Research Article

Modulation of Gamma-Secretase for the Treatment of Alzheimer’s Disease

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The Amyloid Hypothesis states that the cascade of events associated with Alzheimer’s disease (AD)—formation of amyloid plaques, neurofibrillary tangles, synaptic loss, neurodegeneration, and cognitive decline—are triggered by Aβ peptide dysregulation (Kakuda et al., 2006, Sato et al., 2003, Qi-Takahara et al., 2005). Since γ-secretase is critical for Aβ production, many in the biopharmaceutical community focused on γ-secretase as a target for therapeutic approaches for Alzheimer’s disease. However, pharmacological approaches to control γ-secretase activity are challenging because the enzyme has multiple, physiologically critical protein substrates. To lower amyloidogenic Aβ peptides without affecting other γ-secretase substrates, the epsilon (ε) cleavage that is essential for the activity of many substrates must be preserved. Small molecule modulators of γ-secretase activity have been discovered that spare the ε cleavage of APP and other substrates while decreasing the production of Aβ42. Multiple chemical classes of γ-secretase modulators have been identified which differ in the pattern of Aβ peptides produced. Ideally, modulators will allow the ε cleavage of all substrates while shifting APP cleavage from Aβ42 and other highly amyloidogenic Aβ peptides to shorter and less neurotoxic forms of the peptides without altering the total Aβ pool. Here, we compare chemically distinct modulators for effects on APP processing and in vivo activity.

1. Introduction

Gamma-secretase (γ-secretase) is required for the production of amyloid beta peptides (Aβ) and decreasing Aβ production as a disease modifying approach for the treatment of Alzheimer’s disease (AD) has received intense interest. The initial focus was on the discovery of compounds that would decrease γ-secretase activity. γ-Secretase cleaves the membrane bound C-terminal domain (C99) of APP at the ε site to produce the intracellular domain, AICD. The enzyme then makes sequential cuts of the remaining intramembrane APP fragment at each turn of the alpha helix (every 3–4 amino acids) until Aβ peptides are formed and released into the extracellular space [1–3]. This protein processivity produces Aβ peptides that vary in size, from 43–34 amino acids [4, 5]. In Alzheimer’s disease, a greater number of the longer forms of Aβ, including Aβ42 and Aβ43, or a high ratio of the long peptides to the shorter forms, appear to occur [6]. These longer Aβ peptides readily oligomerize, forming toxic species, as well as becoming the seeds for amyloid plaques [7, 8].

The full inhibition of γ-secretase appeared to be a sound approach. However, it was found that γ-secretase plays a broader biological role and cleaves multiple proteins to yield physiologically essential products. Thus, total inhibition results in severe adverse effects in vivo [9–11]. This played out in the clinic in the trial of the γ-secretase inhibitor, semagacestat from Eli Lilly [12–14]. Patients treated with this drug developed skin and gastrointestinal side effects that are characteristic of the inhibition of γ-secretase processing of Notch, leading to the discontinuation of the clinical trial in 2010 [13, 14].

The discovery of compounds that could decrease the production of the more amyloidogenic Aβ42 peptide while preserving total Aβ levels and γ-secretase cleavage of other substrates led to a clinical trial of one of these newly
identified, first generation gamma-secretase modulators (GSMs) [15]. The NSAID-derived, Flurizan from Myriad Genetics was tested in a Phase 3 trial in mild to moderate AD patients. However, Flurizan is a very weak modulator of γ-secretase, with an IC₅₀ of ~250 μM [16]. In addition, this compound has very poor distribution into the central nervous system [16, 17]. Not surprisingly, the combination of these less than desirable properties resulted in no exposure in the brain, and a failed Phase III trial in June 2008 [18].

Second and now third generation GSMs have been discovered and are proceeding toward clinical trials in AD. These GSMs are significantly more potent than Flurizan and appear to have better drug-like properties. However, the majority of these compounds fall into one of two chemical classes, with little structural diversity within each of these classes and the development of some has been discontinued because of toxicities and biopharmaceutical limitations that other class members may also share [19]. A structurally unique GSM derived from a core molecule isolated from a natural product is also moving through preclinical testing [20]. While all three chemical series of GSMs share some common pharmacological properties, they differ in other fundamental ways. Here we present data contrasting the pharmacology of members of several structural series of GSMs and how modulation may avoid the pitfalls associated with γ-secretase inhibitors (GSIs).

2. Materials and Methods

2.1. Test Compounds. LY411575, GSI-953, BMS-708163, MK-GSM1, JNJ-40418677, and E-2012 were prepared according to published methods. SPI-1802 and SPI-1810 were prepared at Satori Pharmaceuticals.

2.2. Cell Culture and Compound Treatment. SUP-T1 cells (ATCC) were cultured in T75 flasks in RPMI media (Mediatech 10-041-CV) supplemented with 10% FBS and penicillin/streptomycin at 37°C in a 5% CO₂ atmosphere. One hour prior to drug treatment, six well plates were seeded with 1.5 mL of media containing 2% FBS and cells at a density of 1.5 × 10⁶ cells/mL. Test compounds in DMSO were diluted 100-fold directly into the media with the cells and incubated for 18 hours at 37°C. After treatment 100 μL aliquots of treated cells were assayed for viability with the Promega Cell Titer Glo assay system. The conditioned media was added and cells were further processed to measure Aβ levels and NICD levels, respectively.

CHO-2B7 cells (Mayo Clinic) are Chinese hamster ovary cells stably transfected with human βAPP 695 wt [38, 39]. The cells were cultured in Ham’s F12 media (Thermo Fisher SH30026.01) supplemented with 10% FBS, 0.25 mg/mL Zeocin and penicillin/streptomycin at 37°C in a 5% CO₂ atmosphere. For compound treatment, cells were plated in 96-well plates at a density of 1.0 × 10⁴ cells/mL and allowed to grow to 100% confluence over two days. Test compounds in DMSO were diluted 100-fold directly into the media before adding to the cells. Immediately prior to adding compound-containing media to the cells, they were washed once with 1XPBS. Conditioned media from CHO-2B7 cells were collected after 5 hours of treatment and the levels of Aβ peptides were assessed as described below.

H4 human neuroglioma cells (ATCC) were cultured in 10% FBS/DMEM (Media Tech) with Pen/Strep (50 units/50 μg/mL; Invitrogen). Human WT APP stably transfected CHO cells were cultured in 10% FBS/HAM’S F-12 growth media (Media Tech) supplemented with Pen/Strep and G418 (500 μg/mL; Promega). Cells were plated and grown to confluency in 96-well plates prior to dosing. Cells were washed with PBS and 100 μL of media containing DMSO alone (vehicle) or test compounds in DMSO at a final DMSO concentration of 1% (v/v). Conditioned media was collected after 18 hours of treatment and diluted 1:1 with MSD blocking buffer (1% BSA in MSD wash buffer).

2.3. Solid Phase Extraction. Wells of 30 mg Oasis HLB 96-well extraction plates (Waters Corporation) were activated by addition of 1 mL of methanol followed by rinsing with 1 mL of water utilizing a vacuum plate manifold. 1 mL of SUP-T1 conditioned media was added and wells were then washed sequentially with 2 mL of 10% methanol and then with 2 mL of 30% methanol. Samples were eluted into sample collection tubes by adding 250 μL of 90% methanol with 2% ammonium hydroxide to each well. Eluted samples were concentrated to dryness under vacuum without heating.

2.4. Aβ In Vitro Assay Measurement. Conditioned media was collected after 5–18 hours of treatment and diluted with 1 volume of MSD blocking buffer (1% BSA in MSD wash buffer). Alternatively, dried films of SUP-T1 conditioned media after solid phase extraction were resuspended with 1 volume of MSD blocking buffer (1% BSA in MSD wash buffer). Samples were transferred to blocked MSD Human (6E10) Aβ 3-Plex plates and incubated for 2 hours at room temperature with orbital shaking followed by washing and reading according to the manufacturer’s instructions (SEC-TOR Imager 2400 Meso Scale Discovery, Gaithersburg MD).

2.5. NICD Assay. The remaining cells were washed twice in PBS and then lysed with Promega reporter lysis buffer containing a complete protease inhibitor cocktail (Roche) for 1 hour at 4°C. Lysates were spun at 5,000 RPM for 5 minutes and supernatants were collected. Total protein levels were measured and adjusted to 1-2 mg/mL total protein using the BCA total protein assay (Thermo Scientific). NICD levels were then measured with a cleavage specific Notch1 sandwich ELISA (Cell Signaling Technologies) according to the manufacturer’s instructions.

2.6. Immunoprecipitation and Matrix-Assisted Laser Desorption/Ionization Time-of-Flight (MALDI-TOF) Mass Spectrometry. Chinese Hamster Ovary cells stably transfected with wild-type human APP were treated for 6 hrs with γ-secretase modulators at an approximate concentration of 10-fold the IC₅₀ (Merck GSM1 at 1 μM, JNJ-40418677 at 2 μM, SPI-1802 at 3 μM, and SPI-1810 at 2 μM). Monoclonal Aβ antibodies 6E10 (specific for amino acids 1–16 of Aβ) and 4G8 (specific for amino acids 17–24 of Aβ; Covance, Dedham, MA) were immobilized with agarose resin using the
AminoLink Plus reagents (Thermo Scientific, Rockport, IL). Conditioned media from treated cells was precleared with agaro-se resin overnight, and the supernatant was incubated with agaro-conjugated 6E10/4G8 for 6 hrs. Immunoprecipitates were washed extensively prior to analysis.

Analyses were performed on a Shimadzu Biotech Axima TOF2 (Shimadzu Instruments) matrix-assisted-laser desorption/ionization time-of-flight (MALDI-TOF) mass spectrometer. Peptides were analyzed in positive ion linear mode. For intact peptide mass measurement the instrument was set with a mass range extending up to 6000 m/z using a pulsed extraction setting of 3500. An average mass external standard was used which consisted of angiotensin II (1047.2), P14R (1534.86) and ACTH clip 18–39 (2465.20), insulin B (3496.67), and insulin (5734.51). For sample preparation, 5 μL aliquots of Aβ containing immunoprecipitates were diluted with 10 μL of 0.1% TFA and then desalted using a C18 Zip Tip (Millipore, Corp.). Samples were directly deposited from the Zip Tip onto the MALDI sample target and then mixed with 0.5 μL of matrix solution which consisted of 5 mg/mL of alpha cyano-4-hydroxy cinnamic acid in acetonitrile: 0.1% TFA (50:50). Data was acquired manually using a set laser power and averaging 1500–2000 laser shots.

2.7. In Vivo Study Methods. All animal handling and procedures were conducted in full compliance to AAALAC International and NIH regulations and guidelines regarding animal care and welfare.

Either transgenic mice (Tg2576, 3 mos; N = 21) or wild-type Sprague Dawley rats (200–225 g; N = 8) were utilized to assess in vivo efficacy. All animals were acclimated to the test facility for a minimum of two days prior to initiation of the study. Compounds were dosed orally in 10:20:70 Ethanol/Solutol/Water via oral gavage. Samples were harvested at 6 hrs after dose for Aβ and compound exposure levels. Blood samples were collected into K2EDTA and stored on wet ice until processed to plasma by centrifugation (3500 rpm at 5 °C) within 30 minutes of collection. Each brain was dissected into three parts: left and right hemispheres and cerebellum. Brain tissues were rinsed with ice cold phosphate buffered saline (without Mg²⁺ or Ca²⁺), blotted dry and weighed. Plasma and cerebella were analyzed for parent drug via LC/MS/MS. Parent drug levels were compared to a standard curve to establish the unknown levels.

2.8. Rodent Aβ Determination. This protocol is a modification of protocols described by Lanz et al. [40] and Rogers et al. [41]. Frozen hemispheres were weighed into tared homogenization tubes (MP Biomedicals#6933050 for rat; MP Biomedicals, Solon, OH) and (Simport#T501-4AT; Simport, Beloeil, Qc, Canada) containing one 5 mm stainless steel bead (Qiagen#69989) for mouse). For every gram of brain, 10 mLs of 6 M guanidine hydrochloride (wild-type rat) or 0.2% diethyl amine in 50 mM NaCl (transgenic mouse) was added to the brain-containing tubes on wet ice. Rat hemispheres were homogenized for one minute and mouse hemispheres were homogenized for 30 seconds at the 6.5 setting using the FastPrep-24 Tissue and Cell homogenizer (MP Biomedicals#116004500). Homogenates were rocked for two hours at 4 °C, then precleared by ultracentrifugation at 100,000 × g for one hour at 4 °C. Precleared wild-type rat homogenates were concentrated over solid phase extraction (SPE) columns (Oasis HLB 96-well SPE plate 30 um, Waters®WAT058951; Waters Corp., Milford, MA). Briefly, SPE columns were prepared by wetting with 1 mL of 100% methanol followed by dH₂O using vacuum to pull liquids through. Brain homogenates were then added to the prepared columns (1.0 mL from rat). Columns were washed twice with 10% methanol followed by two washes with 30% methanol. Labeled eluent collection tubes (Costar cluster tubes #4413; Corning Inc., Corning, NY) were placed under SPE columns and samples were eluted under very mild vacuum with 300 μL of 2% NH₄OH/90% methanol. Eluents were dried to films under vacuum with no heat in a speed vacuum microcentrifuge. Films were resuspended in 150 μL of Meso Scale Discovery (MSD, Gaithersburg MD) blocking buffer (1% BSA in MSD wash buffer) for one hour at room temperature with occasional vortexing. A volume of 45 μL of precleared transgenic mouse brain homogenates were diluted into 450 μL of blocking buffer and were neutralized with 5 μL of 0.5 M Tris pH 6.8. For Aβ38, 40, and 42 measurements, MSD 96 well multipot Human/Rodent (4G8) Aβ triplex ultrasensitive ELISA plates were blocked with MSD blocking buffer for 1 hour at room temperature with orbital shaking. A volume of 25 μL of neat resuspended wild-type rat brain homogenate films or diluted transgenic mouse brain homogenates were added in duplicates to the blocked 3-plex Aβ MSD plates with SULFO-TAG 4G8 antibody (MSD). The Aβ 3-plex plates were incubated for 2 hours at room temperature with orbital shaking followed by washing and reading according to the manufacturer’s instructions (SECTOR Imager 2400, MSD). The average Aβ concentrations from duplicate measurements of each animal were converted to percent vehicle values and the treatment group averages were statistically compared by ANOVA analysis.

3. Results and Discussion

γ-Secretase is a complex enzyme with multiple substrates and multiple cleavage sites on at least some of these substrates, including APP. Complete inhibition of γ-secretase activity by targeting the ε cleavage site prevents the processing of multiple physiologically relevant proteins, leading to the severe side effects reported in AD patients [12, 13]. On the other hand, chemically modulating the enzyme with a GSM is a more precise mechanism to enhance certain cleavage events while preventing the cut that yields the amyloidogenic peptide, Aβ42, which is linked to the pathophysiological initiation of AD (Figure 6). In preclinical toxicological testing, GSIs appear to be free of the mechanism-based toxicities attributed to the inhibition of Notch processing that have plagued the enzyme inhibitors. In vitro data demonstrate that Notch cleavage to NICD is not inhibited by any of several represented GSIs at concentrations that do not disrupt cell viability (Figure 1). Treatment with GSIs in rodents have not shown the classical Notch-related toxicities that are associated with GSIs, suggesting that the complete processing of Notch to NICD can occur in the presence of GSIs.
3.1. GSMs Do Not Show a Potency Shift with Changes in Substrate Concentration. γ-Secretase inhibitors can show a significant shift in potency in APP transfected cell-based assays depending on the level of expression of APP. Table 1 compares the IC$_{50}$ for Δβ$_{42}$ lowering of different GSIs in multiple cell lines with varying levels of APP expression. In the highest expressing cell lines, the GSIs assayed here appear more potent (lower IC$_{50}$s in the high expressing, transfected CHO-2B7 cell line) than when the same compounds are tested in assays using lower expressing cell lines (native H4 cell line). This potency shift seen with GSIs has been attributed to a shift in the enzyme/substrate ratio [42], where there is a higher ratio in wild-type cells versus those that overexpress APP. This higher ratio in turn requires more compounds to inhibit the enzymatic reaction. Interestingly, the potency shift due to substrate concentration that occurs with GSIs does not occur with GSMs. As seen in Table 1, the IC$_{50}$ for all the GSMs shown are consistent across cell lines, regardless of the APP expression levels. While this potency shift is not well understood, these data do suggest that GSIs and GSMs affect the enzyme in fundamentally different ways.

3.2. Structural Diversity of GSMs. Many of the second generation GSMs were inspired by the NSAIDs (Figure 2). For instance, all compounds shown in the figure were initially derived from the aryl acetic acid motif found in 1 and similar NSAIDs [21]. The similarity between these compounds makes for a crowded patent landscape, with some compounds potentially covered in multiple patent applications from different sponsors. In addition, for this structural series, the physicochemical properties indicate that all of these compounds carry an increased safety risk due to the high logP, low PSA [43–45], and high degree of aromaticity [46, 47]. Compounds 7–9 are designed to improve upon these properties, but are still considered high risk due to these same physicochemical based in silico models [27–29]. In vivo preclinical toxicity testing will ultimately be needed to assess the safety profile of these similar structures. However, the lack of progression of compounds from this class into and through clinical development suggests that this scaffold may have challenges that will continue to slow or prevent successful conduct of the studies required for registration.

A second class of GSMs that has received significant attention is summarized in Figure 3. Following initial disclosure of 10 and 11, a number of pharmaceutical companies pursued structurally related chemical series [30, 48]. Although this class offers clear distinction from the NSAID-inspired compounds above, their physicochemical properties also reside outside of the molecular space most frequently affiliated with marketed agents for oral therapy. For example, the number of aromatic rings and logP of the representatives shown are higher than the average for oral compounds on the market [46, 47], leading to a higher probability of safety and biopharmaceutical challenges with this class. Although some groups have been successful in developing promising structural alternatives, as exemplified by 16–18, little has been reported on the development of any representatives from this general scaffold [34–36].

A novel and structurally distinct chemical architecture of a third class of GSMs has been reported by Satori Pharmaceuticals. This scaffold was first isolated from black cohosh, leading to the characterization of initial hit 19 (Figure 4) [20]. A combination of synthetic and medicinal chemistry optimization led to 20, which is reported to have better drug properties than 19 [20]. To date, the compounds reported by Satori also fall outside of the guidelines most typically associated with good in vivo disposition. The group notes, however, that the majority of marketed agents derived from natural products also violate these same guidelines, a trend that has led some to conclude that molecules derived via

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**Figure 1:** Multiple GSI and GSMs were examined in for their ability to inhibit NOTCH cleavage using the SUP-T1 cellular assay.
Table 1: GSMs do not show a potency shift with APP overexpression.

| Cell line | H4 | H4-APP | CHO-SW | CHO-7W | CHO-2B7 |
|-----------|----|--------|--------|--------|--------|
| Aβ42 Levels (pg/mL) | 20 | 110 | 131 | 834 | 200 |
| Inhibitors | | | | | |
| LY411575 | 1.40 | 1.20 | 0.20 | ND | 0.05 |
| GSI-953 | 706 | 52 | 5.20 | 8.70 | 2.50 |
| BMS-708163 | 40 | 8 | 0.98 | 1.10 | 0.20 |
| Modulators | | | | | |
| GSM1 | 54 | 64 | 154 | 73 | 62 |
| JNJ-40418677 | 115 | 133 | 190 | 122 | 172 |
| E-2012 | 42 | 84 | 54 | 36 | 33 |

Figure 2: Representative NSAID-inspired GSMs. Compounds 1–6 are aryl acetic acids [21–26], compounds 7 and 8 are piperidine acetic acids [27, 28] and compound 9 is a cyclohexane acetic acid [29].

3.3. Aβ Peptide Profiling of Structurally Diverse GSMs. When the NSAID-type of GSMs was first described (Figure 2), the changes in Aβ peptides that were seen, a decrease in Aβ42, an increase in Aβ38 and little or no change on Aβ40 or total Aβ were labeled the “modulator profile,” nomenclature that was reinforced by the pharmacology reported for the GSMs represented by the structures in Figure 3. However, since that time, additional Aβ peptide profiles have been reported for the chemically distinct scaffold disclosed by Satori Pharmaceuticals, as well as molecules more closely
related to those in Figure 3. Examples of the variety of Aβ peptide profiles produced by GSMs are shown in Figure 6. Merck GSM1 and JNJ-40418677 lower Aβ42 while increasing Aβ38, with little or no effect on Aβ40. Alternatively, the Satori Pharmaceutical compounds, SPI-1802 and SPI-1810 (structures shown in Figure 5) decrease both Aβ42 and Aβ38, but maintain total Aβ levels by increasing Aβ37 and Aβ39. Yet, all of these compounds can be classified as “gamma secretase modulators” based on the commonality of sparing the ε cleavage of C99 and other substrates (e.g., Notch), decreasing Aβ42, and not affecting total Aβ levels.

Based on these data, modulation of γ-secretase is more accurately defined as a shift of the Aβ pool to shorter, but variable length, Aβ peptides. A physiological role for Aβ peptides has not been discovered, but in vitro studies have demonstrated that shorter peptides are incapable of aggregation and oligomerization [19] and may even prevent the oligomerization of Aβ42 by binding to it, suggesting that preserving the total pool of Aβ may be beneficial.

The molecular mechanism by which GSMs modulate γ-secretase activity is not completely understood. The variety of Aβ peptide profiles that result from treatment of cells

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**Figure 3:** Representative aryl imidazole inspired GSMs [30–33]. Compounds 16–18 are the most unique because the aryl imidazole has been replaced by a bioisostere [34–36].

| Compound | Source | IC50 Aβ42 (nM) |
|----------|--------|----------------|
| 10       | Neurogenetics | 200–1000 |
| 11       | Eisai, E-2012 | 92 |
| 12       | Torreyipes | 5 |
| 13       | Roche | 210 |
| 14       | Bristol-Meyers Squibb | 5 |
| 15       | AstraZeneca | 22 |
| 16       | Amgen | 131 |
| 17       | Janssen | 9 |
| 18       | Merck | 255 |

**Figure 4:** Two examples of Satori GSMs.

19: Satori screening hit
IC50 Aβ42 = 100 nM

20: Merck
IC50 Aβ42 = 91 nM
3.4. In Vivo Activity of GSMs. In vivo, GSMs exhibit the same \( \beta \)-profile as observed in vitro with reductions in \( \beta \) and increases or decreases of other \( \beta \) and no effect on total \( \beta \) levels. However, the amount of compound required to generate these effects has been surprisingly high when compared to the in vitro potency of the molecules. For example, the Eisai GSM E-2012 has an in vitro IC\(_{50}\) of 33 nM in a cell-based assay. In vivo, 4.9 \( \mu \)M plasma concentrations of the compound were required to reduce brain \( \beta \) by 25\%. Similarly, Merck GSM1, JNJ-40418677, and SPI-1810 also require high plasma levels in order to achieve 25\% lowering (Table 2), a factor that does not appear to be readily attributable to plasma protein binding. These high compound levels are not required for in vivo efficacy with GSIs which further suggests that the interaction with \( \gamma \)-secretase by a GSM is different from that of a GSI. Why high concentrations of some GSMs are required to modulate \( \gamma \)-secretase activity in vivo is not completely understood. Less than ideal pharmacokinetic properties, such as low free fraction or poor blood brain barrier permeability may contribute to the need for high plasma concentrations of GSMs to see efficacy in the brain in vivo. In addition, there are data indicating that GSMs can bind to both active and inactive forms of presenilin since the binding site of the modulators is available even prior to endoproteolysis that creates the active form of \( \gamma \)-secretase [51, 52]. Conversely, data indicate that GSIs require complex formation prior to binding [53], which may mean more binding sites are available for GSMs than for GSIs.

4. Conclusions

Small molecule modulators of \( \gamma \)-secretase are now in the early stages of clinical testing. In preclinical toxicology studies, these modulators are free of the mechanism-based toxicities that have been seen with GSIs, most of which appear to be due to inhibition of Notch processing by directly targeting the \( \epsilon \) cleavage site. Both in vitro and in vivo data support the conclusion that GSMs do not interfere with Notch
Table 2: GSM potency versus plasma exposure in mice.

|                  | E-2012 | JNJ-40418677 | GSM1 | SPI-1810 |
|------------------|--------|--------------|------|----------|
| **In vitro** Aβ42 IC50 (nM) CHO-2B7 cells | 33     | 172          | 62   | 114      |
| **Plasma exposure (μM)** | 4923   | 7764         | 2744 | 14638    |
| **Plasma fold over Aβ42 IC50 for 25% reduction** | 149    | 45           | 44   | 128      |
| **Brain exposure (μM)** | 2749   | 7497         | 8681 | 20368    |
| **Brain fold over Aβ42 IC50 for 25% reduction** | 83     | 44           | 140  | 179      |

Figure 7: Diagram highlighting the differences between inhibitors and modulators.

processing, but instead, via slight shifts in the cleavage site on APP, lower Aβ42 production to produce the other normal Aβ peptide products (Figures 6 and 7). Thus, modulation of γ-secretase may represent the most selective approach to treating Alzheimer’s disease via a decrease in production of Aβ42, more selective that an inhibitor of either beta or γ-secretase. Modulators do differ both in chemical structure and in their effects on APP processing, producing the desired decrease in Aβ42 and a variety of changes in other Aβ peptides. Understanding the molecular basis of these varying profiles may shed further light on the biology of γ-secretase.

Disclosures

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