Protective Effects of Positive Lysosomal Modulation in Alzheimer’s Disease Transgenic Mouse Models

David Butler1,2*, Jeannie Hwang1,2,3, Candice Estick1,4, Akiko Nishiyama4, Saranya Santhosh Kumar4,5, Clive Baveghems2, Hollie B. Young-Oxendine3, Meagan L. Wisniewski3, Ana Charalambides2,3, Ben A. Bahr1,2,3,4,5,¤

1Neurosciences Program, University of Connecticut, Storrs, Connecticut, United States of America, 2Department of Pharmaceutical Sciences, University of Connecticut, Storrs, Connecticut, United States of America, 3William C. Friday Laboratory, Biotechnology Research and Training Center, University of North Carolina Pembroke, Pembroke, North Carolina, United States of America, 4Department of Physiology and Neurobiology, University of Connecticut, Storrs, Connecticut, United States of America, 5Department of Molecular and Cell Biology, University of Connecticut, Storrs, Connecticut, United States of America

Abstract

Alzheimer’s disease (AD) is an age-related neurodegenerative pathology in which defects in proteolytic clearance of amyloid β peptide (Aβ) likely contribute to the progressive nature of the disorder. Lysosomal proteases of the cathepsin family exhibit up-regulation in response to accumulating proteins including Aβ1–42. Here, the lysosomal modulator Z-Phe-Ala-diazomethylketone (PADK) was used to test whether proteolytic activity can be enhanced to reduce the accumulation events in AD mouse models expressing different levels of Aβ pathology. Systemic PADK injections in APPSwind and APPsw/P51ΔE9 mice caused 3- to 8-fold increases in cathepsin B protein levels and 3- to 10-fold increases in the enzyme’s activity in lysosomal fractions, while neprilysin and insulin-degrading enzyme remained unchanged. Biochemical analyses indicated the modulation predominantly targeted the active mature forms of cathepsin B and markedly changed Rab proteins but not LAMP1, suggesting the involvement of enhanced trafficking. The modulated lysosomal system led to reductions in both Aβ immunostaining as well as Aβ1–42 sandwich ELISA measures in APPswind mice of 10–11 months. More extensive Aβ deposition in 20-22-month APPsw/P51ΔE9 mice was also reduced by PADK. Selective ELISAs found that a corresponding production of the less pathogenic Aβ1–38 occurs as Aβ1–42 levels decrease in the mouse models, indicating that PADK treatment leads to Aβ truncation. Associated with Aβ clearance was the elimination of behavioral and synaptic protein deficits evident in the two transgenic models. These findings indicate that pharmacologically-controlled lysosomal modulation reduces Aβ1–42 accumulation, possibly through intracellular truncation that also influences extracellular deposition, and in turn offsets the defects in synaptic composition and cognitive functions. The selective modulation promotes clearance at different levels of Aβ pathology and provides proof-of-principle for small molecule therapeautic development for AD and possibly other protein accumulation disorders.

Introduction

Alzheimer’s disease (AD) is the most prevalent form of senile dementia, and is characterized by progressive compromise of synaptic integrity and cognitive functions. Aβ accumulation in the brain is a hallmark of AD pathology, and Aβ1–42 species have been implicated in the disruption of synaptic function and neuronal loss [1]. In addition to extracellular deposition, accumulation of Aβ also occurs intraneuronally [2–8], likely due to defective clearance and in many cases occurring prior to amyloid plaque formation. The clearance rates for Aβ peptides were indeed found to be slower in AD patients as compared to the rates in cognitively normal individuals [9]. One conclusion is an imbalance in Aβ production vs. clearance that implicates a plausible mechanism for the Aβ dysregulation in the more common late-onset AD, and perhaps a contributing factor in familial AD.

Intraneuronal Aβ1–42 is found in the brains of Alzheimer patients and individuals with mild cognitive impairment [2,10], thus there is growing evidence that such intracellular accumulation is an early indicator of neuronal compromise that correlates with cognitive decline. Synaptic dysfunction and deterioration are exhibited by Aβ-containing neurons and the intraneuronal accumulation is associated with cognitive deficits in animal models [2,4,6–8,11]. It is widely held that reducing protein accumulation events is critical for slowing the progression of AD, especially those produced by the aggregation-prone Aβ1–42 peptide. Potential targets include Aβ-degrading enzymes such as neprilysin, insulin-degrading enzyme, and endothelin-converting enzyme, and these proteases may in fact be responsible for Aβ homeostasis in the brain [12–15]. The lysosomal hydrolase cathepsin B has also been found to cleave Aβ1–42 into less amyloidogenic species [16]. This is
of particular interest since extracellular Aβ1–42 can be taken up by neurons in AD-vulnerable subfields and sequestered into lysosomes [11,17,18], thus lysosomal cathepsin activity may be important for the clearance of the peptide.

The endosomal/lysosomal system plays an important role in protein clearance, and its enhancement has been suggested as a strategy to reduce aberrant protein accumulation in age-related neurodegenerative disorders [19–24]. Interestingly, the endosomal/lysosomal system exhibits evidence of regulatory events in response to accumulating proteins found in AD [16,24–27] and areas of the aged brain [20,28,29]. The cathepsin family of lysosomal hydrolases appears to be particularly responsive to AD-related proteins accumulating in neurons. Protein accumulation stress, including that produced by Aβ1–42, markedly up-regulates the message, protein, and activity levels of the cysteine protease cathepsin B (EC 3.4.22.1), the aspartyl protease cathepsin D (EC 3.4.23.5), and other isoforms [16,19,20,24]. These protease responses may be reflective of early compensatory processes that keep protein accumulation events partially in check and account for the gradual nature of AD pathology. The response by cathepsin B in APPswInd mice failed to occur in older animals [30], suggesting that reduced efficiency of this compensatory pathway contributes to the age-related vulnerability of the brain.

To investigate the effects of lysosomal enhancement on protein accumulation pathology, a positive modulator of the lysosomal system was tested in transgenic mouse models of AD for its ability to promote cathepsin activity and protein clearance. The modulator selectively enhanced cathepsin B levels in the CNS, resulting in reduced Aβ1–42 levels and increased measures of a truncated Aβ1–38 peptide. Associated with the enhanced clearance of intra- and extracellular Aβ was the corresponding protection of synaptic integrity and improved cognitive ability. These findings provide further evidence that lysosomal enzymes can regulate the level of Aβ in the brain, and they indicate a minimally invasive approach to enhance lysosomal degradation of Aβ as a treatment for AD.

Materials and Methods

Animals and Injection Schedule

All transgenic mice and non-transgenic littermates were obtained from Jackson Laboratories (Bar Harbor, ME) and housed in vivarium facilities until the desired age. The APPswe/PS1A9E9 mice, strain B6C3-Tg(APPswe,PSEN1dE9)85Dbo/J (APP-PS1; stock number 004462) were used at 20–22 months of age. APPswInd mice received from Jackson laboratories (stock number 004661) expressed only 15% of the transgene copy number normally expressed by the B6.Cg-Tg(PDGFB-APPSwInd)20 Lns/J strain. The mice exhibited lower levels of Aβ deposits compared to the original APPswInd line [30], and were used at 10–11 months as a model of early Aβ pathology. Genotype was confirmed by PCR on tail DNAs. Non-transgenic Sprague-Dawley rats (Charles River Laboratories, Wilmington, MA) were used at 11–12 days postnatal to prepare hippocampal slice cultures, following a routine protocol with Millicell-CM inserts (Millipore, Bedford, MA) [19–21]. All studies were carried out in strict accordance with the recommendations from the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. Animal use and analyses were conducted in accordance with approved protocols from the Animal Care and Use Committees of the University of Connecticut (Protocol A09-008) and the University of North Carolina-Pembroke (Protocols 2009-001 and 2010-003). Mice were handled daily for >1 week and subsequently received daily i.p. injections of 18–20 mg/kg Z-Phe-Ala-diazomethylketone (PADK), obtained from Bachem Americas, Inc. (N-1040; Torrance, CA). PADK solutions were initially prepared at 24 mg/ml in dry DMSO, and slowly diluted with PBS to 12 mg/ml. Control mice were injected with the corresponding amount of vehicle (50% PBS and 50% DMSO). Consistent lysosomal modulation results were obtained from at least three PADK preparations represented by different lot numbers.

Behavioral Paradigms

Mice were handled and familiarized with the T-maze and suspended bar setup prior to the start of PADK injections. A day before the end of the injection schedule, spontaneous alternation behavior was assessed to measure episodic memory deficits in APP-PS1 mice. Transgenic and control mice were placed at the intersection of a T-maze, and entries across each arm’s threshold were observed with a closed-circuit monitor for a 10-min period. A minimum of 15 entries was used to determine percent alternations when compared to total alternations possible. An alternation was a succession of entries into 3 different arms of the maze. Mobility in the T-maze or in a novel open field was assessed by grid crossings in the first 3 min of exploration. APPswInd mice and their respective controls were assessed on the suspended rod test, in which the mice were placed in the middle of a 1-cm diameter rod suspended 40-cm over a padded service with platforms in sight 56 cm apart. Time of maintained uprightness was recorded on the third trial as the mice attempted to reach a platform during a period of 60 sec. They were also tested in an open field of 55 cm x 36 cm to determine exploratory distance during novel exposure to the environment and during subsequent re-exposure 24 h later. Grid crossings were assessed for 5 min using a closed-circuit camera.

Tissue Preparation

Immediately following behavior testing, brain tissue was removed and prepared for analyses. Some animals were anesthetized and perfused with 4% paraformaldehyde prior to dissecting brains for tissue sectioning and subsequent hematoxylineosin staining or immunofluorescence protocols. For the remaining mice, brains were removed and quickly dissected in ice-cold buffer containing 0.32 M sucrose, 5 mM HEPES (pH 7.4), 1 mM EDTA, 1 mM EGTA, and the protease inhibitors aprotinin, leupeptin, bestatin, E-64, peptatin A (each at 2 μg/ml), and 4-(2-aminoethyl)benzenesulfonyl fluoride (0.3 mM) (unless otherwise stated, reagents were from Sigma-Aldrich Co., St. Louis, MO). Regions were separated from one hemi-forebrain and snap-frozen in liquid nitrogen for later homogenization in lysis buffer containing protease inhibitors. The other hemi-forebrain was either 1) fixed in phosphate-buffered 4% paraformaldehyde for immunocytochemistry, 2) mechanically homogenized in appropriate buffer conditions to collect the extracellular-enriched fraction following centrifugation [31], or 3) homogenized in ice-cold 0.3 M sucrose with 1 mM EDTA for the isolation of lysosomes (see below).

Immunoblotting and ELISA

Equal amounts of sample protein were separated on standard or tris-tricine gradient gels, and transferred to nitrocellulose for antibody staining. Antibodies utilized were developed against cathepsin B (1:200; Millipore, Bedford, MA), cathepsin D (1:300; Cortex Biochemicals, San Leandro, CA), neprilysin (1:300; Millipore), insulin-degrading enzyme (1:200; Covance, Princeton, NJ), n-secreasate (1:500, against amino acids 732–748 of human TNF-α converting enzyme; ProSci, Poway, CA), actin (1:1,000, Sigma-Aldrich Co.), amino acids 1–16 of human Aβ (E610, 1:500;
Covance), human sAPPα (2B3, 1:100; IBL International, Hamburg, Germany), human sAPPβ-sw (6A1, 1:200; IBL), Rab5a (1:200; Santa Cruz Biotechnology, Santa Cruz, CA), Rab7 (1:100; Santa Cruz Biotechnology), Lamp1 (1:200; GeneTex Inc., Irvine, CA), synaptophycin (Chemicon, Temecula, CA), and the carboxy termini of AMPA receptor subunit GluA1 [19]. Secondary antibodies were from Bio-Rad (Richmond, CA) and immunoreactive bands were assessed for integrated optical density with BIOQUANT software (Nashville, TN). Equal aliquots of soluble homogenate fractions were also assessed by ELISA protocols for the specific detection of Aβ_{42} and Aβ_{38} species using antibodies (12F4 and BA1–13, respectively) and reagents from Covance. Chemiluminescence detection was measured with a SpectraMax L Luminescence Reader (Molecular Devices, Sunnyvale, CA) and converted to femtomoles per milligram sample protein using standard curves generated with pure Aβ_{42} and Aβ_{38} peptides (Bachem).

Immunocytochemistry

Fixed tissue was cryoprotected and serial sectioned at 20-μm thickness. Immunolabeling followed standard free-floating methods using anti-cathepsin B (Millipore), anti-NeuN (Invitrogen,
PADK selectively enhances cathepsin B levels in two transgenic mouse models. Dissected hemi-brains were quickly separated into regions and each immediately homogenized in ice-cold 0.3 M sucrose with 1 mM EDTA and centrifuged at 750 × g for 10 min. The collected supernatant was incubated with 2 mM CaCl₂ at 37°C for 5 min then layered over 24% Percoll in 0.32 M sucrose containing 1 mM EDTA (pH 5.5). After centrifugation at 20,000 × g for 18 min the gradients were fractionated, and those determined to be negative (upper four fractions) or positive (lower two) for cathepsin B by immunoblot were diluted with 5 volumes of sucrose solution to separate organelles from the Percoll by centrifuging at 40,000 × g for 60 min. The pellets were resuspended and protein content determined with the Pierce BCA assay (Thermo Scientific, Rockford, IL). The resulting fractions were aliquoted, lysed in 18 mM citrate, and assessed for cathepsin B proteolytic activity using the Z-Arg-Arg AMC substrate, the fluorogenic Calbiochem assay kit (EMD Chemicals, Gibbstown, NJ), and the Molecular Devices SpectraMax M3 Microplate Reader. Potential inhibitors were also tested in the cathepsin B activity assay, using equal aliquots of Triton X-100 solubilized brain homogenate pre-treated for 30 min with increasing concentrations of the agents.

Table 1. PADK-mediated enhancement across brain regions of transgenic mouse models.

| Brain Region | APPsweInd | APP-PS1 |
|--------------|-----------|---------|
| neocortex    | 3.0±0.17  | 7.3±0.60|
| frontal cortex| 3.2±0.33  | 6.8±1.3 |
| hippocampus  | 4.3±0.38  | 8.4±0.85|
| mesencephalon| 3.7±0.70  | 4.9±0.21|

Table 2. PADK selectively enhances cathepsin B levels in two transgenic mouse models.

|          | APPsweIndveh | APPsweIndPADK | APP-PS1veh | APP-PS1+PADK |
|----------|--------------|---------------|------------|--------------|
| CB       | 95.3±14.2    | 404.6±36.0*** | 44.5±15.2  | 376.0±38.6***|
| nep      | 89.1±19.2    | 95.3±16.6     | 43.0±6.5   | 31.5±6.6     |
| IDE      | 101.1±11.0   | 76.4±6.5      | 96.1±8.3   | 89.5±6.0     |
| α-secretase | 82.3±6.0    | 92.0±5.2      | 60.1±8.9   | 58.9±6.2     |
| LAMP1    | 98.4±5.4     | 102.2±8.6     | 46.1±9.9   | 45.8±4.6     |

APPsweInd and APP-PS1 mice were injected i.p. daily with PADK (20 mg/kg; n = 11–13) or vehicle (n = 10) for 9–11 days. Hippocampal homogenates were analyzed by immunoblot and mean immunoreactivities are shown for active cathepsin B (CB), nephrilysin (nep), insulin-degrading enzyme (IDE), α-secretase (α-secretase), and LAMP1.

***P<0.0001, unpaired t-test.

doi:10.1371/journal.pone.0020501.t002
correlations between Aβ species and cathepsin B enhancement or between GluA1 immunoreactivity and the lysosomal enhancement.

Results

To elicit lysosomal enhancement in mouse models of AD, we administered Z-Phe-Ala-diazomethylketone (PADK), a weak inhibitor of cathepsin B and L (IC50 = 9.4 ± 2.4 μM) as well as a lysosomal modulator previously shown to cause a feedback response involving marked up-regulation of cathepsin isoform expression in vitro [19–21]. The lysosomal modulator (20 mg/kg) was injected i.p. daily for 9 days into 10–11-month APPswInd mice which express the human APP gene with the Swedish (K670N/M671L) and Indiana (V717F) mutations [30], resulting in a marked increase in the active isoform of cathepsin B in the brain as compared to vehicle-injected transgenic mice (Fig. 1A; ANOVA P < 0.0001, post hoc test P < 0.001; n = 13). Cathepsin B immunoreactivity levels were enhanced >4 fold in hippocampal samples, and 3-fold or greater increases were found in samples from neocortex, frontal cortex, and mesencephalon (Table 1). Measures of Aβ-degrading proteases neprilysin and insulin-degrading enzyme, as well as α-secretase which prevents Aβ production, were not altered (Fig. 1A and Table 2), thus the PADK-mediated lysosomal modulation was produced in a selective manner. Similar selectivity was also evident for the PADK effect in 20–22-month APPswe/PS1DDE9 mice (APP-PS1; Table 2), which express a chimeric mouse/human APP and human presenilin 1 directed to CNS neurons [33]. Significant cathepsin B up-regulation was found in different brain regions of the APP-PS1 mice, with hippocampus exhibiting the largest increase of >8 fold (Table 1).

In immunocytochemistry images, intracellular cathepsin B was revealed as punctate staining (green) characteristic of lysosomal organelles in hippocampal CA1 pyramidal neurons (Fig. 1B), and...
the intensity of such immunostaining was enhanced after PADK treatment (Fig. 1C). Neurons were counterstained with anti-NeuN (red), and PADK elicited no apparent change in neuronal density or morphology. Quantitative analysis of the fluorescence intensity across the stratum pyramidale confirmed an increase in cathepsin B immunoreactivity ($P<0.0001$; Fig. 1D). On the other hand, the number of cathepsin B-positive organelles per pyramidal neuron ($n=62$) was found to be unchanged in the view-fields (Fig. 1E), and the lysosome-associated membrane glycoprotein LAMP1 was also unaltered in blot samples from the different treatment groups (Fig. 2A, Table 2). ANOVA assessment of two different cathepsins (B and D) across all transgenic treatment groups found that only cathepsin B increased with PADK treatment (Fig. 2). Thus, the 2- to 3-fold increase in intracellular cathepsin B staining, in the absence of a change in lysosome number, appears to represent the

![Figure 4. Mature forms of cathepsin B exhibit pronounced modulation by PADK.](image)

![Figure 5. Changes in Rab proteins in cultured hippocampal slices treated with PADK.](image)

Figure 4. Mature forms of cathepsin B exhibit pronounced modulation by PADK. APP$^{SweHBD}$ mice subjected to 9 daily injections of PADK (20 mg/kg; $n=10$) or vehicle ($n=9$) were assessed for the different cathepsin B species in hippocampal samples. Homogenates from the transgenic mice and from vehicle-treated wildtype mice were analyzed by immunoblot to label the approximately 40-kDa procathepsin B species (proCB) and the 25- and 30-kDa active mature forms (arrows) within the same samples (A). Immunoreactivity levels of the proenzyme and mature forms (means±SEM) in vehicle- (-) and PADK-treated transgenic samples (+) were determined by image analysis (B). Unpaired Mann-Whitney U-test: *$P=0.0296$, **$P<0.001$. doi:10.1371/journal.pone.0020501.g004

Figure 5. Changes in Rab proteins in cultured hippocampal slices treated with PADK. Stable slice cultures prepared from rat hippocampus were treated daily with PADK (10 μM; $n=12$ groups of 8 slices each) or with the corresponding vehicle (final condition of 0.02% DMSO; $n=14$ groups of slices) for 4 days. Slice groups were homogenized and equal protein aliquots analyzed by immunoblot for Rab5a, Rab7, active form of cathepsin B (act CB), LAMP1, and actin (A). Rab protein levels (means±SEM) in slices treated without (-) or with PADK (+) were determined by image analysis (B). Unpaired Mann-Whitney U-test: *$P=0.0208$, ***$P<0.0001$. doi:10.1371/journal.pone.0020501.g005
primary PADK effect in neurons. Note that a one-way nonparametric Kruskal-Wallis analysis followed by Dunn's post test of the 33-kDa cathepsin D form alone revealed significant albeit small increases produced by PADK in the APPSwInd and APP-PS1 samples (P, 0.05).

To further test whether the intracellular modulation is actually influencing lysosomal cathepsin B, localization of the PADK effect was evaluated in brain tissue double-stained for cathepsin B and LAMP1 (Fig. 3A). As evident in the merged immunofluorescence image, the PADK-modulated cathepsin B co-localized with LAMP1-positive organelles in CA1 pyramidal neurons. Together, the findings indicate that the lysosomal modulator enhances cathepsin B content in lysosomes. APPSwInd mice injected with vehicle or 18 mg/kg PADK daily for 10 days were also assessed for cathepsin B activity in isolated lysosomes using the Z-Arg-Arg AMC probe (Fig. 3B). Rapidly dissected brain regions were subjected to subcellular fractionation in Percoll gradients, and equal protein aliquots from the lysosomal and non-lysosomal fractions were evaluated for hydrolase activity that was blocked by the cathepsin B inhibitor CA074me. Note that in the different brain regions tested, only the lysosomal fractions exhibited PADK-dependent increases in cathepsin B activity of 3-10 fold (P, 0.0001).

Hippocampal homogenates from the APPSwInd mice were further analyzed in an attempt to understand how PADK influences lysosomal levels of cathepsin B. Cathepsin B belongs to the superfamily of papain-like cysteine proteases and is first synthesized as a proenzyme. The approximately 40-kDa procathepsin B exhibited a marginally significant increase of 93% (mean ± SEM; P, 0.03) in the PADK-treated group as compared to vehicle-injected transgenic mice samples (Fig. 4A and B). Within the same immunoblot samples, the percent change in the 25–30-kDa mature cathepsin B forms was found to be much larger than the PADK effect on the proform of the enzyme (270% ± 31%, P, 0.001) (Fig. 4B). PADK's range of effect in the two transgenic models is 4- to 9-fold changes in the mature active isoforms in hippocampus (Table 1), and even greater fold increases when considering the increase in the 25-kDa species alone (see Fig. 4A). These pronounced changes implicate trafficking as a component of the PADK effect, since inactive procathepsin B is processed to the

Figure 6. PADK treatment reduces accumulation events in 10–11-month APPSwInd mice. APPSwInd mice were treated with 9 daily injections of either PADK (20 mg/kg; n = 13) or vehicle (veh; n = 10), and wildtype mice (wt; n = 13) were injected with vehicle for 9 days. Fixed brain sections were stained with the 6E10 antibody; arrows denote extracellular deposits (A). Size bar: 450 µm. sg, stratum granulosum; sp, stratum pyramidale. Sections were also immunostained with an Aβ1–42–specific monoclonal antibody; arrows denote CA1 cell bodies with intracellular labeling (B; size bar: 15 µm). The remaining tissue was separated for rapid homogenization, and equal protein aliquots of hippocampal samples were assessed by immunoblot. The 6E10 antibody labeled the 4-kDa Aβ species and the parent APP, and selective antibodies were used to label sAPPα, sAPPβ, and actin (C). Positions of molecular weight standards are shown. Mean integrated optical densities ± SEM were normalized to 100% in vehicle-treated transgenic samples (-) to allow comparison of PADK effects (+) across the different antigens (D). The samples were also tested with an Aβ1–42 sandwich ELISA to determine femtomoles of Aβ1–42–related peptides per milligram protein (E). ANOVA: P, 0.0001; post hoc test compared to APPswind+vehicle: *P, 0.01.
doi:10.1371/journal.pone.0020501.g006
active forms in late endosomes and lysosomes. To selectively
examine trafficking markers during defined lysosomal modulation,
hippocampal slice cultures were treated with PADK daily for 4
days, causing a >5-fold increase in active cathepsin B. In
conjunction with enhanced cathepsin B, the PADK-treated slices
exhibited a decrease in Rab5a (Fig. 5A), a marker of early
endosomes [34]. From the analysis of integrated optical densities,
control slices were found to contain more than 2-fold more Rab5a
than the PADK slices, whereas the lysosomal modulator increased
the level of Rab7 which controls the late step of endosomal
trafficking to lysosomes (Fig. 5B). In addition, as found
in vivo
, LAMP1 was unchanged (control, 156 ± 7.2; PADK-treated slices,
140 ± 5.1; NS). Thus, positive modulation of trafficking likely plays
a major role in PADK’s ability to increase the content of active
cathepsin B in lysosomes and/or late endosomes.

We next tested whether PADK’s influence on the lysosomal
system is associated with enhanced Aβ clearance. APPswInd mice
of low transgene copy number were used at 10–11 months of age
in order to assess the PADK effects at an early stage of Aβ
pathology. Tissue sections were stained with the 6E10 antibody, and photomicrographs indicate PADK-
mediated reductions of the intracellular labeling in hippocampal subfields and piriform cortex (A). Image analysis for densitometric quantification was conducted across view-fields of four different neuronal layers (B), and mean integrated optical densities were plotted (± SEM). Individual ANOVAs:
P<0.0001; Tukey’s post hoc tests compared to APPswInd+vehicle: **P<0.001. Size bar: 40 μm, CA1 and DG; 65 μm, piriform cortex. DG, dentate gyrus; sg, stratum granulosum; sp, stratum pyramidale.
doi:10.1371/journal.pone.0020501.g007

In Figure 6A, intracellular staining is evident among neurons of the stratum pyramidale and stratum granulosum in vehicle-
jected APPswInd mice, while none was found in wildtype mice, and the transgenics also exhibited sporadic extracellular deposits. The daily PADK administration resulted in reduced deposits and a marked decrease in the cellular labeling. Serial sections stained with Aβ1–42–specific monoclonal antibody confirmed that Aβ1–42
is the primary species that accumulates intracellularly in the APPswInd brain (Fig. 6B), as previously reported for other AD mouse models [35], and it is the same peptide selectively taken up
by neurons of the hippocampus [11]. The Aβ1–42 staining intensity
in pyramidal neurons was diminished by the PADK treatment,
and similar degrees of reduced 6E10 labeling (~63–73%; post hoc
tests P<0.001) were determined across different neuronal layers in
hippocampus and piriform cortex (Fig. 7).

Brain homogenate samples were also analyzed to determine
which APP fragments recognized by 6E10 were reduced by the
PADK treatment. The lysosomal modulator significantly reduced
a 6E10-labeled species of 4 kDa that coincided with the
electrophoretic migration of pure Aβ1–42 (ANOVA P<0.0001,
post hoc test P<0.01), without affecting the parent hAPP protein

Figure 7. PADK-mediated reductions of intraneuronal accumulation in the APPswInd mice. Non-transgenic control (wt) and APPswInd mice were treated with 9 daily injections of PADK or vehicle. Brain sections were stained with the 6E10 antibody, and photomicrographs indicate PADK-
mediated reductions of the intracellular labeling in hippocampal subfields and piriform cortex (A). Image analysis for densitometric quantification was conducted across view-fields of four different neuronal layers (B), and mean integrated optical densities were plotted (± SEM). Individual ANOVAs:
P<0.0001; Tukey’s post hoc tests compared to APPswInd+vehicle: **P<0.001. Size bar: 40 μm, CA1 and DG; 65 μm, piriform cortex. DG, dentate gyrus; sg, stratum granulosum; sp, stratum pyramidale.
doi:10.1371/journal.pone.0020501.g007
cleavage activity of recombinant human β-secretase, nor did the potent cathepsin B inhibitor CA074me (Fig. 8).

To assess the PADK effects at a more extensive stage of Aβ pathology, we utilized double transgenic APP-PS1 mice of 20–22 months. Their extracellular deposits of 50–90 μm were easily detected in hippocampus and other brain regions in hematoxylin-eosin stained sections (see Fig. 9A) as well as with 6E10 immunolabeling (Fig. 9B). The 6E10 antibody produced very faint background staining in wildtype control mice, whereas pronounced intra- and extracellular labeling was evident in brain sections of the vehicle-treated transgenics. As in the younger APPtmd mice, PADK-mediated lysosomal modulation in APPPS1 mice (20 mg/kg/day × 11 days, i.p.) was associated with smaller and fewer extracellular deposits as well as reduced levels of intraneuronal staining (Fig. 9A and B). The specific immunostaining intensity was decreased 80% in the CA1 pyramidal layer, and in close correspondence were 76–85% reductions in the above-threshold staining area for extracellular deposits in the stratum radiatum and piriform cortex (ANOVA: P<0.0001, post hoc test P<0.001; n = 11) (Table 3).

Brains from the PADK-injected APP-PS1 mice also exhibited a reduction in the 4-kDa Aβ band labeled in 6E10 immunoblots (ANOVA P<0.0001, post hoc test P<0.001), whereas no decrease was evident in the parent APP (Fig. 9C). The lysosomal modulator similarly reduced Aβ species assessed in immunoblot samples of extracellular-enriched brain extracts, without affecting levels of full-length or secreted forms of APP. As conducted in the APPtmd model to determine which Aβ species is influenced by PADK, vehicle- and PADK-treated APP-PS1 samples were also subjected to a sandwich ELISA to selectively assess Aβ1–42 species (Fig. 9D). The measured concentrations of peptide (molecules per milligram homogenate protein) indicate that PADK reduced Aβ1–42 by 44–62% (P<0.01). This level of reduction closely matches the PADK-mediated 40–63% decrease in the 4-kDa peptide labeled by 6E10, and the peptide is the only antigen recognized by 6E10 that exhibits a reduction in PADK-treated mice. Brain sections from the three groups of mice were also double-stained for Aβ1–42 (green) and cathepsin B (red) to specifically assess intracellular Aβ1–42 (Fig. 10). Immunofluorescence images revealed punctate Aβ1–42-positive material within CA1 pyramidal neurons, and the intracellular accumulation was reduced by PADK in correspondence with enhanced labeling intensity of cathepsin B-positive organelles. Note that with PADK treatment, co-localization of anti-Aβ1–42 and anti-cathepsin B staining occurs in several organelle-like structures.

As evidence points to Aβ1–42 being reduced in brains of PADK-treated transgenic mice, the opposite is the case for Aβ1–38, a less pathogenic peptide found previously to be a cathepsin B cleavage product of Aβ1–42 [16]. Brain samples from APPtmd and APP-PS1 mice treated with vehicle or PADK were assessed for Aβ1–42 and the truncated Aβ1–38 species using selective sandwich ELISA protocols (Fig. 11). The Aβ1–42 sandwich ELISA does not recognize Aβ1–38 or Aβ1–40 peptides, and the Aβ1–38 ELISA does not recognize Aβ1–40 or Aβ1–42. The 52% decrease in Aβ1–42 measures in PADK-treated APPtmd mice, as compared to vehicle-treated mice, was matched by the significant 52% increase in Aβ1–38 peptide. Regarding the opposing PADK effects in APP-PS1 mice, the 51% decrease in Aβ1–42 was partially matched by the significant 32% increase in Aβ1–38 species. The transient nature of Aβ1–38 production by cathepsin B [16] may account for the smaller change in the truncated peptide.

Together with our immunocytochemistry results, Aβ1–38 appears to have a lower propensity to resist degradation and accumulate in cells than Aβ1–42, as expected from previous studies [11,17].
The lysosomal modulator’s effects on Aβ clearance were associated with synaptic protection. In Figure 12A, hippocampal samples from the two mouse models were analyzed by immunoblot for the presynaptic protein synapsin II and the postsynaptic glutamatergic marker GluA1. Similar to the extent of synaptic decline seen in related transgenic mice [30,33], the synaptic proteins exhibited deficits of 23–31% in APPswInd and APP-PS1 mice as compared to their respective age-matched wildtypes (P<0.01), while actin levels remained unchanged. PADK treatment significantly reduced the GluA1 deficit in the two mouse models, reaching levels comparable to those found in non-transgenic control mice (Fig. 12B) (ANOVA: P<0.0001; n = 12–20). Similar indications of synaptic protection were found when assessing synapsin II and synaptophysin. The integrity of hippocampal dendritic fields was also found preserved in immunostained tissue sections, and the level of GluA1 immunoreactivity within each transgenic mouse correlated with the respective extent of cathepsin B enhancement in the brain (r = 0.78, P = 0.02).

Lastly, we assessed the mouse models to determine whether the PADK-mediated clearance and synaptic protection translate to improvements in behavioral tests. Although the APPswInd mice used in the present study express low levels of Aβ deposits, their
synaptic decline was associated with significant deficits on the suspended rod test (Fig. 13A) as well as on the exploratory habituation test (Fig. 13B), similar to the type of deficit reported previously for APPswind mice of the same age range [36]. Compared to wildtypes, the vehicle-injected APPswind mice exhibited a marked reduction in balance time while coordinating to reach a safe platform, whereas the PADK-treated transgenics exhibited a marked reduction in balance time while coordinating.

APP-PS1 mice were injected i.p. daily with PADK (20 mg/kg; n = 11) or vehicle (n = 10) for 11 days. Fixed tissue was sectioned and stained with the 6E10 antibody. Image analysis for densitometric quantification of the immunostaining (mean integrated optical density±SEM) was conducted across view-fields of the hippocampal CA1 stratum pyramidale (sp). Area of deposit labeling above background was also measured for view-fields of the hippocampal stratum radiatum (sr) and piriform cortex (area). Area of deposit was calculated for the hippocampal CA1 stratum pyramidale (sp). The area of deposit was measured using Adobe Photoshop CS3 software (area)

Table 3. PADK decreases 6E10 immunostaining in APP-PS1 mouse brain.

|            | wt       | APP-PS1+veh | APP-PS1+PADK |
|------------|----------|-------------|--------------|
| hippocampal sp (IOD) | 129±15.1 | 672±58.9 | 241±15.0** |
| hippocampal sr (area) | 0.07±0.02 | 2.90±0.71 | 0.49±0.08** |
| piriform cortex (area) | 0.11±0.03 | 3.89±0.36 | 1.04±0.21** |

APP-PS1 mice were injected i.p. daily with PADK (20 mg/kg; n = 11) or vehicle (n = 10) for 11 days. Fixed tissue was sectioned and stained with the 6E10 antibody. Image analysis for densitometric quantification of the immunostaining (mean integrated optical density±SEM) was conducted across view-fields of the hippocampal CA1 stratum pyramidale (sp). Area of deposit labeling above background was also measured for view-fields of the hippocampal stratum radiatum (sr) and piriform cortex (area). Area of deposit was calculated for the hippocampal CA1 stratum pyramidale (sp). The area of deposit was measured using Adobe Photoshop CS3 software (area)

**P<0.001.

PADK decreases 6E10 immunostaining in APP-PS1 mouse brain.

Discussion

Our study indicates that lysosomes are a site of modulation for the targeted enhancement of cathepsin function to promote clearance of the Aβ peptide. The endosomal-lysosomal pathway has been implicated as playing a role in the degradation of Aβ, and the impairment of this pathway likely contributes to the accumulation of intraneuronal Aβ and other proteins linked to AD. Enhancement of the lysosomal system was achieved with the modulatory agent PADK, resulting in the selective increase of cathepsin B activity in lysosomes. The lysosomal modulation led to reductions in both intracellular and extracellular Aβ, to a degree that ameliorates the phenotype of two AD mouse models with distinctly different levels of Aβ pathology. At the analogous stages of the human disorder, reducing Aβ accumulations is thought to be essential for slowing AD progression.

When administered systemically, the PADK modulator increased cathepsin B levels 4- to 8-fold across different brain regions in APPswind and APP-PS1 mice. From the analysis of immunofluorescence images, enhanced intracellular cathepsin B staining was found to represent the primary PADK effect in neurons. As cathepsin B increased in the transgenic mice, there was a corresponding decrease in Aβ1-42 measured by selective ELISA. Brain immunocytochemistry with anti-Aβ1-42 and 6E10 antibodies suggests that lysosomal modulation leads to decreased levels of intraneuronal deposition. Interestingly, reduction in extracellular deposits occurred in correspondence with the reduced labeling of intracellular Aβ, providing evidence of an equilibrium relationship between secreted Aβ peptides and the extracellularly deposited material. This fits well with Oddo et al. [35] which described the accumulation of intracellular Aβ as a source of extracellular amyloid deposits. Note that PADK had no effect on hAPP expression levels in the transgenic brains, whereas the 4-kDa Aβ species was markedly reduced. This 44–63% decrease in Aβ likely contributes to the reduced extracellular deposition as multiphoton microscopy and microdialysis studies showed that reduction of Aβ by as little as 20–25% was capable of diminishing plaque deposits [38]. The actions of PADK on cathepsin B and Aβ indicate the intraneuronal clearance process as an appropriate target for offsetting protein accumulation pathology. Since extracellular peptide taken up by vulnerable neurons is thought to contribute significantly to the accumulation of intracellular Aβ1-42 [11,17,18,39], the modulated lysosomal system may be exerting its clearance effects directly on the internalized Aβ1-42 peptide as well as by increasing the turnover of Aβ-containing APP fragments. Lysosomal modulation in other cell types will be the subject of follow-up studies since in microglia, internalized oligomeric Aβ has recently been shown to be trafficked to lysosomes and degraded by cysteine proteases including cathepsin B [40].

PADK-mediated lysosomal modulation appears to involve both increased expression of cathepsin B and enhanced trafficking of the enzyme. The modulation consisted of a modest increase in procathepsin B as compared to larger 4- to 9-fold increases in the mature active forms of the enzyme. Further evidence that PADK has a positive effect on endo-lysosomal trafficking includes i) cathepsin B hydrolytic activity was enhanced in isolated lysosomes to a much greater degree than the enhancement of proenzyme in brain samples, ii) active cathepsin B forms were enhanced in the absence of any increase in the lysosomal marker LAMP1, iii) the modulated cathepsin B co-localized with LAMP1 in neurons, and iv) decreases in the Rab5a early endosome marker corresponded with the enhanced levels of mature cathepsin B in hippocampal tissue. Elevated expression of early endosome regulators, on the other hand, occur in hippocampal neurons from individuals with AD and mild cognitive impairment, suggesting that early endosomal dysfunction contributes to AD progression [27,41].

The action of PADK may offset endocytic dysfunction and promote protein clearance through enhanced trafficking of cathepsin B and/or Aβ-containing species from early endosomes to late endosomes and lysosomes. Perhaps enhanced cathepsin B maturation is facilitated by the small PADK-mediated increase in mature cathepsin D, since previous studies found the latter to activate procathepsin B into mature forms [42,43]. Regarding PADK’s lack of influence on LAMP1, this would appear to rule out any broad effect on lysosome production or the modulation of TFEB, a transcription factor that regulates lysosomal biogenesis.
TFEB does share with PADK the ability to elicit protection in models of pathogenic accumulation events, as shown in TFEB-transfected cells insulted with a parkinsonian neurotoxin [45].

An obvious paradox follows from PADK promoting lysosomal function while historically being classified as an inhibitor of the thiol proteases cathepsins B and L. Indeed, extended exposure of hippocampal slice cultures to 40 μM PADK causes multiple signs of lysosomal dysfunction and the characteristic proliferation of a lysosomal marker [46,47]. On the other hand, low concentrations of 1–10 μM significantly increased cathepsin enzymes without producing lysosomal pathology or synaptic decline over extended treatments of 5–20 days [19,20,46], and without any obvious axonopathy, swelling of axonal initial segment, or transport failure. In fact, the low-level PADK treatment led to enhanced clearance in the hippocampal slice model of protein accumulation pathology, as well as to recovered levels of microtubule stability markers, transport function, tubulin-binding proteins, and pre- and postsynaptic proteins. Here, comparable effects on Aβ clearance and synaptic protection were produced by PADK in transgenic mice.

The pronounced up-regulation of cathepsin B and the changes in Rab proteins are not the first paradoxical effects found produced by a cathepsin inhibitor. PADK causes increased proenzyme expression in the transgenic mouse brains, likely related to the paradoxical findings in which the protease inhibitor leupeptin increased cathepsin activities in different tissues. The broad-acting inhibitor, as high as 200 mg/kg, caused significant inhibition of cathepsins B and L, but after clearance of the administered leupeptin not only did cathepsin activities recover to control levels they continued to increase more than two-fold [48–51]. Note that high levels of PADK administered to mice (60 mg/kg i.p. in 100% DMSO) or to 3T3 fibroblasts (100 μM applied daily) found no such increases in cathepsin B and L activities [48,51]. However, the study that tested lower PADK concentrations reported increased amounts of the 39-kDa procathepsin L [51], perhaps related to the increase in procathepsin B in the present study. Unique compared to leupeptin, PADK had a long-term effect on increasing the active intermediate of a cathepsin. In fact, they showed that their lowest concentration tested, 3 μM, led to pronounced accumulation of the active 29-kDa lysosomal form. The buildup of cathepsin intermediates may signify modulated enzyme trafficking, resulting in a larger pool of accessible isoforms for more efficient maturation, and thereby improving lysosomal efficiency.

The enhanced lysosomal capacity described here is through targeted enzyme modulation, producing selective enhancement of a clearance pathway since PADK was found to increase cathepsin B activity whereas neprilysin and insulin-degrading enzyme levels were unchanged in the APPSwe/m1 and AP-PS1 mice. In the normal brain, cathepsin B may act together with identified Aβ-degrading enzymes to provide efficient Aβ clearance. Extracellular and, in some cases, intracellular Aβ levels have been shown to be decreased by neprilysin, insulin-degrading enzyme, and endothelin-converting enzyme [12–15]. Note that correlational analyses found a negative correlation between neprilysin expression and Aβ...
The PADK-mediated changes in Aβ effects are expressed as mean percent change compared to the corresponding vehicle-treated transgenic data: 

\[ \frac{\text{PADK effects}}{\text{Vehicle effects}} = \frac{\text{PADK}}{\text{Vehicle}} \]



In brain samples (n = 9 per group), using selective sandwich ELISAs. The PADD-mediated changes in Aβ were determined from data presented in Figures 6E and 9D. In the APPSwInd and APP-PS1 samples, PADD increased Aβ, species from 60.6±7.7 to 922±5.7 fmol/mg and from 104±7.0 to 137±18 fmol/mg, respectively. The plotted PADD effects are expressed as mean percent change±SEM. Post hoc tests compared to the vehicle-treated transgenic data: *p<0.01.

** doi:10.1371/journal.pone.0020501.g011

Figure 11. Lysosomal modulator treatment promotes truncation of the Aβ1-42 peptide. APPSwInd and APP-PS1 mice treated with vehicle or PADD were assessed for Aβ1-42 and truncated Aβ1-38 species in brain samples (n = 9–11 per group), using selective sandwich ELISAs. The PADD-mediated changes in Aβ1-42 were determined from data presented in Figures 6E and 9D. In the APPSwInd and APP-PS1 samples, PADD increased Aβ1-38 species from 60.6±7.7 to 922±5.7 fmol/mg and from 104±7.0 to 137±18 fmol/mg, respectively. The plotted PADD effects are expressed as mean percent change±SEM. Post hoc tests compared to the vehicle-treated transgenic data: *p<0.01.

** doi:10.1371/journal.pone.0020501.g012

Figure 12. Lysosomal modulation is associated with preservation of synaptic markers in APPSwInd and APP-PS1 mice. Transgenic and wildtype (wt) mice were injected daily with PADD (+) or vehicle (-) for 9–11 days. Equal protein aliquots of hippocampal homogenates were analyzed by immunoblot for synaptic markers and actin, showing PADD-improved levels of GluA1 and synapsin II (syn II) in transgenic mice (A). The mean GluA1 immunoreactivities±SEM are shown for vehicle-treated wildtypes and for the vehicle- and PADD-treated transgenics (B). Post hoc tests compared to vehicle-treated transgenics: **p<0.001.

** doi:10.1371/journal.pone.0020501.g012

Accumulation as well as clinical diagnosis, but this was not the case regarding the expression of other Aβ-degrading enzymes [52]. Lysosomal integrity and cathepsin B regulatory responses are perturbed in the aged brain [16,33], suggesting that age-related disruptions of effective cathepsin B and neprilysin activities together contribute to the aging risk factor of AD.

The lack of change regarding sAPPα and sAPPβ in both transgenic models also rules out PADD's modulation of z- or β-secretase. Positive modulation of β-secretase would promote Aβ production in contrast to the reduction of Aβ species found elicited by PADD. PADD was absent of any inhibitory action on β-secretase activity as well, thus ruling out the weak cathepsin inhibitor's ability to influence Aβ levels by blocking β-site cleavage of APP. Positive regulation of z-secretase has the potential to preclude Aβ production. The z-secretase enzyme has also been implicated in an alternative route of clearance in mice in which cathepsin-mediated elevation of z-secretase activity regulates Aβ production [34,55]. PADD's influence on cathepsins may indirectly utilize z-secretase, however, PADD effects did not include altered z-secretase or sAPPα levels.

A defect in proteolytic mechanisms that can degrade Aβ or its precursors may constitute a major determinant of AD pathogenesis and progression, especially in late-onset sporadic AD [56]. Perhaps related to this, reduced efficiency of the lysosomal system leads to the buildup of amyloidogenic species [46,57–59] and other proteins linked to aggregation events and synaptotoxicity [19,22,60–62]. Of particular interest in regards to the present study, genetic inactivation of cathepsin B resulted in increased abundance of Aβ1-42 and the worsening of Aβ pathology [16]. Corresponingly, elevating cathepsin B expression or net cathepsin activity leads to protection against Aβ pathology [16,59], and purified cathepsin B was found to cleave Aβ1-42 into shorter less pathogenic peptides [16]. Here, the induced increase in cathepsin B activity in lysosomes points to this enzyme's modulation as being involved in PADD's protective clearance of Aβ1-42. In fact, increases in the active form of cathepsin B correlated with the degree of Aβ1-42 reduction, and co-localization of Aβ1-42 and cathepsin B occurred in lysosome structures. In contrast to the reduced levels of Aβ1-42 peptide, PADD-treated transgenic mice exhibited an increase in the shorter Aβ1-38 species, indicating that the protective modulation involves the truncation of Aβ1-42. The corresponding increase in the Aβ1-38 peptide as Aβ1-42 was reduced supports the direct intracellular processing by modulated lysosomes as a route of Aβ detoxification.

Reducing the accumulation of Aβ peptides, especially the aggregation-prone species Aβ1-42, is the main objective of many approaches attempting to reduce synaptopathogenesis and associated cognitive defects in AD. Aβ species are known to disrupt neurotransmission, synaptic plasticity, and memory encoding [6,31,63–67], and cause the loss of synaptic integrity
APP-PS1 mice exhibited distinct declines in pre- and postsynaptic proteins as well. PADK-mediated lysosomal modulation and Aβ clearance translated into restoration of synaptic components, perhaps by reducing specific accumulation events that have been implicated in transport failure and axonopathy. Associated with the synaptic decline, APPswInd mice displayed balance and coordination defects and failed to recognize a familiar environment, and the older APP-PS1 mice exhibited deficits in hippocampal-dependent spontaneous alternation behavior. These data provide evidence of intracellular Aβ1-42 accumulation correlating with functional compromise. Corresponding with the synaptic protection, the lysosomal modulator treatment significantly improved performance in both mouse models, in which behavioral measures in PADK-treated transgenics were comparable to those found in non-transgenic control mice.

The opposing effects of lysosomal modulation on cathepsin B vs. Aβ1-42 levels support the idea that the lysosomal pathway works, at least in part, towards the clearing of toxic Aβ peptides. The lysosomal system responds to a pharmacologically plausible enhancement strategy that is mechanism based, resulting in intracellular Aβ clearance that also leads to reduction of extracellular deposits. These findings support new ideas regarding Aβ metabolism and the equilibrium events that influence extracellular deposition. Future work has the potential to identify lysosomal modulatory agents that have specific actions to best alleviate protein accumulation pathology. For instance, modulators of lysosomal biogenesis could tap into the TFEB pathway, thereby eliciting broad action across several lysosomal enzyme systems. Decoupling the inhibitory nature of PADK from its trafficking modulation effects may provide an important direction to develop agents that specifically enhance cathepsin maturations for promoting protein clearance. PADK’s weak inhibitory potency may explain its effect on procathepsin B as in the case of previously reported leupeptin effects, whereas the influence on mature cathepsin forms is most apparent with PADK treatment. The corresponding amelioration of synaptic and behavioral deficits in transgenic models suggests that lysosomal enhancement can provide effective protection at different stages of AD pathogenesis, thus having important implications for the development of disease-modifying therapies.

Acknowledgments
The authors thank Lara Batey, David Karanian, Robert Kwon, Dawn Jellison, and Arthur Colon for excellent assistance.

Author Contributions
Conceived and designed the experiments: DB BAB. Performed the experiments: DB JH CE SSK HBY-O MLW AC BAB. Analyzed the data: DB JH CE AN SSK HBY-O MLW AC BAB. Contributed reagents/materials/analysis tools: AN SSK HBY-O MLW AC BAB. Wrote the paper: DB JH BAB.

References
1. Walsh DM, Selkoe DJ (2007) Aβ oligomers: A decade of discovery. J Neurochem 101: 1172–1184.
2. Takahashi RH, Milner TA, Li F, Nam EE, Edgar MA, et al. (2002) Intraneuronal Alzheimer Aβ42 accumulates in multisynaptic bodies and is associated with synaptic pathology. Am J Pathol 161: 1869–1879.
3. Takahashi RH, Almeida CG, Kearney PF, Yu F, Lin MT, et al. (2004) Oligomerization of Alzheimer's β-amyloid within processes and synapses of cultured neurons and brain. J Neurosci 24: 3592–3599.
4. Oddo S, Caccamo A, Shephard JD, Murphy MP, Golde TE, et al. (2003) Triple-transgenic model of Alzheimer’s disease with plaques and tangles: Intracellular Aβ and synaptic dysfunction. Neuron 39: 409–421.
5. Li M, Chen L, Lee DHS, Yu LC, Zhang Y (2007) The role of intracellular amyloid β in Alzheimer’s disease. Prog Neurobiol 83: 131–139.
6. Billings LM, Oddo S, Green KN, McGaugh JL, LaFerla FM (2005) Intraneuronal Aβ causes the onset of early Alzheimer’s disease-related cognitive deficits in transgenic mice. Neuron 45: 675–688.
7. Bayer TA, Breyhan H, Duan K, Rettig J, Wirths O (2008) Intraneuronal Aβ-amyloid is a major risk factor–novel evidence from the APP/PS1KI mouse model. Neurodegener Dis 5: 140–142.
8. Gouras GK, Tampellini D, Takahashi RH, Capetillo-Zarate E (2010) Intraneuronal Aβ-amyloid accumulation and synapse pathology in Alzheimer’s disease. Arq Neuropsiquiatr 68: 523–541.
9. Maslennikova KG, Sugishita W, Ovod V, Munsell L, Kasten T, et al. (2010) Decreased clearance of CNS β-amyloid in Alzheimer’s disease. Science 330: 1774.
10. Gouras GK, Tsai J, Nashlund J, Vincent B, Edgar M, et al. (2000) Intraneuronal Aβ42 accumulation in human brain. Am J Pathol 156: 15–20.

Figure 13. PADK reduces behavioral deficits in APPswInd and APP-PS1 mice. In the first model, vehicle-treated wildtype mice (n = 11) were tested with groups of vehicle- (n = 10) and PADK-treated APPswInd mice (n = 13) across trials on the suspended rod test (A), and time maintained on the rod during the third trial was plotted (means±SEM). The animal groups were also tested across consecutive days in the same novel field, and the percent change±SEM in exploratory distance on the second day compared to the first was determined (B). In the second model, age-matched vehicle-treated wildtypes were tested with groups of vehicle- (n = 10) and PADK-treated APP-PS1 mice (n = 11) for spontaneous alternation behavior in a T-maze (C); data are plotted as percent of maximum alternations possible (mean±SEM). Open field mobility assessment confirmed no change in mean grid crossings±SEM across the three groups of mice (D). Post hoc tests compared to vehicle-treated transgensics: *P<0.01, **P<0.001. doi:10.1371/journal.pone.0020501.g013

[2,4,8,11,68,69]. Loss of synaptic markers was evident even at early stage Aβ pathology in the 10–11-month APPswInd mice, and
11. Bahr BA, Hoffman KB, Yang AJ, Hess US, Glabe CG, et al. (1998) Amyloid β protein is internalized selectively by hippocampal field CA1 and causes neurons to accumulate amyloidogenic carboxyterminal fragments of the amyloid precursor protein. J Comp Neurol 397: 139–147.

12. Higuchi M, Iwata N, Saido TC (2005) Understanding molecular mechanisms of proteolysis in Alzheimer's disease: Progress toward therapeutic interventions. J Neurobiol Acta 1731: 60–67.

13. Eckman EA, Eckman CB (2005) A\ldots

14. Higuchi M, Iwata N, Saido TC (2005) Understanding molecular mechanisms of proteolysis in Alzheimer's disease: Progress toward therapeutic interventions. J Neurobiol Acta 1731: 60–67.

15. Fuentealba RA, Liu Q, Zhang J, Kanekiyo T, Hu X, et al. (2010) Low-density lipoprotein receptor-related protein 1 (LRP1) mediates neuronal Aβ1–42 uptake and lysosomal trafficking. PLoS ONE 5: e11884. doi:10.1371.168: 233–234.

16. Bahr BA, Hoffman KB, Yang AJ, Hess US, Glabe CG, et al. (1998) Amyloid β protein is internalized selectively by hippocampal field CA1 and causes neurons to accumulate amyloidogenic carboxyterminal fragments of the amyloid precursor protein. J Comp Neurol 397: 139–147.

17. Ryzhikov S, Bahr BA (2008) Gephyrin alterations due to protein accumulation and lysosomal stress are reduced by the lysosomal modulator Z-Phe-Ala-diazomethylketone. J Mol Neurosci 34: 131–139.

18. Lee HJ, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

19. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

20. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

21. Ryzhikov S, Bahr BA (2008) Gephyrin alterations due to protein accumulation and lysosomal stress are reduced by the lysosomal modulator Z-Phe-Ala-diazomethylketone. J Mol Neurosci 34: 131–139.

22. Lee HJ, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

23. Lee HJ, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

24. Lee HJ, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

25. Lee HJ, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

26. Ginsberg SD, Hemby SE, Lee VM, Eberwine JH, Trojanowski JQ (2000) Pathogenic lysosomal dehydropeptidase in Parkinson's disease. J Neurosci 20: 12535–12544.

27. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

28. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

29. Bi X, Yong AP, Zhou J, Gall CM, Lynch G (2000) Regionally selective changes in N-methyl-D-aspartate receptor currents in rat brain neurons after ventricular infusion of leupeptin. Brain Res 640: 25–32.

30. Mucke L, Masliah E, Yu GQ, Mallory M, Rockenstein EM, et al. (2000) High-expression of the amyloid precursor protein potently inhibits hippocampal long-term potentiation. J Neurosci 20: 12535–12544.

31. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

32. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

33. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

34. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

35. Oddo S, Caccamo A, Smith IF, Green KN, LaFerla FM (2006) A dynamic gene network regulating lysosomal biogenesis and function. Science 325: 177–180.

36. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

37. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

38. Yan P, Bero AW, Cirrito JR, Xiao Q, Hu X, et al. (2009) Characterizing the appearance and growth of amyloid plaques in APP/PS1 mice. Neuroscience 29: 10706–10714.

39. Yu H, Crik SL, Bu G, Frieden C, Pappu RV, et al. (2009) Amyloid seeds formed by cellular uptake, concentration, and aggregation of the amyloid-β peptide. Proc Natl Acad Sci USA 106: 20224–20229.

40. Yang CN, Shiao VJ, Shee FS, Guo BS, Chen PH, et al. (2011) Mechanism mediating oligomer Aβ clearance by native primary microglia. Neurobiol Dis 42: 221–230.

41. Ginsberg SD, Alldred MJ, Coussns SE, Cataldo AM, Neve RL, et al. (2010) Microarray analysis of hippocampal CA1 neurons implicates early endosomal dysfunction during Alzheimer's disease progression. Biol Psychiatry 68: 885–893.

42. Tanaka K, Isegaki A, Ichihara A (1984) Purification and characterization of hemoglobin-hydrolyzing acid thiol protease induced by leupeptin in rat liver. J Biol Chem 259: 5937–5940.

43. van der Stappen JW, Williams AC, Maciewiz RA, Parazakev C (1996) Activation of cathespin B, secreted by a colorectal cancer cell line requires low pH and is mediated by cathespin D. Int J Cancer 67: 547–554.

44. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

45. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

46. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

47. Bahr BA (1995) Long-term hippocampal slice neuroglycination a\ldots

48. Sutherland JH, Khoshaghideh F, Patel S, Lee SJ (2004) Clearance of a\ldots

49. Butler D, Brown QB, Chin DJ, Batey L, Karim S, et al. (2005) Cellular proteolytic processing\ldots

50. Tanaka K, Isegaki A, Ichihara A (1984) Purification and characterization of hemoglobin-hydrolyzing acid thiol protease induced by leupeptin in rat liver. J Biol Chem 259: 5937–5940.

51. Salminen A, Gottesman MM (1999) Inhibitor studies indicate that active cathespin B in inflammatory cells is not responsible for the tiss\ldots

52. Mucke L, Mathews PM, Cataldo AM (2001) The neuronal endosomal-lysosomal system in Alzheimer's disease. J Alzheimers Dis 3: 97–107.

53. Butler D, Nixon RA, Bahr BA (2006) Potential compensatory responses through autophagic/lysosomal pathways in neurodegenerative diseases. Autophagy 2: 234–237.

54. Hook G, Hook VY, Kindy M (2007) Cysteine protease inhibitors reduce brain β-amyloid and β-secretase activity in vivo and are potential Alzheimer's disease therapeutics. Biol Chem 388: 979–983.

55. Klein DM, Fehrenstein KM, Bremenren D (2009) Cathespin B and L differentially regulate amyloid precursor protein processing. J Pharmacol Exp Ther 328: 813–821.

56. Selkoe DJ (2001) Clearing the brain's amyloid cobwebs. Neurosci 32: 177–180.

57. Mielke JG, Murphy MP, Maritz J, Bengualid KM, Ivy GO (1997) Chloroquine administration in mice increases \ldots

58. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.

59. Selkoe DJ (2001) Clearing the brain's amyloid cobwebs. Neurosci 32: 177–180.

60. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.

61. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.

62. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.

63. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.

64. Walsh DM, Klyubin I, Fadeeva JV, Cullen WK, Anwyl R, et al. (2002) Naturally occurring Lysosomal Modulation and therapeutic intervention. PLoS ONE 32: 813–821.
65. Cleary JP, Walsh DM, Hofmeister JJ, Shankar GM, Kuskowski MA, et al. (2005) Natural oligomers of the amyloid-β protein specifically disrupt cognitive function. Nature Neurosci 8: 79–84.

66. Shankar GM, Li S, Mehta TH, Garcia-Munoz A, Shepardson NE, et al. (2008) Amyloid-β protein dimers isolated directly from Alzheimer’s brains impair synaptic plasticity and memory. Nature Med 14: 837–842.

67. O’Nuallain B, Freir DB, Nicoll AJ, Risse E, Ferguson N, et al. (2010) Amyloid β-protein dimers rapidly form stable synaptotoxic protofibrils. J Neurosci 30: 14411–14419.

68. Lacor PN, Buniel MC, Furlow PW, Clemente AS, Velasco PT, et al. (2007) Aβ oligomer-induced aberrations in synapse composition, shape, and density provide a molecular basis for loss of connectivity in Alzheimer’s disease. J Neurosci 27: 796–807.

69. Shankar GM, Bloodgood BL, Townsend M, Walsh DM, Selkoe DJ, et al. (2007) Natural oligomers of the Alzheimer amyloid-β protein induce reversible synapse loss by modulating an NMDA-type glutamate receptor-dependent signaling pathway. J Neurosci 27: 2866–2875.