The Cysteine Protease Inhibitor, E64d, Reduces Brain Amyloid-β and Improves Memory Deficits in Alzheimer’s Disease Animal Models by Inhibiting Cathepsin B, but not BACE1, β-Secretase Activity

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Abstract. The cysteine protease cathepsin B is a potential drug target for reducing brain amyloid-β (Aβ) and improving memory in Alzheimer’s disease (AD), as reduction of cathepsin B in transgenic mice expressing human wild-type amyloid-β protein precursor (AβPP) results in significantly decreased brain Aβ. Cathepsin B cleaves the wild-type β-secretase site sequence in AβPP to produce Aβ, and cathepsin B inhibitors administered to animal models expressing AβPP containing the wild-type β-secretase site sequence reduce brain Aβ in a manner consistent with β-secretase inhibition. But such inhibitors could act either by direct inhibition of cathepsin B β-secretase activity or by off-target inhibition of the other β-secretase, the aspartyl protease BACE1. To evaluate that issue, we orally administered a cysteine protease inhibitor, E64d, to normal guinea pigs or transgenic mice expressing human AβPP, both of which express the human wild-type β-secretase site sequence. In guinea pigs, oral E64d administration caused a dose-dependent reduction of up to 92% in brain, CSF, and plasma of Aβ40 and Aβ42, a reduction of up to 50% in the C-terminal β-secretase fragment (CTFβ), and a 91% reduction in brain cathepsin B activity, but increased brain BACE1 activity by 20%. In transgenic AD mice, oral E64d administration improved memory deficits and reduced brain Aβ40 and Aβ42, amyloid plaque, brain CTFβ, and brain cathepsin B activity, but increased brain BACE1 activity. We conclude that E64d likely reduces brain Aβ by inhibiting cathepsin B and not BACE1 β-secretase activity and that E64d therefore may have potential for treating AD patients.

Keywords: Alzheimer disease, amyloid-β, BACE1, cathepsin B, guinea pig, memory deficits, oral E64d, transgenic mice

INTRODUCTION

Alzheimer’s disease (AD) results in the progressive loss of cognitive function and the accumulation of neurotoxic amyloid-β peptides (Aβ) in the brain.
The abnormal accumulation of Aβ has been demonstrated as a major disease factor based largely on studies in transgenic mouse models in which the overproduction of brain Aβ results in memory deficits and brain amyloid plaque, which mimics that seen in AD patients [4–6]. Compounds which reduce brain Aβ accumulation in transgenic mouse models have been shown as a major disease factor based largely on studies in transgenic mouse models in which the abnormal accumulation of Aβ was demonstrated as a major disease factor based largely on studies in transgenic mouse models 

The abnormal accumulation of Aβ is a hallmark of AD and they may represent asymptomatic AD [20]. AD and they may represent asymptomatic AD [20].

Brain Aβ production and its neurotoxicity, as well as the potential methods of pharmaceutical intervention to reduce that production and toxicity, have been the subject of many reviews [7–12]. Briefly, Aβ peptides are cleaved out of the amyloid-β protein precursor (AβPP) by proteases, functionally called β- and γ-secretase, which cleave AβPP at the N- and C-termini of the Aβ peptides, respectively. β-secretase first cleaves AβPP to produce the C-terminal β-secretase fragment (CTFβ) from which the γ-secretase produces Aβ of varying lengths, the most common of which contain 40 (Aβ40) or 42 (Aβ42) amino acids. Soluble Aβ monomers form into soluble oligomers, which can then form fibrils and eventually into insoluble brain amyloid plaque, but soluble Aβ oligomers appear to correlate with the cognitive decline seen in AD [13]. Another protease, which is functionally called the α-secretase, competes with the β-secretase for AβPP cleavage and produces soluble AβPP α (sAβPPα), which is neuroprotective [10, 14, 15]. A β-secretase inhibitor is a particularly attractive means of reducing brain Aβ because it would reduce both forms of neurotoxic Aβ and increase neuroprotective sAβPPα.

The vast majority of AD patients develops sporadic AD and produces Aβ by β-secretase cleavage of the normal wild-type (wt) β-secretase cleavage site sequence in AβPP [8, 16]. Thus, a β-secretase inhibitor must be able to inhibit wt β-secretase site cleavage if it is to be effective in treating most AD patients. A good model for evaluating the effects of β-secretase inhibitors on natural human wt β-secretase production of Aβ is the normal guinea pig, which expresses AβPP containing the human wt β-secretase site sequence [17–19]. But guinea pigs do not develop brain plaque or memory deficits and thus cannot be used to evaluate the pathology or behavior of AD. Transgenic mice expressing human AβPP having mutations that cause overexpression of brain Aβ, develop brain plaque and memory deficits, but none of these models perfectly reflects AD and they may represent asymptomatic AD [20]. AβPP having mutations at the β-secretase cleavage site sequence, called the Swedish mutation (Swedish), is widely used in transgenic mouse models [21]. Only one family expresses AβPP Swedish and those individuals develop familial AD, which is characterized by more brain Aβ accumulation and a much earlier onset of the disease than in sporadic AD [22]. But γ-secretase cleavage of the wt and Swe β-secretase sites may differ and thus inhibitors of the Swe β-secretase site cleavage may or may not be effective for treating most AD patients [23]. The effects of β-secretase inhibitors on wt β-secretase activity, brain plaque and memory deficits can be studied in transgenic mice expressing AβPP containing a mutation near the γ-secretase site sequence but which has the wt β-secretase site sequence. One such mutant AβPP is known as the London mutant (AβPPLondon) and is found in 30 families, which develop familial AD [24–26] (AD&FTD Mutation Database http://www.molgen.s.u.ac.be/ADMutations).

Two proteases identified as β-secretases exhibit very different cleavage efficiencies for the Swe mutant versus wt β-secretase site sequences [23]. The aspartyl protease, BACE1, is the most well known β-secretase and was originally identified as such in transfected cells and in transgenic animal models containing Swe β-secretase site sequence [27–31]. BACE1 cleaves the Swe β-secretase site sequence but barely cleaves the wt β-secretase site sequence [32–34]. The cysteine protease cathepsin B has also been identified as a β-secretase in models expressing the wt β-secretase site sequence, but in contrast to BACE1, cathepsin B readily cleaves the wt β-secretase site but does not cut the Swe mutant β-secretase site [35–37].

Although gene knockout and siRNA studies have clearly shown that cathepsin B is a valid target for reducing Aβ in models expressing the wt β-secretase site sequence [36, 38], cathepsin B’s role as a β-secretase is controversial in large part because the relationship between cathepsin B and BACE1 β-secretase activity is unclear. For example, it is possible that the reduction in Aβ, which results from deleting or suppressing the cathepsin B gene, may be due to an indirect inhibition of BACE1 β-secretase activity and cathepsin B β-secretase inhibition may be of only minor importance.

Previously, we showed that cathepsin B inhibitors reduce brain Aβ in the guinea pig and transgenic AβPPLondon mouse models [39–41] and improve memory deficits and reduce brain amyloid plaque in the transgenic AβPPLondon mouse model [41]. But in those studies, cathepsin B inhibitors were delivered directly to the brain by intracerebroventricular (icv) administration and no evaluation of brain BACE1 and cathepsin
B activities were made. Thus, those studies established the proof of principle that cathepsin B inhibitors can reduce brain Aβ plaques, and memory deficits. The studies did not show that cathepsin B inhibitors could be therapeutically developed because icv administration is not an acceptable means of treating AD patients. Moreover, they did not determine the relative importance in vivo of brain BACE1 and cathepsin B β-secretase inhibition in reducing brain Aβ by a cathepsin B inhibitor.

To evaluate these issues, we studied the effects of oral administration of a small molecule, cysteine protease inhibitor E64d (2S,3S)-trans-epoxysuccinyl-L-leucylamido-3-methylbutane ethyl ester, also known as EST and loxistatin, 342.2 mol wt) in either the guinea pig or transgenic AβPPLon mouse model. Oral E64d administration to either model caused a large dose-dependent reduction in brain Aβ, CTFF, and cathepsin B activity, but increased sAβPPu and BACE1 activity relative to controls. Importantly, feeding E64d chow to transgenic AβPPLon mice also improved spatial memory deficits, and reduced brain Aβ and amyloid plaque relative to controls. These data are consistent with E64d inhibiting β-secretase activity and show that the compound does not act by inhibiting BACE1 β-secretase activity, but most likely acts by inhibiting cathepsin B β-secretase activity. These data show that oral E64d administration is efficacious in AD animal models expressing the AβPP containing the human wt β-secretase site sequence found in most AD patients and, thus, it may have potential as an oral AD therapeutic agent.

MATERIALS AND METHODS

E64d

E64d is a prodrug ethyl ester of its biologically active acid form, E64c, and is rapidly hydrolyzed in the gut to E64c when it is orally administered. E64c then circulates systemically [42]. E64c permanently inhibits cysteine proteases by covalently binding to the sulphydryl groups in the active site of the protease [42]. Thus, E64c irreversibly binds to the active site of human papain-like cysteine proteases, B, C, F, H, K, L, O, S, W, V, and Z [43, 44], and calcium ion-dependent papain-like cysteine proteases (calpain) [45]. E64c does not inhibit BACE1 activity [23].

E64d was synthesized by American Life Science Pharmaceuticals (San Diego, CA) using methods developed and modified from those previously described [46]. Purity was determined to be 99% by reverse-phase high-pressure liquid chromatograph (RP-HPLC) assay, which was confirmed against a qualified reference standard. The identity of the compound was confirmed by 1H nuclear magnetic resonance (NMR), melting point, elemental analysis and liquid chromatography mass spectroscopy (LCMS) (data not shown). E64d exhibited excellent stability with 99% of the compound remaining after a 5-week stability test at 60°C (data not shown). However, E64d is also commercially available from various suppliers (e.g., Calbiochem, San Diego).

Animal models

Guinea pigs

Guinea Pigs (male, Hartley strain, average weight 400 g corresponding to animals about 6 weeks old) were obtained from Charles River (Wilmington, MA). The animals were given free access to standard chow and water before and during the experiment.

Transgenic AD mice

Male transgenic mice expressing human AβPP containing the wt β-secretase site and the London mutant β-secretase site sequences were generated by Applied Neurotechnology, Inc. (Charleston, SC) by methods previously described [41]. Briefly, AβPPLon was generated using site-directed mutagenesis to insert the V7171 London mutation into human AβPP cDNA. The mouse strain was created in a C57Bl/6 mouse background using the Thy-1.2 expression cassette driven by the Thy-1 promoter containing an SV40 polyadenylation site [47]. Mice were given free access to food and water before and during the experiment.

From about 3 months of age onwards, AβPPLon mice express soluble Aβ aggregates and develop cognitive impairment but they do not appreciably develop brain plaque until at least about 12 months of age and thus their cognitive impairment is dissociated from plaque formation [25, 26]. In order to evaluate the effects of E64d before and after brain plaque formation occurs, two age groups of transgenic AβPPLon mice were studied, young (6 to 9 months of age) and old (12 to 15 months of age).

Oral formulations

Gavage formulation

Delivering a drug by gavage offers the advantage of accurate dosing but is traumatic and thus only suitable for relatively short dosing periods (up to about a week). Delivery by gavage was used for the guinea pig stud-
E64d was suspended in Me2SO at the indicated concentrations (0.1, 1.0, 5, and 10 mg/kg) and administered by gavage daily using a feeding tube. Vehicle control animals were treated by gavage of Me2SO alone.

**Mouse chow formulation**
Delivering a drug by doped chow feeding allows for extended dosing periods without trauma but results in less accurate dosing because individual animals consume variable amounts of chow. Doped chow feeding was used for the transgenic AD mice experiments. A 10 mg/kg/day E64d oral dose was maximally effective at lowering brain Aβ and cathepsin B activity in guinea pigs and thus a calculated dose of that amount was administered to the transgenic mice in doped chow. E64d doped mouse chow was prepared in the mouse chow routinely used to feed the transgenic AD mice (2018, Teklad Global 18% Protein Rodent Diet, Harlan Laboratories, Inc. of Madison, WI). Briefly, 40 milligrams (mg) of E64d per kilogram (kg) of chow was processed into standard 12.7 millimeter pellets according to the manufacturer’s procedure. The resulting doped chow was analyzed by RP-HPLC and found to contain 20 mg of E64d per kg of chow. RP-HPLC studies showed that only 50% of the E64d extracted from spiked chow is detected (data not shown). Thus the doped chow actually contained about 40 mg E64d per kg of chow. The mice consume on average about 5 grams (gm) of chow per day and had an average body weight of about 20 gm. Thus, on average, mice ate a daily E64d dose of 10 mg/kg.

**Oral dosing**

**Guinea pig gavage dosing**
Guinea pigs were treated by gavage daily for one week. Guinea pigs were lightly anesthetized with isoflurane prior to treatment. 250 μl of compound solution per animal per day was administered and the animals were sacrificed 4 h after the last dosing.

**Mouse chow dosing**
Two dose regimes, 1 month and 3 months, were used for both young (6 months old) and old (12 months old) transgenic mice. All animals were fed normal chow up to the appropriate age (6 or 12 months) at which age the experimental animals were switched to feeding on E64d doped chow for 1 or 3 months. Thus, the experimental young mice were fed doped chow beginning at 6 months of age and continued with that fed until they were either 7 or 9 months old. Similarly, old mice were fed doped chow beginning at 12 months of age and continued until they were 13 or 15 months old. The experimental mice were fed doped chow while training for and being tested in the Morris water maze test. Control mice received normal chow throughout their life. All animals were fed ad libitum. No difference was observed in the amount of E64d doped and normal chow consumed by the different groups (data not shown).

**Analyses**

**Spatial memory deficits and memory retention**
The memory deficits in transgenic AβPP1-23 mice was measured using the Morris water maze test [48, 49]. Briefly, the spatial memory capability of each animal was assessed by the Morris water maze swimming test (700-0718-4 W SD Instruments)[41]. Mice were individually trained in a 1.2-m open field water maze in a pool filled with water to a depth of 30 cm and maintained at 25°C and rendered opaque by the addition of a small amount of nontoxic white powder. Four positions around the edge of the tank were arbitrarily designated north (N), south (S), east (E), and west (W); to provide four alternative start positions and define the division of tank into four quadrants: NE, SE, SW, and NW. An escape platform (10 cm square and white) was placed 1 cm below the surface of the water and invisible to the mice. All animals underwent nonspatial pretraining for 5 consecutive days, which prepared the animals for the final behavioral test to determine the retention of memory to find the platform. The day following the nonspatial pretraining, the hidden platform was placed in the center of one quadrant of the pool. The animal was released into the pool facing the pool wall in a random fashion and the time (latency period) and the distance traveled from the release to the animal standing on the platform was measured using video recording (Smart Video Tracking System; SD Instruments). On the day after the last training session, the platform was removed and a spatial probe test conducted in which each mouse was allowed to search for the platform for 60 seconds (memory retention) and the percent time spent in the NE target quadrant where the platform had been was determined.

**Brain amyloid plaque**
Amyloid load was assessed in brain sections from mice by methods previously used [41]. Amyloid plaques were located by immunostaining for Aβ (Aβ antibody 10D5, Elan Pharmaceuticals). Brain tissues were fixed in 4% paraformaldehyde and then in 4% par-
formaldehyde and 30% sucrose for 24 h each at 4°C. Tissues were washed in buffered saline and transferred to an optimum cutting temperature (O.C.T.) medium. Cryosections were cut and blocked with normal serum, incubated with anti-Aβ and stained with diaminobenzic acid (Vector ABC Elite kit, Vector Laboratories). Bright field light microscopy imaged brain areas from which stained amyloid areas were quantitated using image analysis (NIH Image software).

**Brain Aβ analysis**
The brain Aβ analysis method was that previously described for guinea pigs and transgenic AD mice [39–41]. Animals were sacrificed and brain extracts were prepared as previously described. Briefly, brain extracts were homogenized (1:3 weight/volume of buffer) in buffer of 5 M guanidine HCl in 50 mM Tris–HCl, pH 7.6, 150 mM NaCl, plus protease inhibitors (Sigma). Homogenates were diluted to 0.5 M guanidine and centrifuged (200,000 g for 20 min), and supernatant and pellet fractions were collected. Protein content was determined by the Bradford method [50].

Enzyme-linked immunosorbent assays (ELISAs) measured Aβ peptides by methods previously described [51, 52]. The pellet from the brain extract procedure was sonicated in 6 M guanidine and centrifuged at 200,000 g for 20 min at 4°C, and the supernatant was diluted to 0.5 M guanidine. The two supernatants were combined, and Aβ40 and Aβ42 were determined using ELISA kits specific for each peptide (IBL, JP27718 and JP27711).

**CSF Aβ**
CSF was collected from the guinea pig brains and protease inhibitors were added and Aβ40 and Aβ42 were determined using ELISA kits specific for each peptide (IBL, JP27718 and JP27711).

**Plasma Aβ**
Plasma samples were collected from the guinea pigs following sacrifice, protease inhibitors were added and Aβ40 and Aβ42 were determined using ELISA kits specific for each peptide (IBL, JP27718 and JP27711).

**Brain CTFβ and sAβPPα**
CTFβ and sAβPPα were determined by methods previously described [39–41]. Briefly, CTFβ and sAβPPα were determined in the pellet and supernatant fractions, respectively, from the brain extract by Western blot (antibody A8717 and antibody 6E10, respectively, Sigma). Relative amounts of CTFβ and sAβPPα were assessed by densitometry of the Western blots.

**Brain cathepsin B activity analysis**
Cathepsin B activity was determined using a fluorometric assay kit from Abcam (ab65300) as described by the manufacturer.

**Brain BACE1 activity analysis**
The brain BACE1 activity was determined using a fluorometric assay kit from Abcam (ab65357) as described by the manufacturer.

**Animal treatment**

**Humane animal use**
Animal studies were conducted according to regulations by the NIH and as approved by the IACUC at the Medical University of South Carolina and Ralph H. Johnson VA Medical Center.

**Statistics**
10 guinea pigs and transgenic AβPP mice were studied per group. Numerical data were collected and entered into GraphPad Prizm 5 software, which was used to graphically display and analyze the results. The means of the guinea pig data were compared using ANOVA and Dunnett’s multiple comparison tests, which compared the control group with the E64d treated groups. The means of the transgenic AβPP mouse data were compared using unpaired t-test comparisons of age-matched animals receiving normal or E64d doped chow.

The guinea pig dose-response reduction curves were determined using linear regression two-phase exponential decay, one fast and on slow, least squares fit. The paired guinea pig and transgenic AβPP mouse data were analyzed using linear regression analysis.

**RESULTS**

**Guinea pig data**

**Guinea pig Aβ**
Oral administration of E64d to guinea pigs resulted in a dose-dependent reduction in brain, CSF and plasma Aβ40 and Aβ42 (Fig. 1 A–F). All three compartments and both Aβ forms had similar biphasic dose responses with a sharp reduction in Aβ occurring between 0.0 to 1 mg/kg/day dose with a more gradual
Fig. 1. E64d reduces brain, CSF, and plasma Aβ1-40 and Aβ1-42 in guinea pigs. Once-a-day oral administration of E64d for one week to guinea pigs results in similar biphasic dose-dependent reductions in (A) brain Aβ1-40, (B) brain Aβ1-42, (C) CSF Aβ1-40, (D) CSF Aβ1-42, (E) plasma Aβ1-40, and (F) plasma Aβ1-42. The minimum effective dose was about 1 mg/kg/day. The maximum dose (10 mg/kg/day) caused a 90% reduction or more in both Aβ1-40 and Aβ1-42 in all compartments. The guinea pig is a natural model of human wild-type AβPP processing and, thus, the major reduction in Aβ caused by oral E64d administration in that model suggests that the compound may have similar effects in humans. The fact that both Aβ1-40 and Aβ1-42 were similarly affected in all compartments is consistent with the compound inhibiting Aβ-secretase activity because Aβ-secretase produces both Aβ1-40 and Aβ1-42 and both forms were similarly reduced.

The mean Aβ1-40 and Aβ1-42 in all compartments and was significantly less than the control group (no E64d) for all E64d doses equal or greater than 1 mg/kg/day. The highest dose, 10 mg/kg/day, achieved the maximum reduction of about 92% lower than the control group (no E64d) in Aβ1-40 and Aβ1-42 in all compartments (brain, CSF, and plasma). These data are consistent with the inhibitor reducing β-secretase activity because β-secretase produces both Aβ1-40 and Aβ1-42 and both forms were similarly reduced.
Guinea pig brain CTFβ
E64d also caused a biphasic dose-dependent reduction in brain CTFβ (Fig. 2A, B). The mean CTFβ levels were significantly lower than the control group (no E64d) for E64d doses of 1 mg/kg/day or greater with the highest E64d dose resulting in the maximal reduction in CTFβ of about 50% lower than that of the control group. These data are also consistent with E64d reducing β-secretase activity because CTFβ is the cleavage product of β-secretase.

Guinea pig brain sAβPPα
E64d caused a dose-dependent increase in brain sAβPPα (Fig. 2C, D). The mean sAβPPα levels were significantly higher than the no dose group for E64d doses of 5 mg/kg/day or greater with the highest E64d dose resulting in the maximum increase in sAβPPα of about 54% more than the control group. These data also support the hypothesis that E64d is reducing β-secretase activity because sAβPPα is derived from AβPP α-secretase cleavage, which competes with β-secretase cleavage of AβPP, and thus the inhibition of β-secretase activity provides more AβPP for α-secretase to produce more sAβPPα.

Guinea pig brain BACE1 activity data
In contrast, oral E64d administration produced a slight dose-dependent increase in brain BACE1 activity (Fig. 3B). The increase in BACE1 activity rises as the dose increases to 5.0 mg/kg/day and levels off.

Fig. 2. E64d lowers and increases brain CTFβ and sAβPPα, respectively, in guinea pigs. Oral administration of E64d to guinea pigs caused a dose-dependent reduction in brain CTFβ with the maximum dose reducing CTFβ by more than 50% (Fig. 2A, B). CTFβ is a proteolytic product of β-secretase cleavage of AβPP and thus these data are also consistent with the compound acting by inhibiting β-secretase activity. In contrast, the oral E64d treatment caused a dose-dependent increase in brain sAβPPα with the maximum dose increasing sAβPPα by about 54% (Fig. 2C, D). These data are consistent with E64d inhibiting β-secretase activity because sAβPPα is derived from AβPP α-secretase cleavage, which competes with β-secretase cleavage of AβPP, and thus the inhibition of β-secretase activity provides more AβPP for β-secretase to produce more sAβPPα. The mean and SEM are displayed at each data point (Dunnett’s multiple comparison test ***p < 0.05, n = 10 animals/group).
Fig. 3. E64d reduces and increases brain cathepsin B and BACE1 activities, respectively, in guinea pigs. Oral E64d administration to guinea pigs caused a biphasic dose-dependent reduction in brain cathepsin B activity with a minimum effective dose of about 1 mg/kg/day and the maximum dose lowering cathepsin B activity by 90% (Fig. 3A). As such, the brain cathepsin B effect is very similar to reduction in Aβ caused by E64d. Cathepsin B inhibition by E64d is required if the compound acts by inhibiting cathepsin B-secretase activity and thus these data support that mechanism of action. In contrast, that treatment caused an increase in brain BACE1 activity with the maximum dose increasing BACE1 activity by almost 20% (Fig. 3B). The increase in BACE1 activity rules out the hypothesis that E64d reduces brain Aβ by inhibiting BACE1-secretase activity because BACE1 activity was increased. The mean and SEM are displayed at each data point (Dunnett’s multiple comparison test ***p<0.05, n=10 animals/group).

5.0 and 10.0 mg/kg/day doses resulted in the maximum increase in brain BACE1 activity of about 20% more than the control group. Thus, E64d increases guinea pig brain BACE1 activity, which rules out E64d acting by inhibiting BACE1-secretase activity in this model.

Guinea pig brain Aβ40 or Aβ42 positively correlated with brain cathepsin B activity

Paired analysis of brain Aβ40 or Aβ42 and brain cathepsin B activity from individual guinea pigs showed a slight negative correlation for Aβ40 or Aβ42, respectively (Fig. 4A, B). Thus, brain cathepsin B activity is strongly predictive of brain Aβ40 and Aβ42 levels, which is consistent with E64d reducing Aβ by inhibiting cathepsin B β-secretase activity.

Guinea pig brain Aβ40 or Aβ42 negatively correlated with brain BACE1 activity

Paired analysis of brain Aβ40 or Aβ42 and brain BACE1 activity from individual guinea pigs showed a slight negative correlation for Aβ40 or Aβ42, respectively (Fig. 4C, D). Thus, brain BACE1 activity is not correlated with brain Aβ, which is inconsistent with E64d reducing brain Aβ by inhibiting BACE1 β-secretase activity.

Transgenic AβPP671/71 mice data

Mouse brain Aβ data

Young (6–9 months of age) and old (12–15 months of age) transgenic mice fed E64d doped chow for 1 or 3 months had significantly lower brain levels of Aβ40 or Aβ42 (Fig. 5A, B). The brain Aβ40 or Aβ42 reductions as a percentage of control for each age group and feeding schedule are summarized in Tables 1 and 2. The average brain Aβ40 and Aβ42 reduction for both treatments in both age groups was 62% and 58%, respectively. Again, this data are consistent with the inhibitor reducing β-secretase activity because both Aβ40 and Aβ42 were similarly reduced.

Mouse brain CTFβ

E64d feeding caused a significant 50% reduction in brain CTFβ for both feeding schedules in both age groups relative to controls (Fig. 6A, B, C). These data are also consistent with E64d reducing β-secretase activity because CTFβ is the cleavage product of β-secretase and it is reduced with treatment.

Mouse brain sAβPPα

E64d feeding caused a significant increase in brain sAβPPα for both feedings and in both age groups (Fig. 6D, E, F). Treatment for one or 3 months in the young or old animals resulted in a 21% or 34% increase, respectfully, in brain sAβPPα relative to controls. As discussed above, these data also support the hypothesis that E64d is reducing β-secretase activity because β-secretase inhibition increases sAβPPα.

Mouse brain cathepsin B activity

The E64d treatment for one or 3 months in the young and old mice caused an average a 77% reduction in
Cathepsin B and Aβ are positively correlated, BACE1 and Aβ are negatively correlated in guinea pigs. The individual paired data from all guinea pigs for brain Aβ and Aβ data versus brain cathepsin B activity were plotted (Fig. 4A, B). Linear regression analysis for brain Aβ versus brain cathepsin B activity finds a $r^2 = 0.98$ and a 95% confidence interval for a 0.21 to 0.22 slope, which is a significantly non-zero slope at $p < 0.0001$. For brain Aβ versus brain cathepsin B activity, linear regression analysis of finds a $r^2 = 0.87$ and a 95% confidence interval for a 0.45 to 0.56 slope, which is a significant non-zero slope at $p < 0.0001$. Thus, brain cathepsin B activity is a strong predictor of brain Aβ and these data are consistent with E64d reducing brain Aβ by inhibiting cathepsin B β-secretase activity. In contrast, paired data from individual guinea pigs for brain Aβ and Aβ vs. brain BACE1 activity at the various E64d doses shows a very weak, slightly negative correlation (Fig. 4C, D). Linear regression analysis of brain Aβ versus brain BACE1 activity finds a $r^2 = 0.45$ and a a 95% confidence interval for a $-0.06$ to $-0.03$ slope, which is a significant non-zero slope at $p < 0.0001$. Linear regression analysis of brain Aβ versus brain BACE1 activity finds $r^2 = 0.36$ and a 95% confidence interval for a $-0.14$ to $-0.06$ slope, which is a significant non-zero slope at $p < 0.001$. These data show that brain BACE1 activity is not predictive of brain Aβ and argue against E64d reducing brain Aβ by inhibiting BACE1 β-secretase activity.

Mouse brain BACE1 activity

E64d feeding for one or 3 months caused higher brain BACE1 activity in all mice, but a statistically significant increase averaging only 17% occurred in the old mice (Fig. 7B). Thus, E64d treatment increased brain BACE1 activity in the APP/PLoN mice. These data rule out E64d as inhibiting brain BACE1 β-secretase activity and thereby lowering brain Aβ because brain BACE1 activity would have to decrease, not increase as was observed, for that to be the mechanism of action.

Mouse memory deficits – latency period

Feeding E64d for one or 3 months significantly improved the memory deficits of both age groups as measured by the latency period in the Morris water maze swimming test (Fig. 8A). The percent improvement in the latency period for each age group and feeding schedule is summarized in Table 3 and is an average improvement of 35%. The average swimming
Fig. 5. Feeding E64d chow to transgenic AD mice reduces brain Aβ40 and Aβ42. Brain Aβ is shown as a function of the mouse age with young (6 – 9 months of age) and old (12 – 15 months of age) mice boxed separately. The arrows indicate the age at which the experimental animals were switched from normal to E64d chow and fed that chow for either one or 3 months. In other words, the young animals were fed the E64d chow until they were 7 or 9 months old and the old animals were fed the E64d chow until they were 13 or 15 months old. Brain Aβ40 was significantly reduced in both young and old mice fed for either one or 3 months (Fig. 5A, arrows indicate age at which E64d feeding began, and Table 1). Brain Aβ42 was also significantly reduced in both young and old mice fed E64d chow for either one or 3 months (Table 2). Thus, even though brain Aβ40 and Aβ42 increased with increasing age in control animals, the Aβ40 and Aβ42 in the E64d treated animals was consistently reduced at both ages and feeding schedules. Particularly significant is that E64d was efficacious at reducing Aβ40 and Aβ42 in the old animals even though feeding was not initiated until after those animals had accumulated abnormal levels of Aβ40 and Aβ42. This suggests that the compound is effective at arresting further brain Aβ40 and Aβ42 accumulation even after abnormal levels have developed. The mean and SEM are displayed at each data point. Stars represent statistically significant differences between the means of age-matched E64d and normal chow fed mice (t-test comparison ***p<0.001 10 animals/group).

Table 1
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<tr>
<th>Mouse Age</th>
<th>E64d feeding treatment (months)</th>
<th>Reduction in brain Aβ40 relative to controls (%)</th>
</tr>
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<tr>
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<td>50</td>
</tr>
<tr>
<td>Young†</td>
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<td>Old‡</td>
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<td>Average for all ages and treatments</td>
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†Young mice began E64d feeding at 6 months of age; ‡Old mice began E64d feeding at 12 months of age.

Table 2
<table>
<thead>
<tr>
<th>Mouse Age</th>
<th>E64d feeding treatment (months)</th>
<th>Reduction in brain Aβ42 relative to controls (%)</th>
</tr>
</thead>
<tbody>
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<td>Young†</td>
<td>1</td>
<td>52</td>
</tr>
<tr>
<td>Young†</td>
<td>3</td>
<td>72</td>
</tr>
<tr>
<td>Old‡</td>
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<td>46</td>
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<td>61</td>
</tr>
<tr>
<td>Average for all ages and treatments</td>
<td>undefined</td>
<td>58</td>
</tr>
</tbody>
</table>

†Young mice began E64d feeding at 6 months of age; ‡Old mice began E64d feeding at 12 months of age.

Mouse memory deficits – distance traveled

Feeding E64d for one or 3 months caused a significant reduction in the distance traveled in the Morris water maze test (Fig. 8B). The percent improvement in the latency period for each age group and feeding schedule is summarized in Table 4 and is an average improvement of 34%. Significantly, the E64d treatment for one and 3 months resulted in a marked improvement in memory deficits for the old mice, which had already developed impairment. Again, these data support the potential of oral E64d for improving memory deficits in AD patients, possibly even after it has developed.

Mouse memory retention

In the spatial probe trial, the platform was removed from the pool and memory retention was measured by determining the percent of a 60 second time period in which the animal swam in the NE quadrant where the platform had previously been. The mice fed the E64d
Fig. 6. Feeding E64d chow to transgenic AβPP/APP mice decreases brain CTFβ and increases sAβPPα. Western blots show that feeding E64d chow results in qualitatively less brain CTFβ than controls (Fig. 6A-C, arrows indicate the age at which E64d feeding began). Quantitative densitometry of the Western CTFβ blots shows that brain CTFβ is reduced in young and old animals fed E64d for one or 3 months by about 50% (Fig. 6C, arrows show age at which E64d chow feeding began). As CTFβ is a proteolytic fragment of β-secretase cleavage, these data are consistent with E64d acting by inhibiting β-secretase activity. In contrast, Western blots of sAβPPα qualitatively show that E64d feeding increased brain sAβPPα relative to controls (Fig. 6D-F). Quantitative densitometry of the Western sAβPPα blots shows that there is consistently more brain sAβPPα in young and old animals fed E64d for one or 3 months than in controls (Fig. 6F). Increased sAβPPα is consistent with E64d inhibiting β-secretase activity because that inhibition would allow more production of sAβPPα. The mean and SEM are displayed at each data point (t-test comparison *p < 0.02, **p < 0.004, ***p < 0.0001, n = 10 animals/group).

Mouse brain amyloid plaque
As discussed above transgenic AβPP/APP mice accumulate brain Aβ and amyloid plaque as they age. Young mice fed either the control or E64d chow for one month (7 months old) had no brain amyloid plaque (Fig. 9). But young mice fed the control for 3 months (until they were 9 months old) had only a small amount of plaque (0.12% area), which the E64d diet completely prevented. Consistent with previous transgenic AβPP/APP mouse studies [25, 26], only...
Feeding E64d chow to transgenic AβPP695AD mice reduces brain cathepsin B activity and either has no effect or increases brain BACE1 activity. Feeding E64d chow to either young or old mice for one or 3 months reduced brain cathepsin B activity by about 83% (Fig. 7A, arrows show age at which E64d chow feeding began). Brain cathepsin B inhibition is required if E64d acts by suppressing cathepsin B/b-secretase activity and thus these data are consistent with the compound acting by that mechanism of drug action. In contrast, feeding E64d chow for one or 3 months to young mice increased brain BACE1 activity but the increase was not statistically significant (Fig. 7B). Feeding E64d chow for one or 3 months to old mice significantly increased brain BACE1 activity relative to animals fed normal chow by about 16%. These data show that the effects of E64d on brain Aβ42 are not due to off-target inhibition of brain BACE1/b-secretase activity because E64d treatment did not inhibit brain BACE1 activity, but, rather, increased brain BACE1 activity. The mean and SEM are displayed at each data point (t-test comparison *p < 0.04, n = 10 animals/group).

Older (>12 months of age) animals had significant brain plaque. Old animals fed the control or the E64d diet for one month (until they were 13 months old) had 0.35 or 0.18% plaque areas, respectively, with the control chow E64d chow

Young mice Old mice

Age (months)

6 7 9 12 13 15

Brain Cathepsin B Activity (Relative to Units)

Distance Traveled (Centimeters)

Latency Period (sec)

Fig. 8. Feeding E64d chow to transgenic AβPP695AD mice improves memory deficits and memory retention. As they age, transgenic AβPP695AD mice mimic the behavior of AD patients in that they progressively develop memory deficits, which can be assessed using the Morris water maze test by measuring the time it takes the animals to find an invisible, submerged platform (latency period) and distance traveled to find that platform. Feeding E64d chow to young (6–9 months) or old (12–15 months of age) mice for either one or 3 months significantly improved the animals’ ability to remember the location of the platform as measured by the latency period (Fig. 8A, arrows indicate age at which E64d feeding began, and Table 3). Significantly, E64d treatment improved memory deficits in 15-month-old animals, which had significant memory deficits, suggesting that the compound may be able to restore memory after a deficit has occurred. Similarly, feeding E64d chow to young or old mice for either one or 3 months reduced the distance traveled (Fig. 8B and Table 4). Again, this data shows that oral administration of E64d is able to improve memory deficits that otherwise will occur in this transgenic AD animal model. To test memory retention, mice were also tested for the percentage of time spent in the northeast (NE) target quadrant after the platform was removed from that quadrant (Fig. 8C and Table 5). All E64d treated groups spent more time in the target quadrant than did age-matched untreated animals showing that the E64d resulted in greater memory retention. Together, these data suggest that oral E64d administration may also be able to prevent or reverse the memory decline seen in AD patients. The mean and SEM are displayed at each data point (t-test comparison ***p < 0.0001, n = 10 animals/group).
Table 3
Improvement of memory deficits as measured by the latency period in the Morris water maze test for young and old transgenic AD mice fed E64d for one or 3 months

<table>
<thead>
<tr>
<th>Mouse Age</th>
<th>E64d feeding treatment (months)</th>
<th>Improvement in latency period relative to controls (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Young†</td>
<td>1</td>
<td>25</td>
</tr>
<tr>
<td>Young‡</td>
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<td>41</td>
</tr>
<tr>
<td>Old†</td>
<td>1</td>
<td>35</td>
</tr>
<tr>
<td>Old‡</td>
<td>3</td>
<td>38</td>
</tr>
<tr>
<td>Average for all ages and treatments</td>
<td>35</td>
<td></td>
</tr>
</tbody>
</table>

†Young mice began E64d feeding at 6 months of age; ‡Old mice began E64d feeding at 12 months of age.

Table 4
Improvement of memory deficits as measured by the distance traveled in the Morris water maze test for young and old transgenic AD mice fed E64d for one or 3 months

<table>
<thead>
<tr>
<th>Mouse Age</th>
<th>E64d feeding treatment (months)</th>
<th>Improvement in distance traveled relative to controls (%)</th>
</tr>
</thead>
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<tr>
<td>Young†</td>
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<td>25</td>
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<td>Young‡</td>
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<td>41</td>
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<td>35</td>
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<tr>
<td>Old‡</td>
<td>3</td>
<td>38</td>
</tr>
<tr>
<td>Average for all ages and treatments</td>
<td>35</td>
<td></td>
</tr>
</tbody>
</table>

†Young mice began E64d feeding at 6 months of age; ‡Old mice began E64d feeding at 12 months of age.

Table 5
Improvement of memory retention as measured by the percentage of time spent in the quadrant, which had previously contained a submerged platform, for young and old transgenic AD mice fed E64d for one or 3 months

<table>
<thead>
<tr>
<th>Mouse Age</th>
<th>E64d feeding treatment (months)</th>
<th>Improvement in percentage of time spent in NE quadrant (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Young†</td>
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<td>35</td>
</tr>
<tr>
<td>Young‡</td>
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<td>3</td>
<td>207</td>
</tr>
<tr>
<td>Average for all ages and treatments</td>
<td>105</td>
<td></td>
</tr>
</tbody>
</table>

†Young mice began E64d feeding at 6 months of age; ‡Old mice began E64d feeding at 12 months of age.

One-month E64d treatment causing a 49% reduction in brain plaque. Old animals fed the control or the E64d diet for 3 months (until they were 15 months old) had 0.68 or 0.27% plaque areas, respectively, with the 3-month treatment causing a 60% reduction in brain plaque. Thus, the E64d treatment not only reduced brain Aβ but it also reduced the brain amyloid plaque load, which is a pathological hallmark of AD.

Mouse latency period is positively correlated with brain Aβ

Paired analyses of the latency period and brain Aβ data from individual mice showed strong positive correlations (Fig. 10A–D). The one and 3-month feeding schedule data was combined and the control and treated data separately analyzed. Interestingly, in the older animals, the slope for the E64d treated animals was about 2 times steeper than that of the control for both Aβ40 and Aβ42. The steeper slopes for the treated animals suggest that E64d may affect other factors in addition to brain Aβ and thereby improve memory.

DISCUSSION

A significant result of this study is that simply feeding E64d chow to transgenic AβPPLon mice improved the memory deficits and memory retention in these mice relative to age-matched transgenic mice fed control chow (Fig. 8). On average, young and old mice fed E64d chow for one or 3 months had about a 35% improvement in their latency period and their distance traveled swimming to the submerged platform in the...
Fig. 9. E64d treatment of transgenic AβPP/lon AD mice reduces brain amyloid plaque. Transgenic AβPP/lon mice also mimic the pathology seen in AD patients in that they develop brain amyloid plaque as they age. Quantitative image analysis of brain sections stained for amyloid plaque in young mice show that control and treated mice have no amyloid plaque at 7 months of age and that control and E64d treated mice have very little amyloid plaque and no amyloid plaque, respectively, at 9 months of age (Fig. 9A arrows show age at which E64d chow feeding began). Quantitative image analysis of brain sections from old mice show that control mice have significant amyloid plaque at 13 and 15 months of age and that E64d significantly reduced the amyloid plaque at both ages (Fig. 9A). Histological micrographs of a brain region from a 15 month-old mouse fed control chow for 3 months shows numerous plaques, including one in the hippocampus pointed to by the arrow (Fig. 9B). A corresponding histological micrograph from a 15 month-old animal fed E64d chow for 3 months shows very little plaque (Fig. 9C). These data suggest that oral E64d administration may also be able to reduce the brain plaque seen in AD patients. The mean and SEM are displayed at each data point (t test comparison **p < 0.002, ***p < 0.0001, n = 10 animals/group).

Morris water maze test relative to animals fed control chow. These data show that the E64d improved memory deficits. Moreover, in the memory retention test in which the platform was removed, E64d fed animals spent an average 105% greater percent time in the quadrant where the platform had been relative to controls. These data show that E64d improved memory retention. Importantly, beginning E64d feeding when mice were old (12 months of age) and had already developed significant memory impairment, improved the memory deficits and memory retention relative to age-matched controls. That shows that in this model the compound retards or arrests the memory decline even after it has manifested itself. Young mice also have memory deficits, although not as severe as that in old mice, and feeding E64d to young mice beginning at 6 months of age significantly improved their memory deficits and memory retention relative to age-matched controls. Thus, the data show that oral E64d administration improves in young and old animal models the primary problem in AD, memory decline, which suggests that the compound may have potential for treating early and late stages of AD.

Feeding E64d to transgenic AβPP/lon mice also reduces brain amyloid plaque, which develops in these animals and mimics that pathology seen in AD patients (Fig. 9). Feeding E64d to mice that are old and have significant brain plaque reduced the brain plaque by 49% on average relative to age-matched controls. Although the young animals had not yet manifested significant amounts of brain plaque, the small amount found in the 9 month old controls was completely absent in
the age-matched E64d treated mice. Thus, oral E64d administration reduces a characteristic AD pathology, brain amyloid plaque, in this animal model and that also suggests E64d may have AD therapeutic potential.

The brain amyloid plaque and memory deficits observed in the transgenic AβPPLon mice are likely due to the abnormal accumulation of brain Aβ because the major difference between the transgenic AβPPLon mice and their non-transgenic strain matched controls is brain Aβ accumulation. The brain amyloid plaque that develops in these animals is undoubtedly caused by the abnormal brain Aβ accumulation because brain Aβ peptides are the primary component of brain amyloid plaque. In regard to the memory deficits, the fact that E64d treated animals had significantly reduced brain Aβ and improved memory deficits relative to age-matched controls not fed E64d is consistent with brain Aβ causing the memory deficits (Figs. 5 and 8). Furthermore, the strong positive correlation found between brain Aβ40 and Aβ42 and increasing memory deficits also supports this hypothesis (Fig. 10). In young mice, the correlation slopes between Aβ40 or Aβ42 and the latency period were about the same for E64d treated and control transgenic mice showing that the brain Aβ effect on memory was proportionally the same for both groups. In old mice, on the other hand, the correlation slope between the brain Aβ40 or Aβ42 and the latency period was significantly more positive...
for the E64d treated than for the old control transgenic mice showing that brain Aβ effect on memory was not proportionally the same for both groups, which suggests that additional factor(s) may also contribute to the improved memory seen in the old animals treated with E64d. As discussed below, such additional factors may be due to the neuroprotective effects of E64d.

Another important result of this study is that oral administration of E64d to transgenic AβPP/Lon mice or normal guinea pigs results in an approximately equal reduction in brain Aβ40 and Aβ42, a concomitant reduction in brain CTFβ and an increase in brain sAβPPs. In the transgenic AβPP/Lon mice, E64d treatment for all ages and treatments reduced brain Aβ40 and Aβ42 by about 62% and 58%, respectively, brain CTFβ by about 50% and increased brain sAβPPs by about 29%. In guinea pigs, E64d treatment produced dose-dependent reductions in brain Aβ40 and Aβ42, brain CTFβ and a dose-dependent increase in brain sAβPPs. These data agree with our previous work, which showed that icv E64d administration to either transgenic AβPP/Lon mice or normal guinea pigs also resulted in the same brain Aβ peptide pattern [39–41]. That pattern is consistent with the compound reducing brain Aβ by lowering its production through inhibiting β-secretase activity [23]. Thus, the memory improvement combined with reduced brain Aβ and amyloid plaque, which occurred upon treating the transgenic AβPP/Lon mice, is likely due to a decrease in brain Aβ production caused by inhibition of β-secretase activity.

But the question is which β-secretase activity is inhibited by E64d? Two proteases are currently proposed to have that activity, the aspartyl protease BACE1 and the cysteine protease cathepsin B. Deleting the BACE1 gene in transgenic mice expressing AβPP containing the Swe β-secretase site sequence results in the characteristic peptide pattern expected of β-secretase inhibition [53], but detailed evaluation of the BACE1 gene knockout studies indicate that they do not preclude participation of cathepsin B in Aβ production (see review in [23]). Likewise, deleting the cathepsin B gene in transgenic mice expressing human wt AβPP also results in that peptide pattern of β-secretase inhibition [36]. The data here clearly show that E64d administration caused an increase in brain BACE1 activity and a decrease in brain cathepsin B activity in both the guinea pigs and the transgenic AβPP/Lon mice. BACE1 activity was slightly negatively correlated whereas brain cathepsin B activity was highly positively correlated with brain Aβ40 or Aβ42, versus brain Aβ40 or Aβ42 in guinea pigs. Thus, brain BACE1 activity had no predictive power whereas brain cathepsin B activity had very high predictive power for determining brain Aβ40 or Aβ42 levels in guinea pigs. These data are consistent with in vitro data, which has shown that E64d has no effect on BACE1 activity but is a potent cathepsin B inhibitor [23]. Thus, the data show that E64d is not reducing brain Aβ through BACE1 β-secretase inhibition but that it is likely inhibiting cathepsin B β-secretase activity. It is not clear why E64d causes an increase in brain BACE1 activity but it may be due to a compensatory mechanism by which the brain is attempting to augment the reduced cathepsin B β-secretase by increasing BACE1 activity. But if that is the case, the compensation is ineffective with respect to Aβ because E64d treatment results in significantly reduced brain Aβ. Much more research is needed to understand the relationship between cathepsin B and BACE1 β-secretase activity.

Both the transgenic AβPP/Lon mice and the guinea pigs express AβPP, which contain the human wt β-secretase site sequence, thus the β-secretase activity that occurs in these animal models is that found in the vast majority of AD patients. E64d’s ability to inhibit wt β-secretase activity and thereby reduce brain Aβ in these animal models suggests that E64d is likely to do so in most AD patients. Moreover, guinea pigs and AD patients express physiologically normal levels of AβPP and thus the fact that E64d reduced brain Aβ in guinea pigs argues that it also will have a similar effect on processing in AD patients.

The strong positive correlation between increasing Aβ40 or Aβ42 and worsening memory as measured by the latency period in the transgenic AβPP/Lon mice (Fig. 10) is similar to the positive correlation observed between Aβ40 or Aβ42 and cognitive decline in AD patients [54]. As previously reported [25, 26], the young (6–9 months of age) transgenic AβPP/Lon mice developed memory deficits prior to the formation of significant brain amyloid plaque (Figs. 8 and 9A). Young transgenic AβPP/Lon mice are known to form soluble Aβ aggregates [26], which are similar to the soluble Aβ oligomer species correlated with cognitive decline in AD [13], and are thought to model mild cognitively impaired (MCI) patients [26]. As such, the improvement in memory deficits resulting from E64d treatment of young transgenic AβPP/Lon mice bodes well for the compound being effective at treating MCI or early stage AD. Also as previously reported [25, 26], the old (12–15 months of age) transgenic AβPP/Lon mice developed memory deficits and significant brain amyloid plaque (Figs. 8 and 9A). Our finding that amyloid plaque did not correlate with memory...
deficits in old mice is consistent with earlier AD studies, which also found very little correlation between amyloid deposits and dementia [55–57]. Nonetheless, the fact that E64d treatment improved memory deficits and reduced brain amyloid plaque, a hallmark of AD, in old animals, suggests that the compound may be effective for advanced AD. Moreover, the brain amyloid plaque in old transgenic AβPP/PP mice is known to contain post-translationally modified Aβ forms called pyroglutamate Aβ (pyroGluAβ) [26], which may be critical in late stage disease development [58] because they are highly neurotoxic [59], greatly accelerate Aβ aggregation [60, 61], and are much more prevalent in AD brain than age-matched controls [62]. Thus, E64d may be able to reduce amyloid plaque containing the particularly pernicious pyroGluAβ form found in AD brain. Together, the data support the possibility that E64d may be successful in treating MCI or advanced AD patients.

On the other hand, treating transgenic mice expressing AβPP containing the Swe mutant or CTFβ levels [41, 63]. This is consistent with data showing that deleting the cathepsin B gene in such animals reduces brain Aβ levels [41, 63]. The lack of effect is likely due to the fact that cathepsin B does not cleave the Swe β-secretase site sequence to produce Aβ and thus inhibiting cathepsin B activity or deleting the cathepsin B gene does not reduce brain Aβ or CTFβ in animals expressing AβPP/Swe [23].

The ability to inhibit other cysteine proteases in addition to cathepsin B provides the E64 family of compounds the added advantage of also being potent neuroprotectants. For example, as discussed above, chronic E64 injection (ip) improved memory deficits in an AD transgenic AβPP/Swe mouse model relative to controls even though brain Aβ levels were not affected [63]. This memory improvement was attributed to E64’s inhibition of calpain, another cysteine protease. Increased Aβ causes increased calpain activity, which causes synaptic dysfunction and, thus, E64 is thought to reduce the synaptic dysfunction and thereby improve memory deficits by inhibiting calpain activity. AD results in widespread brain calpain activation and has long been considered a molecular basis of AD [65]. E64d and E64-c have also been shown to have significant neuroprotective effects in ischemic primate and rat models, with up to about 84% of the CA1 hippocampal neurons being retained upon treatment relative to ischemic controls [66–69]. In these models, the neuroprotective effect is attributed to E64-c inhibiting cathepsin B and L, which is another cysteine protease. Thus, the fact that E64d inhibits a range of cysteine proteases may be advantageous in treating AD patients by providing neuroprotection.

Another mechanism by which E64d is likely to cause neuroprotection is by increasing brain sAβPPs. Increasing sAβPPs is generally beneficial to neurons since its administration to mice and rats enhances learning and memory [14, 15]. Thus, the increased brain sAβPPs caused by oral E64d demonstrated here or by icv administration as previously shown [40, 41] also likely confers neuroprotection.

The neuroprotective effects may explain why in old AβPP/PP mice E64d administration improved memory deficits more than could be attributed to brain Aβ lowering alone. In most AD patients who express AβPP containing the wt β-secretase site sequence, E64d may have multiple beneficial effects by both lowering toxic brain Aβ levels and by providing neuroprotection against that toxicity.

However, one report speculated that cathepsin B may be important for the extracellular degradation of brain Aβ in vivo and that inhibiting cathepsin B may increase brain Aβ and thereby promote memory loss and AD [64]. That hypothesis was based in large part on data showing that doubling brain cathepsin B activity by injecting a lent-cathepsin B vector into the brains of transgenic AβPP/Swe mice reduced brain plaque in the area receiving the injection. Moreover, our previous data showed that deleting the cathepsin B gene in transgenic mice expressing human AβPP/Swe reduces brain Aβ [36]. These data also are contrary to the hypothesis that inhibiting cathepsin B may increase brain Aβ because brain Aβ decreased upon completely eliminating brain cathepsin B activity. The data of this study and the knockout data can be explained by the fact that the animal models used in those experiments expressed AβPP containing wt β-secretase site sequence, which cathepsin B β-secretase activity cleaves to produce brain Aβ. Thus, chemically inhibiting cathepsin B activity or ablating the cathepsin B gene in such animals reduces brain Aβ production, making Aβ degradation unnecessary because...
brain Aβ need not be degraded if it is not produced. But even when cathepsin B inhibition does not reduce brain Aβ, cathepsin B inhibition probably does not significantly impact the degradation of brain Aβ. As discussed above, chronic E64 treatment of transgenic AβPPSwe mice had no effect on brain Aβ levels but did improve memory deficits [63]. If cathepsin B degradation of Aβ is significant in vivo, E64 treatment would be predicted to increase brain Aβ and worsen memory deficits because E64 is a potent cathepsin B inhibitor. But since that did not occur, these data too do not support the hypothesis. While it may be that inducing viral over expression of cathepsin B can reduce brain plaque under certain conditions, the in vivo inhibitor reported here do not support the conjecture that cathepsin B inhibition would so reduce brain Aβ degradation that brain Aβ levels would increase and memory would worsen. On the contrary, the data presented here show that inhibiting brain cathepsin B activity reduces brain Aβ and brain plaque and improves memory deficits.

Studies in human AD brains support a role for cathepsin B in the “disease situation.” Cathepsin B is colocalized with Aβ in amyloid plaques of AD brains [70–72] and age-related changes in cathepsin B are consistent with the late age of onset of AD [73, 74]. Proteomic analysis of AD and age-matched controls showed AD patients have significantly higher CSF cathepsin B precursor levels [75] and higher plasma cathepsin B levels [76]. These findings indicate a role for cathepsin B in AD.

Cathepsin B belongs to the C1 (papain) family of CA clan consisting of 11 human cysteine proteases [43, 77, 78]. The normal and pathological roles of cysteine cathepsins have been reviewed recently [79]. The cathepsins B, C, F, H, L, O, and Z are ubiquitously but differentially expressed in mammalian tissues, whereas cathepsin K is mainly found in osteoclasts and cathepsin S is in peripheral antigen-presenting cells [80]. Traditionally, cysteine cathepsins function as lysosomal proteases for protein degradation. However, they also have other intra- and extracellular functions, many of which have been revealed by gene-knockout studies (see review [78]). Cysteine proteases activate proenzymes and prohormones, mediate MHC-II-mediated antigen presentation, remodel bone, differentiate keratinocytes, regulate hair growth, and control apoptosis. They also participate in tumor metastasis, inflammatory rheumatoid arthritis, arthritis, atherosclerosis and periodontitis. Cathepsin B is a lysosomal protease but is also found in regulated secretory vesicles of chromaffin cells that produce Aβ peptides [35] and secretory vesicles of pancreatic beta-cells [81]. In brain, cathepsin B, as well as several cysteine cathepsins, show differences in neuroanatomical distribution [72, 82]. Cathepsin B deficient (non-transgenic) mice appear normal and cannot be distinguished from their wild-type counterparts [83] but have reduced trypsinogen activation in experimental pancreatitis, reduced tumor necrosis factor-α induced apoptosis, delayed mammary cancer and reduced tumor vascularization and growth [79].

Surprisingly, despite inhibiting this spectrum of proteases, E64d has been shown to be safe in man. E64d was originally developed by the Japanese Ministry of Health, which called it EST, in the 1980’s to treat muscular dystrophy and EST completed Phase III trials but did not become a drug due to a lack of sufficient efficacy for that disease [84]. Nonetheless, extensive pharmacokinetic, adsorption, metabolism, distribution, elimination, toxicology, and clinical data on EST were published in peer-reviewed Japanese journals. The details of that data cannot be fully discussed here, but in summary, they show that the compound has good oral bioavailability, a very wide therapeutic window [85–88], is safe for clinical use and was chronically administered to pediatric patients [89–92]. The low toxicity and safe clinical use may be due to the compound attenuating, but not eliminating, protease activity. Nonetheless, the safe use of E64d plus the oral efficacy demonstrated here in AD animal models gives E64d real potential as a candidate for subsequent AD clinical trials.

In conclusion, this study shows that oral administration of the cysteine protease inhibitor, E64d, to transgenic AβPPLOL mice improves the memory deficits, with reduction of brain Aβ peptides and amyloid plaque, that develop in this AD model and mimics that seen in AD patients. These effects are likely due, in large part, to E64d’s ability to reduce brain Aβ. In transgenic AβPPLOL mice and guinea pigs, oral E64d administration reduces brain Aβ40, Aβ42, CTFβ, and cathepsin B activity while increasing brain sAβPPβ and BACE1 activity, which is consistent with the compound reducing brain Aβ production by inhibiting cathepsin B, and not BACE1, β-secretase activity. Because E64d reduces brain Aβ in animal models having the wt β-secretase activity, it is likely to also do so in most AD patients, who also have that activity. Independent of its Aβ lowering effect, E64d has also been shown to be a potent neuroprotectant in AD and thus can be expected to provide that benefit in treating AD as well. Finally, the fact that the compound is orally efficacious in AD animal models and is clini-
cally safe to use makes E64d an excellent candidate for advancement to clinical AD trials.

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