Novel Selective Calpain 1 Inhibitors as Potential Therapeutics in Alzheimer’s Disease

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Abstract. Alzheimer’s disease, one of the most important brain pathologies associated with neurodegenerative processes, is related to overactivation of calpain-mediated proteolysis. Previous data showed a compelling efficacy of calpain inhibition against abnormal synaptic plasticity and memory produced by the excess of amyloid-β, a distinctive marker of the disease. Moreover, a beneficial effect of calpain inhibitors in Alzheimer’s disease is predictable by the occurrence of calpain hyperactivation leading to impairment of memory-related pathways following abnormal calcium influxes that might ensue independently of amyloid-β elevation. However, molecules currently available as effective calpain inhibitors lack adequate selectivity. This work is aimed at characterizing the efficacy of a novel class of epoxide-based inhibitors, synthesized to display improved selectivity and potency towards calpain 1 compared to the prototype epoxide-based generic calpain inhibitor E64. Both functional and preliminary toxicological investigations proved the efficacy, potency, and safety of the novel and selective calpain inhibitors NYC438 and NYC488 as possible therapeutics against the disease.

Keywords: Alzheimer’s disease, amyloid-β, calpain, learning, long-term potentiation, memory

INTRODUCTION

Calcium-activated neutral cysteine proteases (calpains) are a variegated cluster of calcium-dependent proteases, able to modify the function of several target proteins by partial truncation. This limited non-digestive proteolysis is a particular form of post-translational modification that changes physiological activity and translocation of the target proteins [1], including calpains themselves [2]. Calpains regulate through proteolysis several cellular functions, including cytoskeleton assembly and disassembly.

In the central nervous system (CNS), where calpain I and calpain II are the main calpain isoforms,
their activation is to synaptic plasticity and memory as well as to neurodegeneration [3, 4]. Events that have been proposed to participate in synaptic plasticity and memory, including cytoskeletal regulation, AMPA receptor trafficking, actin polymerization, and regulation of local protein synthesis are regulated by calpains [5] through a plethora of protein targets like CaMKIIα, protein kinase C and PP1/alpha/calcineurin [6–10] and transcription factors such as the CAMP response element-binding protein (CREB) [11–16]. Abnormal calcium influxes intensify calpain activity at supra-physiological levels that are evident in a number of neurological disorders (i.e., Alzheimer’s disease (AD) [4, 17, 18]), generating a variety of detrimental effects in pathways related to synaptic plasticity and memory, including the decrease in CREB phosphorylation and activation [19, 20], and accompanied by synaptic dysfunction, which is a robust predictor of cognitive impairment in AD [21, 22].

Despite the intense effort in the field of AD directed to study and to provide a neuropathological substrate for an effective therapy toward AD, the current therapeutic arsenal (such as galantamine, rivastigmine, donepezil, and memantine) is at best symptomatic and provides solely temporary relief without a real causative breakthrough [23, 24]. There are no FDA-approved drugs that can delay or halt the progression of the disease. Given the perspective epidemiological AD and other neurodegenerative diseases [25], it is rather urgent to develop a proficient line of therapeutics with high translational potential and optimal therapeutic index.

The vast majority of failing clinical efforts deal with the problem of amyloidogenic protein deposition, because proteinaceous aggregates consisting of deposition of extracellular amyloid plaques and intracellular neurofibrillary tangles are the major histopathological hallmarks of the disease. Our approach, instead, focuses on preserving synaptic functionality. This is justified by ample evidence suggesting that AD starts as a synaptic disorder [26]. It is likely that the very fine and variable amnesic symptoms occurring at the beginning of the disease in the absence of any other clinical signs of brain injury, are caused by discrete changes in synaptic function, produced at least in part, by amyloid-β (Aβ) species (e.g., Aβ40 and Aβ42) [27–30], peptides derived from processing of amyloid-β protein precursor (AβPP). Previously, we validated the inhibition of calpains as a therapeutic target against their overactivation in AD toward the recovery of synaptic dysfunctions induced by Aβ [19, 31]. These findings led to an effort aimed to discovering novel calpain inhibitors that might be utilized against AD. Here we report findings from a phenotypic screening of three generations of peptidomimetic epoxide warhead containing molecules that have been previously proved to be unreactive toward reaction with free thiols while displaying irreversible active site calpain 1 inhibition with sub-micromolar potency [32]. We designed our drug screening for calpain inhibitors using a phenotypical modality combined with medicinal chemistry refined through target-based computational approach [32, 33], focusing on the capability of our candidate molecules to protect from the detrimental effect of oligomerized Aβ42 on hippocampal long-term potentiation (LTP), a type of synaptic plasticity thought to underlie learning and memory. Following this screening, the last generation of leads was further tested for pharmacokinetic and toxicological features, and then for the recovery of cognitive impairments in a mouse model of amyloid deposition, the AβPP/PS1 mouse [32].

MATERIAL AND METHODS

Animals

All experiments were performed with the approval of the Columbia University Animal Care and Use Committee in accordance with the guidelines for the humane treatment of animals (protocol #AC-AAAB9126). Hemizygous transgenic (HuAPP985SWE) 2576 mice expressing mutant human AβPP (K670N, M671L) [34] were crossed with hemizygous PS1 mice that express mutant human PS1 (M146V; Line 6.2) [35]. The offspring, double-transgenic mice overexpressing AβPP/PS1, were compared with their wild type (WT) littermates so that age and background strain were comparable. To identify the genotype of the animals, we used DNA extracted from tail tissue as previously described [35, 36]. For pharmacokinetic testing, we used instead ICR mice.

Aβ peptide oligomerization

Recombinant Human Aβ42 peptide (American Peptides) was oligomerized as previously described [37]. Briefly, crude lyophilized Aβ peptide was re-suspended in cold 1:1.1,3,3,3-hexafluoro-2-propanol (HFIP, Sigma) and aliquoted in polypropylene vials. After 24 h, the HFIP solution was allowed to evaporate in a fume hood until a thin film of monomeric peptide is formed on the bottom of the vials. Peptide films were dried under gentle vacuum and stored in sealed vials at –20 ºC. Prior to use, anhydrous DMSO (Sigma) was
added to obtain a pure monomeric Aβ/DMSO solution and then sonicated for 10 min [37]. Low-order oligomer-enriched Aβ42 was obtained by incubating an aliquot of monomeric Aβ/DMSO solution in sterile artificial cerebrospinal fluid (ACSF) phosphate buffer at 4°C overnight. Oligomerized Aβ peptide was then further diluted up to 200 nM concentration with vehicle right before the experiments.

### Drug administration

Et46 from Sigma-Aldrich and all candidate compounds were solubilized in 100 μl Tween-80 + 100 μl dimethylsulfoxide (DMSO) and then diluted with vehicle solution to the appropriate experimental concentration for in vitro experiments or experimental dose for in vivo experiments. Oligomerized Aβ42 was either administered in vitro alone or co-administered in vitro with Et46 or one of the candidate compounds.

### Pharmacokinetic assessment

A total of 8-10 male ICR mice were divided into 3 dosing groups (18 mice in each group), which were administrated by intraperitoneal route at the same equimolar dose (NYC215, 7.57 mg/kg; NYC438, 7.86 mg/kg; NYC488, 7.83 mg/kg). Test article solutions were prepared by dissolving in 4% DMSO/4% Tween 80/92% deionized water to yield final concentrations as showed in the above table. Dose volume for each test animal was determined based on the most recent body weight. Blood (approximately 250 μl) was collected via retro-orbital puncture into tubes containing sodium heparin anticoagulant at 7.5, 15, 30, 60, 120, and 240 min post-dosing. Mice were sacrificed by cervical dislocation after blood harvest. The plasma were separated via centrifugation (11,000 rpm, 5 min) and stored in –80°C before analysis. Frozen unidentified plasma samples were thawed at room temperature and vortexed thoroughly. With a pipette, 25 μl of plasma was transferred into a 1.5 ml Eppendorf tube. To each sample, 25 μl of methanol/methanol-water (1:1, v/v) and 25 μl of internal standard (IS) (100 ng/ml NYC488 for NYC215, 50 ng/ml YFP for NYC438, and no IS was used for NYC488) were added, followed by the addition of 100 μl acetonitrile. The sample mixture was vortexed for approximately 1 min. After centrifugation at 11,000 rpm for 5 min, the upper layer was vaporized under nitrogen stream. The residue was dissolved with mobile phase and 20-μl aliquot was injected onto the LC/MS/MS system for analysis. Calibration standards were prepared by spiking 25 μl of the analyte standard solutions into 25 μl of heparinized blank mice plasma. The nominal standard concentrations in plasma were 3.0, 10.0, 30.0, 100, 300, 1,000, 3,000, and 10,000 ng/ml for each analyte. Quantification was achieved by the internal standard method using peak area ratios of the analyte.
to IS in plasma for NYC215 and NYC438. For NYC488, external standard method was used. Concentrations were calculated using a weighted least-squares linear regression (W = 1/x2). The assay was performed using an LC/MS/MS system consisting of the following components: HPLC system, GI319A vacuum degasser, GI311A quaternary pump, GI316A column oven (Agilent, Waldbronn, Germany) and NANOSPACE SI-2 HTS autosampler Z 3133 (Shiseido, Tokyo, Japan); MS/MS system, API 4000 triple quadrupole mass spectrometer, equipped with a TurboIonSpray (ESI) Interface (Applied Biosystems, Concord, Ontario, Canada). For the NYC215 and NYC438, we used a Capcell C18 column (100 mm x 4.6 mm I.D., 5 μm, Shiseido, Japan) while for the NYC488 we used a Synergi 4 μm Hydro-RP 80A (150 mm x 4.6 mm I.D., Phenomenex, Torrance, CA, USA). Mobile phase was acetonitrile versus 0.2% formic acid in 5 mM ammonium acetate at different mix depending on the candidate compound. The major pharmacokinetic parameters were calculated by non-compartmental analysis using WinNonlin 5.3 (Pharsight USA).

**Spectrin western blotting**

Hippocampal lysates for immunoblotting were prepared as previously described [19] with slight modifications. Hippocampal tissue was homogenized in lysis buffer (62.5 mM Tris-HCl pH 6.8, 3% LDS, 1 mM DTT) and incubated at 4°C for 10 min, then sonicated before centrifugation at 20,000 rpm for 5 min. Whole cell extracts were electrophoresed on 3–8% gradient Tris-Acetate PAGE gel (Invitrogen) and then transferred to PVDF membrane. The membrane was blocked with 5% non-fat milk in TBS (50 mM Tris-HCl, 150 mM NaCl, pH 7.4) for 1 h, then incubated overnight at 4°C with spectrin antibody (1:1,000 concentration for immunoblotting). Spectrin antibody was from Millipore, β-III-Tubulin antibody was purchased from Promega.

**Histologic analysis**

Mouse organ (liver, heart, muscle, stomach, kidney, brain) samples collected at the end of the chronic treatment with either vehicle or candidate compounds were fixed in 10% buffered paraformaldehyde, processed through conventional histological techniques, and stained with hematoxylin and eosin. Microscopy was performed using an optical microscope (Olympus BX51) equipped with a camera (Olympus Q-Color 5), and the images were recorded in a computer using the Image Pro-Express software.

**Behavioral assessment**

A) Associative contextual memory

Associative memory was probed through fear conditioning in either vehicle or transgenic AβPP/PS1 mice, according to previously proposed method [39]. Our conditioning chamber was located inside a sound-attenuating box (72 x 54 x 48 cm). A clear Plexiglas window (2 x 12 x 20 cm) allowed the experimenter to film the mouse performance with a camera placed on a tripod and connected to FreezeFrame software (MDS Associates Inc.). To provide background white noise (72 dB), a single computer fan was installed in one of the sides of the sound-attenuating chamber. The conditioning chamber (33 x 20 x 22 cm) was made of transparent Plexiglas on two sides and metal on the other two. One of the metal sides had a speaker and the other had a 24 V light. The chamber had a 36-bar insulated shock grid floor. The floor was removable and after each use we cleaned it with 75% ethanol and then with water. Only one animal at a time was present in the experimentation room. The other mice remained in their home cages. During the contextual conditioning experiment, mice were placed in the conditioning chamber for 2 min. In the last 2 s of the 2 min, mice were given a foot shock of 0.50 mA for 2 s through the bars of the floor, and left in the conditioning chamber for another 30 s before being placed back in their home cages. “Freezing” behavior, defined as the absence of all movements except for that necessitated by breathing, was assigned scores using FreezeView software (MDS Associates Inc.). For evaluation of contextual fear learning, freezing at 24 h post-training was measured for 5 consecutive minutes in the chamber in which the mice were trained. Twenty-four hours after the contextual testing, cued fear conditioning was evaluated by placing the mice in a novel context (triangular cage with a smooth flat floor) for 2 min (pre-CS test), after which they were exposed to the CS for 3 min (CS test), and freezing was measured. In a separate set of experiments, we tested whether the four different experimental groups of mice had similar exploratory behavior and anxiety by carrying out the open field test. Animals were positioned in an open arena with a floor that was divided into compartments. The internal dimensions of the arena were 72 x 72 x 33 cm. An area measuring 36 x 36 cm in the center of the open field was defined as the “central compartment”. Behavioral scoring was evaluated by the percentage of time spent in the center compartment and the number of entries into the center compartment. No differences were found among the four groups of mice (data not shown).
Statistical analysis

For all experiments, mice were coded by “blind” investigators with respect to treatment and genotype. Data are expressed as mean ± SEM. Statistical analysis was performed with one-way ANOVA (for fear conditioning experiments), two-way ANOVA with repeated measures (for LTP and 2 day RAWM experiments) and Student’s t test (pairwise comparisons). The level of significance was set for p < 0.05.

RESULTS

Phenotypic screening of novel calpain 1 inhibitors using Aβ-induced reduction of LTP

For the present studies, we synthesized three generations of calpain inhibitors by modifying mostly the P2 and P3 domains of the inhibitor E64 backbone (Fig. 1). The P2 recognition group is important for the selectivity towards calpain 1 versus other proteases such as cathepsin B [32]. E64 was used as our benchmark compound for assessing assay in synaptic plasticity. As a fundamental requirement, the screening of candidate compounds aimed to verify whether the new derivatives possessed at least equal capabilities as E64 to rescue the Aβ-induced defect in LTP of hippocampal slices [19].

Acute exposure to 200 nM Aβ42 blocks LTP induction [29] (Fig. 2A). A dose-response curve for E64, co-applied with oligomerized Aβ42 (200 nM) prior to the induction of LTP by tetanic stimulation showed that E64 restored LTP to the vehicle-treated level. The EC50 for E64 was 650 nM (Fig. 2B). Therefore, we evaluated all candidate lead compounds at a concentration corresponding to the EC50 for E64.

Fig. 1. Chemical structure of the prototype epoxide compound E64. The structure of E64 can be functionally divided into three main domains: the epoxide warhead that interacts with an enzymatic pocket of calpains producing the protease inhibition, a peptidomimetic leucin domain P2 that it is important to modulate the selectivity of any derivative compound for calpains, and finally a P3 compound cap that is useful for drugability development and for improving the pharmacological potency.

B) Spatial memory

Spatial memory was assessed through the 2 day RAWM as previously described [40, 41]. Transgenic AβPP/PS1 mice were trained in fifteen daily sessions to identify a platform location by alternating between a visible and a hidden platform in the goal arm. The final three trials on day 1 and all fifteen trials on day 2 used a hidden escape platform to probe the ability of the mouse to find the goal arm location. The reaching of the learning criterion (max 1 arm error average in three consecutive trials) was obtained by WT vehicle-treated animals. The visible platform testing was observed among the different groups of mice indicating that visual, motor, and motivation skills were not affected by the experimental procedure (data not shown). Higher the scoring, less efficacy is associated to the experimental drug (by definition the vehicle-treated AβPP/PS1 display the worst score in criterion).

Amyloid-β assessment

Aβ content was assessed both in mouse hippocampi and plasma collected at the end of the chronic treatments, as described [42]. Hippocampi were homogenized in 880 μl of tissue lysate buffer (20 mM Tris-HCl (pH 7.4), 1 mM ethylenediaminetetraacetic acid, 1 mM ethyleneglycoltetraacetic acid, 250 mM sucrose) supplemented with 3X protease inhibitors (Roche). Blood samples were instead collected in EDTA-treated tubes, centrifuged to obtain the plasmatic fraction, and then mixed with an aliquot of the lysate buffer to maintain the matrix. ELISA assay (EZBRAIN-SET, Millipore, USA) was performed according to the manufacturer’s protocol on a Costar-like 96-well plates were incubated overnight at 4°C with capture antibody (in 0.1 M sodium bicarbonate, pH 8.2) at a dilution of 4 μg/ml. Upon blocking, plate wells were incubated overnight with 30 μl of brain lysate and then washed with PBS and incubated with the antibody at a 1 μg/ml. Values were read at 620 nM wavelength 30 min after adding 100 μl of colorimetric buffer. The signal was normalized to the protein concentration for each sample.

For all experiments, mice were coded by “blind” investigators with respect to treatment and genotype. Data are expressed as mean ± SEM. Statistical analysis was performed with one-way ANOVA (for fear conditioning experiments), two-way ANOVA with repeated measures (for LTP and 2 day RAWM experiments) and Student’s t test (pairwise comparisons). The level of significance was set for p < 0.05.
Fig. 2. Electrophysiological screening of three generations of calpain inhibitors through evaluation of their ability to protect against the Aβ42-induced LTP impairment. A) Example of effect produced by the co-administration of E64, the prototype epoxide compound with calpain inhibitory activity, together with Aβ42 (200 nM, 20 min). E64 rescues the detrimental effect induced by Aβ42 onto LTP, recovering the potentiation values up to the levels observed upon perfusion with vehicle alone. B) Dose-response curve of LTP levels observed upon co-administration of different concentrations of E64 with Aβ42 (200 nM, 20 min). The ED50 for E64 is 650 nM. Boxes represent the S.E.M range. C) Phenotypical screening of three generations of new E64 derivatives as calpain inhibitors based on the ability to rescue the detrimental effect of Aβ42 (200 nM, 20 min) onto LTP in vitro when co-administered at 650 nM, the ED50 for E64. Compounds are grouped according to the progressive optimization throughout the drug discovery process: 1st generation compounds (1st Gen) are compounds with potency equal or slightly higher versus E64; 2nd generation compounds (2nd Gen) have higher potency than E64 in inhibiting calpain; and 3rd generation (3rd Gen) compounds, instead, are more potent than E64 displaying improved selectivity toward calpain 1 inhibition.

Novel calpain inhibitors developed throughout the three synthesis generation (1st generation: NYC165, NYC131; 2nd generation: NYC215, NYC227, NYC103; 3rd generation: NYC438, NYC488, and NYC106) provided robust results, except compounds NYC227 and NYC106. Slices...
perfused with our candidate compounds were able to preserve synaptic plasticity in the presence of 200 nM Aβ42, confirming the validity of the phenotypical drug screening [32] (Fig. 2C).

These results further validate the use of calpain inhibitors to recover synaptic plasticity in AD. All molecules were calpain inhibitors with equal or superior potency with respect to E64, while 2nd and 3rd generation molecules were developed to increase selectivity for calpain 1, as previously described [32].

Pharmacokinetic (PK) profile and brain drug activity of novel calpain 1 inhibitors

Next, we assessed the PK profile of the best 2nd and 3rd generation compounds obtained from the functional screening on synaptic plasticity (NYC215, NYC438, and NYC488) via LC-MS/MS determination of plasma concentrations. We treated i.p. three sets of mice for the PK assessment of different candidate compounds at equimolar concentrations: NYC215 (7.57 mg/kg), NYC438 (7.86 mg/kg), and NYC488 (7.83 mg/kg). The analysis of kinetics indicated that all three candidate compounds NYC215, NYC438, and NYC488 were rapidly absorbed. The peak plasma concentration occurred at 0.25, 0.5, and 1.25 h after dosing, respectively. Figure 3 shows the plasma concentrations at each sampling time. The absolute bioavailabilities of NYC215, NYC438, and NYC488 were 80.4%, 87.3%, and 41.3%, respectively. Their 1/2 lives were 0.6 h, 1.1 h, and 0.6 h.

Our next goal was to determine if the inhibitor candidate is capable of lowering levels of spectrin proteolytic degradation products in hippocampi from adult animals upon in vivo administration. Spectrin is a cytoskeleton protein target for calpain cleavage. The presence of a specific calpain fragment around 145 kDa is an index of calpain activity [43] and a decreased immunoreactivity in western blotting assays would instead indicate low activity of calpain inhibitors [44]. This kind of investigation offers insights on specific calpain inhibition and brain penetration at the dose used in the efficacy studies in vivo. Using western blot analysis, we checked the prevention of calpain-generated spectrin fragments following i.p. treatment for 12 days with NYC215, the best 2nd generation compound, and NYC438 and NYC488, the two 3rd generation compounds that surpassed E64 benchmark in the LTP rescue assay, at the same concentrations used for PK assessment. The compound NYC215 was slightly less efficient at preventing the spectrin cleavage by calpains while the remaining two compounds NYC438 and NYC488 dramatically reduced the amount of fragments (Fig. 4). This result confirmed the ability of NYC215, NYC438, and NYC488 to cross the blood-brain barrier (BBB) and inhibit calpain in the brain. The brain penetration could result from either passive diffusion or active transport through a BBB transporter. Future experiments should involve in vitro studies with various BBB transporters to understand the transport modalities.

Preliminary toxicity profile of optimized compounds

Our next goal was to have a preliminary assessment of the toxicity profile of NYC215, NYC438, and NYC488 in vivo. Using western blot analysis, we checked the prevention of calpain-generated spectrin fragments following i.p. treatment for 12 days with NYC215, the best 2nd generation compound, and NYC438 and NYC488, the two 3rd generation compounds that surpassed E64 benchmark in the LTP rescue assay, at the same concentrations used for PK assessment. The compound NYC215 was slightly less efficient at preventing the spectrin cleavage by calpains while the remaining two compounds NYC438 and NYC488 were absolutely more active in decreasing the spectrin fragment at around 145 kDa that is generated specifically by calpain. The decrease of the specific calpain-generated fragment demonstrates that the drugs can reach the brain upon systemic administration, overcoming the problems related to first-pass metabolism and blood-brain barrier penetration.
and NYC488. Typically, toxicity studies would help in predicting possible side effects and deciding the safe dose of drugs to be administered during clinical studies [45, 46]. In a series of experiments, we determined the maximum tolerated dose (MTD) following NYC215, NYC438, and NYC488 administration in mice. MTD was computed as the maximum administered dose that does not produce any toxicity effect in terms of malaise (i.e., immobility, altered gait, hunched posture, spiky coat/stops, grooming, altered urination, and/or defecation, porphyrin staining around eyes and nose, vocalization, decrease access rate to food and water, PICA behavior) or death. This restrictive definition is actually superimposable with the one of “no observed adverse effect level” (NOAEL) dose [47].

From the NOAEL, it can be calculated the human equivalent dose (HED) and maximum recommending starting dose (MRSD) as prescribed by current FDA indications [48].

For assessment of the MTD, WT mice (3–5 month old) were acutely injected either with NYC215, NYC438, and NYC488 in an exploratory challenging dose-response treatment designed as sequential acute i.p. injections to establish the dose that produces a marked malaise. MTD was experimentally obtained as the dose immediately antecedent the one inducing malaise, and was found to be at 100 mg/kg i.p. for NYC215, at 150 mg/kg i.p. for NYC438, and at 200 mg/kg i.p. for NYC488. All these doses were >10 times higher than the concentration used in the efficacy study.

The evaluation of acute toxicity at MTD doses was then carried out in another set of animals (3–5 month old), acutely injected either with vehicle, NYC215, NYC438, or NYC488. No clinical sign of toxicity (as measured through food and liquid intake, weight change, locomotion and exploratory behavior, as well as mortality) were observed during the first 24 h with continuous monitoring given in the first 4 h, as well as for 14 days after acutely single dose administration. This observation was supported by the necropsy performed on the treated animals at 14 days after the acute treatment. Necropsy included weights and measurements of organs, appearance of organs (fat deposition, hemorrhage, pigment deposition or other changes, lesion, consistency), and examination of specific macro lesions such as abnormal growths, fibrosis, and necrosis. We did not observe signs of anatomical modifications.

Next, we performed the evaluation of chronic toxicity. An additional set of experiments was performed with treatment for 15 days at the respective MTD for each drug or vehicle. Body weight, fluid and food intake, as well as any sign of behavioral distress, were continuously monitored throughout the treatment. No physical/behavioral distress or death was observed at the end of the chronic treatment, animals were sacrificed and necropsy was carried out in all the animals. Finally, in a separate set of experiments, treatment was perform for 15 days at the MTD and animals were monitored afterwards for additional 15 days to examine possible delayed signs of toxicity. Again, we did not observe any signs of toxicity.

In the absence of gross abnormalities, histopathological evaluation after necropsy was limited to organs that have reported pathology linked to calpain inhibition or loss of function (for a review on the role of calpains in pathology see [49]). In particular, we focused on the condition of the liver (hepatotoxicity has been reported in association with protease inhibitors used in the treatment of HIV that inhibit calpain activity, hepatic steatosis and fibrosis, elevated free fatty acid levels and insulin resistance are associated with decreased activity of calpain 10), the kidney (looking for signs of diabetic nephropathy because of the possible inhibition of calpain 10, cellularity, inflammation, collagen deposition/fibrosis/sclerosis on glomeruli, vessels, tubules, collecting ducts and interstitium, common nephrotoxic effects as proximal tubular epithelial cell damage or renal papillary necrosis), the muscle (checking for myofiber size and fibrosis, necrotic fibers and dystrophy because the loss of function mutations in calpain 3 results in Limb Girdle Muscular Dystrophy Type 2A, regenerative fibers, fat deposition, inflammation), the stomach (assessing the possible presence of gastric cancer because calpain 9 has been proposed to act as a gastric cancer suppressor), and finally the brain (assessing cytoarchitecture, neuronal loss including both apoptosis and necrosis, inflammation, axonal degeneration, gliosis, myelinlation, body inclusions, neurotoxicity in neocortex, striatum, thalamus, hippocampus, brain stem, and cerebellum). Overall, the histopathological evidence did not reflect any generalized toxicity induced by the chronic treatment at MTD for the three candidate inhibitors (Table 1). However, it is noteworthy that potential nephrotoxicity, induced by the 2nd generation compound NYC215, was suggested by the isometric nephrotoxic effects as proximal tubular epithelial cell damage or renal papillary necrosis.
Table 1
Assessment of the toxicity profile of new calpain inhibitors in vivo. Compounds NYC215, NYC438, and NYC488 were chronically administered in vivo at the respective MTDs (that are over 10 times higher than the expected therapeutic dose). Histological evaluation for probing gross modifications were carried out in different target organs known to be a possible source of concern in epoxide-based and calpain inhibition treatments (liver, kidney, muscle, stomach, and brain). The 3rd generation compounds NYC438 and NYC488 showed no signs of toxicity whereas the 2nd generation compound NYC215 induced a discrete isometric tubular epithelial vacuolization in the kidney, probably associated to osmolarity adjustment.

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Phenotypic screening of novel calpain 1 inhibitors with cognitive tests

Second and third generation lead compounds were then tested for the ability to ameliorate the defect in associative memory through fear conditioning assessment [39] and in short-term reference memory using the 2 day radial-arm water maze (RAWM) test [40, 41] in double transgenic AβPP/PS1 mice both at early stages of Aβ deposition (3 months) and late stages (7 months). Both associative memory and short-term spatial memory are early signs of cognitive decline in AD [51, 52].

We first examined contextual fear learning, a hippocampus-dependent task, in both AβPP/PS1 and WT littermates treated with either vehicle or NYC215, NYC438, and NYC488 at equimolar doses (7.57 mg/kg, 7.86 mg/kg, and 7.83 mg/kg, respectively) from the age of 2 months until 3 months. Both associative memory and short-term spatial memory are early signs of cognitive decline in AD [51, 52].

We assessed 24 h later by measuring freezing behavior in response to presentation of the context (contextual conditioning) or of the auditory cue within a completely different context (cued conditioning). We found no difference in the freezing behavior of the vehicle and inhibitor-treated AβPP/PS1 mice compared with vehicle- and inhibitor-treated WT littermates during the training phase of the fear-conditioning test (data not shown). Twenty-four hours later, we found decreased freezing behavior in vehicle-treated AβPP/PS1 mice compared with vehicle-treated WT littermates in the analysis of contextual learning. However, treatment with our lead molecules restored freezing in AβPP/PS1 mice and did not affect the performance of WT mice (Fig. 6A). Treatment with the compounds did not affect the performance of WT mice, further suggesting that they do not induce toxicity (Fig. 6A). Moreover, vehicle-treated WT mice showed similar freezing time as control untreated WT mice (data not shown). We also did not find a difference in freezing behavior during cued learning (data not shown). These results indicate that the impairment in contextual fear learning in AβPP/PS1 mice can be rescued by treatment with a calpain inhibitor. Testing of the effects of the novel inhibitors in older mice that were treated from the age of 2 months until 7 months confirmed the results of younger mice (Fig. 6B). The novel calpain inhibitors were able to prevent the cognitive disturbance.
in associative memory in double transgenic AβPP/PS1 mice at 7 month of age. AβPP/PS1 mice treated with the novel calpain inhibitors showed consistently more freezing in fear conditioning assessment than vehicle-treated mice, demonstrating the possibility to prevent the occurrence of cognitive disturbances in AβPP/PS1 mice even at ages when typically learning and memory is severely impaired.

Next, we aimed to verify whether the same treatment could reverse the spatial learning impairment in AβPP/PS1 mice. The 2-day RAWM task was performed as previously described [41]. The mouse had to swim in a 6 arm maze filled with milky water until it was able to find a hidden platform at the end of one of the arms (submerged platform) using the visual orientation cues placed within sight above the maze. During the first day (training), mice were trained to identify the platform location by alternating between a visible and a hidden platform in a specific maze arm (goal arm) during several consecutive training trials. We evaluated the number of mouse entries in an arm with no platform (incorrect arm entries). Failure to select an arm after 15 s was counted as an error. Each trial lasted up to 1 min. After 1 min, if the platform had not been located, the mouse was then directed towards the platform while swimming.

NYC215 or NYC438 or NYC488 improved the performance of AβPP/PS1 mice with the 2-day RAWM task without affecting the performance of WT littermates (Fig. 6C). Indeed, vehicle-treated AβPP/PS1 failed to reach the learning criterion (1 error) in the 2-day RAWM task by block 9 and 10 of day 2 whereas vehicle-injected WT littermate mice succeeded. NYC215, NYC438, and NYC488 ameliorated the deficit in RAWM performance in transgenic mice on the last block. NYC215, NYC438, and NYC488 did not affect the WT littermate performance with the RAWM. Vehicle-treated WT mice had similar performance as control untreated WT mice (data not shown).

To test for visual, motor, and motivational deficits, all mice also underwent visible platform task after performing the RAWM test. We found no difference in speed and latency period to the platform for the various groups of mice (data not shown).

Testing of the effects of the novel inhibitors in older mice that were treated from the age of 2 months until 7 months confirmed the results in younger mice (Fig. 6D). Our novel calpain inhibitors prevented deficits in spatial memory in double transgenic AβPP/PS1 mice at 7 months of age.

Overall, AβPP/PS1 mice treated with the novel calpain inhibitors showed consistently more freezing in fear conditioning assessment and fewer errors in 2-days RAWM than vehicle–treated mice, demonstrating the possibility to prevent the occurrence of cognitive disturbances in AβPP/PS1 mice even at ages when typically learning and memory is severely impaired.
Fig. 6. Behavioral evaluation of the ability of novel calpain inhibitor to rescue memory defects in AβPP/PS1 mice. A) Daily treatment with 2nd generation NYC215, and 3rd generation NYC438 and NYC488 from the age of 2 months until 3 months ameliorated the defect in contextual fear memory in AβPP/PS1 mice. WT-vehicle: n = 15, WT-NYC215: n = 8, WT-NYC438: n = 8, AβPP/PS1-vehicle: n = 15, AβPP/PS1-NYC215: n = 10, AβPP/PS1-NYC438: n = 11, AβPP/PS1-NYC488: n = 10, p < 0.05 in all transgenic groups treated with compound compared to their respective vehicle-treated transgenics. B) Daily treatment with NYC215, NYC438 and NYC488 from the age of 2 months until 7 months ameliorated the defect in contextual fear memory. WT-vehicle: n = 18, WT-NYC215: n = 9, WT-NYC438: n = 9, WT-NYC488: n = 9, AβPP/PS1-vehicle: n = 17, AβPP/PS1-NYC215: n = 10, AβPP/PS1-NYC438: n = 10, AβPP/PS1-NYC488: n = 11, p < 0.05 in all transgenic groups treated with compound compared to their respective vehicle-treated transgenics. C) Daily treatment with NYC215, NYC438, and NYC488 from the age of 2 months until 3 months ameliorated the defect in spatial memory in AβPP/PS1 mice. WT-vehicle: n = 15, WT-NYC215: n = 8, WT-NYC438: n = 8, WT-NYC488: n = 8, AβPP/PS1-vehicle: n = 15, AβPP/PS1-NYC215: n = 10, AβPP/PS1-NYC438: n = 11, AβPP/PS1-NYC488: n = 10, p < 0.05 in all transgenic groups treated with compound compared to their respective vehicle-treated transgenics. D) Daily treatment with NYC215, NYC438, and NYC488 from the age of 2 months until 7 months ameliorated the defect in spatial memory in AβPP/PS1 mice. WT-vehicle: n = 18, WT-NYC215: n = 9, WT-NYC438: n = 9, WT-NYC488: n = 9, AβPP/PS1-vehicle: n = 17, AβPP/PS1-NYC215: n = 10, AβPP/PS1-NYC438: n = 10, AβPP/PS1-NYC488: n = 11, p < 0.05 in all transgenic groups treated with compound compared to their respective vehicle-treated transgenics.

Taken together, these data suggest that the novel calpain inhibitors are capable and quite effective in restoring the cognitive abilities in the AβPP/PS1 AD mouse model. This implies that early treatments with calpain inhibitors may prevent from the progressive cognitive sequelae already established in the early stage of the disease. These results have significant translational value in further reinforcing the need for clinical trials of calpain inhibitors in AD.

Nevertheless, in the current study we verified whether the 3rd generation compound NYC438, which has longer half-life and better bioavailability, altered Aβ levels. ELISA analysis of Aβ40 and Aβ42 levels in hippocampal and plasma samples revealed readily quantifiable levels of these peptides in the hippocampus and blood following 5-month treatment with NYC438 or vehicle in double transgenic AβPP/PS1 mice. Daily treatment with NYC438 did not reduce hippocampal or blood levels of Aβ40 and Aβ42 in AβPP/PS1 mice (Fig. 7). Overall, these data support the hypothesis that the beneficial effects of calpain inhibition are produced by mechanisms downstream of Aβ production, while still counteracting the Aβ-induced detrimental effects.

Novel calpain 1 inhibitors do not alter cerebral and plasma Aβ content

We previously demonstrated that both E64 and BDA-410 do not have any effect on Aβ levels [19].
AD is a neurological, multifactorial illness, of epidemic proportions, which is characterized by a number of neuropathological features, including the steady presence of diffuse proteinaceous aggregates in the brain diffused along brain areas, neuronal death, and synaptic changes leading to dementia [53, 54]. AD incidence in the population is growing together with progressively increasing lifespans [25, 55]. Thus, there is an undisputable need for robust and safe therapeutic tactics to treat the disease. One of the putative causes of AD is attributable to the increased presence or circulation of soluble oligomers of Aβ provoking both a substrate for synaptic degeneration and diffusion of the pathology along brain areas [30]. Substantial evidence supports the ability of Aβ oligomers to reduce plasticity and memory both in translational experimental models of the disease (for a review see [56]) and in humans [57–59]. It is therefore conceivable to counteract the detrimental effects of oligomeric Aβ exposure as a possible target for the development of a causal therapy in AD. Indeed, the potential therapeutic value in AD of our novel calpain inhibitors was demonstrated in their ability to ameliorate deficits in LTP and memory induced by soluble Aβ-induced, overcoming the possible criticism of drugs developed solely to block protein aggregation and deposition [48].

Another important feature of our approach is targeting calpain inhibition. Over-activation of calpains, whether as a direct consequence of Aβ activity or due to other factors affecting calcium signaling (for a review, see [60, 61]), is one of the culprits of AD [62]. Moreover, calpain hyperactivation has been shown in other proteinopathies [63–65]. Therefore, the identification of calpain inhibitors might prove to attenuate elevated calpain activity while preserving physiological levels of calpain activation [66, 67]. Among the known calpain inhibitors which have been studied in AD is A-705253, developed by Abbvie Pharma, a notable ketoamide-based calpain inhibitor that is active in the 3xTgAD mouse model rescuing memory defects, and reducing levels of BACE enzyme, Aβ deposits, and overall neuroinflammation [68, 69]. We designed a novel series of epoxide-based calpain inhibitors using E64 as a lead. Two 3rd generation inhibitors, NYC438 and NYC488, were potent inhibitors of calpain 1 (IC50 <100 nM) with improved selectivity and easy synthetic scalability [72]. Inhibition of spectrin cleavage provided evidence of functional brain bioavailability, and both inhibitors were able to recover both the plasticity and the memory impairment associated with the exposure to Aβ42.

Fig. 7. ELISA analyses of Aβ40 and Aβ42 levels after treatment with the novel calpain inhibitor NYC438. 3rd generation compound NYC438 did not affect Aβ40 and Aβ42 levels in hippocampal of 7-8 month old AβPP/PS1 mice (n = 5 per group).
not show reactivity toward non-specific targets [32]. BDA-410 and E64 bear cyclopropenone and epoxide warheads, respectively [19]. Both cyclopropenone and epoxide warheads are electrophilic, with the potential to covalently modify the cysteine active site of calpain 1, or to non-selectively modify other protein-thiols, or even to form glutathione conjugates and therefore be rapidly cleared [72]. In our drug discovery design, therefore, we considered the knowledge of reactivity toward proteins and free thiols as a specific requirement early in the drug development project, in addition to the simple structure-based, "rational" drug design strategy towards selective calpain 1 inhibitors using phenotypical screening.

Both of our 3rd generation compounds, NYC438 and NYC488, had very good PK characteristics with a slightly longer half-life for NYC438 versus NYC488 (∼1.1 h and ∼0.6 h). The inhibitors, moreover, showed no overall toxicity, despite the general idea that epoxides are liable because of non-specific interaction with thiol groups in off-targets [71]. This is important as our toxicological tests were performed at the MTD, which is at least ten-times higher than specific therapeutic doses [48]. This signifies an intrinsic safety of our lead 3rd generation compounds [73]. Epoxide inhibitors might display a number of advantages over other types of cysteine protease inhibitors. In fact, epoxides are less reactive than other classes of inhibitors, such as halomethylketones or aldehydes, showing to be more directed to a single target (whether calpains or cathepsins) and therefore they do not modify the levels of Aβ [42]. As recently remarked by the Alzheimer’s Association [25], the slow, insidious nature of AD progression, the staggering impoverishment of the quality life for AD patients, and the destructive public health outcomes such as family disintegration/impoveryments, and cost of caregiving supports the quest for new symptom- and disease-modifying treatments. The availability of a new series of effective, non-toxic therapeutics, exerting their action on the molecular mechanisms of the disease, is a positive step toward a better cure for AD and any other disease associated with calpain overactivation.

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