived βAPP fragment (C99) in two in vivo models, 3xTgAD mice and adeno-associated viral-mediated C99-infected mice. We present a pathological loop in which the accumulation of C99 is both the effect and causality of impaired lysosomal-autophagic function. The deleterious effect of C99 was found to be linked to its aggregation within EAL-vesicle membranes leading to disrupted lysosomal proteolysis and autophagic impairment. This effect was Aβ independent and was even exacerbated when γ-secretase was pharmacologically inhibited. No effect was observed in inhibitor-treated wild-type animals suggesting that lysosomal dysfunction was indeed directly linked to C99 accumulation. In some brain areas, strong C99 expression also led to inflammatory responses and synaptic dysfunction. Taken together, this work demonstrates a toxic effect of C99 which could underlie some of the early-stage anatomical hallmarks of Alzheimer’s disease pathology. Our work also proposes molecular mechanisms likely explaining some of the unfavorable side-effects associated with γ-secretase inhibitor-directed therapies.

Keywords Alzheimer · C99 · Lysosomes · Autophagy · Aggregation · Triple-transgenic mouse · γ-Secretase inhibition

Introduction

Alzheimer’s disease (AD) is the most prevalent neurodegenerative disease and cause of dementia in elderly. Histopathologically, it is characterized by the deposition of extracellular amyloid plaques and intraneuronal neurofibrillary tangles, as well as synaptic pathology and neurodegeneration [7]. Growing evidence supports that these alterations are preceded and may be even caused by a defective endolysosomal/autophagic degradation [33, 36, 39, 43, 50]. Indeed, endolysosomal/autophagic anomalies are well-recognized early neuropathological features of AD, marked by enlarged endosomes, lysosomal alterations and progressive accumulation of autophagic vacuoles (AVs) [33–35, 39, 43, 50]. It is also well established that endosomal-autophagic-lysosomal (EAL) compartments constitute the main sites of proteolytic processing of the amyloid-β precursor protein βAPP [43] and that amyloidogenic species can accumulate inside organelles of the EAL machinery [12, 20, 24, 26, 62]. Moreover, the accumulation of the aggregate-prone Aβ42 within EAL organelles was found to disturb normal EAL function, suggesting that it could be both a
consequence and a cause of endolysosomal-autophagic dysfunction [43].

EAL dysfunction has been extensively described in transgenic mice displaying AD-like anatomo-pathology [11, 13, 47, 51, 55, 61] including in the widely used 3xTgAD mouse (APPswe, TauP301L, PS1M146V) [10, 11, 37]. It is also well known that this mouse model displays an early accumulating and particularly high intraneuronal amyloid-like immunostaining within the subiculum of the hippocampus, which was firstly accounted for Aβ [37]. Yet, we recently demonstrated that this staining corresponded to the β-secretase-derived fragment, C99 (βCTF) rather than to Aβ, which was only detected at low levels and at late stages in these mice [24]. We found that C99 accumulated in enlarged cathepsin B-positive structures suggesting a link between C99 and EAL pathology [24]. In this work, we establish that C99 accumulation is the consequence of impaired lysosomal-autophagic degradation but that C99, in turn, also contributes to lysosomal dysfunction. Interestingly, these effects of C99 were observed in the 3xTgAD mouse, in which C99 is generated by proteolytically APP processing, but also in a direct transgenic C99-expressing mouse. C99 fragments were found to aggregate within membranes of EAL vesicles, proposing that C99 induces autophagic-lysosomal dysfunction by interfering with EAL membrane integrity. The pharmacological inhibition of γ-secretase increased the levels of EAL-associated C99 and exacerbated pathology, clearly demonstrating a C99 dose-dependent but Aβ-independent effect. Overall, our work demonstrates a detrimental loop in which aggregated C99 contributes to anatomical hallmarks reminiscent of those occurring early in Alzheimer’s disease.

Materials and methods

Viral constructions

Virus production was performed following a protocol previously described [4]. Briefly, HEK293 cells were transfected with the adenovirus helper plasmid (pXG6), the AAV packaging plasmid (rAAV2-rh10), and the AAV10 plasmid empty vector or encoding either human C99 or GFP under control of the synapsin-1 promoter (AAV-empty, AAV-synC99 and AAV-synGFP). Viruses were produced, purified and vector titers were determined by real-time PCR and expressed as viral genomes per ml (vg/ml) [4].

Animals and in vivo drug treatment

3xTgAD (harboring PS1M146V, βAPPswe, and TauP301L transgenes) and non-transgenic (nonTg) mice [37] were generated from breeding pairs provided by Dr. LaFerla (Irvine, USA). 2xTgAD (PS1wt, βAPPswe and TauP301L) were produced by crossing 3xTgAD with nonTg mice, as described [38]. For AAV-mediated in vivo delivery, 1-day-old C57BL6 mice (Janvier Labs., France) were injected with 4 µl of AAV virus (5.5 × 10^12 vg/ml (viral genomes per ml)) into the left lateral ventricle, as described [21] and mice were analyzed at 2 months post-AAV delivery. 5-month-old nonTg and 3xTgAD males or 2-month-old AAV-infected mice (males and females) were treated daily for 12 or 30 days with the γ-secretase inhibitor ELND006, referred to as D6 hereafter (30 mg/kg, Elan Pharmaceuticals, San Francisco, USA) [9, 45] or with vehicle alone (methylcellulose/polyosorbate 80, Sigma) via oral gavage, as described [24]. All experimental procedures were in accordance with the European Communities Council Directive of 24 November 1986 (86/609/EEC) and local French legislation.

Immunohistochemical analysis

Animals were deeply anesthetized with a lethal dose of pentobarbital and perfused transcardially with cold phosphate-buffered saline (PBS). Brains were fixed in 4 % Paraformaldehyde/PBS then embedded in paraffin and sliced (8 µm) or cut on a vibratome (50 µm). For FCA18, 82E1, NU1, 4G8 and 6E10 staining, sections were treated with formic acid (90 % or 50 %/5 min for paraffin and vibratome sections, respectively). Sections were then incubated at 4 °C overnight with primary antibodies (see supplementary table) followed by Alexa Fluor-conjugated antibodies (Molecular Probes, 1:1000). Cathepsin B labeling was amplified using the Vectorstain ABC kit (Vector) and streptavidin-Alexa594 (Molecular Probes, 1:1000). Nuclei were stained with DAPI (Roche, 1:20,000) and fluorescence was visualized using a confocal microscope (Fluoview10, Olympus). For DAB development, sections were incubated with HRP-conjugated secondary antibodies (Jackson ImmunoResearch, 1/1000) or with vehicle alone (methylcellulose/polysorbate 80, Sigma) via oral gavage. For DAB development, sections were incubated with HRP-conjugated secondary antibodies (Jackson ImmunoResearch, 1/1000) followed by DAB substrate (DAB impact, Vector). Neuronal nuclei were stained with cresyl violet. For quantitative analysis of lamp1 staining in AA V-empty and AA V-C99 brain slices, images were thresholded, converted to mask and analyzed using Particle Analysis ImageJ plugin.

Cell culture and experimental treatments

The human neuroblastoma cell line SH-SY5Y, naive or stably expressing APPswe or pcDNA3 (mock), previously described [40] was exposed to the following drugs for 16–20 h: D6 (50 nM to 5 µM, Elan Pharmaceuticals, San Francisco) in vehicle (methylcellulose/polysorbate 80, Sigma), DAPT (5 µM in DMSO, Sigma), leupeptin (10 µM in H₂O, Sigma), PADK (5 µM in DMSO,
Bachem), NH₄Cl (10 mM in H₂O, Sigma), pepstatin A (10 μM in EtOH, Enzo Life sciences), smear28 (50 μM in DMSO, Sigma) and bafilomycin A1 (50 nM in H₂O, Sigma). Some cells were transfected with GFP-LC3 (Addgene) using Lipofectamine 2000 according to standard protocols. Other cells were infected with lentiviruses (LV) encoding mouse cathepsin B (LV-mCatB-FUGW2) or control (LV-FUGW2) [32] and polyclonal cell lines stably expressing mCatB (naive SH-SY5Y or APPswe-SH-SY5Y cells) were obtained. For immunocytochemical experiments of C99 expression, COS-7 cells were transfected with C99 using Lipofectamine 2000.

Immunocytochemistry

Cells were immunostained with α-APPct (1:5000) and α-lamp-1 (E-5, 1:100) or α-cis-golgi (GM130, Cell signaling, 1:400) followed by Alexa Fluor-488 or Alexa Fluor-594 conjugated antibodies (1:1000, Molecular Probes). Nuclei were stained with DAPI (1:20,000, Molecular probes). Other cells were incubated 30 min with Lysotracker-Red DND-99 (1:20,000, Invitrogen) at 37 °C, fixed with PFA 4 % and stained with DAPI. Images were acquired using an inverted confocal microscope (Fluoview10, Olympus, France).

In vitro degradation assay

Recombinant C99 (C100-flag), previously described [49], was incubated in the absence or presence of purified cathepsin B from human placenta (5, 10 or 50 ng, Sigma) in 100 mM sodium acetate pH 5 buffer containing 1 mM EDTA and 8 mM cysteine at 37 °C during 60 min, then analyzed on Tris-Tricine 16 % acrylamide gels as described below. C100flag was revealed using an anti-flag antibody (M2, 1:5000, Sigma).

Electron microscopy

Cells were fixed 20 min with 2.5 % glutaraldehyde/phosphate buffer 0.1 N (pH 7.4). Mice were anesthetized with a lethal dose of pentobarbital and perfused transcardially with ice cold physiological saline followed by 2 % glutaraldehyde/cacodylate buffer 0.1 M (pH 7.4). The brain was sliced (200 μm) on a vibratome and 2 mm cubes from the subiculum were microdissected under binoculars and post-fixed in osmium tetroxide (1 % in cacodylate buffer 0.1 M). The tissue was embedded in Epon resin (EMS) and 80 nm ultrathin sections were contrasted with uranyl acetate and lead citrate and visualized using a JEM 1400 electron microscope operating at 100 kV equipped with a Morada SIS camera.

Preparation of brain fractions

Dissected hippocampi of vehicle or D6-treated 3xTgAD or nonTg mice, or hemispheres from AAV-injected mice, were homogenized in RIPA buffer (Tris 50 mM; pH 7.4 containing NaCl (150 mM), EDTA (1 mM), Triton X100 (1 %), deoxycholate (0.5 %), SDS (0.1 %) and protease inhibitor cocktail (Sigma) and soluble and insoluble fractions were prepared as described [24]. Synaptosomal and microsomal enriched fractions were prepared from hippocampi of vehicle or D6-treated 3xTgAD mice. Tissue was homogenized in 0.32 M sucrose containing protease inhibitor cocktail (Sigma), centrifuged twice at 850 × g/5 min and the collected supernatants were centrifuged twice at 12,000 × g/20 min. The obtained pellet (“synaptosomal fraction”) was resuspended in Heps 4 mM/EDTA. The supernatant was centrifuged at 20,000 × g/1 h to obtain the “microsomal fraction”.

SDS/PAGE and western blot analyses

Proteins were separated on 16 % Tris-Tricine gels (for APP-CTFs, cathepsin B and LC3) or 10 % Tris–glycine gels (for βAPP p62 and actin) and transferred to nitrocellulose membranes. After probing with primary antibodies (see supplementary table), immunological complexes were revealed with HRP-conjugated antibodies (Jackson Immunoresearch, 1/10,000) followed by electrochemiluminescence (Supersignal West Pico/West Maximum Sensitivity chemiluminescence substrate, Thermo Scientific, France). Peak height of signal intensities from protein bands were quantified with ImageJ software.

Quantification of human Aβ

Aβ40 and Aβ42 levels were quantitatively detected in soluble and insoluble fractions (see above) of mice hippocampi (for 3xTgAD mice) or hemispheres (for AAV-synC99 mice) using sandwich ELISA kits detecting human Aβ40 and Aβ42, respectively ( Biosource, Invitrogen, France). Absorbance was read at 450 nm using a spectrophotometer.

In vitro cathepsin B activity assay

Cells were lysed mechanically in homogenization buffer (250 mM sucrose, 1 mM EDTA, 5 mM Heps pH 7.4). The cell suspension was centrifuged at 850 × g/5 min and the supernatant was centrifuged at 20,000 × g/90 min. The pellet (membrane enriched fraction) was resuspended in Tris–HCl (10 mM, pH 7.5). To monitor cathepsin B (CatB) activity, 60 μg of protein extracts were incubated in acetate buffer (25 mM, pH 5.5, 100 μl and 8 mM l-cysteine HCl) containing CatB substrate
Figure 1c, d) increased similarly with age, while only cellular staining (Suppl. Figure 1a) and C99 levels (Suppl. Figure 1a). Interestingly, in the two strains this intracellular C99 accumulation in double transgenic (2xTgAD) mice harboring APPswe and TauP301L but wild-type presenilin1 [24] (Suppl. Figure 1e). Overall, these data indicated that C99 accumulation did not correlate with Aβ load and therefore could not be accounted for altered γ-secretase processing alone. In situ immunolabeling with 82E1 revealed that C99 accumulated in intraneuronal organelles positive for the lysosomal enzyme cathepsin B (Fig. 1a) and suggested a link between C99 accumulation and defective lysosomal clearance. In young 3xTgAD mice (3 months of age), the majority of the neurons displayed numerous small-sized cathepsin B puncta (Fig. 1a, arrows), but some neurons also presented enlarged cathepsin B structures which were positive for C99 (Fig. 1a, arrowheads). Similar results were obtained with an antibody directed against the endosomal-lysosomal membrane-associated protein lamp1, and the number and size of these structures were high in both 14-month-old 2AD and 3AD mice (Fig. 1b). The large size of these structures proposed that many of them corresponded to endo- or autolysosomes rather than to lysosomes and would support impaired lysosomal-autophagic-mediated C99 degradation in both 2xTgAD and 3xTgAD mice. In agreement, electron microscopy revealed an increased number of dense autophagic vesicles in brains from both of these mice, as compared to non-transgenic mice (Fig. 1c, red arrowheads). These autophagic vacuoles corresponded to both small and very dark autolysosomes (<1.5 µm, examples in lower middle and right images) containing various compositions of densely packed undigested material and lipid droplets (clear round structures).

Electrophysiological recordings

LTP measurements were performed by E-Phy-Science, Sophia Antipolis, France. 2-month-old AAV-empty, vehicle or D6-treated AAV-C99 females, were anesthetized with isoflurane and decapitated. Stimulations were delivered in CA1 fibers in the alveus and field excitatory postsynaptic potential (f-EPSP) were recorded from the subicular pyramidal neurons in the middle portion of the subiculum. The LTP-induction protocol consisted of four trains of 100 stimuli at 100 Hz repeated every 20 s, as described [41].

Statistical analysis

Statistical analysis was performed with PRISM software (Graph-Pad Software, San Diego, CA, USA) by using a non-parametric two-tailed Mann–Whitney U test for pairwise comparisons or the two-way ANOVA test followed by either the Tukey’s post hoc test for multiple comparisons or the Dunnett post hoc test when comparison to controls. All data are expressed as the mean ± SEM. Differences were considered significant at *p < 0.05, **p < 0.01, ***p < 0.001.

Results

3xTgAD and 2xTgAD mice display early and progressive C99 accumulation within enlarged cathepsin B and lamp1 positive structures

We previously demonstrated that the β-secretase-derived APP C-terminal stub C99 corresponded to the early appearing and accumulating intracellular label in the subiculum of 3xTgAD mice [24]. This label could be detected with N-terminal directed antibodies including the β-secretase-mediated cleavage-specific antibodies FCA18 [3] and 82E1 [19] known to recognize both Aβ and C99. A similar accumulating intraneuronal FCA18-immunostaining was observed in double transgenic (2xTgAD) mice harboring APPswe and TauP301L but wild-type presenilin1 [24] (Suppl. Figure 1a). Interestingly, in the two strains this intracellular labeling (Suppl. Figure 1a) and C99 levels (Suppl. Figure 1c, d) increased similarly with age, while only 3xTgAD mice produced extracellular plaques at late ages that were stained with both FCA18 (Suppl. Figure 1a) and anti-Aβ42 (a-Aβ42) (Suppl. Figure 1b). The detection of very few extracellular Aβ plaques in situ in aged 2xTgAD mice was corroborated by the absence or very low detectable levels of both Aβ40 and Aβ42 measured by ELISA (Suppl. Figure 1e). These data indicated that C99 accumulation did not correlate with Aβ load and therefore could not be accounted for altered γ-secretase processing alone. In situ immunolabeling with 82E1 revealed that C99 accumulated in intraneuronal organelles positive for the lysosomal enzyme cathepsin B (Fig. 1a) and suggested a link between C99 accumulation and defective lysosomal clearance. In young 3xTgAD mice (3 months of age), the majority of the neurons displayed numerous small-sized cathepsin B puncta (Fig. 1a, arrows), but some neurons also presented enlarged cathepsin B structures which were positive for C99 (Fig. 1a, arrowheads). Similar results were obtained with an antibody directed against the endosomal-lysosomal membrane-associated protein lamp1, and the number and size of these structures were high in both 14-month-old 2AD and 3AD mice (Fig. 1b). The large size of these structures proposed that many of them corresponded to endo- or autolysosomes rather than to lysosomes and would support impaired lysosomal-autophagic-mediated C99 degradation in both 2xTgAD and 3xTgAD mice. In agreement, electron microscopy revealed an increased number of dense autophagic vesicles in brains from both of these mice, as compared to non-transgenic mice (Fig. 1c, red arrowheads). These autophagic vacuoles corresponded to both small and very dark autolysosomes (<1.5 µm, examples in lower middle and right images) containing various compositions of densely packed undigested material and lipid droplets (clear round structures).

Cellular analysis shows that APP-CTFs are cleared through the autophagic-lysosomal degradation pathway

The above data suggested a role of lysosomal-autophagic degradation in C99 accumulation and would agree with earlier data proposing a major contribution of this degradation pathway in APP-CTF clearance [1, 8, 54, 57]. To confirm the role of lysosomal enzymes in the fate of C99, we used SH-SY5Y neuroblastoma overexpressing APPswe (known to potentiate β-secretase-mediated C99 and Aβ production [29] and harbored by the βAPP transgene in both 2xTgAD and 3xTgAD mice [37]) (SH-APPswe). Clearly, treatments of SH-APPswe with either the lysosomotropic weak base NH4Cl or specific lysosomal protease inhibitors, such as the cathepsin B inhibitors leupeptin and PADK or the cathepsin D inhibitor pepstatin A, all led to large increases in APP-CTFs, including the most abundantly expressed C83,
C99 and AICD (Fig. 2a, b). These treatments also led to the accumulation of other higher molecular mass APP-CTFs, notably one about a size of 25 kDa (Fig. 2a, arrows), whose levels were very low in non-treated cells, showing its particularly high sensitivity to lysosomal degradation. Although, we did not determine the exact identity of...
Fig. 2 In vitro analysis shows that APP-CTFs are degraded by cathepsins through autophagy. **a**, **b** SH-APPswe cells were treated with NH₄Cl, leupeptin (Leu), pepstatin A (PepA) or PADK and analyzed by western blot using α-APPct. Arrows correspond to a non-identified 25 kDa APP-CTF fragment. **Bars** in **b** correspond to quantification of C99 and C83 immunoreactivities obtained in **a** and expressed relative to expressions measured in DMSO-treated cells normalized to actin. Data are represented as mean ± SEM, as determined by ANOVA one-way Dunnett post hoc test, **p** < 0.01 and *p* < 0.05 indicate significant differences relative to control cells (**n** = 10–16, from four independent experiments). **c** Immunocytochemical analysis using α-APPct and α-lamp1 on H₂O or NH₄Cl treated SH-APPswe cells. Note the labeling of α-APPct in enlarged α-lamp1 vesicles in merged images (arrow). **d** SH-APPswe cells were infected with lenti-cathepsin B (CatB) or empty vector (mock) virus at different concentrations (1, 5 or 10 μl of a stock corresponding to 3.75 × 10⁸ TU/ml) and CatB and APP-CTF levels were analyzed after 48 h by western blot. α-cathepsin B revealed immature (proCatB) and mature (mCatB) cathepsin B. **e**, **f** CatB and APP-CTF levels were analyzed by western blot in CatB stable cell lines after six passages. **Bars** in **f** represent the quantification of mCatB and C83 and C99 immunoreactivities obtained in **e**, each expressed as respective expressions measured in mock cells. Data are represented as mean ± SEM, Mann–Whitney, ***p** < 0.001 and **p** < 0.01 (**n** = 7 from three independent experiments). **g** Recombinant C100-flag was incubated in the absence or presence of increasing concentrations of purified CatB at 37 °C during 0 (To, 50 ng) or 60 min (5, 10 or 50 ng) and C100 levels were detected by western blot using α-Flag antibody. **h**, **i** SH-APPswe cells were treated with smer-28 (smer), PADK or bafilomycin A1 (BafA1), and APP-CTF levels were analyzed by western blot using α-APPct. Arrow corresponds to a non-identified 25 kDa APP-CTF fragment. **Bars** in **i** correspond to quantification of C83 and C99 immunoreactivities obtained in **h** and expressed as the percentage of the expressions in DMSO-treated cells normalized to actin. Data are represented as mean ± SEM, as determined by ANOVA one-way Dunnett post hoc test, **p** < 0.01 and *p* < 0.05 indicate significant differences relative to control cells **n** = 8, from three independent experiments.
Fig. 3 In vivo treatment of 3xTgAD mice with the γ-secretase inhibitor ELND006 (D6) leads to increased APP-CTFs localized to large cathepsin B-positive structures and to autophagic impairment. 5-month-old non-transgenic (nonTg) or 3xTgAD males were treated daily with D6 or vehicle by oral gavage during 1 month. a Immunostaining with FCA18 in the subiculum (see insert) of 3xTgAD vehicle- (AD-CT) or D6-treated (AD-D6) mice. Scale bar 125 and 20 μm, respectively. b, c RIPA-soluble (sol.) fractions from hippocampi of AD-CT or AD-D6 mice were analyzed for βAPP, APP-CTF, LC3-I, LC3-II and p62 expression by western blot. LC3-II was revealed after long time exposure of the same blot revealed for LC3-I. RIPA insoluble acid formic retrieved fractions (is) were analyzed for APP-CTF expression. Bars in e are the mean ± SEM of 12 animals of each treatment and represent the quantification of βAPP, C83, C99 and p62 immunoreactivities expressed as the percentage measured in AD-CT mice normalized to actin, and the LC3-II to LC3-I ratio normalized to AD-CT. Statistical analysis was performed by Mann–Whitney and **p < 0.01 indicate significant differences relative to AD-CT. d, e RIPA-soluble fractions from hippocampi of non-transgenic vehicle- (nonTg-CT) or D6-treated (nonTg-D6) mice were analyzed for APP-CTF, LC3-I, LC3-II and p62 expression by western blot. Right lane corresponds to AD-D6 mice. Bars in e are the mean ± SEM of six animals of each treatment and represent the quantification of C83 and p62 immunoreactivities expressed as the percentage measured in nonTg-CT mice. Statistical analysis was performed by Mann–Whitney and ***p < 0.001 indicate significant differences relative to nonTg-CT. f Images correspond to co-immunohistochemical staining with 82E1 and α-cathepsin B in subiculum of AD-D6 mice. Scale bar 20 μm.
should conversely expect to reduce C99 levels by enhancing lysosomal enzyme expression. Indeed, reduced APP-CTF levels were observed in both lentiviral-induced cathepsin B overexpressing cells (Fig. 2d) and in cells stably expressing this enzyme (Fig. 2e, f), as compared to mock-transfected cells. Furthermore, to demonstrate the ability of cathepsin B to catabolize C99 directly, we designed a bimolecular enzymatic assay in which various amounts of recombinant cathepsin B were incubated with recombinant C99 (C100-flag). This experiment confirmed the direct degradation of C99 by cathepsin B (Fig. 2g). Finally, we used a pharmacological approach to either activate or inhibit autophagy by using the small molecule enhancer of rapamycin, smer-28 [54] or the vacuolar-type H+ ATPase inhibitor, bafilomycin A1, respectively. As expected, APP-CTFs levels were reduced by smer-28 and enhanced by both bafilomycin A1 and the cathepsin B inhibitor PAKD (Fig. 2h, i). All together, these findings highlighted a critical role of lysosomal-autophagic function in APP-CTF clearance and clearly supported our hypothesis connecting impaired lysosomal-autophagic degradation and C99 accumulation.

**In vivo treatment of young 3xTgAD mice with the γ-secretase inhibitor ELND006 triggers massive increases in EAL-associated APP-CTFs and leads to autophagic pathology**

We then questioned whether C99 per se could contribute to lysosomal-autophagic dysfunction. To this end, we took advantage of a pharmacological approach in which we used the γ-secretase inhibitor ELND006 (D6) found to induce enhanced C99 (and C99-derived C83) levels [24]. Young mice were treated daily for 1 month with D6 and analyzed for C99 expression by immunohistochemistry (Fig. 3a, f) or western blot (Fig. 3b, c) as well as for lysosomal function (Fig. 3b, c). As expected, FCA18-associated immunolabeling showed a remarkable increase in lysosomal-associated C99 staining in brains of D6-treated mice (AD-D6) as compared to those of vehicle-treated mice (AD-CT) (Fig. 3a). This pharmacological treatment also drastically enhanced the number of enlarged cathepsin B-positive structures in C99-containing neurons indicating concomitant C99 accumulation and exacerbated lysosomal pathology (Fig. 3f). These observations were confirmed by western blot analysis revealing significant increases in the autophagic marker LC3-II and the autophagic substrate p62/SQSTM1 (Fig. 3b, c), thus indicating autophagic impairment [6]. In wild-type (nonTg) mice, the treatment with the inhibitor also led to increased levels of C83, which however remained very low as compared to those observed in AD-D6 mice (right lane in the western blot) (Fig. 3d, e). In both vehicle and D6-treated nonTg mice, LC3-II was not detected and p62 levels were unchanged (Fig. 3d, e) indicating the absence of effect of D6 on autophagic function in these mice.

In light of these observations, electron microscopy analysis (EM) was performed to confirm the above-described autophagic alterations. The subiculum of young AD vehicle-treated mice (AD-CT) displayed some small dense bodies (Fig. 4a, blue arrows) and various-sized dense lipid-containing autophagic vacuoles (Fig. 4a, red arrows) localized to perikarya, while age-matched vehicle-treated (nonTg-CT) (Suppl. Figure 2a) or D6-treated nonTg (nonTg-D6) neurons (Suppl. Figure 2b) presented very few of such structures. In D6-treated AD animals (AD-D6), EM revealed a strong autophagy-related pathology and intraneuronal damage (Fig. 4b, d–i and Suppl. Figure 2e–j). The number of small dense structures (<0.5 μm, blue arrows “1” in Fig. 4i) was identical in AD-CT and AD-D6 mice, but the number of larger sized autophagic vesicles, many containing lipids (>0.5 μm, red arrows “2” in Fig. 4i), was significantly increased in AD-D6 mice. However, the most striking feature of the autophagy-related pathology was the presence of multiple large vacuoles filled with heterogenous non-digested cargoes and multimellar bodies (blue ML) (Fig. 4d, e–i and Suppl. Figure 2e–j—red arrowheads), (red arrowheads “3” in Fig. 4i), probably corresponding to immature autophagic vesicles [62]. Moreover, many AD-D6-treated neurons exhibited large electron-lucent areas of the cytoplasm localizing close to autophagic vesicles, suggesting a link between autophagic leakage and...
erosive destruction (Fig. 4h, and Suppl. Figure 2f, j—blue star). D6-treated neurons also presented a high number of damaged mitochondria (see examples in Fig. 4d, g and Suppl. Figure 2g, i—black arrows) and reduced synaptic contacts within the neuropil (Fig. 4d, yellow arrows), as compared to control mice (Fig. 4c—yellow arrows).

We next compared the effect of D6-treatment on APP-CTF accumulation in 2xTgAD and 3xTgAD mice. This treatment led to same increases in C99 (and C99-derived C83) (Suppl. Figure 3a, b) and in punctiform staining (Suppl. Figure 3d) in the two strains, whereas the effect of D6 on Aβ load was detected only in 3xTgAD mice displaying enough Aβ to detect a decrease (Suppl. Figure 3c). These data fully agreed with our proposal of a dual effect of D6 on APP-CTF accumulation, one linked to its canonical inhibitory effect of γ-secretase and one to its inhibitory
effect on lysosomal degradation. In agreement with defective lysosomal degradation, both D6-treated 2xTgAD and 3xTgAD animals displayed enhanced, rather than lowered levels of the other γ-secretase cleavage-derived metabolite AICD (Suppl. Figure 3a, b) another lysosomal substrate (Fig. 2a) (see also [56]).

The γ-secretase inhibitor ELND006 triggers lysosomal APP-CTF accumulation, reduced cathepsin B activity and autophagic impairment in SH-APPswe cells, but not in SH-mock cells

To more precisely dissect the C99-mediated molecular mechanisms, we studied the effect of D6 treatment on lysosomal function in SH-APPswe cells. Firstly as expected, D6 triggered a drastic decrease in extracellular Aβ (not shown) and concomitant increase in C99 and C83 (Fig. 5a). Immunocytochemical analysis showed that these APP-CTFs were particularly abundant in lamp1-positive structures (Fig. 5b), which were enlarged and more numerous than those observed in control cells, similarly to cells treated with lysosomal inhibitors (Fig. 2c). These data were confirmed by Lysotracker-red, which labeled many and mostly large-sized puncta in D6-treated cells as compared to vehicle-treated cells (Fig. 5c). No increase in Lysotracker-red staining was seen in D6-treated SH-mock cells (i.e., cells only expressing endogenous levels of wild-type APP), suggesting a strong link between lysosomal dysfunction and APP-CTF accumulation. In addition, four independent lines of data supported our hypothesis, when we compared the effects of D6 and DAPT (another well-known γ-secretase inhibitor [58]) in SH-APPswe cells to SH-mock cells. Firstly, D6 and DAPT reduced in vitro cathepsin B activity (Fig. 5d) and lowered the levels of mature cathepsin B (Fig. 5e, f) in SH-APPswe but not in SH-mock cells (Fig. 5e), while NH4Cl strongly inhibited cathepsin B in both cell types (Fig. 5d–f). Moreover, in SH-APPswe cells both γ-secretase inhibitors led to the accumulation of a low-molecular weight (12 kDa) cathepsin B immunopositive product, probably corresponding to a degradation product of this enzyme (CatBdg) and supporting defective lysosomal clearance (Fig. 5e, f). Secondly, D6 and DAPT increased the levels of the autophagic markers LC3-II and p62 in only APPswe cells and the levels of these markers correlated with those of APP-CTFs (Fig. 5e, f). Thirdly, D6 treatment significantly increased LC3-positive vesicular profiles in SH-APPswe cells, whereas little if any LC3-GFP puncta were found in D6-treated SH-mock cells (Fig. 5g). Fourth, electron microscopy revealed the presence of large (0.5–2 μm) abnormal dense autolysosomal structures in D6-treated SH-APPswe cells, which structurally looked similar to those observed in leupeptin-treated cells (Fig. 5h).

Adeno-associated viral vector (AAV)-mediated expression of C99 in wild-type mice leads to the accumulation of aggregated C99 in enlarged cathepsin B and lamp1-positive structures

In order to definitely demonstrate that C99 per se is sufficient to induce lysosomal pathology, we used an AAV-mediated strategy to express C99 under the transcriptional control of the neuron-specific synapsin I in wild-type animals. AAV-C99, AAV-GFP- or AAV-empty was injected into the ventricle of newborn mice and C99 expression was analyzed at 2 months post-AAV delivery. Western blot revealed a significant and specific increase of C99 in AAV-C99 injected brains (Fig. 6a). As compared to AAV-GFP mice, in which GFP was observed in many neurons and throughout the brain (Fig. 6b, left panels), immunohistochemical analysis of C99 in AAV-C99 mice using α-APPct showed a more restricted expression that was particularly high in the hippocampus and neocortex (Fig. 6b, right panels). Within the neurons of these areas, the staining was intraneuronal and membrane-associated giving rise to a distinct immunostaining when compared to that of the cytoplasmic protein GFP (Fig. 6b). To discriminate between overexpressed C99 and endogenous APP and APP-CTFs that are also detected with the α-APPct antibody (recognizing both human and mouse APP, see supplementary table), we compared the immunostainings in AAV-C99 and AAV-empty mice. As expected, α-APPct detected membranous staining in many neurons and throughout the brain in AAV-empty mice; however, the intensity of this staining was much lower than that obtained in the AAV-C99 mice, indicating that the largest part of this staining corresponded to exogenously expressed C99. To determine the exact intraneuronal staining of C99, we performed immunocytochemical staining of C99 in cultured transfected cells that revealed that C99 was localized almost exclusively to the Golgi apparatus (Suppl. Figure 4a), whereas plasma membrane-associated C99 was also evident in few cells (Suppl. Figure 4b). As observed in SH-APPswe cells, the treatment with NH4Cl or D6 both led to the redistribution of APP-CTFs into EAL-associated structures and very little or no co-localization with GM130 was seen in these cells (Suppl. Figure 4c, d). In AAV-C99 brains, α-APPct did not detect EAL-associated staining, but interestingly when we used FCA18 or 82E1, the staining was punctiform and looked similar to that observed in the 3xTgAD mice. This labeling was present in the same brain regions labeled with α-APPct, but was localized to a much more restricted number of neurons and most strongly in the subiculum (Fig. 6d) and in some layers of the cortex. Interestingly, the other N-terminal-directed antibodies 6E10 and 4G8 recognized both the membrane-associated labeling and puncta (Fig. 6d,
Fig. 5 γ-Secretase inhibitor treatment triggers autophagic dysfunction in APPswe-expressing SHSY-5Y but not mock-transfected cells. a SH-APPswe cells were treated with ELND006 (D6) and analyzed for APP-CTF levels by western blot. APP-CTFs were detected using α-APPct, 4G8 or 6E10 (see diagram for epitope recognition) and compared to C99-flag expression in HEK transfected cells. b Co-immunocytochemical staining of vehicle (Veh) or D6-treated APP-swe cells using α-APPct and α-lamp1 shows a high overlap of these labelings. Scale bar 2 μm. c Vehicle (V) or D6-treated SH-mock or SH-APPswe cells were stained with Lysotracker red and DAPI. Scale bar 5 μm. d Cathepsin B (CatB) activity was monitored in vitro from microsomal fractions of SH-APPswe cells treated with vehicle (V), D6, DAPT (DT) or NH₄Cl and western blot analysis was used to determine the levels of βAPP and APP-CTFs, LC3-I, LC3-II, p62, CatB and actin. Besides the detection of the 30 kDa immunoreactivity corresponding to mature CatB, the CatB antibody revealed immunoreactivity at about 12 kDa most probably corresponding to a degradation product of this enzyme (CatB-dg). e, f Stably CatB expressing mock or APPswe cells were treated with vehicle (V), D6, DAPT (DT) or NH₄Cl and western blot analysis was used to determine the levels of βAPP and APP-CTFs, LC3-I, LC3-II, p62, CatB and actin. Data are represented as mean ± SEM, as determined by ANOVA one-way Dunnett post hoc test, ***p < 0.001 indicate significant differences relative to control cells (vehicle-treated cells), n = 8–14 from 2 independent experiments. e, f Stably CatB expressing mock or APPswe cells were treated with vehicle (V), D6, DAPT (DT) or NH₄Cl and western blot analysis was used to determine the levels of βAPP and APP-CTFs, LC3-I, LC3-II, p62, CatB and actin. Besides the detection of the 30 kDa immunoreactivity corresponding to mature CatB, the CatB antibody revealed immunoreactivity at about 12 kDa most probably corresponding to a degradation product of this enzyme (CatB-dg). f Quantification of immunoreactivities obtained in e for APPswe cells were normalized to actin and expressed as the percent of expression levels obtained in control treated cells. g SH-mock or SH-APPswe cells were transfected with LC3-GFP, treated with vehicle or D6 and was visualized by fluorescence microscopy. Scale bar 5 μm. h Electron microphotographs show representative cells from each condition. Scale bars are 1 or 2 μm, as indicated.
e—arrow and arrowhead, respectively) suggesting the presence of two distinct C99 species. Since, the lysosomal-associated staining required formic acid retrieval and since C99 has been shown to have a high tendency to self-aggregate and form amyloid-like fibrils in vitro [30, 60], we hypothesized that the C99 accumulating in lysosomes could correspond to aggregated or misfolded forms that would not be detected with C-terminal-directed antibodies because of aggregation-dependent structural or conformational changes of the C-terminal moiety. In agreement with this hypothesis, we found that the punctiform staining was also detected with the aggregate-specific antibody NU1 [23] (Fig. 6d). As observed in the 3xTgAD mice, these aggregates were localized to enlarged cathepsin B- and
for 3xTgAD mice, C99-expressing mice were treated with D6. As described above, C99 accumulated in enlarged cathepsin B-positive structures. 2-month-old AAV-empty, AAV-GFP or AAV-C99 were analyzed by western blot (a) or immunohistochemistry (b). a RIPA-soluble fractions from forebrain hemispheres were analyzed for APP-CTF expression using either α-APPct or 6E10. b GFP fluorescence was observed in AVV-GFP mice using a confocal microscope and C99 expression was visualized after immunohistochemical staining with α-APPct and using AAV-empty mice as a negative control. The lower panel shows higher magnification images at the level of the boxed areas highlighting the different subcellular labeling of GFP and C99. c Bars correspond to ELISA measurement of Aβ40 and Aβ42 in soluble (sol) and insoluble (is) fractions, respectively, and are represented as mean ± SEM n = 6 animals. d Images from immunohistochemical analysis of C99-associated expression in C99-AAV mice at the level of the subiculum using α-APPct, FCA18, 6E10, 4G8 or NU1. Scale bar 25 μm. e High-magnification images of co-immunolabeling with α-APPct and 4G8 illustrating the intraneuronal membrane-associated (arrow) and punctiform (arrowhead) stainings. Scale bar 5 μm. f, g Co-labeling of 82E1 and cathepsin B (f) or 4G8 and lamp1 (g) shows that C99 localizes to enlarged EAL-associated structures (arrowhead in merged images). Scale bar 25 and 5 μm, respectively. h Quantitative analysis of lamp1 structures in the subiculum of AAV-empty or AAV-C99 mice. Data are represented as the relative number per cell (counts (a.u)/cell) and average size of lamp1-positive puncta expressed as arbitrary units (a.u.). Data are represented as mean ± SEM and are from two AAV-empty and two AAV-C99 mice, three brain slices for each animal and six images each slice. Images show representative images of lamp1 staining.

lamp1-positive structures (Fig. 6f, g) and the number and size of such enlarged structures was enhanced in the AAV-C99 mice, as compared to AAV-empty mice (Fig. 6h).

ELISA showed that C99-expressing mice do produce Aβ, corresponding mainly to Aβ40 but only very low levels of the aggregation-prone Aβ42 (Fig. 6c). Therefore, one could not exclude the possibility, that in this model, the punctiform staining could correspond to aggregated Aβ, instead of C99. Thus, to rule out this possibility, the C99-expressing mice were treated with D6. As described for 3xTgAD mice, γ-secretase inhibition enhanced the levels of C99 and C99-derived C83 [16] (Fig. 7a, but see also Fig. 8b) and strongly reduced Aβ levels (Fig. 7b). Immunohistochemical analysis using NU1 showed an increased rather than decreased punctiform immunostaining in D6-treated mice, clearly indicating that this staining should be accounted for C99 and not Aβ (Fig. 7c). Similar enhanced punctiform staining was observed with other N-terminal directed antibodies, 82E1 (Fig. 8a), FCA18, 6E10 (not shown) and 4G8 (Fig. 7d). In D6-treated mice, in areas expressing particular high punctiform staining, NU1 and 4G8 also gave rise to an extracellular diffuse staining surrounding the neurons (Fig. 7c, d, respectively), probably corresponding to the release of intraneuronal material from dying neurons (see also below). We also used α-APPct to analyze the effect of D6-treatment (Fig. 7c, e) and found that, within the subiculum, D6 also led to a strong APP-CTF associated staining localizing to synaptic regions (note the high overlap of α-APPct and the presynaptic marker synaptophysin in D6 treated AAV-C99 mice (Fig. 7e). The low intensity of α-APPct labeling in D6-treated AAV-empty mice showed that these synaptic localized APP-CTFs were mostly linked to viral-mediated C99. These observations confirmed the presence of two species of C99 (APP-CTFs), one non-aggregated and recognized by α-APPct and one aggregated and detected with N-terminal- and aggregate-specific antibodies. D6 treatment led to increased levels of both non-aggregated APP-CTFs localizing in presynaptic regions and of aggregated APP-CTFs accumulating in lysosomal-derived structures. It is noteworthy that similar D6-induced increases in both synaptophysin-positive APP-CTFs (Suppl. Figure 5a, d–e) and NU1-labeled aggregated APP-CTFs (Suppl. Figure 5b) was found in 3xTgAD mice, thus ruling out the possibility of artificial routing of C99 in the C99-expressing mice model. Moreover, western blot analysis confirmed the accumulation of C99 and C83 in both microsome- and synaptosome-enriched fractions of hippocampi from D6 treated 3xTgAD mice (suppl. Figure 5c).

Furthermore, and more surprisingly, in the C99-expressing mouse, D6 not only led to an enhanced staining in the cortex and subiculum, the two main regions expressing C99, but it also induced a strong immunostaining in various other brain regions, in which very low or no C99 staining was seen in vehicle-treated AAV-C99 mice (Fig. 7f), but in which GFP expression was high in AAV-GFP mice (Fig. 6b, left panels). Again, the labeling obtained with N-terminal antibodies was punctiform and in some regions also diffuse and extracellular. Taken together, these findings clearly demonstrated that this labeling should be ascribed to C99 and not to Aβ. However, they also suggested that these regions display either higher γ-secretase activity or alternative D6-sensitive C99 degradation, explaining the absence of C99 labeling in vehicle-treated C99-expressing mice.

**D6 treatment not only leads to autophagic dysfunction but also to inflammatory responses and synaptic defects in C99-expressing mice, but not in wild-type mice**

As described above, C99 accumulated in enlarged cathepsin B- and lamp1-positive structures and particularly within the subiculum in C99-expressing mice. In brains from D6-treated AAV-C99 mice, the number of neurons displaying such enlarged structures was increased rather than decreased (Fig. 8a). Furthermore, these brains displayed elevated levels of the autophagic marker LC3-II (Fig. 8b, c), while this was not the case in those from D6-treated AAV-GFP mice (Fig. 8d), again indicating that these lysosomal alterations were linked to C99 expression.
Moreover, as described above, in some regions in which lysosomal-associated C99 was abundant, 4G8 also unraveled an important and diffuse extracellular staining likely corresponding to the release of intracellular material from dying neurons (Fig. 8e, f, but see also Fig. 7c, d). In agreement with necrotic cell death, nuclear staining with DAPI or Cresyl violet revealed the presence of many pycnotic nuclei (Fig. 8e, f). The same regions also displayed an increased number of both Iba-1-positive microglia (Fig. 8g) and GFAP-positive astrocytes (Fig. 8h). Iba-1 labeling was enhanced by the sole expression of C99 (compare AAV-C99 and AAV-empty) and further enhanced by D6 treatment, indicating concomitant C99 accumulation and inflammatory responses (Fig. 8g). Importantly, microglial...
and astrocytic activation (Fig. 8h) were not induced by D6 in AAV-empty mice. Finally, we assessed the effects of C99 expression on synaptic function by measuring long-term potentiation (LTP) at the CA1 pyramidal neurons in subicular neurons in AAV-empty or AAV-C99 vehicle or D6-treated mice (Fig. 8i–k). Whereas no differences were seen in basal synaptic transmission between AAV-empty, AAV-C99 vehicle or D6-treated mice (Fig. 8i), only the AAV-empty mice displayed a robust induction in LTP (Fig. 8j, k). D6 did not worsen LTP alterations, but this could be due to the drastic floor effect of C99 over-expression alone. However, the fact that D6 did not reverse synaptic dysfunction indicated that it was not caused by Aβ, but should be directly linked to C99.

Discussion

Recently, we showed that the 3xTgAD mouse model develops an early, age-dependent intraneuronal accumulation of the β-secretase-derived βAPP fragment C99 [24]. In the present work, we demonstrate a link between this accumulating C99 and endosomal-lysosomal dysfunction (EAL) pathology. Firstly, the comparison of 2xTgAD (harboring wild-type PS1) and 3xTgAD mice (harboring PS1M146V) revealed that C99 accumulation was the consequence of impaired lysosomal-lysosomal fusion rather than of altered γ-secretase processing and in agreement with this, C99 was found to accumulate within endolysosomes in both of these mice. Moreover, our in vitro data and earlier work [1, 8, 57] demonstrated a major role of the lysosomal-lysosomal pathway in C99 degradation. The 3xTgAD mice harbor PS1 and tau mutations, and the 2xTgAD mice harbor mutated tau, which possibly could have interfered with C99/APP trafficking and degradation, as described for some PS1 and tau mutants [25, 63]. Therefore, to determine whether EAL-associated C99 accumulation could be seen also in a model devoid of any mutated transgenes, we generated a viral-based C99-expressing transgenic model. Interestingly, in these mice immunohistochemical studies revealed the presence of two clearly distinct C99 species. One of it was intraneuronal membrane-associated and mainly localizing to the golgi apparatus. This C99 was recognized by α-APPct and representing the major part of C99. Another one was localized to EAL structures and restricted to a subgroup of the neurons. This C99 was aggregated or misfolded and recognized by both N-terminal-directed and aggregate-specific antibodies. Also in these mice, many of the C99-accumulating EAL vesicles were enlarged, again implying a link between C99 and EAL pathology. Taken together, these data ruled out the possibility that C99 aggregation and lysosomal localization was linked to artificial cellular trafficking caused by the expression of other mutated transgenes or to a putative lack of adequate targeting of exogenously expressed C99. We cannot exclude that the overexpression of these proteins itself could also have influenced degradation, but importantly elevated C99 levels have also been reported in human AD post-mortem brains [18, 22, 42], indicating that C99 accumulation is not specific to mouse AD models, but also do occur in human AD.

The EAL-associated C99 was recognized by the aggregate-specific antibody NU1 and localized to the membranes of these vesicles. We therefore hypothesized that this aggregated or misfolded C99 itself could also be a trigger of lysosomal-autophagic dysfunction. To validate this hypothesis, we took advantage of a pharmacological approach in which 3xTgAD or C99-expressing mice were treated with the γ-secretase inhibitor, ELND006 (D6) that enhances C99 levels [24]. Indeed, in both mouse models, D6 treatment increased EAL-associated C99 levels and led to exacerbated pathology: elevated levels of the autophagic markers LC3-II and p62, increased number of abnormalized C99-positive cathepsin B- and lamp1-structures and a higher density of non-digested autophagic vesicles highlighted by electron microscopy. Ultrastructural analysis also revealed the presence in neuropil of large autophagic vesicles filled with aberrant storage material likely corresponding to immature AVs, suggesting that, in addition to the effect on lysosomal proteolysis, γ-secretase inhibition also led to disturbed AV maturation and/or transport. Importantly, these effects of D6 were not observed in wild-type animals, strongly suggesting that they were dependent
Figure 1: Examples of morphological changes and protein expression changes in AAV-C99-D6 versus AAV-C99-Veh.

**a)** Representative images showing the effect of AAV-C99-D6 treatment on cellular morphology compared to AAV-C99-Veh.

**b)** Western blot analysis showing differences in protein expression between AAV-C99-Veh and AAV-C99-D6.

**c)** Graph depicting the LC3-I/LC3-II ratio, with a significant increase in AAV-C99-D6 compared to AAV-C99-Veh.

**d)**Additional Western blot analysis showing protein expression changes.

**e)** Immunofluorescence images showing cellular localization of specific proteins in AAV-C99-D6 versus AAV-C99-Veh.

**f)** Additional immunofluorescence images with different staining conditions.

**g)** Further immunofluorescence images with Iba-1 staining for macrophage/microglia activation.

**h)** GFAP staining to assess astrocyte activation.

**i)** Graph showing changes in EPSP slope over time with various treatments.

**j)** Additional graph showing EPSP slope changes under different conditions.

**k)** Bar graph comparing average EPSP slope over the last 40-60 minutes, with statistical significance indicated.

All images are labeled with appropriate scales and markers to ensure clarity and precision in the representation of the experimental results.
Fig. 8 γ-Secretase inhibitor treatment of AAV-C99 expressing mice leads to exacerbated autophagic dysfunction, inflammatory responses and synaptic dysfunction. a Brain slices at the levels of the subiculum from vehicle- (Veh) or D6-treated AAV-C99 mice were immunostained with 82E1 and α-CatB. Scale bar 25 μm. b–d Western blot analysis of RIPA-soluble fractions from AAV-C99 injected mice (b, e) or AAV-GFP mice (d) treated with D6 or vehicle (Veh). Brains were analyzed for APP-CTF (using either α-APPct or 6E10) or LC3-II/LC3-I expression. Bars in c correspond to the quantitative analysis of LC3-II, expressed as the LC3-II/LC3-I ratio, and are relative to control (AAV-C99-Veh). Mann–Whitney test, p < 0.001, n = 8. Right lane corresponds to AAV-C99 mice. e, f Immunostaining with 4G8 or NU1 of D6-treated AAV-C99 brains using fluorescence (e) or peroxidase-DAB labeling (f). Note the extracellular staining (white arrows in e and black arrows in f) surrounding pycnotic nuclei visualized by DAPI (e, white arrowheads) or Cresyl violet (f, black arrowheads). Scale bars 50 or 10 μm, respectively. g, h Immunostaining with 4G8, Iba1 (g) or GFAP (h) reveals both microglial and astrocytic activation in D6-treated mice within brain regions expressing important APP-CTF levels. Scale bar 300 and 50 μm, respectively. i Basal synaptic transmission in AAV-empty, vehicle-treated AAV-C99 or D6-treated AAV-C99 mice (n = 2–3 slices per mouse from 6 to 7 mice per group. All values are mean ± SEM. j Field excitatory postsynaptic potential (fEPSP) slopes in subiculum (n = 2–3 slices per mouse from 6 to 7 mice per group). k Summary graph of LTP magnitudes calculated 40 to 60 min after high-frequency stimulation from graphs in (j) with statistical analysis (*p < 0.05; one-way ANOVA with the Tukey’s post hoc test). Error bars represent SEM

Overall, the above set of data unveiled a pathological loop in which C99 accumulation was not only the result of altered autophagy, but could also be the cause of it. The observations that aggregated C99 accumulated within the membranes of the EAL vesicles suggested that it likely interfered directly with EAL function by perturbing membrane integrity and inhibiting normal lysosomal proteolysis. Possibly exacerbated C99 accumulation could also lead to the leakage of catabolic contents, lysosomal enzymes and acidity into the cytoplasm and to resulting neuronal cell death [17, 28]. Indeed, both electron microscopy and immunohistochemical analysis revealed clear signs of neuronal necrosis and release of aggregated C99 (and C99-derived C83) to extracellular spaces. Strikingly, recent papers described similar deleterious effects of EAL-associated Aβ42 in a transgenic Aβ42-expressing drosophila model [26–28], suggesting that whatever the nature of the aggregates, their accumulation within EAL-vesicle membranes could disturb lysosomal-autophagic function. In the D6-treated AAV-C99 mice, C99 accumulation was also associated with strong inflammatory responses including both microglial activation and astrogliosis. These findings showed that the pathology observed in the C99 mice resembled that seen in lysosomal storage disorders, in which a primary defect of lysosomal function induces both autophagic impairment and the activation of the immune system as a response to the accumulation of storage material and/or dying neurons [44]. Interestingly, we also observed that C99 expression strongly affected synaptic function. Long-term potentiation recordings in the subiculum of the AAV-injected mice, showed an almost complete absence of LTP in the subiculum of C99-expressing mice. D6 did not reverse this effect clearly indicating that it was linked to C99 expression itself and not to Aβ.

Altogether, our data showed a direct and clearly Aβ-independent detrimental effect of C99 and provide new molecular explanations for earlier mechanistically unsolved studies describing APP-CTF-linked toxicity. For instance, it was reported that the conditional knock-out of presenilin1 in mice led to a progressive development of synaptic and cognitive alterations that were temporally correlated with APP-CTF accumulation [48]. Moreover, the treatment with γ-secretase inhibitors was found to induce both long-term deficits and APP-CTF accumulation in mice [31, 52] and to accelerate cognitive decline in human [15]. Consistent with these findings, the work from the group of Dr d’Adamo showed that the inhibition of β-secretase but not of γ-secretase led to reduced synaptic/memory defects in a mouse model of familial dementia [53]. Finally, the transgenic overexpression of BACE-1 in mice was observed to induce a pathogenic pathway involving the accumulation of APP-CTFs but not of Aβ [46]. Recent work from the group of Dr Nixon proposed a direct link between C99 accumulation and endocytic abnormalities in fibroblasts derived from Down’s syndrome patients [20, 22], but these studies did not investigate the downstream effects of C99 accumulation on lysosomal and autophagic function.

This work is the first to demonstrate the presence of two structurally and immunologically distinguishable C99 species, one corresponding to non-aggregated and intraneuronal membrane-bound C99 and one to aggregated or misfolded C99 localized to EAL-vesicle membranes. The pharmacological inhibition of γ-secretase led to drastic increases in both non-aggregated C99 accumulating within synaptic regions, and in aggregated EAL-associated C99. These findings therefore reconcile previous works demonstrating different localizations of APP-CTFs. Whereas some studies reported presynaptic APP-CTF accumulation after γ-secretase inhibition [5, 31] or in presenilin knock-out mice [48], others found lysosomal APP-CTF accumulation in presenilin-invalidated cultured cells [14].

Altogether, our findings add a new toxic trigger to the amyloid cascade, which could underlie some of the AD-related hallmarks occurring especially in the early stages of the disease. In the 3xTgAD mouse, the intraneuronal C99 accumulation occurs earlier and is more important than that
of $\beta$ and in the AAV-C99 mice, the aggregation-prone A$\beta_{42}$ is only produced in very low amounts, compared to that of $\beta_{40}$. These findings might explain the major contribution of C99 to the observed phenotypes in our models, but of course we do not exclude the contribution of $\beta$ to the pathology. Finally, our data also show a region-specific accumulation of C99, indicating differences in proteolytic processing and degradative mechanisms within neuronal subtypes. These differences could explain the accumulation of distinct catabolites in specific neuronal populations and underlie mechanisms contributing to neuronal susceptibility to AD-related pathology. Taken together, our study demonstrates that C99 can participate to lysosomal pathology in the absence of $\beta$.

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Compliance with ethical standards

Ethical approval All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted.

Conflict of interest The authors declare no competing financial interests.

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References

1. Asai M, Yagishita S, Iwata N, Saito TC, Ishiura S, Maruyama K (2011) An alternative metabolic pathway of amyloid precursor protein C-terminal fragments via cathepsin B in a human neuro-gliaoma model. FASEB J 25:3720–3730
2. Baranger K, Marchalant Y, Bonnet AE, Crouzin N, Carrette A, Paumier JM, Py NA, Bernard A, Bauer C, Charrat E, Moschke K, Seiki M, Vignes M, Lichtenhaler SF, Checler F, Khrestchatisky M, Rivera S (2015) MT5-MMP is a new pro-amyloidogenic proteinase that promotes amyloid pathology and cognitive decline in a transgenic mouse model of Alzheimer’s disease. Cell Mol Life Sci 73:217–236
3. Barelli H, Lebeau A, Vizzavona J, Delaere P, Chevallier N, Drouot C, Marambaud P, Ancolio K, Buxbaum JD, Khorkova O, Heroux J, Sahasrabudhe S, Martinez J, Wartier JM, Mohr M, Checler F (1997) Characterization of new polyclonal antibodies specific for 40 and 42 amino acid-long amyloid beta peptides: their use to examine the cell biology of presenilins and the immunohistochemistry of sporadic Alzheimer’s disease and cerebral amyloid angiopathy cases. Mol Med 3:695–707
4. Benkhelifa-Ziyat S, Besse A, Roda M, Duque S, Astord S, Carron M, Marais T, Barkats M (2013) Intramuscular scAAV9-SMN injection mediates widespread gene delivery to the spinal cord and decreases disease severity in SMA mice. Mol Ther 21:282–290
5. Bittner T, Fuhrmann M, Burgold S, Jung CK, Volbracht C, Steiner H, Mittwegger G, Kretzschmar HA, Haass C, Herm J (2009) Gamma-secretase inhibition reduces spine density in vivo via an amyloid precursor protein-dependent pathway. J Neurosci 29:10405–10409
6. Bjorkoy G, Lamark T, Brech A, Outzen H, Perander M, Overvatn A, Stenmark H, Johansen T (2005) p62/SQSTM1 forms protein aggregates degraded by autophagy and has a protective effect on huntingtin-induced cell death. J Cell Biol 171:603–614
7. Blennow K, de Leon MJ, Zetterberg H (2006) Alzheimer’s disease. Lancet 368:387–403
8. Boland B, Smith DA, Mooney D, Jung SS, Walsh DM, Platt FM (2010) Macroautophagy is not directly involved in the metabolism of amyloid precursor protein. J Biol Chem 285:37415–37426
9. Brigham E, Quinn K, Kwong G, Willits C, Goldbach E, Motter R, Lee M, Ho K, Wallace, Khodolenko D, Tanaka P, Ni H, Hemphill S, Chen X, Eichenbaum T, Ruslin L, Nguyen L, Santiago P, Liao A, Soriano F, Boja M, Probst G, Dappen M, Latimer L, Jagodzinski J, Konrady A, Garofalo A, Webb S, Sham H, Wehner N, Tonn G, Sauer J, Basi G, Ness D (2010) Pharmacokinetic and pharmacodynamic investigation of ELND006, a novel APP-selective gamma-secretase inhibitor, on amyloid concentrations in the brain, CSF and plasma of multiple nonclinical species following oral administration. In: International Conference on Alzheimer’s Disease
10. Caccamo A, De Pinto V, Messina A, Branca C, Oddo S (2014) Genetic reduction of mammalian target of rapamycin ameliorates Alzheimer’s disease-like cognitive and pathological deficits by restoring hippocampal gene expression signature. J Neurosci 34:7988–7998
11. Caccamo A, Majumder S, Richardson A, Strong R, Oddo S (2010) Molecular interplay between mammalian target of rapamycin (mTOR), amyloid-beta, and Tau: effects on cognitive impairments. J Biol Chem 285:13107–13120
12. Cataldo AM, Petanceska S, Terio NB, Peterhoff CM, Durham R, Mercken M, Mehta PD, Buxbaum J, Haroutunian V, Nixon RA (2004) Abeta localization in abnormal endosomes: association with earliest Abeta elevations in AD and Down syndrome. Neurobiol Aging 25:1263–1272
13. Cataldo AM, Peterhoff CM, Schmidt SD, Terio NB, Duff K, Beard M, Mathews PM, Nixon RA (2004) Presenilin mutations in familial Alzheimer disease and transgenic mouse models accelerate neuronal lysosomal pathology. J Neuraphathol Exp Neurol 63:821–830
14. Chen F, Yang DS, Petanceska S, Yang A, Tandon A, Yu G, Rozmahel R, Ghiso J, Nishimura M, Zhang DM, Kawarai T, Levesque G, Mills J, Levesque L, Song YQ, Rogaeva E, Westaway D, Mount H, Gandy S, St George-Hyslop P, Fraser PE (2000) Carboxyl-terminal fragments of Alzheimer beta-amyloid precursor protein accumulate in restricted and unpredicted intracellular compartments in presenilin 1-deficient cells. J Biol Chem 275:36794–36802
15. Doody RS, Raman R, Farlow M, Iwatsubo T, Velas B, Joffe S, Kieburtz K, He F, Sun X, Thomas RG, Aisen PS, Siemens E,
Sethuraman G, Mohs R (2013) A phase 3 trial of semagacestat for treatment of Alzheimer’s disease. N Engl J Med 369:341–350

16. Flammang B, Pardossi-Piquard R, Sevalle J, Debaye D, Dabet-Gay AS, Thevenet A, Lauritzen I, Checler F (2012) Evidence that the amyloid-beta protein precursor intracellular domain, AICD, derives from beta-secretase-generated C-terminal fragment. J Alzheimers Dis 30:145–153

17. Glabe C (2001) Intracellular mechanisms of amyloid accumulation and pathogenesis in Alzheimer’s disease. J Mol Neurosci 17:137–145

18. Hebert SS, Horre K, Nicolai L, Papadopoulou AS, Mandemakers W, Silhataroglu AN, Kauppinen S, Delacourte A, De Strooper B (2008) Loss of microRNA cluster miR-29a/b-1 in sporadic Alzheimer’s disease correlates with increased BACE1/beta-secretase expression. Proc Natl Acad Sci USA 105:6415–6420

19. Horikoshi Y, Sakaguchi G, Becker AG, Gray AJ, Duff K, Aisen PS, Yamaguchi H, Maeda M, Kinoshita N, Matsuoka Y (2004) Development of Abeta terminal end-specific antibodies and sensitive ELISA for Abeta variant. Biochem Biophys Res Commun 319:733–737

20. Jiang Y, Mullaney KA, Peterhoff CM, Che S, Schmidt SD, Boyer-Boiteau A, Ginsberg SD, Cataldo AM, Mathews PM, Nixon RA (2010) Alzheimer’s-related endosome dysfunction in Down syndrome is Abeta-independent but requires APP and is reversed by BACE-1 inhibition. Proc Natl Acad Sci USA 107:1630–1635

21. Kim JY, Ash RT, Ceballos-Diaz C, Levites Y, Golde TE, Smirnakis SM, Jankowsky JL (2013) Viral transduction of the neuronal brain delivers controllable genetic mosaicism for visualising and manipulating neuronal circuits in vivo. Eur J Neuroscience 37:1203–1220

22. Kim S, Sato Y, Mohan PS, Peterhoff C, Pensalini A, Rigoglioso A, Jiang Y, Nixon RA (2016) Evidence that the rab5 effector APPL1 mediates APP-betaCTF-induced dysfunction of endosomes in Down syndrome and Alzheimer’s disease. Mol Psychiatry 21:707–716

23. Lambert MP, Velasco PT, Chang L, Viola KL, Fernandez S, Lacor PN, Khuon D, Gong Y, Bigio EH, Shaw P, De Felice FG, Kraft GA, Klein WL (2007) Monoclonal antibodies that target pathological assemblies of Abeta. J Neurochem 100:23–35

24. Lauritzen I, Pardossi-Piquard R, Bauer C, Brigham E, Abraham JD, Ranaldi S, Fraser P, St-George-Hyslop P, Le Thuc O, Espin V, Chami L, Dunys J, Checler F (2012) The beta-secretase-derived C-terminal fragment of betaAPP, C99, but not Abeta, is a key contributor to early intraneuronal lesions in triple-transgenic mouse hippocampus. J Neurosci 32:16243–16255

25. Lazarov O, Morfini GA, Pignino G, Gadadhar A, Chen X, Robinson J, Ho H, Brady ST, Sisodia SS (2007) Impairments in fast axonal transport and motor neuron deficits in transgenic mice expressing familial Alzheimer’s disease-linked mutant presenilin 1. J Neurosci 27:7011–7020

26. Ling D, Magallanes M, Salvaterra PM (2014) Accumulation of amyloid-like fibrils in COS cells overexpressing part of the Alzheimer amyloid protein precursor. Nature 347:566–569

27. Mitani Y, Yarimizu J, Saita K, Uchino H, Akashiba H, Shitaka Y, Ni K, Matsuoka N (2012) Differential effects between gamma-secretase inhibitors and modulators on cognitive function in amyloid precursor protein-transgenic and nontransgenic mice. J Neurosci 32:2037–2050

28. Mueller-Steiner S, Zhou Y, Arai H, Roberson ED, Sun B, Chen J, Wang X, Yu G, Esposito L, Mucke L, Gan L (2006) Antiamyloidogenic and neuroprotective functions of cathepsin B: implications for Alzheimer’s disease. Neurology 51:703–714

29. Nixon RA (2007) Autophagy, amyloidogenesis and Alzheimer disease. J Cell Sci 120:4081–4091

30. Nixon RA, Cataldo AM, Mathews PM (2000) The endosomal-lysosomal system of neurons in Alzheimer’s disease: a review. Neurochem Res 25:1161–1172

31. Nixon RA, Yang DS (2011) Autophagy failure in Alzheimer’s disease—locating the primary defect. Neurobiol Dis 43:38–45

32. Nixon RA, Yang DS (2012) Autophagy and neuronal cell death in neurological disorders. Cold Spring Harb Perspect Biol 4(10). doi:10.1101/cshperspect.a008839

33. Oddo S, Caccamo A, Shepherd JD, Murphy MP, Golde TE, Kayed R, Metherate R, Mattson MP, Akbari Y, LaFerla FM (2003) Triple-transgenic model of Alzheimer’s disease with plaques and tangles: intracellular Abeta and synaptic dysfunction. Neuron 39:409–421

34. Oddo S, Caccamo A, Tseng B, Cheng D, Vasilevko V, Cribbs DH, LaFerla FM (2008) Blocking Abeta42 accumulation delays the onset and progression of tau pathology via the C terminus of heat shock protein70-interacting protein: a mechanistic link between Abeta and tau pathology. J Neurosci 28:12163–12175

35. Orr ME, Oddo S (2013) Autophagic/lysosomal dysfunction in Alzheimer’s disease. Alzheimers Res Ther 5:53

36. Oules B, Del Prette D, Greco B, Zhang X, Lauritzen I, Sevalle J, Moreno S, Paterlini-Brechot P, Trebak M, Checler F, Benfenati F, Chami M (2012) Ryanodine receptor blockade reduces amyloid-beta load and memory impairments in Tg2576 mouse model of Alzheimer disease. J Neurosci 32:11820–11834

37. Pardossi-Piquard R, Lauritzen I, Bauer C, Sacco G, Robert P, Checler F (2016) Influence of genetic background on apathy-like behavior in triple transgenic AD mice. Acta Neuropathol 123:1–8. doi:10.1186/s13287-015-0203-3

38. Pera M, Alcolea D, Sanchez-Valle R, Guardia-Laguarta C, Colom-Cadena M, Badiola N, Suarez-Calvet M, Llado A, Barrera-Ocampo AA, Sepulveda-Falla D, Blesa R, Molinuevo JL, Clarinjon I, Ferrer I, Lelief E, Lleo A (2013) Distinct patterns of APP processing in the CNS in autosomal-dominant and sporadic Alzheimer disease. Acta Neuropathol 125:201–213

39. Peric A, Annaert W (2015) Early etiology of Alzheimer’s disease: tipping the balance toward autophagy or endosomal dysfunction? Acta Neuropathol 129:363–381

40. Platt N, Speak AO, Colaco A, Gray J, Smith DA, Williams IM, Wallom KL, Platt FM (2016) Immune dysfunction in Niemann-Pick disease type C. J Neurochem 136(Suppl 1):74–80

41. Probst G, Aubele DL, Bowers S, Dressen D, Bova MP, Hemphill SS, Nguyen L, Ruslim L, Tanaka F, Wallace Y, Yednock TA, Basi GS (2013) Discovery of (R)-4-cyclopropyl-7,8-difluoro-5-(4-(trifluoromethyl)
phenylsulfonyl)-4,5-dihydro-1H-pyrazolo[4,3-c]quinoline (ELND006) and (R)-4-cyclopropyl-8-fluoro-5-[(6-(trifluoromethyl)pyridin-3-yl)sulfonyl]-4,5-dihydro-2H-pyrazolo[4,3-c]quinoline (ELND007): metabolically stable gamma-secretase Inhibitors that selectively inhibit the production of amyloid-beta over Notch. J Med Chem 56:5261–5274

46. Rockenstein E, Mante M, Alford M, Adame A, Crews L, Hashimoto M, Esposito L, Mucke L, Masliah E (2005) High beta-secretase activity elicits neurodegeneration in transgenic mice despite reductions in amyloid-beta levels: implications for the treatment of Alzheimer disease. J Biol Chem 280:32957–32967

47. Sanchez-Varo R, Trujillo-Estrada L, Sanchez-Mejias E, Torres M, Baglietto-Vargas D, Moreno-Gonzalez I, De Castro V, Jimenez S, Ruano D, Vizuete M, Davila JC, Garcia-Verdugo JM, Jimenez AJ, Vitorica J, Gutierrez A (2011) Abnormal accumulation of autophagic vesicles correlates with axonal and synaptic pathology in young Alzheimer’s mice hippocampus. Acta Neuropathol 123:53–70

48. Saura CA, Chen G, Malkani S, Choi SY, Takahashi RH, Zhang D, Gouras GK, Kirkwood A, Morris RG, Shen J (2005) Conditional inactivation of presenilin 1 prevents amyloid accumulation and temporarily rescues contextual and spatial working memory impairments in amyloid precursor transgenic mice. J Neurosci 25:6755–6764

49. Sevalle J, Amoyel A, Robert P, Fournie-Zaluski MC, Roques B, Checler F (2009) Aminopeptidase A contributes to the N-terminal truncation of amyloid beta-peptide. J Neurochem 109:248–256

50. Shacka JJ, Roth KA, Zhang J (2011) The autophagy-lysosomal degradation pathway: role in neurodegenerative disease and therapy. Front Biosci 13:718–736

51. Spilman P, Podlutskaya N, Hart MJ, Debnath J, Gorostiza O, Martin D, Gouras GK, Kirkwood A, Morris RG, Shen J (2005) Conditional inactivation of presenilin 1 prevents amyloid accumulation and temporarily rescues contextual and spatial working memory impairments in amyloid precursor transgenic mice. J Neurosci 25:6755–6764

52. Tamayev R, D’Adamo L (2012) Inhibition of gamma-secretase worsens memory deficits in a genetically congruous mouse model of Danish dementia. Mol Neurodegener 7:19

53. Tamayev R, Matsuda S, Arancio O, D’Adamo L (2012) beta-but not gamma-secretase proteolysis of APP causes synaptic and memory deficits in a mouse model of dementia. EMBO Mol Med 4:171–179

54. Tian Y, Bustos V, Flajolet M, Greengard P (2011) A small-molecule enhancer of autophagy decreases levels of Abeta and APP-CTF via Atg5-dependent autophagy pathway. FASEB J 25:1934–1942

55. Torres M, Jimenez S, Sanchez-Varo R, Navarro V, Trujillo-Estrada L, Sanchez-Mejias E, Carmona I, Davila JC, Vizuete M, Gutierrez A, Vitorica J (2012) Defective lysosomal proteolysis and axonal transport are early pathogenic events that worsen with age leading to increased APP metabolism and synaptic Abeta in transgenic APP/PS1 hippocampus. Mol Neurodegener 7:59

56. Vingtdeux V, Hamdane M, Begard S, Loyens A, Delacourte A, Beauvillain JC, Buee L, Marambaud P, Sergeant N (2007) Intra-cellular pH regulates amyloid precursor protein intracellular domain accumulation. Neurobiol Dis 25:686–696

57. Vingtdeux V, Hamdane M, Loyens A, Gele P, Drobeck H, Begard S, Galas MC, Delacourte A, Beauvillain JC, Buee L, Sergeant N (2007) Alkalizing drugs induce accumulation of amyloid precursor protein by-products in luminal vesicles of multivesicular bodies. J Biol Chem 282:18197–18205

58. Weihofen A, Lemberg MK, Friedmann E, Rueeger H, Schmitz D, Gouras GK, Kirkwood A, Morris RG, Shen J (2005) Conditional inactivation of presenilin 1 prevents amyloid accumulation and temporarily rescues contextual and spatial working memory impairments in amyloid precursor transgenic mice. J Neurosci 25:6755–6764

59. Willem M, Tahirovic S, Busche MA, Ovespian SW, Chafai M, Koort S, Hornburg D, Evans LD, Moore S, Daria A, Hampel H, Muller V, Giudici C, Nuscher B, Wenninger-Weinzierl A, Kremmer E, Heneka MT, Thal DR, Giedraitis V, Lannfelt L, Muller U, Livesey FJ, Meissner F, Herms J, Konnerth A, Marie H, Haass C (2015) eta-Secretase processing of APP inhibits neuronal activity in the hippocampus. Nature 526:443–447

60. Wolf D, Quon D, Wang Y, Cordell B (1990) Identification and characterization of C-terminal fragments of the beta-amyloid precursor produced in cell culture. EMBO J 9:2079–2084

61. Yang DS, Stavrides P, Mohan PS, Kaushik S, Kumar A, Ohno M, Schmidt SD, Wesson D, Bandypadhyay U, Jiang Y, Pawlik M, Peterhoff CM, Yang AJ, Wilson DA, St George-Hyslop P, Westaway D, Mathews PM, Levy E, Cuervo AM, Nixon RA (2010) Reversal of autophagy dysfunction in the TgCRND8 mouse model of Alzheimer’s disease ameliorates amyloid pathologies and memory deficits. Brain 134:258–277

62. Yu WH, Cuervo AM, Kumar A, Peterhoff CM, Schmidt SD, Lee JH, Mohan PS, Mercken M, Farnemy MR, Tjernberg LO, Jiang Y, Duff K, Uchiyama Y, Naslund J, Mathews PM, Cataldo AM, Nixon RA (2005) Macroautophagy—a novel Beta-amyloid peptide-generating pathway activated in Alzheimer’s disease. J Cell Biol 171:87–98

63. Zhang B, Higuchi M, Yoshiyama Y, Ishihara T, Forman MS, Stavrides P, Mohan PS, Kaushik S, Kumar A, Ohno M, Schmidt SD, Wesson D, Bandypadhyay U, Jiang Y, Pawlik M, Peterhoff CM, Yang AJ, Wilson DA, St George-Hyslop P, Westaway D, Mathews PM, Levy E, Cuervo AM, Nixon RA (2010) Reversal of autophagy dysfunction in the TgCRND8 mouse model of Alzheimer’s disease ameliorates amyloid pathologies and memory deficits. Brain 134:258–277

64. Yu WH, Cuervo AM, Kumar A, Peterhoff CM, Schmidt SD, Lee JH, Mohan PS, Mercken M, Farmery MR, Tjernberg LO, Jiang Y, Duff K, Uchiyama Y, Naslund J, Mathews PM, Cataldo AM, Nixon RA (2005) Macroautophagy—a novel Beta-amyloid peptide-generating pathway activated in Alzheimer’s disease. J Cell Biol 171:87–98

65. Zhang B, Higuchi M, Yoshiyama Y, Ishihara T, Forman MS, Martínez D, Joyce S, Trojanowski JQ, Lee VM (2004) Retarded axonal transport of R406W mutant tau in transgenic mice with a neurodegenerative tauopathy. J Neurosci 24:4657–4667