Increasing Cu bioavailability inhibits Aβ oligomers and tau phosphorylation

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Edited by Peter J. Sadler, University of Warwick, Coventry, United Kingdom, and accepted by the Editorial Board November 21, 2008 (received for review September 11, 2008)

Cognitive decline in Alzheimer’s disease (AD) involves pathological accumulation of synaptotoxic amyloid-β (Aβ) oligomers and hyperphosphorylated tau. Because recent evidence indicates that glycogen synthase kinase 3β (GSK3β) activity regulates these neurotoxic pathways, we developed an AD therapeutic strategy to target GSK3β. The strategy involves the use of copper-bis(thiosemicarbazone) complexes to increase intracellular copper bioavailability and inhibit GSK3β through activation of an Akt signaling pathway. Our lead compound CuII(gtsm) significantly inhibited GSK3β in the brains of APP/PS1 transgenic AD model mice. CuII(gtsm) also decreased the abundance of Aβ trimers and phosphorylated tau, and restored performance of AD mice in the Y-maze test to levels expected for cognitively normal animals. Improvement in the Y-maze correlated directly with decreased Aβ and tau levels. This study demonstrates that increasing intracellular copper bioavailability can restore cognitive function by inhibiting the accumulation of neurotoxic Aβ trimers and phosphorylated tau.

Alzheimer’s disease | bioinorganic chemistry | glycogen synthase kinase | therapeutic | animal model

Alzheimer’s disease (AD) is a neurodegenerative disorder characterized clinically by impaired cognitive performance and pathologically by cerebral deposition of extracellular amyloid plaques and intracellular neurofibrillary tangles. Amyloid plaques in AD contain aggregated forms of the 39- to 43-aa amyloid-β peptide (Aβ) and Aβ is strongly implicated as a causative agent responsible for cognitive failure in AD. A diverse range of mechanisms for Aβ toxicity has been reported (1). Aβ is produced from the amyloid precursor protein (APP) (2–5) and Aβ aggregation pathway, primarily low-molecular-mass oligomers such as dimers and trimers, exhibit the greatest neurotoxicity (6–8). In addition to Aβ oligomers, aberrantly phosphorylated microtubule-associated protein tau is also associated with cognitive decline in AD (9). Intracellular neurofibrillary tangles in the AD brain contain hyperphosphorylated tau, and Aβ induced cognitive deficits characteristic of AD transgenic mice are attenuated by decreasing levels of endogenous tau (10).

It is now widely recognized that a truly effective therapeutic compound for treating AD needs to attenuate both the Aβ- and tau-mediated pathologies. Recent positive outcomes for PBT2 in clinical and preclinical trials are therefore pertinent. Lannfelt et al. (11) demonstrated in phase IIa clinical trials that PBT2 lowers plasma Aβ levels and attenuates cognitive decline, and Adlard et al. (12) have shown that PBT2 decreases interstitial Aβ and phosphorylated tau in the brains of AD model mice. PBT2 is a second-generation 8-OH quinoline, which, unlike its predecessor clioquinol, lacks iodine and was selected for clinical development because of its easier chemical synthesis, higher solubility, and increased blood–brain barrier permeability. Its clinical application was developed based on the potential to inhibit Aβ toxicity through modulation of Aβ-metal interactions (13–16), but the in vivo mechanism of action for PBT2 still requires elucidation. Cell culture studies have provided mechanism of action data to show that membrane-permeable compounds with the potential to form metal complexes, such as PBT2, can inhibit Aβ toxicity by increasing cellular metal ion bioavailability and activating neuroprotective cell signaling pathways (17–20). Central to these pathways is inhibition of glycogen synthase kinase 3β (GSK3β). Inhibition of this kinase in vitro increases expression of Aβ-degrading proteases (20), and GSK3β is a kinase capable of phosphorylating tau (21, 22). In this study, we tested the hypothesis that accumulation of Aβ and phosphorylated tau could be prevented in vivo by inhibiting GSK3β with an agent that increases intracellular bioavailability of copper. To do this we used the bis(thiosemicarbazone) compounds CuII(gtsm) and CuII(atsm) (Fig. 1). We were able to demonstrate that inhibition of GSK3β activity with CuII(gtsm), but not the control compound CuII(atsm), decreased abundance of Aβ trimers and phosphorylated tau in vivo. Importantly, treatment with CuII(gtsm) reversed cognitive deficits in APP/PS1 transgenic AD mice.

Results

CuII(gtsm) Inhibits GSK3β and Decreases Tau Phosphorylation in Vitro

SH-SY5Y cells were treated with CuII(gtsm) or CuII(atsm) (Fig. 1) or the vehicle control dimethyl sulfoxide (DMSO). These metal complexes were used because both compounds are cell permeable and deliver Cu into the cell. Importantly, once CuII(gtsm) is exposed to the intracellular reducing environment, CuII is released to CuI, causing it to dissociate from the ligand and to increase Cu bioavailability. The negative control CuII(atsm) does not reduce and dissociate under normal intracellular conditions, and therefore does not increase Cu bioavailability (18, 23–26). We found both metal complexes increased cellular Cu levels several hundredfold.
CuII(gtsm) modulates GSK3β signaling pathways and inhibits tau phosphorylation in SH-SYSY cells. (A) ICP-MS data showing CuII(gtsm) and CuII(atsm) both dramatically increase cellular levels of Cu in SH-SYSY cells. Cu content of DMSO treated cells was 34 nmol of Cu per milligram of protein. (B) Western blot analyses reveal that CuII(gtsm), but not CuII(atsm), induced GSK3β phosphorylation at serine-9 (ser9) via its upstream kinase, protein kinase B (Akt), and that phosphorylation of the associated extracellular signal-related kinase 1/2 (ERK1/2) was also increased (Fig. 2B). These data demonstrate that increased bioavailability of intracellular Cu rather than elevated cellular Cu per se is required to induce cellular pathways that lead to increased GSK3β phosphorylation at ser9 (i.e., GSK3β inhibition). We have shown previously that induction of these cellular pathways using metal complexes leads to decreased levels of Aβ in cell cultures because of increased expression of Aβ-degrading proteases (17, 18, 20).

To determine whether CuII(gtsm)-mediated inhibition of GSK3β in SH-SYSY cells inhibited activity of the kinase toward its substrate tau, we determined levels of tau phosphorylation by Western blot analysis. Tau phosphorylation at ser404 was decreased in CuII(gtsm)-treated cells by 64% (Fig. 2C, *P = 0.0002*), but unaltered in CuII(atsm)-treated cells (*P = 0.376*). Phosphorylation of tau at ser396 was also decreased by treating with CuII(gtsm) [supporting information (SI) Fig. S1]. The abundance of total tau was unchanged for all treatments (Fig. 2C).

CuII(gtsm) Rescues Aβ-Induced Inhibition of LTP. Aβ has strong inhibitory potential toward hippocampal long-term potentiation (LTP) (7, 27), and metal complex-mediated activation of neuroprotective pathways can decrease Aβ levels in vitro (17, 18, 20). Therefore, we determined whether CuII(gtsm) was able to rescue Aβ-induced inhibition of LTP. When included in the artificial cerebrospinal fluid (ACSF) at 2 μM, synthetic Aβ42 inhibited LTP in hippocampal slices from wild-type C57/bl6 mice by 28% (DMSO treatment fEPSP = 130.4%, Aβ treatment fEPSP = 94.5%, Fig. 3A, *P = 0.01*). Including 1 μM CuII(gtsm) in the ACSF prevented Aβ toxicity and returned LTP to levels that were not significantly different to controls (Aβ + CuII(gtsm) treatment fEPSP = 121.4%, *P = 0.308*). CuII(gtsm) alone had no effect on LTP (CuII(gtsm) treatment fEPSP = 127.6%, Fig. 3A, *P = 0.4557*).

CuII(gtsm) Improves Cognitive Performance of AD Mice in Y-Maze. Based on our in vitro data showing that CuII(gtsm) activates potentially neuroprotective cell signaling pathways (Fig. 2) and prevents Aβ-mediated inhibition of LTP (Fig. 3A) we treated APP/PS1 transgenic AD mice with CuII(gtsm) at 10 mg/kg of body weight (*n = 15*) or vehicle control (*n = 14*) by daily gavage. Sham- and CuII(gtsm)-treated AD mice were analyzed for cognitive performance by using a 3-arm radial maze (Y-maze) for spatial working memory and learning (28). Sham-treated AD mice exhibited poor recognition of the Y-maze novel arm (33% entry to novel arm, Fig. 3B), demonstrating impaired cognitive performance consistent with previous studies (29, 30). By contrast, the CuII(gtsm)-treated mice showed significantly improved cognitive performance with 48% entry to the novel arm (Fig. 3B, *P = 0.0002*). These Y-maze data indicate CuII(gtsm) restored cognitive performance of the AD mice to levels expected for healthy, cognitively normal mice. In contrast to CuII(gtsm), and consistent with our hypothesis that the therapeutic effects of CuII(gtsm) are based on the compound’s capacity to increase intracellular Cu bioavailability, treating the AD mice with CuII(atsm) did not affect cognitive performance in the Y-maze test, even at an elevated dose of 100 mg/kg of body weight (Fig. 3B, *P = 0.280*).
CuII(gtsm) Inhibits GSK3β in Vivo. Biochemical examination of brain tissue from sham- and CuII(gtsm)-treated AD mice revealed strong consistency with data from experiments using SH-SY5Y cells. CuII(gtsm) induced increased ser9 phosphorylation of GSK3β via its upstream kinase Akt, induced phosphorylation of ERK1/2 (Fig. 4A, P < 0.018, 0.018, and 0.049, respectively). Concomitant with the CuII(gtsm)-mediated inhibition of GSK3β was a decrease in detectable levels of phosphorylated tau (Fig. 4B, P < 0.030). Abundance of total tau was not altered in the CuII(gtsm)-treated mice (Fig. 4B, P = 0.2248).

In addition to GSK3β, cyclin-dependent kinase 5 (cdk5) with its regulatory cofactor P35 can also phosphorylate tau (31). We therefore examined whether decreased phosphorylation of tau in AD mice involved decreased cdk5/P35. Neither cdk5 nor P35 levels were altered in the brains of AD mice after treatment with CuII(gtsm) (Fig. 4B, P = 0.49 and 0.47, respectively), indicating that decreased tau phosphorylation in vivo is more likely the result of GSK3β inhibition.

CuII(gtsm) Decreases Brain Aβ Trimer Levels in AD Mice. Because increasing cellular metal ion bioavailability prevents Aβ accumulation in vitro through inhibition of GSK3β (20), we examined whether CuII(gtsm)-mediated inhibition of GSK3β in vivo (Fig. 4A) affected Aβ levels in the brains of AD mice. ELISA analyses demonstrated that Aβ in the PBS soluble fraction of the brain from CuII(gtsm)-treated mice was unaltered, but that Aβ in the PBS-insoluble fraction was decreased by 24% (Fig. 5A, P = 0.033). More

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**Fig. 4.** CuII(gtsm) modulates GSK3β signaling pathways and inhibits tau phosphorylation in the brains of AD mice. (A) Western blot and densitometry analyses showing CuII(gtsm) increases phosphorylation of GSK3β, Akt and ERK1/2 in brains of AD mice. ERK1/2 could not be detected in PBS extracts of mouse brains and data for ERK1/2 was therefore obtained by using cell lysis buffer as described in the methods. Data for all other proteins was obtained by using PBS extracts. (B) Western blot and densitometry analysis of brain samples shows tau phosphorylation at ser404 is decreased after treatment with CuII(gtsm) in the absence of effects on total tau, cdk5, or P35. All Western blot densitometry data are normalized for the loading control GAPDH. P values shown indicate that treatments induced significant effects compared with the sham control. Densitometry data in A and B are mean values ± SEM.

**Fig. 5.** CuII(gtsm) inhibits accumulation of PBS insoluble Aβ trimers in the brains of AD mice. (A) ELISA data showing that overall levels of PBS insoluble Aβ, but not PBS soluble Aβ, are decreased in the brains of CuII(gtsm)-treated AD mice. (B) SELDI-TOF MS analysis showing Aβ trimers are decreased in the PBS-insoluble fraction of CuII(gtsm)-treated mice, but monomer levels are unaffected. All SELDI-TOF MS data are normalized for total brain protein and are expressed as Aβ peak intensity per milligram of protein. (C) Correlation analysis showing cognitive performance in Y-maze testing of AD mice improves with decreasing Aβ trimer levels in the PBS-insoluble fraction of the brain. P values shown indicate that treatments induced significant effects compared with the sham control. Data in A and Aβ peak intensity data in B are mean values ± SEM.
rigorous examination of PBS-insoluble brain samples using surface enhanced laser desorption ionization time-of-flight (SELDI-TOF) mass spectrometry (MS) was performed to establish whether the ELISA data were due to an overall decrease in total brain Aβ or due to decreased abundance of specific Aβ species. The SELDI-TOF MS data show that although monomeric forms of Aβ40, Aβ42, and Aβ43 resolved discretely with accurate molecular masses, their levels were not altered by CuII(gtsm) (P = 0.43, 0.28 and 0.47, respectively). By contrast, a readily detectable Aβ species with a molecular mass of 13,801 Da, equivalent to an Aβ trimer, was significantly decreased in CuII(gtsm)-treated mice (Fig. S5B, P = 0.02). A mass spectrometry technique has not previously supported the existence of an Aβ trimer in vivo. The second peak shown in Fig. 5B at 14,018 Da is consistent with the Aβ trimer containing a sinapinic acid adduct; sinapinic acid is part of the SELDI-TOF matrix. Other Aβ oligomer species such as dimers, tetramers, and Aβ56* were not detected when using SELDI-TOF MS. Similarly, trimers containing Aβ40 were also not detected. Examination via immunohistochemistry revealed treatment with CuII(gtsm) did not affect Aβ plaque deposition in the brains of AD mice (Fig. S2, P = 0.39). Although numerous studies demonstrate the neurotoxic potential of Aβ trimers (32) there is little in vivo evidence to show specific reduction of their abundance in the brain correlates directly with improved cognitive performance. To address this, we performed correlation analyses of brain Aβ trimer levels and performance of mice in the Y-maze test. These analyses revealed that mice with lower levels of trimeric Aβ in the brain performed better in the Y-maze test (R² = 0.18, P = 0.034) (Fig. 5C). These data, therefore, provide in vivo evidence for the role of Aβ trimers in impaired cognitive performance in AD mice and indicate that therapeutic inhibition of Aβ trimer accumulation by increasing cellular metal bioavailability can rescue impaired cognition.

Discussion
Therapeutic inhibition of Aβ accumulation and tau phosphorylation in the brain is a primary target in AD research because aberrant Aβ accumulation and tau phosphorylation are now widely recognized as the main contributors to the disease-associated cognitive decline. Recently a therapeutic compound, PBT2, successfully completed phase IIa clinical trials, and was also shown to decrease Aβ levels in cerebrospinal fluid and attenuate cognitive decline in AD patients (11). Importantly, a prescience analysis of PBT2 demonstrated that the compound decreased interstitial Aβ as well as phosphorylated tau in the brains of AD model mice, leading to their improved cognitive performance (12). Based on previous cell culture studies (18, 20) it was speculated that the mechanism of action for PBT2 involves sequestration of Cu and Zn from extracellular Aβ aggregates and subsequent PBT2-mediated delivery of the metal ions intracellularly (12). Data to show the mechanism of action through which a PBT2-mediated increase in cellular metal ions can inhibit accumulation of Aβ and phosphorylated tau are yet to be presented. We used the tbc metal complexes CuII(gtsm) and CuII(atsm) because these complexes can cross the blood–brain barrier (33), and as described above, they exhibit remarkably different chemical properties that are determined by the substituents attached to the diimine backbone of the ligand and enable controlled regulation of cellular metal ion bioavailability.

The possibility that metal complexes with the capacity to increase intracellular metal bioavailability can activate neuroprotective cell signaling pathways pertinent to AD is based on cell culture studies using a range of metal complexes. In particular, treating cell cultures with specific Cu complexes can activate a cell signaling pathway that leads to increased expression of Aβ-degrading proteases and, therefore, decreased Aβ levels (18, 20). The signaling pathway involves phosphorylation of ERK1/2, JNK, and GSK3β, and metal complex-mediated phosphorylation of PI3K/Akt because of activation of the epidermal growth factor receptor is an important upstream event (17–20, 34). In vivo support for targeting Aβ and tau in the AD by modulating metal bioavailability comes from a small number of studies. When administered to AD model mice that develop cognitive deficits, clioquinol and PBT2 both decreased the brain Aβ burden and improved cognitive performance (12, 14). Furthermore, PBT2 and clioquinol both provided promising cognitive outcomes in clinical trials (11, 16). In a related study, AD mice were treated for 7 months with the metal-ligating compound pyrrolidine dithiocarbamate (PDTC), and the treatment was shown to induce phosphorylation of Akt and GSK3β, decrease tau phosphorylation, and improve cognitive performance of the mice (35). Surprisingly, however, PDTC did not change Aβ levels as determined by ELISA. Common to the beneficial effects of clioquinol, PBT2, and PDTC is that the compounds were all administered without metal ions already bound to the ligand. Therefore, if their capacity to inhibit tau phosphorylation and Aβ accumulation involved the formation of a metal complex, the activity relied on the compounds interacting with metal ions adventitiously in vivo. Such strategies therefore do not rule out the possibility of nonspecific sequestration of metal ions. We believe this is central to our observed effects for CuII(gtsm) on both Aβ trimers (Fig. 5) and phosphorylated tau (Fig. 4) in the brains of AD mice, because our cell culture work shows metal-free compounds that rely on adventitious binding of metal ions are considerably less potent than metal complexes where the compound is already loaded with Cu (18, 20).

Aβ potently inhibits hippocampal LTP (32), an in vitro assay used to determine the cellular mechanisms of memory and learning. Studies on Aβ-mediated inhibition of LTP indicate Aβ trimers as the main toxic Aβ species (7). Here, we have shown that CuII(gtsm) attenuates Aβ-mediated inhibition of LTP. More importantly, we have shown that treatment with CuII(gtsm) decreases the abundance of Aβ trimers in the brains of AD mice and that decreased Aβ trimer levels correlate directly with improved cognitive performance in the Y-maze test. The significance of the data shown in Fig. 5C is that mass spectrometry techniques have not previously been used to verify that apparent Aβ oligomer species seen on a Western blot have a molecular mass consistent with the Aβ oligomer. Our SELDI-TOF MS data therefore differentiates between Aβ trimers and APP C-terminal fragments that have the potential to resolve on SDS/PAGE gels in the region expected for Aβ oligomers. Previous studies have reported an apparent correlation between decreased abundance of the Aβ species termed Aβ56* and improved cognition based on Western blot methods (36), but we propose that more rigorous Aβ identification techniques are essential before performing correlation analyses (see Table S1).

CuII(gtsm)-mediated inhibition of GSK3β is central to our mechanism of action pathway (Fig. 6) and our observed effects for CuII(gtsm) on Aβ trimers, tau phosphorylation, and cognitive improvement. Originally described for its capacity to phosphorylate glycogen synthase kinase (37), GSK3β is somewhat unique because its capacity to phosphorylate a substrate is inhibited when the kinase itself is phosphorylated at the regulatory site ser9 (38). Phosphorylation of GSK3β therefore inhibits its kinase activity, and the inhibition of GSK3β gained recognition as a potential therapeutic target in AD once tau was identified as a GSK3β substrate (21, 22). Hyperphosphorylated tau is abundant within neurofibrillary tangles of the AD brain, indicating a loss of tau kinase/phosphatase homeostasis. Promoting a substrate-specific increase in phosphatase activity is practically impossible because of relatively poor phosphatase-substrate specificity; therefore targeting tau kinases is more
deficits are only observed in AD mice with elevated Aβ for tau in cognitive decline, but also demonstrates that cognitive deficits of AD and that GSK3β activity is an important intermediate. In this study, we have shown that inhibition of GSK3β in vivo by therapeutically increasing Cu bioavailability decreases Aβ trimer levels and tau phosphorylation in AD mice. Importantly, the CuII(gtsm)-mediated decrease in Aβ trimer levels correlated directly with improved cognitive performance in the Y-maze test, and therefore provides strong support for increasing cellular metal ion bioavailability as a valid therapeutic strategy for treating AD.

Materials and Methods
The following is a brief summary. Full details are provided in SI Text.

CuII(gtsm) and CuII(atsm). The btsc compounds CuII(gtsm) and CuII(atsm) were synthesized as described previously (42, 43).

Cell Culture Experiments. SH-SY5Y cells at ~80% confluency were treated with 25 μM CuII(gtsm), 25 μM CuII(atsm) or the vehicle control DMSO in serum-free media for 5 h. Cells collected by centrifugation were resuspended in 50 μL of Cytobuster Protein Extraction reagent (Novagen) supplemented with protease and phosphatase inhibitors. Protein content of the cell extracts was determined by using a protein content determination kit (Pierce).

Inductively Coupled Plasma Mass-Spectrometry Determination of Metals Ions. After treating and collecting cells as described above, pelleted cells were rinsed with PBS then analyzed for Cu, Fe, Zn, and Mn by using inductively coupled plasma mass-spectrometry (ICP-MS) as described previously (20).

AD Mice and Treatments. The AD mice used in this study expressed mutant human APP (K670N, M671L) and mutant human PS1 (E9) (44). Treatments were administered daily by oral gavage and began when the mice were 5–6 months old. Acid stability studies have shown dissociation of Cu–btsc complexes is unlikely to occur in the gut (45). Mice were tested for cognitive performance by using the Y-maze test, and the brains collected after culling were used for biochemical and immunohistochemical analyses.

Y-Maze Test for Spatial Memory and Learning. Mice were subjected to a 2-trial Y-maze test, with all testing performed during the light phase of the circadian cycle. Three identical arms of the maze were randomly designated start arm, novel arm, and other arm. The Y-maze tests were performed after 6 weeks of treatment and consisted of 2 trials separated by a 1-h intertrial interval to assess spatial recognition memory. The first trial (training) was for 10 min, and the mice were allowed to explore only 2 arms (starting arm and other arm). For the second trial (retention), mice were placed back in the maze in the same starting arm, and allowed to explore for 5 min with free access to all 3 arms. By using a ceiling-mounted CCD camera, all trials were analyzed for the number of entries the mice made into each arm. Data are expressed as the percentage of novel arm entries made during the 5-min second trial.

Preparation of Mouse Brain Samples for Biochemical Analyses. Mice were anesthetized, dissected to expose the chest cavity, and then perfused with ~10 mL of Dulbecco’s PBS (Invitrogen) supplemented with heparin, butylated hydroxytolulene, and phosphatase inhibitors. Brains were removed and stored frozen. Right hemispheres were homogenized by sonication in 1 mL of 10 mM of Dulbecco’s PBS (Invitrogen) supplemented with phosphatase inhibitors. Left hemispheres were homogenized by sonication in 1 mL of PBS supplemented with phosphatase inhibitors. Left hemispheres were homogenized mechanicially in Protein Extraction Buffer as per SH-SY5Y samples. Soluble and insoluble fractions were separated by centrifugation and protein content determined by using a protein assay kit (Pierce).

Mass Spectrometry Analysis of Aβ in Brain Tissues. Analysis of Aβ from brain tissues was performed with surface enhanced laser desorption ionisation time-of-flight (SELDI-TOF) mass spectrometry (MS) using the ProteinChip system (Ciphergen Biosystems). This method enables direct mass-spectrometry identification of individual proteins after immunoaffinity capture purification of the proteins of interest from tissue samples. PBS-soluble and -insoluble extracts of AD mouse brains were subjected to antibody capture using the WO2 antibody for human Aβ (epitope = Aβ residues 5–8), and the mass of each Aβ species was determined by using the ProteinChip reader. Data generated by the ProteinChip software was analyzed by using Biomarker Patterns Software for differential peak intensities of each peptide. All SELDI-TOF MS brain Aβ data are expressed as log-normalized Aβ peak intensity per milligram of brain protein.

Feasible. Here, we have shown that therapeutic inhibition of GSK3β by treating with CuII(gtsm) correlated with decreased tau phosphorylation in vitro and in vivo. Considered together with our observations for decreased Aβ trimer levels, our data for CuII(gtsm)-mediated decreases in tau phosphorylation raise direct questions regarding whether Aβ trimer levels or aberrantly phosphorylated tau contributes most significantly to effects on cognition. There is conjecture in the literature on this point but growing evidence indicates that although tau is an essential mediator of cognitive decline, Aβ accumulation is a critical upstream event (39, 40). A pertinent study to support this shows that cognitive deficits in an AD mouse model are attenuated by crossing AD mice overexpressing APP with tau−/− mice: AD/tau−/− mice exhibit unaltered Aβ levels compared with AD/tau+/+ mice (10). This work demonstrates a critical role for tau in cognitive decline, but also demonstrates that cognitive deficits are only observed in AD mice with elevated Aβ levels. In a more recent study, it was shown that exposure to Aβ induces tau phosphorylation by promoting GSK3β activity (41). These studies therefore demonstrate an intimate relationship between Aβ accumulation and tau phosphorylation in the cognitive

Fig. 6. Schematic overview of proposed mechanism by which CuII(gtsm) decreases Aβ oligomer burden (trimers) and tau phosphorylation in the brains of AD mice. Cell-permeable CuII(gtsm) dissociates in the presence of intracellular reductants to release CuI and increase Cu bioavailability, thereby activating cell signaling pathways that lead to decreased Aβ trimer levels, decreased tau phosphorylation, and improved performance of AD mice in the Y-maze test for spatial learning and memory.


