N-Methylated Peptide Inhibitors of β -Amyloid Aggregation and Toxicity. Optimization of the Inhibitor Structure[†]

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ABSTRACT: The key pathogenic event in the onset of Alzheimer's disease (AD) is believed to be the aggregation of the β -amyloid (A β) peptide into toxic oligomers. Molecules that interfere with this process may therefore act as therapeutic agents for the treatment of AD. N-Methylated peptides (meptides) are a general class of peptide aggregation inhibitors that act by binding to one face of the aggregating peptide but are unable to hydrogen bond on the other face, because of the *N*-methyl group replacing a backbone NH group. Here, we optimize the structure of meptide inhibitors of A β aggregation, starting with the KLVFF sequence that is known to bind to A β . We varied the meptide length, N-methylation sites, acetylation, and amidation of the N and C termini, side-chain identity, and chirality, via five compound libraries. Inhibitor activity was tested by thioflavin T binding, affinity chromatography, electron microscopy, and an 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide toxicity assay. We found that inhibitors should have all D chirality, have a free N terminus but an amidated C terminus, and have large, branched hydrophobic side chains at positions 1–4, while the side chain at position 5 was less important. A single *N*-methyl group was necessary and sufficient. The most active compound, D-[(chGly)-(Tyr)-(chGly)-(chGly)-(mLeu)]-NH₂, was more active than all previously reported peptide inhibitors. Its related non-N-methylated analogues were insoluble and toxic.

Alzheimer's disease (AD)¹ is the most common form of senile dementia, affecting more than 15 million people worldwide. With an increased life expectancy and aging population, this number is set to rise considerably. At least 20 diseases are caused by proteins or peptides folding incorrectly and aggregating into fibrils or plaques, including AD, type-II diabetes, Parkinson's disease, Huntington's disease, and the spongiform encephalopathies (1). They accumulate either intra- or intercellularly in a variety of organs, including the liver, spleen, and, most importantly, the brain, causing severe neurological disorders. The aggregates are often in the form of regular amyloid fibrils. Amyloid is rich in β -sheet structure, with hydrogen bonding between monomers parallel to the fibril axis, leading to fibrils of indefinite length. The best studied amyloid-based disease is AD, characterized pathologically by abnormally high levels of brain lesions (senile plaques) and neurofibrillary tangles in dead and dying neurons and by elevated numbers of amyloid deposits in the walls of cerebral blood vessels (2).

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The major component of senile plaques is a small peptide of 39–43 amino acids called β -amyloid (A β). A β (1–40) is the most prevelant species, while A β (1–42) is more toxic; other lengths are rare. A β is produced through endoproteolysis of the amyloid precursor protein. Compelling evidence indicates that factors that increase the production of A β , particularly its more amyloidogenic variants, or that facilitate deposition or inhibit elimination of amyloid deposits cause AD or are risk factors for the disease (3).

Controversy has raged over whether the fibrils are an epiphenomenon linked to disease or whether fibril formation causes disease (4). For example, while many mutations in the A β precursor protein gene are linked to premature onset of AD, the amount of amyloid deposited in the brain does not correlate to severity. A resolution of this apparent paradox from in vitro and in vivo evidence is that soluble, oligomeric forms on A β have potent neurotoxic activity and are the primary causes of neuronal injury and cell death occurring in AD (5, 6).

Numerous groups are developing treatments designed to block various key steps in the amyloidosis process. Specific therapeutic strategies currently being pursued include (7-9) (1) inhibiting the expression of the amyloidogenic protein or stabilizing its native form using small organic ligands, (2) inhibiting the release of the amyloidogenic peptide from its parent protein using protease inhibitors, (3) inhibiting the aggregation of the protein or peptide directly using small ligands or indirectly by vaccination, (4) inhibiting other effects of the disease, which may or may not be directly associated with amyloidosis (e.g., inflammation and oxidative

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¹ Abbreviations: AD, Alzheimer's disease; A β , β -amyloid; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; ThT, thioflavin T; EM, electron microscopy; HFIP, hexafluoroisopropanol; TFA, trifluoroacetic acid.

stress), (5) replacing cells that have been killed by the disease (e.g., by cell or gene therapy), and (6) alleviating the symptoms of the disease but without necessarily blocking the pathogenic process.

The most effective treatments may be those designed to inhibit steps that precede protein/peptide aggregation, by blocking production of the amyloidogenic protein or peptide in the first place. However, this requires blocking the expression or activity of a natural protein or peptide that has presumably evolved to perform some other, important biological function in vivo. For example, many groups are currently developing inhibitors of β - or γ -secretase as potential drugs for AD. These two enzymes cleave amyloid precursor protein to produce the A β peptide associated with this disease, but they have also been shown to perform other, important biological functions (10). On the other hand, treatments designed to target steps that follow protein/peptide aggregation are less likely to be effective because they would not prevent the formation of toxic soluble oligomers or insoluble fibers, which could continue to kill cells. Thus, an attractive therapeutic strategy in principle is to inhibit protein/ peptide aggregation itself, because this appears to be the first step in the pathogenic process of amyloidosis, which is not associated with some natural biological function (11, 12).

An attractive strategy to develop amyloid aggregation inhibitors is to use the wild-type peptide, because it is already known to bind to itself, thus avoiding the usual necessity in drug discovery of finding a starting compound with some activity. The first group to make use of a core section of $A\beta$ as a structural starting point was Tjernberg and co-workers, who showed that $A\beta(16-20)$ was able to bind full-length A β and thus prevent its assembly into fibrils (13). Despite being shown to form fibrils in isolation, $A\beta(16-20)$ (KLVFF) was proposed as being a key region from which a lead compound could be created against amyloid. Soto et al. also began work on inhibitors aimed at the core region of A β , in this case, residues 17-21 (LVFFA) (14). The strategy is based on substituting key residues for prolines in a bid to reduce the β propensity of the peptide, while retaining its hydrophobicity. A lead 11 amino acid inhibitor was reduced to five residues with greater ability to prevent fibril formation. All D analogues were found to be as effective but with increased protease resistance. These so-called β -sheet breaker peptides were shown not only to be stable in vivo but also to have blood-brain-barrier permeability (15, 16). Initially on the basis of the 15–25 region of A β , Murphy and workers designed a peptide with a "recognition element" homologous to the A β peptide but with a disrupting element of at least three lysines tagged to the C terminus, designed to interfere with A β aggregation (17–19). Sato et al. designed heptapeptide inhibitors of A β toxicity based on β -sheet packing (20). New types of small molecule inhibitors of A β aggregation have also recently been reported (21-23).

Here, we use N-methylated peptides (meptides) as inhibitors of amyloidosis. One side presents a hydrogen-bonding "complementary" face to the protein, with the other having *N*-methyl groups in place of backbone NH groups, thus presenting a "blocking" face. Substituents larger than a methyl group are also effective (24). We have previously shown that *N*-methyl derivatives of $A\beta(25-35)$ are able to prevent aggregation and inhibit toxicity in PC12 cells (25). Meredith and co-workers (26, 27) investigated N-methylated peptides of a region corresponding to 16–22 and later 16– 20 of the amyloid "core domain" region. They can prevent $A\beta$ fibrils from forming and break down preformed fibrils. These peptides have the added advantages of high proteolytic resistance, solubility, membrane permeability (27), and high propensity to form β structure at the N-methylated site. Cruz et al. found that single *N*-methyl amino-acid-containing peptides related to 16–20 of $A\beta$ were able to reduce the cytotoxicity of $A\beta(1-42)$ (28).

MATERIALS AND METHODS

Stock Preparation. $A\beta(1-40)$ and $A\beta(1-42)$ [(Cl) salt] was purchased from the American Peptide Company. A 10 mM stock solution of each target peptide in 1,1,1,3,3,3-hexafluoroisopropanol (HFIP) was prepared according to the Zagorski method to ensure that the peptide is initially in a monomeric state (29) as follows: (i) 3 cycles of dissolution with vortexing and sonication in high-grade trifluoroacetic acid (TFA, from Fluka) and drying under a slow stream of dry N₂, (ii) 3 cycles of dissolution (with vortexing and sonication) in high-grade HFIP and drying under a slow stream of dry N₂, (iii) dissolution of the target to 10 mM in HFIP, and (iv) storage of the 10 mM target stock solution in HFIP at 4 °C. TFA/HFIP treatment ensures that the amyloidogenic peptide is in a monomeric, soluble state.

Inhibitors (prepared by solid-phase Fmoc chemistry using *O*-(7-azabenzotriazol-1-yl)-1,1,3,3-tetramethyluronium as the coupling agent) were supplied in lyophilized form at >95% purity, verified by high-performance liquid chromatography (HPLC) and mass spectrometry, by Peptide Protein Research Ltd. (Southampton, U.K.). Inhibitors were dissolved to 1 mM (or in a few cases 0.5 or 2 mM) in 50% acetonitrile. In a few cases, some inhibitors were dissolved in 100% H₂O. The concentration of each A β -derived inhibitor was measured by its Phe absorbance at 257.4 nm ($\epsilon = 197 \text{ M}^{-1} \text{ cm}^{-1}$ per Phe) or by weighing. The inhibitor solutions were stored at 4 °C prior to lyophilization of individual aliquots for each assay.

Thioflavin T (ThT). Inhibition assays were performed with 100 µM target peptide in 200 µL Dulbecco A phosphatebuffered saline (PBS) buffer (Oxoid), with or without each inhibitor at a concentration of $10 \,\mu\text{M}$ (for 1:0.1 molar ratio), 100 µM (for 1:1 molar ratio), or 1 mM (for 1:10 molar ratio). Sufficient target peptide was lyophilized, redissolved, and thoroughly vortexed as one single batch (for immediate use in all target-inhibitor mixes) to a concentration of 200 μ M in pure, sterile-filtered ddH₂O to delay aggregation until the addition of each inhibitor. In parallel, 2 nmol ($20 \times 100 \,\mu\text{L}$ stock), 20 nmol (20 μ L × 1 mM stock), or 200 nmol (200 μ L × 1 mM stock) of each inhibitor was lyophilized and redissolved in a 1 mL Eppendorf tube to a concentration of 20 µM (1:0.1), 200 µM (1:1) or 2 mM (1:10), respectively, in 100 μ L sterile-filtered 2× Dulbecco A PBS buffer (Oxoid: 320 mM NaCl, 6 mM KCl, 16 mM Na₂HPO₄, and 2 mM KH₂PO₄ at pH 7.4) All of the inhibitor solutions were then thoroughly vortexed to ensure complete dissolution. Finally, a 100 μ L aliquot of the target solution was added to each 100 μ L aliquot of inhibitor to give a total assay volume of 200 μ L containing 100 μ M target and either 10 μ M, 100 μ M, or 1 mM inhibitor in 1× Dulbecco PBS buffer. The assay mixtures were vortexed and held in a 37 °C incubator for 4 days to induce aggregation in the presence of each inhibitor.

The ThT assay solution was prepared from stock containing 500 μ M ThT in 50 mM glycine buffer at pH 8.5. The stock was aliquoted into 1 mL Eppendorfs and kept frozen until it was needed. It was then allowed to thaw at room temperature for 10 min before $25 \times$ dilution into the appropriate glycine buffer, giving the required freshly prepared ThT assay solution containing 20 µM ThT in 50 mM glycine buffer at pH 8.5. A total of 180 µL of the ThT assay solution was then added to each well of a 96-well black-bottomed microplate per desired repeat, using a multichannel pipet. Next, 20 µL aliquots of each inhibition/ reversal assay mixture (thoroughly vortexed) were added to individual wells containing the ThT solution using an automated micropipet. The fluorescence of amyloid-bound ThT was measured in a Twinkle LB970 Fluorescence Plate Reader (Berthold Technologies) using the "top reading" setting with the lamp intensity set at 10 000 and a 3 s "shake" prior to each fluorescence reading, which was recorded over a period of 1 s using appropriate excitation and emission filters. Bound ThT excites at 450 nm and emits at 482 nm (30).

For the inhibition assays, the target-inhibitor mixtures were incubated at 37 °C. Single ThT readings were taken on days 1, 2, and 3, and five repeat readings were taken on day 4. For the reversal assays, the target was incubated alone at 37 °C for 3 days before the addition of 200 μ L to each lyophilized inhibitor. The vortexed target-inhibitor solutions were then incubated at 37 °C for a further 4 days, during which time single ThT readings were taken on days 1, 2, and 3 and then five readings were taken on day 4 because maximal ThT binding was found after 4 days of aggregation (Figure A in the Supporting Information). A decrease in ThT fluoresence after 4 days may be due to the formation of larger fibrils that are able to bind a smaller amount of ThT. Inhibition/disaggregation activity was calculated by the percent reduction in the amyloid-bound ThT fluorescence of the target-inhibitor mixture compared with that of the target alone. The percentage amyloid is given by $(F_i - F_b)/$ $(F_{\rm a} - F_{\rm b}) \times 100\%$, where the percentage of amyloid is the percent reduction in the amyloid-bound ThT fluorescence, F_i is the ThT fluorescence of the target-inhibitor mixture, $F_{\rm a}$ is the ThT fluorescence of the target alone, and $F_{\rm b}$ is the background ThT fluorescence with no target or inhibitor present.

Affinity Chromatography. Two copies of the A β core aggregation sequence were attached to the resin in tandem, linked by a pair of glycine residues for flexibility to allow for the formation of a β -turn/hairpin (sequence L-Ac-[(KLVFFAE)-GG]₂-resin). The affinity columns were synthesized by Peptide Protein Research Ltd. on a CP glass resin. The glass resin was chosen because of the inability of the inhibitor peptides to bind to the resin and for its incompressibility, allowing use in an HPLC system. All inhibitors were diluted to 100 μ M stock solutions, containing 5% acetonitrile from their 1 mM stocks in 50% acetonitrile. A few inhibitor peptides were kept as 1 mM stocks in 100% H₂O and were diluted to 100 μ M in 100% H₂O. A total of 180 μ L of the 100 μ M inhibitor stocks was transferred to specific inserts, which were then placed into vials. These specific HPLC vials

were positioned on the autosampler of the Agilent Technologies HPLC instrument, where before the start of an affinity chromatography run, 50 μ L would be collected from the insert and injected into the affinity column automatically, as controlled by the HPLC software (LC/MSD Chemstation, version A. 09.03). Each inhibitor was run through each affinity column 3 times using a linear gradient of 5–95% acetonitrile/0.1% TFA in ddH₂O/0.1% TFA for 33 min. The column was then washed with 95% acetonitrile/0.1% TFA for 2 min, decreased to 5% acetonitrile/0.1% TFA for 2 min, and equilibrated at 5% acetonitrile/0.1% TFA for 3 min, at a flow rate of 1 mL/min. The same cycle was repeated for each affinity run.

Electron Microscopy (EM). The same samples that were prepared for ThT fluorescence were also used for preparing samples for the EM assay. From the 200 μ L that was prepared, 180 μ L was used for the ThT assay and the remaining 20 µL was used for preparing EM samples. A total of 10 μ L drops of the peptide solution samples were placed side by side on a clean strip of Parafilm along with 10 μ L drops of 2% uranyl acetate (w/v in water) on another strip of Parafilm. Carbon-coated copper grids (400 mesh/ inch) were then glow-discharged for 30 s to render them hydrophilic. The glow-discharged EM grids were then inverted on the peptide solution drops for 30 s, carefully blotted by touching the grid edge to the filter paper and negatively stained by inverting them on the uranyl acetated drop for 30 s. The grids were allowed to dry overnight. They were then ready for viewing under the microscope. Electron micrographs were recorded on a Philips EM 301 electron microscope at 100 kV at a magnification of 44000×.

3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium Bromide (MTT) Cell-Toxicity Assay. Rat phaeochromocytoma (PC12) cells were chosen for assessing toxicity because they have been identified as sensitive to the toxic effects of $A\beta$ using MTT (31). The MTT cell titer 96 proliferation assay (Promega) is an in vitro assay measuring the conversion of the MTT tetrazolium component of the CellTitre 96 dye solution into a formazan product by the living PC12 cells, where the formazan product becomes solubilized and the absorbance is read at 570 nm. A color change takes place, and a measure of the percentage of MTT reduction is calculated, where the percentage of cellular MTT reduction provides an indication of the cell condition or health.

Briefly, 10 μ M inhibitors were screened in the presence of 10 μ M A β (1-42). The required volume from inhibitor and target stock solutions was freeze-dried overnight. The freeze-dried inhibitors and target peptide were resuspended in 100% dimethyl sulfoxide (DMSO), each at 2 mM concentration. A total of 5 μ L from each of the resuspended inhibitor/DMSO and target/DMSO was mixed in a well of a 96-well preparation plate, thus giving 10 μ L of 1:1 mM inhibitor/target concentration ratio in 100% DMSO. A total of 90 μ L of Optimem media was added to the 10 μ L inhibitor/target mixture (100:100 µM inhibitor/target ratio in 10% DMSO). A total of 10 μ L of the 100:100 μ M inhibitor/target mixture in 10% DMSO was then dispensed into 90 μ L of media/PC12 cells (n = 8), at final inhibitor and target concentrations of 10 μ M. These were incubated for 48 h at 37 °C, 5% CO₂, prior to the addition of the MTT dye. A total of 15 μ L of the dye was added to each well and incubated for a further 4 h at 37 °C, 5% CO₂. A total of 100 μ L of the stop/solubilization solution was then added to each well and was allowed to stand for 1 h. The absorbance was measured at 595 nm using a 96-well Tecan Ultra 384 plate reader. The percent MTT reduction was calculated as percent MTT reduction = $((x - A)/(B - A)) \times 100\%$, where *x* is the absorbance value in each well under study, *A* is the mean absorbance of the negative control, which corresponds to 0% MTT reduction because of the presence of dead cells (0.1% Triton X-100 with cells), and *B* is the positive control, which corresponds to 100% MTT reduction (live cells in 1% DMSO).

While the MTT assay with PC12 cells is the most widely used assay for measuring $A\beta$ toxicity, we were concerned not to use this as the sole measure of toxicity. We therefore verified our MTT results with the SHSY5Y cell line and other indicators of cell viability, namely, calcein, lactate dehydrogenase, cell titer blue, and the tetrazolium salts XTT and MTS. We found that the MTT data were the most reproducible and required the lowest concentration of $A\beta$, giving a more useful concentration range for assaying inhibitors. The MTT results correlated well with all other methods (not shown), giving confidence in using MTT as our main toxicity assay.

RESULTS

Initial Library. The initial inhibitors were named SEN001-SEN021 (Table 1). These were mostly based around Nmethylating the LVFF sequence, known to be key for amyloid formation in A β ; in addition, KLVFF has been shown to act as an inhibitor of A β aggregation (13). Most were acetylated at the N terminus, and all were amidated at the C terminus, to more closely mimic the polypeptide backbone and provide additional potential hydrogen-bond acceptors and donors. Some peptides introduced specific amino acid substitutions to explore the effect of different residues on their ability to induce an inhibitory effect: peptides SEN001, SEN002, SEN011, SEN012, SEN014, SEN015, and SEN020 incorporated Lys at the N terminus, to include residue 16 of the A β wild-type sequence. SEN004-SEN010 had an additional glutamine at the N terminus, thus incorporating residue 15 of A β , and SEN020 included Glu at the C terminus, residue 22 of the wild-type sequence. SEN013 incorporated an Ala instead of Leu¹⁷, while Lys¹⁶ changed positions with Phe¹⁹, thus substituting a charged, hydrophilic side chain with an aromatic hydrophobic side chain at the N terminus. Reference peptides that had already been published as inhibitors of A $\beta(1-40)$ and/ or A $\beta(1-42)$ amyloid formation were used as controls. These included SEN001 (13), SEN202 (26), PPI-1019 (32), and $iA\beta 5p$ (14). D- $iA\beta 5p$ was the enantiomer of $iA\beta 5p$, while C1 and C2 were control peptides anticipated to have no effect on A β .

The ThT results for inhibition of both $A\beta(1-40)$ and $A\beta(1-42)$ for these peptides are shown in Figures A and B in the Supporting Information. Figure A in the Supporting Information shows that the inhibitors affect $A\beta$ aggregation kinetics. The results for SEN001 and SEN002 confirm the results of Tjernberg et al. that nonmethylated KLVFF can act as an inhibitor. However, N-methylation reduced inhibitor activity for all of these L peptides. The most potent inhibitor in this round was the Praecis compound, PPI-1019, sequence

D-[H-[(mL)-V-F-F-L]-NH₂], where methylation is on the N-terminal amine group, rather than a secondary amide in the backbone. We found no activity in iA β 5p, sequence L-[Ac-[L-P-F-F-D]-NH₂], previously reported as an $A\beta$ aggregation inhibitor by Soto et al. (14). No activity was found for the control peptides, C1 and C2, as expected. These results were confirmed by EM, where $A\beta$ fibril morphology was unchanged (data not shown). PPI-1019, the most potent inhibitor in the initial round, is a D amino acid peptide. ThT fluorescent studies by Chalifour et al. indicated that the D enantiomers of peptide-based inhibitors containing the KLVFF A β -recognition motif were more potent inhibitors of the natural L-A β (1-40)-induced aggregation and neurotoxicity than their natural L peptide counterparts (33). Similarly, the L peptides were more effective at inhibiting D-A β (1-40) aggregation than the D peptides. The results therefore indicate that these peptides can have an inhibitory effect of aggregation through a heterochiral type of stereospecificity. The inhibitors in the next round were therefore all D.

Round-1 Library. The round-1 library, sequences SEN101– SEN120 (Table 1), was based on the sequences of SEN018, SEN019, and PPI-1019. Specifically, the new library was composed of acetylated and corresponding nonacetylated peptides, all composed of the LVFFL domain, that were singly, doubly, or triply methylated. The peptides were all D enantiomers and were amidated at the C terminus. The aim of the round-1 library was to assess the effect of acetylation or nonacetylation and to identify the best sites for N-methylation. All inhibitors were assayed by binding affinity, ThT, and EM and compared to C1, C2, SEN018, SEN019, PPI-1019, and $iA\beta5p$. The rest of the library was derived from SEN019 and composed of the LVFFL hydrophobic domain, incorporating either single, double, or triple methylations. Results are shown in Figures 1–3.

As shown in Figure Aa in the Supporting Information, the range of activities of the round-1 inhibitors on $A\beta(1-$ 40) is from 10 to 100% when assayed with ThT, with errors of typically 15%, showing that reliable ranking of inhibitors was possible. The best round-1 inhibitors reduce amyloid formation by $A\beta(1-40)$ to 24, 23, and 13% for SEN101, SEN113, and SEN116, respectively (compared with 64-70 and 44-80% for SEN018 and SEN019, respectively, and 35-42% for PPI-1019). SEN113 and SEN116 are more active against $A\beta(1-40)$ than PPI-1019, as well as $iA\beta 5p$, which had 61% amyloid formation. SEN101 is an acetylated peptide, singly N-methylated at the N-terminal Leu residue (X1), while SEN113 is nonacetylated, singly N-methylated at the C-terminal Leu (X5). SEN116, the best inhibitor studied to this point, is also nonacetylated but doublemethylated at both the N- and C-terminal leucines (X1 and X5). Other peptides, including SEN105 and SEN117, had no effect.

Structure/activity relationships could be inferred by comparing inhibitors differing by a single substituent: the acetylated inhibitors were generally less potent than their nonacetylated counterparts; methylation of the X3 position causes a very negative inhibitory effect, especially when X1 and/or X5 are/is also methylated; methylation of X1 and/or X5 is very favorable; methylation of X2 and/or X4 are/is better than methylation of X3 but not as good as methylation of X1 and/or X5. Overall, the best inhibitor is SEN116,

name	chirality	sequence	pame	chirality	sequence
SEN001	T		SEN220	D	H-[(m])_(chGly)_F F (m])] NH-
SEN001 SEN002	L L	Ac-[K-L-V-F-F]-NH $_{2}$	SEN230 SEN231	D	$H^{(mL)}(chOry)^{-r}-F^{(mL)}$
SEN003	L	$Ac-[L-V-F-F-A]-NH_2$	SEN232	D	$H_{(mL)}(mL) + (mL) +$
SEN004	L	Ac-[Q-K-L-V-F-F]-NH ₂	SEN233	D	$H-[(mL)-(2nAla)-F-F-(mL)]-NH_2$
SEN005	L	$Ac-[Q-K-(mL)-V-F-F]-NH_2$	SEN241	D	$H-[(mL)-V-A-F-(mL)]-NH_2$
SEN006	L	Ac-[Q-K-L-(mV)-F-F]-NH ₂	SEN242	D	$H-[(mL)-V-V-F-(mL)]-NH_2$
SEN007	L	Ac-[QK-L-V-(mF)-F]-NH ₂	SEN243	D	$H-[(mL)-V-I-F-(mL)]-NH_2$
SEN008	L	$AC-[Q-K-L-V-F-(IIIF)]-NH_2$ $AC-[O-K-(mL)-V-(mE)-F]-NH_2$	SEN244 SEN245	D	$H_{(mL)} - V - L - \Gamma - (mL) - N H_2$ $H_{(mL)} - V - M - F_{(mL)} - N H_2$
SEN010	L	$Ac-[O-K-L-(mV)-F-(mF)]-NH_2$	SEN116	D	$H_{[(mL)} = (mL) - V - F - F - (mL) - NH_2$
SEN011	L	Ac-[K-L-(mV)-F-(mF)-A]-NH ₂	SEN247	D	$H-[(mL)-V-Y-F-(mL)]-NH_2$
SEN012	L	Ac-[K-(mL)-V-(mF)-F-(mA)]-NH2	SEN248	D	$H-[(mL)-V-W-F-(mL)]-NH_2$
SEN013	L	Ac-[F-A-(mF)-K-(mV)-L]-NH ₂	SEN249	D	$H-[(mL)-V-(tbGly)-F-(mL)]-NH_2$
SEN014	L	Ac-[K-L-(mV)-F-(mF)]-NH ₂	SEN250	D	H-[(mL)-V-(chGly)-F-(mL)]-NH ₂ H [(mL)-V (chAlo) F (mL)] NH
SEN015 SEN016	L	AC-[\mathbf{K} -(IIIL)-V-(IIIF)-F]-INH ₂ Δc -[I -(mV)-E-(mE)- Δ]-NH ₂	SEN251 SEN252	D	$H_{(mL)-V_{(mL)}-F_{(mL)}-F_{(mL)}}$ $H_{(mL)-V_{(hPhe)-F_{(mL)}}-H_{(mL)}$
SEN017	L	$Ac_{[m]} - (m) -$	SEN252	D	$H^{-1}(mL) - V^{-1}(mL) - V^{$
SEN018	D	$Ac-[L-(mV)-F-(mF)-L]-NH_2$	SEN261	D	$H^{-1}(mL)-V-F-A^{-1}(mL)]-NH_2$
SEN019	D	Ac- $[(mL)-V-(mF)-F-(mL)]-NH_2$	SEN262	D	$H-[(mL)-V-F-V-(mL)]-NH_2$
SEN020	L	Ac-[K-(mL)-V-(mF)-F-(mA)-E]-NH ₂ ^b	SEN263	D	$H-[(mL)-V-F-I-(mL)]-NH_2$
SEN021	L	Ac- $[P-L-V-F-F-A]-NH_2$	SEN264	D	$H-[(mL)-V-F-L-(mL)]-NH_2$
PPI-1019	D	$H_{(mL)} - V - F - F - L - NH_2^{\circ}$	SEN265	D	$H-[(mL)-V-F-M-(mL)]-NH_2$
$iA\beta 5n$	L	AC-[L-P-F-F-A]-NH $_2$ Ac-[L-P-F-F-D]-NH $_2^d$	SEN110 SEN267	D	$H_{I}(mI) - V - F - Y_{-}(mI) - NH_{2}$
$D-iA\beta 5p$	D	Ac-[L-P-F-F-D]-NH ₂	SEN268	D	$H_{[(mL)} V F W_{(mL)]} NH_2$
C1	L	Ac-[S-K-S-G-Y]-NH ₂	SEN269	D	H-[(mL)-V-F-(tbGly)-(mL)]-NH ₂
C2	L	Ac-[Y-G-S-K-S]-NH ₂	SEN270	D	H-[(mL)-V-F-(chGly)-(mL)]-NH ₂
SEN101	D	Ac-[(mL)-V-F-F-L]-NH ₂	SEN271	D	$H-[(mL)-V-F-(chAla)-(mL)]-NH_2$
SEN102	D	Ac-[L-V-(mF)-F-L]-NH ₂	SEN272	D	$H-[(mL)-V-F-(hPhe)-(mL)]-NH_2$
SEN103 SEN104	D	$AC-[L-V-F-F-(ML)]-NH_2$ $Ac-[(mL)-V-(mE)-E-L]-NH_2$	SEN273 SEN280	D	$H_{(mL)-V-F-(2nAla)-(mL)]-NH_2}$ $H_{(mL)-V-F-F-(none)]-NH_2}$
SEN104 SEN105	D	$Ac_{[(IIIL)-V-(IIII)-IV-L]-IVII_2}$ Ac-[L-V-(IIIL)-F-(IIIL)]-NH ₂	SEN280	D	$H_{mL} - V - F_{mL} - NH_{2}$
SEN106	D	$Ac-[(mL)-V-F-F-(mL)]-NH_2$	SEN282	D	$H^{-1}(mL) - V - F - F - V - NH_2$
SEN019	D	Ac- $[(mL)-V-(mF)-F-(mL)]-NH_2$	SEN283	D	H-[(mL)-V-F-F-I]-NH ₂
SEN108	D	Ac-[L-(mV)-F-F-L]-NH ₂	PPI-1019	D	$H-[(mL)-V-F-F-L]-NH_2^c$
SEN109	D	Ac-[L-V-F-(mF)-L]-NH ₂	SEN285	D	H-[(mL)-V-F-F-M]-NH ₂
SEN018 DDI 1010	D	$Ac-[L-(mV)-F-(mF)-L]-NH_2$	SEN289	D	$H-[(mL)-V-F-F-(tbGly)]-NH_2$
SEN112	D	$H_{1}(ML) - V_{1} - L_{1} - L_{1} - M_{12}$ $H_{1} - V_{1}(mF) - F_{1} - L_{1} - M_{12}$	SEN290	D	$H_{mL} - V - F - F_{mL} - C - C - C - C - C - C - C - C - C - $
SEN112	D	$H = [L - V - F - F - (mL)] - NH_2$	SEN292	D	$H_{[(mL)-V-F-F-(hPhe)]-NH_2}$
SEN114	D	$H-[(mL)-V-(mF)-F-L]-NH_2$	SEN293	D	$H-[(mL)-V-F-F-(2nAla)]-NH_2$
SEN115	D	$H-[L-V-(mF)-F-(mL)]-NH_2$	SEN301	D	H-[(chGly)-(chAla)-(chGly)-(chGly)-(mL)]-NH ₂
SEN116	D	$H-[(mL)-V-F-F-(mL)]-NH_2$	SEN302	D	H-[(chAla)-(chGly)-(chGly)-(mL)]-NH ₂
SENII/	D	$H_{(mL)} - V_{(mF)} - F_{(mL)} - NH_2$	SEN303	D	H-[($chAla$)-($chAla$)-($chGly$)-($chGly$)-(mL)]-NH ₂ H [($chGly$) (X) ($chGly$) ($chGly$) (mL)] NH
SEN110	D	$H_{1} - [L_{1} + [mv] - F_{1} - F_{2} - [mv] - F_{1} - [mv] - [$	SEN304	D	$H_{(chGly)-(chAla)-(chGly)-(chGly)-(mL)]-NH_2}$
SEN120	D	$H_{1} = (mV) - F_{1} = (mF) - L_{1} - NH_{2}$	SEN306	D	H-[(chGly)-(chAla)-(chAla)-(chGly)-(mL)]-NH ₂
SEN200	D	H-[(none)-V-F-F-(mL)]-NH ₂	SEN307	D	H-[(chGly)-(chAla)-(chGly)-(chGly)-(mA)]-NH ₂
SEN201	D	$H-[A-V-F-F-(mL)]-NH_2$	SEN313	D	Ac-[V-I-L-(mL)]-NH ₂
SEN202	D	$H-[V-V-F-F-(mL)]-NH_2$	SEN314	D	H-[V-I-L-(mL)]-NH ₂
SEN203	D	$H-[I-V-F-F-(mL)]-NH_2$	SEN315	D	Ac- $[K-V-I-L-(mL)]-NH_2$
SEN115 SEN205	D	$H_{L} = V - F - F - (IIIL) - NH_2$ $H_{L} = M - V - F - F - (mL) - NH_2$	SEN310 SEN317	D	$\Pi - [I - V - I - L - (IIIL)] - I \Pi - I = I = I = I = I = I = I = I = I = I$
SEN206	D	$H = [F - V - F - F - (mL)] - NH_2$	SEN318	D	$H-[I-V-I-L-(mL)]-NH_2$
SEN207	D	H-[Y-V-F-F-(mL)]-NH ₂	SEN319	D	$H-[I-V-L-L-(mL)]-NH_2$
SEN208	D	$H-[W-V-F-F-(mL)]-NH_2$	SEN320	D	H-[I-V-L-I-(mL)]-NH ₂
SEN209	D	$H-[(tbGly)-V-F-F-(mL)]-NH_2$	SEN321	D	$H-[I-V-I-I-(mL)]-NH_2$
SEN210	D	$H_{(chGly)} - V_{F} - F_{(mL)} - NH_2$	SEN322	D	H-[I-V-I-L-(mA)]-NH ₂
SEN211 SEN212	D	$H_{(n)} = H_{2}$ $H_{(n)} = H_{2}$ H_{2}	SEN401 SEN402	D	$H_{(ChOIy)-I}(chOIy)-(chOIy)-L_{1}NH_{2}$ $H_{(Y_{(chOIy)-(chOIy)-I})-NH_{2}$
SEN212	D	$H^{-1}(2nA a)-V-F-F^{-1}(mL)-NH_{2}$	SEN402	D	H-[(chGly)-Y-(chGly)-(chGly)-(chGly)]-NH ₂
SEN221	D	$H_{(mL)}-A-F-F_{(mL)}-NH_2$	SEN404	D	H-[Y-(chGly)-(chGly)-(chGly)]-NH ₂
SEN116	D	$H-[(mL)-V-F-F-(mL)]-NH_2$	SEN405	D	H-[(chGly)-(chGly)-(chGly)-(chGly)-(chGly)]-NH ₂
SEN223	D	$H-[(mL)-I-F-F-(mL)]-NH_2$	SEN406	D	H-[(chGly)-(chGly)-(chGly)-(chGly)]-NH ₂
SEN224	D	$H-[(mL)-L-F-F-(mL)]-NH_2$	SEN407	D	H-[I-V-I-L-L]-NH ₂
SEN225 SEN224	D	$H_{(mL)}-M_{F-F-(mL)}-NH_2$	SEN408 SEN400	D	$H = [V = I - L - L] = NH_2$
SEN220 SEN227	ע	$H_{(mL)} - Y_{F} - F_{(mL)} - NH_{2}$	SEN409 SEN410	ע	11-[1- v -1-1-1]-1N112 H-[V-I-I-I]-NH2
SEN228	D	$H_{(mL)} - W_{F-F-(mL)}$	SEN411	D	H-[(chGly)-(chGly)-(chGlv)]-NH ₂
SEN229	D	H-[(mL)-(tbGly)-F-F-(mL)]-NH ₂			

^{*a*} Abbrevations: *tert*-butyl-Gly (tbGly), cyclohexyl-Gly (chGly), cyclohexyl-Ala (chAla), homo-Phe (hPhe), and 2-naphthyl-Ala (2nAla). ^{*b*} From ref 26. ^{*c*} From ref 32. ^{*d*} From ref 14.



FIGURE 1: Round-1 ThT assay. $A\beta(1-40)$, 100 μ M; inhibitor, 100 μ M. Error = 1 standard deviation; n = 3. $A\beta(1-40)$ (100 μ M) in isolation is scaled to 100%.

which fits with these conclusions, being nonacetylated and having methylated residues at the X1 and X5 positions.

A selection of the best inhibitors were then tested against $A\beta(1-42)$ to determine their effect on this more amyloidogenic peptide (Figure C in the Supporting Information). Amyloid formation was reduced to 52, 51-70, and 62% for SEN101, PPI-1019, and SEN116, respectively. These were smaller reductions than with A β (1-40), confirming that the longer peptide is a more challenging target. Interestingly, $iA\beta 5p$ induces $A\beta(1-42)$ aggregation, shown by the higher ThT fluorescence than that of the target alone. Overall, SEN101 and PPI-1019, both methylated at the N-terminal Leu, as well as SEN116, methylated at both the N- and C-terminal leucines, are the best modulators of $A\beta(1-42)$ aggregation. In contrast to $A\beta(1-40)$, SEN113, methylated only at the C-terminal Leu of the LVFFL domain, did not have high inhibitor activity, suggesting that, when methylating at the fifth (X5) position, inhibition is decreased. Structure/activity relationships indicated that acetylated inhibitors were less potent than their nonacetylated counterparts, methylation at the first (X1) position was only favored when X5 was methylated (as in SEN113 and SEN116). Overall, only methylation of X1 is undoubtedly favorable.

Selected round-1 inhibitors were assayed for their ability to reverse aggregation by $A\beta(1-40)$ and $A\beta(1-42)$ (Figure D in the Supporting Information). Each target was allowed to aggregate for 3 days at 37 °C prior to the addition of 100 μ M of each inhibitor for a subsequent period of 4 days. C1, C2, and iA β 5p could not reverse aggregation of A $\beta(1-40)$, while all of the all-D-peptide inhibitors tested gave about 20-40% reversal (as opposed to 60-80% inhibition at the same inhibitor/target concentrations). PPI-1019 had the highest potency in reversing A $\beta(1-40)$ fibril formation by reducing aggregation to 59%. SEN101, SEN113, and SEN116 were able to reverse A $\beta(1-40)$ aggregation to 65-71%. Only PPI-1019 was able to reverse A $\beta(1-42)$ aggregation.

The round-1 inhibitors were all tested 3 times on the A β affinity column, giving highly reproducible results with a standard deviation of less than 0.2 min. Peptides that were found to bind tightly to the columns did not bind to the CPG glass resin, confirming that the inhibitors do not bind to any other component of the system besides the target peptide attached to the resin. Results are shown in Figure 2. The acetylated peptides bind to the column more tightly than the



FIGURE 2: Round-1 affinity chromatography.

nonacetylated peptides. Within each acetylated or nonacetylated series, however, structure/activity relationships could be inferred that agree well with ThT results. The control peptides, C1 and C2, did not bind at all. $iA\beta 5p$ bound poorly to the A β column, in agreement with ThT results. All of the methylated inhibitors bound to the columns, indicating that methylation increases the binding affinity of the peptides, unlike proline (as in iA β 5p). Specifically, PPI-1019, being methylated at the X1 position only, reported strong binding affinity, thus suggesting that the first residue should be methylated in A β inhibitors. Moreover, SEN113 and SEN116, methylated at X5 and X1/X5, respectively, were tightly bound peptides, correlating with ThT activity, which suggested that methylation of X1 and/or X5 is favorable. These results suggested that the affinity chromatography method could give valuable results, provided that all of the inhibitors are N-methylated and unacetylated, so that they are soluble and do not aggregate.

EM was performed on selected round-1 inhibitors to assess fibril morphology of A $\beta(1-40)$ and A $\beta(1-42)$ in the presence or absence of inhibitors (Figure 3 and Table 2). Samples were prepared from the same sample used for ThT fluorescence spectroscopy to directly compare results. A β -(1-40) is characterized by large fibril networks, composed of long, single, straight, unbranched fibers of varying length and 9-14 nm in diameter. The fibril diameter in the presence of PPI-1019 and SEN113 increases to 14-17 and 8-20 nm, respectively, while remaining unaltered in the presence of SEN116. There is a reduction in the number and length of fibers present with PPI-1019, SEN113, and SEN116. The most obvious morphological change took place in the presence of SEN116, where the fibril length was considerably reduced to 200-500 nm. SEN116 caused changes in fibril association, where two fibrils were laterally associated; a few twisted ribbons were also present with a helical repeat of 100–200 nm. A β (1–42) formed a similar morphology to $A\beta(1-40)$, although the fibril diameter was slightly larger, forming long twisted ribbons with no apparent lateral association. PPI-1019 and SEN116 caused a reduction in fibril formation as well as altering fibril length in $A\beta(1-$ 42), supporting the ThT results.

Round-2 Library. The round-1 results suggested the following preferred features for an A β aggregation inhibitor: derivatives of LVFFL, all D peptide enantiomers, nonacetylated, C-terminally amidated, and single methylations at residues of the first or fifth positions or double methylations at both the first and fifth positions.

To identify the optimal side chain at each position, 13 different residues were substituted at each position in turn,

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FIGURE 3: Electron micrographs. All at 0.1:1 inhibitor/target ratio (10 μ M inhibitor/100 μ M target). (i) (A) A β (1–40), (B) A β (1–40)/ PPI-1019, (C) A β (1–40)/SEN113, (D) A β (1–40)/SEN116, (E) A β (1–42), (F) A β (1–42)/PPI-1019, and (G) A β (1–42)/SEN116. (ii) (A) A β (1–40)/C2, (B) A β (1–40)/SEN200, (C) A β (1–40)/SEN211, (D) A β (1–40)/SEN221, (E) A β (1–40)/SEN231, (F) A β (1–40)/SEN250, (G) A β (1–40)/SEN270, and (H) A β (1–40)/SEN281. (iii) A β (1–40) in the presence of (A) SEN301, (B) SEN 303, (C) SEN304, (D) SEN313, (E) SEN318, (F) SEN319, (G) SEN321, and (H) SEN323.

Table 2: Fibril Morphology of $A\beta(1-40)$ and $A\beta(1-42)$ in the Presence of Selected Inhibitors as Detected by EM

target (100 µM)	inhibitor name	inhibitor concentration (µM)	diameter (nm)	length (nm)	fibril morphology
$\begin{array}{c} A\beta(1{-}40) \\ A\beta(1{-}42) \\ A\beta(1{-}40) \end{array}$	N/A N/A PPI-1019	N/A N/A 100	9-14 10-17 14-17	N/A N/A 600-700	long, single fibers without obvious twists or lateral association long twisted ribbons, no apparent lateral association lateral fibril association composed of two fibers, few twisted ribbons with believel repeat of 100–200 Å
$\begin{array}{c} \mathrm{A}\beta(1{-}40)\\ \mathrm{A}\beta(1{-}40) \end{array}$	SEN113 SEN116	100 100	8-20 7-15	500-700 200-500	straight unbranched fibers lateral fibril association composed of two fibers, single parrower fibers also present, no twisted ribbons
$A\beta(1-42)$	PPI-1019	100	10-14	300-600	twisted ribbons and single fibers present, no apparent lateral association
$A\beta(1-42) \\ A\beta(1-40)$	SEN116 C2	100 10	10-13 8-11	100–250 N/A	unable to distinguish lateral or twisted fibril morphology long, single fibers without obvious twists or lateral association, similar to target alone
A $\beta(1-40)$	SEN200	10	11-18	N/A	long, single fibers without obvious twists or lateral association, similar to target alone
A $\beta(1-40)$	SEN211	10	15-19	300-1150	twisted ribbons usually composed of two-three fibers and few fibers arranged by lateral association
$A\beta(1-40)$	SEN221	10	8-12	N/A	long, single fibers without obvious twists or lateral association,
$A\beta(1-40)$	SEN231	10	12-24	500-950	short, unbranched twisted ribbons composed of two fibers,
$A\beta(1-40)$	SEN250	10	6-8	800-1200	few hoers associating laterary, less norm networks man the while type few but very long, straight twisted ribbons composed of two fibers
$A\beta(1-40)$	SEN270	10	14-19	200-300,	small networks of fibers present, few twisted ribbons composed
$A\beta(1-40)$	SEN281	10	12-17	100 - 600	single twisted fibers forming large networks similar to the wild type, fau straight short fibers present
Α $β$ (1-40)	SEN301	10	8-16	180–190, 600–700,	long, single fibers and twisted ribbons composed of two or more fibers, more than one fibril population present
Α $β$ (1-40)	SEN303	10	11-18	1500-1600 400-500, 800-1200, 1600-1700	long, single fibers with bends and twisted ribbons composed of two fibers
$A\beta(1-40)$	SEN304	10	13-23	200-300	twisted ribbons usually composed of two fibers with helical repeat
$A\beta(1-40)$	SEN313	10	10-20	N/A	long, single fibers without obvious twists or lateral association, similar to target alone
$A\beta(1-40)$	SEN318	10	12-21	900-1100	very long twisted ribbons composed of two fibers,
$A\beta(1-40)$	SEN319	10	18-22	1200-1800	twisted ribbons composed of more than two fibers with helical
$A\beta(1-40)$	SEN321	10	10-22	1800-1900	very long twisted ribbons composed of two fibers with helical
$A\beta(1-40)$	SEN323	10	10-16	N/A	long, single fibers without obvious twists or lateral association, similar to target alone, few large fibril networks present

while taking the remaining four positions from H-[(mL)-V-F-F-(mL)]-NH₂ (SEN116). The side chains tested were A, V, I, L, M, F, Y, W, tert-butyl-Gly (tbGly), cyclohexyl-Gly (chGly), cyclohexyl-Ala (chAla), homo-Phe (hPhe), and 2-naphthyl-Ala (2nAla). Uncharged and nonpolar side chains were chosen because the target region of $A\beta$ is assumed to also be nonpolar. Nonnatural amino acids were included to test many potentially interesting derivatives. The N- and C-terminal residues were also deleted. The entire round-2 library is listed in Table 1 and named SEN200-SEN293, where SEN200-SEN213 are modified at position 1, SEN221-SEN233 are modified at position 2, SEN241-SEN253 are modified at position 3, SEN261-SEN273 are modified at position 4, and SEN270-SEN283 are modified at position 2. The side chains are numbered in the same order within each of the five series; e.g., SEN205, SEN225, SEN245, SEN265, and SEN285 have Met at positions 1-5, respectively. Peptides that had been previously used retained their former names. The control peptides, C1 and C2, $iA\beta 5p$, and the three best peptide inhibitors identified in round 1,

namely, PPI-1019, SEN113, and SEN116, were also included.

With the improved activity of the inhibitors, the inhibitor/ A β stoichiometry was reduced to 0.1:1 (10 μ M inhibitor/ 100 μ M A β) to maximize the range of results. The best round-2 inhibitors now reduced amyloid formation by A β -(1-40) to 50% (Figure 4), showing the best side chains at each position to be (1) (chGly/2nAla/chAla) > (hPhe/Trp/ Leu) > (Val/Ile/Phe); (2) (chAla/Tyr) > (chGly/Phe/Trp/ Leu/2nAla) > (Val/Ile); (3) (chGly/chAla/tbGly) > (2nAla/ Leu/hPhe/Trp) > (Ile/Phe); (4) (chGly/2nAla/Ile) > (chAla/ Leu/tbGly) > (Val/Met); and (5) (Ala/Val) > (tbGly/Met).

All round-2 peptides were tested 3 times on A β -affinity columns, giving data that support the ThT results (Figure 5). In general, the binding affinity of the SEN200–SEN293 peptide libraries increased with the hydrophobicity of the alternative substituted residue, although some smaller and less hydrophobic side chains did increase the binding affinity in certain positions, such as Val at position 2 (SEN222) and Ile at the second, third, and fourth positions (SEN223,



FIGURE 4: Round-2 ThT assay. $A\beta(1-40)$, 100 μ M; inhibitor, 10 μ M. Error = 1 standard deviation; n = 3. $A\beta(1-40)$ (100 μ M) in isolation is scaled to 100%.

SEN243, and SEN263, respectively). These results therefore provided an opportunity to design peptides with increased binding affinity while reducing the molecular weight. It should also be noted that the SEN280-SEN293 library, where the last residue is unmethylated, bound very tightly to the affinity column. This could be due to the aggregation of the inhibitors on the affinity column because of nonmethylation of the last residue. Figure E in the Supporting Information shows good correlations between affinity binding and ThT results for each of the SEN200-SEN273 series but no correlation for the SEN280-SEN293 series.

Electron micrographs showing A $\beta(1-40)$ in the presence of selected round-2 inhibitors are shown in Figure 3 and

summarized in Table 2. SEN211, SEN231, SEN250, SEN270, and SEN281 were chosen to study because they were identified as among the most active inhibitors against A β -(1-40) aggregation by ThT. SEN201 and SEN221 were chosen because of their inability to inhibit amyloid formation by A β (1-40). C2, SEN200, and SEN221 had no effect on fibril formation. In the presence of the active selected round-2 inhibitors, a clear change in fibril morphology was detected with a reduction in the fibril length, while the number of large fibril networks throughout the EM grid was considerably reduced. In general, the fibril diameter ranged from 12 to 18 nm, although there were cases where the fibril diameter was either increased or reduced, depending upon the inhibitor



FIGURE 5: Round-2 affinity chromatography.

effect. For example, SEN250 caused the most obvious morphology change, where the fibril diameter was reduced to 6–8 nm, while the fibril length was relatively longer than other inhibitor effects (up to 1200 nm). SEN231, however, which was also found to have a similar inhibitory effect to SEN250 against $A\beta(1-40)$ (by ThT), caused a different morphology change: the fibril diameter was thicker than $A\beta$ -(1–40) alone, reaching 24 nm in diameter, while the fibril length was reduced to 500–950 nm. Moreover, inhibitors appeared to induce fibril association, where a small number of twisted ribbons were formed (as opposed to single, long fibers forming complex fibril networks by the amyloidogenic target), usually composed of two–three fibers with a helical repeat of 200 nm.

Round-3 Library. A total of 18 new inhibitors, combining the best side chains, were designed, synthesized, and studied (SEN301-SEN324; Table 1). All peptides were based on combinations of the best side chains at each position, with one or two alternative side chains substituted at each position. Larger aromatic side chains (Trp and 2nAla) were avoided, where possible, to maintain a lower molecular weight and avoid potential oxidation in vivo. N-Terminal truncations of combined sequences were also included to determine whether the weight can be reduced without the loss of activity. The round-3 peptide inhibitors were compared directly to the standard control peptides, namely, C1, C2, PPI-1019, and $iA\beta 5p$, as well as the best compounds from round 1 (PPI-1019, SEN113, and SEN116) and the most active inhibitors from round 2 (SEN211, SEN231, SEN250, SEN271, SEN208, and SEN228).

ThT results are shown in Figure 6. The inhibitor/A β stoichiometry was again 0.1:1 [10 μ M inhibitor/100 μ M A β -(1-40)]. SEN304 D-(H-[(chGly)-(Tyr)-(chGly)-(chGly)-(mLeu)]-NH₂) was the most active compound, being able to inhibit A β (1-40) aggregation to 43%. Figure 7 shows A β -affinity chromatography results. SEN301 had the strongest binding affinity to the A β column. SEN302, SEN305, and SEN306 also bound tightly to the A β column. SEN304 bound relatively tightly to the A β column, although not as tightly as SEN301. When the N-terminal residue was missing or polar, as in SEN314, SEN315, and SEN316, there was no binding. The correlation between the ThT and affinity results was weak within this library.

Figure 3 presents the fibril morphology of a selection of the most active round-3 inhibitors in the presence of A β -(1-40), namely, SEN303, SEN304, SEN318, SEN319, and



FIGURE 6: Round-3 ThT assay. $A\beta(1-42)$, 10 μ M; inhibitor, 5 μ M. Error = 1 standard deviation; n = 3. $A\beta(1-42)$ (10 μ M) in isolation is scaled to 100%.



FIGURE 7: Round-3 affinity chromatography.

SEN321, as well as three of the least active inhibitor peptides against A $\beta(1-40)$ aggregation, which include SEN301, SEN313, and SEN323. Fibers formed by inhibitor/target mixtures were of varying length and morphology. Specifically, the peptides that had been identified as poor inhibitors of amyloid formation by A β (1–40) (SEN301, SEN313, and SEN323), all exhibited a similar morphology, which was closely related to the target alone. They were characterized by the formation of large fibril networks throughout the EM grid, composed of very long single fibers with a relatively small diameter of 8–16 nm. SEN301/A β (1–40) also formed more than one fibril population in terms of length, ranging from 180 to 1600 nm. A selection of the best inhibitors of $A\beta(1-40)$ fibrillogenesis, which included SEN303, SEN318, SEN319, and SEN321, were characterized by the formation of thicker fibers with the average diameter ranging from 10 to 22 nm and the formation of straight but twisted ribbons usually composed of two-three fibers with a helical repeat of 100-120 or even 350 nm (as was the case with SEN319 and SEN321). It was evident, however, that the presence of the inhibitors not only changed the morphology of the fibers but also reduced the abundance of fibrils in solution, and consequently, only a few and small clustered fibril aggregates could be identified. The effect of the good inhibitors on fibril length varied. For example, SEN303 exhibited more than one fibril length, while SEN318, SEN319, and SEN321 formed long fibers that were shorter than the A β alone (because they could be measured) but were still longer than the average length of fibril formation in inhibitor/target mixtures. The most striking evidence of inhibitor activity

against $A\beta(1-40)$ aggregation was the effect of SEN304. SEN304/ $A\beta(1-40)$ formed very short (200-300 nm) twisted ribbons composed of two fibers with a helical repeat of 100 nm and a diameter length of 13-23 nm. The most obvious effect on inhibition was the absence of large fibril networks or any formation of clustered aggregates (as detected by all of the previously mentioned mixtures). Instead, it was characterized by small isolated fibers that were present throughout the EM grid (and thus throughout the solution) that could only be detected at high magnification. These isolated fibers may have contributed to the 40% amyloid formation observed by ThT. These results support the ThT assay, which indicated that SEN304 was the most potent inhibitor of $A\beta(1-40)$ aggregation.

Round-4 Library. The round-4 peptides (SEN401– SEN411) were all nonmethylated, to test the hypothesis that N-methylation is useful for inhibitor activity. SEN401 is nonmethylated SEN304. SEN402–SEN406 and SEN411 contained multiple chGly residues to test whether the activity in the best round-3 inhibitors was a nonspecific property of this side chain. SEN407–SEN410 were nonmethylated derivatives of promising compounds from round 3 that were based on Val, Ile, and Leu side chains.

All of the round-4 peptides proved to be very difficult to work with, because of the low solubility in aqueous solvents, precluding the collection of ThT data. Because it was often not possible to achieve an adequate concentration of peptide, it was difficult to assess their efficacy. In the affinity assay, SEN402 eluted at 17.49 min, while SEN404 did not bind at all; affinity data on the other round-4 peptides could not be acquired.

MTT Toxicity Assay. Figure 8 shows results from the MTT toxicity assay using A $\beta(1-42)$ and selected inhibitors. The assay was performed with 10 μ M A β (1-42) and 10 μ M inhibitor (Figure 8A) or 5 μ M inhibitor (Figure 8B). The MTT reduction with MTT alone was 25-30%. None of the inhibitors studied here increased toxicity, while many partially reversed A β toxicity, shown by an increase in the percentage of MTT reduction. In general, the MTT results agreed well with those from other assays. The control peptides C1 and C2 had no effect nor did iA β 5p. PPI-1019 successfully reduced A β toxicity. Some round-1 inhibitors had a weak beneficial effect, which was improved by some round-2 inhibitors. Round-3 inhibitors were the most effective, especially SEN303 and SEN304. Data for the 400 series of non-N-methylated peptides show that most have some toxicity in isolation, in contrast to all of the N-methylated peptides, which were always nontoxic (Figure F in the Supporting Information). SEN401 is the non-N-methylated analogue of SEN304, and it was similarly active, despite its toxicity.

DISCUSSION

Our initial inhibitor was a core amyloidogenic fragment from the full-length $A\beta$, namely, L-Ac-[KLVFF]-NH₂. After four rounds of optimization, we reached a series of compounds with much higher activity, with SEN304 (D-[H-[(chGly)-(Tyr)-(chGly)-(chGly)-(mLeu)]-NH₂]) being the most effective inhibitor. Every side chain was thus changed from the starting compound, and the chirality was inverted. It is not clear why the D peptide should be more effective



FIGURE 8: MTT toxicity. (A) Screened at 1:1 [10 μ M inhibitors versus 10 μ M A β (1-42)]. (B) Screened at 0.5:1 [5 μ M inhibitors versus 10 μ M A β (1-42)].

against an L peptide target, but similar results have been reported by other groups (33). This may be a general result, because Dzwolak et al. reported that mixtures of L- and D-polylysine formed more stable β sheets than the pure L or D peptides (34). The 400 series of inhibitors were designed to probe the effect of N-methylation on activity. They proved to be very difficult to work with, however, because of the low solubility, and were often toxic. We can conclude that a single N-methyl group greatly increases aqueous solubility, in agreement with previous work (27), and reduces inhibitor toxicity, even if its effect on inhibitor efficacy is difficult to assess. Only a single N-methyl group is necessary, and there was little or no gain in activity from multiple N-methylation.

Compounds were optimized primarily using the affinity and ThT data. These results correlated well with reducing A β toxicity, as measured by the MTT assay. The affinity assay gave data that correlated with ThT binding within a series of similar compounds, such as when just one side chain was altered. When large multiple changes were made, such as acetylation, changing chirality, and changing side chains, the results were more difficult to interpret. While the affinity assay is attractive, because dozens of compounds can be accurately measured in a day without using any A β , its results need to be interpreted cautiously. EM was less useful, because while some inhibitors did reduce or abolish fibril formation, others altered fibril morphology. How these EM data relate to toxicity is unclear. Our primary toxicity assay was MTT with PC12 cells, because it gave reproducible data at low concentrations and is widely used to assay A β toxicity. We verified it by a comparison with other toxicity assays and a different cell line. It was reassuring, however, that the results from all of the assays correlated well and we were able to achieve significant increases in efficacy with each round. In addition, while the compounds were optimized primarily using the affinity and ThT data, these results paralleled with reducing $A\beta$ toxicity, as measured by the MTT assay.

The best inhibitors never gave less than a 40% signal with the ThT assay, even if toxicity was reduced much below this in cell-based assays, while EM often showed a change in fibril morphology, rather than complete inhibition of fibril formation. We suggest that many inhibitors, including notably SEN304, may act by causing toxic A β oligomers to aggregate into nontoxic fibrils that weakly bind ThT. One concern at the start of this project was that if inhibitors compete for binding to aggregating A β with the wild-type peptide, they may increase the oligomer population, thus increasing toxicity. One could imagine that inhibitors could bind to both sides of a growing β sheet, blocking further aggregation and keeping the $A\beta$ in an oligometric form. Indeed, any interference with A β aggregation could increase the amount in shorter, more toxic forms. Reassuringly, no significant increase in toxicity was found with any Nmethylated peptide inhibitor. Toxicity problems were only encountered with the rare cases when the inhibitors were toxic in isolation, as shown within the non-N-methylated round-4 library. Further work is clearly needed on the mode of action of the inhibitors.

It was also unclear whether inhibitors should be targeted at $A\beta(1-40)$ or $A\beta(1-42)$. While $A\beta(1-40)$ is 10 times more abundant than $A\beta(1-42)$ in vivo, $A\beta(1-42)$ is more toxic. We addressed this by often assaying inhibitors against both target peptides. The results correlated very well, suggesting that optimizing the inhibitor structure against either target is acceptable. This may be because the binding site for an inhibitor is common to both target peptides.

In conclusion, we have found that N-methylated peptides can be powerful inhibitors of $A\beta$ toxicity in vitro. Our best compounds had a higher activity than previously reported peptides, although D-[(mL)VFFL]-NH₂ (29) was also effective. We found no activity for Ac-[LPFFD]-NH₂ (15), with repeated measurements using a range of assays, despite previous reports. This may be because incorporating a Pro group prevents the adoption of the β conformation required for binding.

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SUPPORTING INFORMATION AVAILABLE

Figure A, aggregation kinetics of $A\beta(1-40)$ and $A\beta(1-42)$ with and without selected round-1 inhibitors; Figure B, initial round ThT assay; Figure C, round-1 ThT assay; Figure D, $A\beta(1-40)$ versus $A\beta(1-42)$ reversal; Figure E, $A\beta(1-40)$ inhibition by ThT versus $A\beta$ -binding affinity; Figure F, MTT toxicity assay for round-4 inhibitors. This material is available free of charge via the Internet at http://pubs.acs.org.

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