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1 **Identification of a sub-micromolar, Nonpeptide Inhibitor of β -secretase**
2 **with low neural cytotoxicity through *In-silico* Screening**

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2 Nowadays identification of novel non-peptide β -secretase (BACE-1, hereinafter) inhibitors with
3 low cytotoxicity and good blood-brain barrier (BBB) property holds common interest of drug
4 discovery for Alzheimer's disease. 20 SPECS compounds were tested in BACE-1 FRET assays
5 and methylthiazoletetrazolium (MTT) cytotoxicity experiment. Two compounds: **2** and **15**
6 demonstrated IC₅₀ values of 0.53 and 9.4 μ M. In addition, **2** showed least toxic effect to the
7 neuroblastoma cells. The results from both *in silico* and *in vitro* studies provided new
8 pharmacophoric entities for chemical synthesis and optimization on the current discovered
9 BACE-1 small molecule inhibitors.

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2 Alzheimer's disease (AD) is a chronic neurodegenerative disorder of the central nervous
3 system, resulting in severe cognitive deficits along with psychiatric complications. More than 37
4 million individuals suffer from this condition and the number is expected to continue growing
5 dramatically^{1, 2}. Five drugs that are currently on the market – the cholinesterase inhibitors
6 donepezil, rivastigmine, galantamine, and tacrine, the N-methyl-D-aspartate (NMDA) receptor
7 modulator memantine approved for the treatment of symptoms of AD.³ However, they are unable
8 to halt or reverse the disease progression. The major pathological hallmark of AD is the
9 deposition and aggregation of amyloid β -peptide ($A\beta$) in the brain tissues and hence leads to
10 intracellular formation of neurofibrillary tangles. This small peptide consists of 39-43 residues
11 that are endo-proteolytically derived from a transmembrane amyloid precursor glycoprotein
12 (APP)⁴. $A\beta$ production involves the sequential actions of two proteases, the BACE-1 and γ -
13 secretase⁵. Inhibition of these enzymes by small molecule interference is a promising strategy to
14 prioritize pharmaceutical candidates and such approach was highly encouraged by the discovery
15 in 2001 that no $A\beta$ were generated in BACE-1 knockout mice which were devoid of BACE-1
16 activity and displayed a normal phenotype⁶. In the past, the major effort in designing BACE-1
17 inhibitors was the production of transition state isosteres such as hydroxyethylamines, reduced
18 amides, statine-based peptidomimetic inhibitors countering the catalytic aspartyl groups.
19 Although large number of potent peptidomimetic inhibitors can be generated, their relatively
20 large size and number of hydrogen bond donors and acceptors make it challenging to achieve
21 high oral bioavailability^{7, 8}. Therefore, identification of novel small molecule non-peptide
22 inhibitors with balanced ADMET properties will make the chemicals more favorable for further
23 development into leads and drugs. Some non-peptide compounds were identified as inhibitors of

1 BACE-1. Astex researchers highlighted their work in discovering aminopyridine and cyclic
2 amidine classes as BACE-1 inhibitors⁹. Barrow *et al* reported the identification of
3 spiro-piperidine inhibitor template for BACE-1¹⁰. More recently, researchers from Weyth
4 described their identification of aminohydantoin as potent BACE-1 inhibitors¹¹. The most
5 prominent inhibitor was from GSK group which generated GSK188909 as the first orally
6 available BACE-1 inhibitor showing its therapeutic effect in an animal model¹². More
7 information on non-peptide inhibitors of BACE-1 was reviewed by Huang *et al*¹³. Despite the
8 painstaking effort put in discovering drug candidates for AD, no truly disease modifying agents
9 are yet available¹⁴. Hence, investigations on new drug discovery and development for AD draw a
10 great deal of interest to the medicinal chemists.

11 Herein, we report the discovery of two novel BACE-1 small organic inhibitors via a receptor-
12 based virtual screening followed by compound drug-likeness analysis, cytotoxicity prediction,
13 and bioassays. Scheme 1 shows the overall approach adopted in this study. The preparation of
14 protein structure was done according to our previously reported protocol¹⁵. The SPECS chemical
15 library (~280,000 compounds) was screened using GOLD 3.0.1 with default parameters. The
16 active site radius is 10Å from atom 1869 (OG1) of amino acid Thr232 which was one of the key
17 amino acid residues in BACE-1. The GoldScore fitness function was applied and top 5000
18 molecules with the highest scores from initial virtual screening were then re-submitted for
19 multiple docking of 10 conformations for each ligand. Finally, the top 200 hits were kept for
20 further selection. To select which of the *in silico* compounds would be purchased and tested in
21 vitro, several factors were taken account in this step and these included: 1) Information on
22 molecular weight, predicted LogP, LogS and toxicity values of the compounds, toxicity
23 prediction by Pallas (Compudrug Pty Ltd) software; 2) Our knowledge in the molecular

1 interactions between BACE-1 and its substrate, together with the predicted putative H-bonds
2 formed by the hits and active site residues of BACE-1, potential hydrophobic interaction and
3 aromatic-aromatic interactions. Based on the factors stated above, the consensus judgment from
4 both of the selection criteria led us to purchase 20 compounds for enzymatic assays. These
5 compounds mostly have molecular weights < 550 Da, $\text{LogP} < 8$, hydrogen bond donor < 3 , and
6 hydrogen bond acceptor < 7 . The features were considered as favourable physicochemical
7 properties of CNS drugs as highlighted in Pardridge's review¹⁶. Finally twenty SPECS
8 compounds were purchased and tested for their inhibitory activities against BACE-1 in FRET
9 assays. BACE-1 inhibition assays were carried out using [Fluorescence Resonance Energy](#)
10 [Transfer](#) (FRET) assay kits according to the protocol described previously^[15]. 6 compounds
11 exhibited over 50% inhibition of BACE-1 activity at $100\mu\text{M}$. Among them, **2** and **15** had IC_{50}
12 values of $0.53\mu\text{M}$ (Figure 1) and $9.4\mu\text{M}$ respectively. Table 1 shows the structures of the
13 inhibitors and their enzyme assay results. To elucidate the possible interactions between the
14 inhibitors and their active binding site, the docked conformations of **2** and **15** were superimposed
15 in the binding site and the docked poses enabled us to understand the main protein-inhibitor
16 interactions. **The top scored conformations were selected and showed from all conformations**
17 **generated by GOLD.** As shown in Figure 2, **2** and **15** fell into two different interaction patterns
18 with BACE-1. Compound **15** was docked into only half of the BACE-1 active pocket (S1', S1,
19 S2 and S3). Figure 3 depicts the detailed binding modes of the inhibitors in BACE-1. The bulky
20 aromatic groups in **15** made it mainly interact via hydrophobic contacts with the surrounding
21 active residues such as Thr 231, Thr 232, and Tyr71 (Figure 3(b)). In contrast, compound **2**
22 spanned the binding with the extension to the prime side of the active site, as seen by the
23 occupancy of the S4' sub-site (Figure 3(a)). In particular (refer to Figure 4), the ethoxyphenyl

1 occupies the intermediate space between S3 and S4, forming Van der Waals interaction with Leu
2 30 and Ile 110. The phenyl pyrozol group is more exposed to the solvent but a π -cation
3 interaction could be present between Arg 235 and this group and further stabilizing the binding
4 of **2** entropically. Interestingly, unlike other reported inhibitors which are capable of forming
5 multiple hydrogen bonds with the two aspartic acid residues in BACE-1, **2** does not contain a
6 moiety to form hydrogen bonds with either of the aspartic acid residues (Asp 32 or Asp 228) in
7 the S1 catalytic hole. Instead, a benzene ring occupies the S1 region. The aromatic-aromatic
8 interaction with Tyr 71 and Phe 108 could be the reason for an aromatic group to be localised at
9 the S1 of BACE-1 by GOLD. In addition, due to the hydrophobic nature in the S1-S3 region, it is
10 reasonable that a large hydrophobic or aromatic group should be well tolerated. The overall
11 comparison in the binding modes of compound **2** and **15** derived from docking were hence in
12 good agreement with the difference in their inhibitory potencies from enzyme assays.

13 To validate the toxicity prediction from Pallas, MTT assays¹⁷ were conducted on the
14 neuroblastoma cell line SH-SY5Y. MTT reacted with the mitochondrial succinic dehydrogenase
15 in the living cells to produce a dark-purple formazan and the reaction was indicative of the
16 presence of viable cells¹⁷. Neuroblastoma cell lines, SH-SY5Y, were purchased from ATCC
17 through local vendor. The base medium mixture was created using F12 medium and ATCC-
18 formulated Eagles's minimum Essential medium with a ratio of 1:1. The complete medium was
19 formed by adding FBS, fetal bovine serum into the base medium with the final concentration of
20 10%. Cells were cultured in T25 flasks containing 7ml of complete medium. The flask was
21 incubated for 5 to 7 days at 37°C in 5% CO₂ incubator. The SH-SY5Y cells were sub-cultured
22 and fed with fresh medium every 5 to 7 days. Briefly, the test compounds were serially diluted
23 by complete medium to achieve final concentrations ranging from 12.5 μ M to 1.56 μ M in the 96-

1 well plate. 100 μ l of test compound was added into the neuroblastoma cells pre-seeded at the
2 concentration of 3×10^5 cells/ml and incubated for 4 days. Total DMSO concentration in the well
3 is less than 2%. Next, 100 μ l of MTT solution was introduced to each well (final MTT
4 concentration = 0.5mg/ml) and incubated for 4 h at 37°C + 5% CO₂. Finally, 100 μ l of detergent
5 (sodium dodecyl sulfate + isobutanol + 36.5% hydrochloric acid) was added into each well and
6 the absorbance values of 570nm and 630nm (reference) was taken after overnight incubation.
7 The culture medium was used as control in the MTT experiment. The cell viability percentage
8 was calculated using the formula $(A_{570}-A_{630})_{\text{sample}} / (A_{570}-A_{630})_{\text{control}} \times 100\%$.

9 Compound **2** was tolerable by the cells at a concentration of up to 25 μ M whereas compound
10 **15** exhibited a 50% reduction in cell viability at 12.5 μ M. Hence, compound **2** showed a 50-fold
11 selectivity in inhibitory potency over toxic effect. Meanwhile, **2** has a molecular weight of 489,
12 predicted LogP of 4.93, LogS of -8.4, 6 hydrogen bond acceptors and 1 hydrogen bond donor.
13 These features make it nearly fit the criteria for being CNS drugs especially for the BBB
14 permeation¹⁶. The overall “portfolio” of **2** suggested it is an attractive hit and worth taking
15 downstream steps to make it into lead compounds against AD.

16 Albeit **2** showed BACE-1 inhibition at sub-micromolar, there is still no enough information
17 whether it can reduce the A β from the cells. Future work on the measurement of the secreted
18 amount of A β from the current cell line will help us understand more on the medicinal potential
19 of compound **2**.

20 In conclusion, an *in-silico* virtual screening of commercial database SPECS together with
21 enzyme assays and cell-based toxicity testings resulted in the identification of two novel non-
22 peptide inhibitors of human BACE-1. **2** emerged as a sub-micromolar inhibitor (IC₅₀ 0.53 μ M)
23 with low cytotoxicity. Docking of compound **2** in the BACE-1 active site revealed that a central

1 hydrophobic benzene ring occupied the aspartic acid catalytic holes and also suggested the
2 putative binding and interactions with other sub-sites. MTT assays indicated compound **2** was
3 well tolerated by the brain cell line SH-SY5Y. Therefore, we believe that future effort in
4 optimizing present sub-micromolar hit into more potent nanomolar leads based on the molecular
5 clues from this research will lead the drug discovery for Alzheimer's disease to another luminous
6 milestone.

7

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14 opportunity to thank Compudrug for allowing us using the Pallas demo software for predictions
15 of features on compounds.

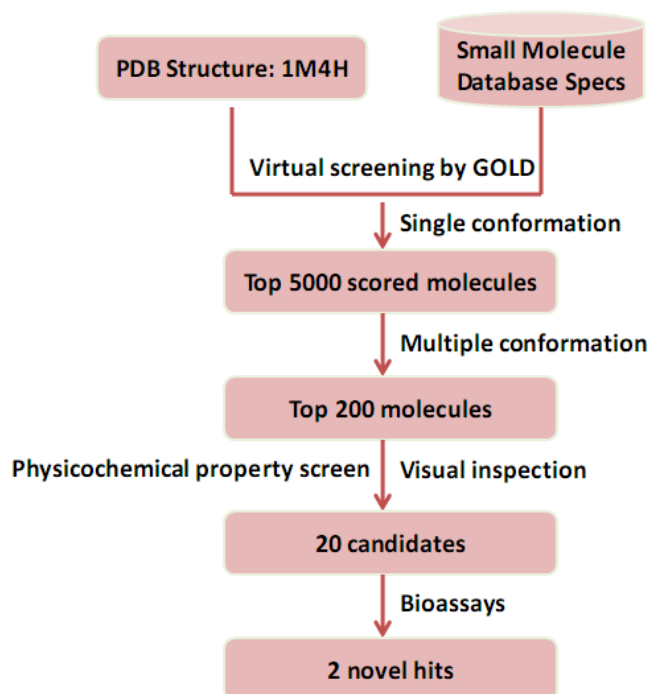
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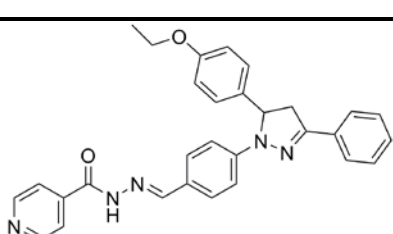
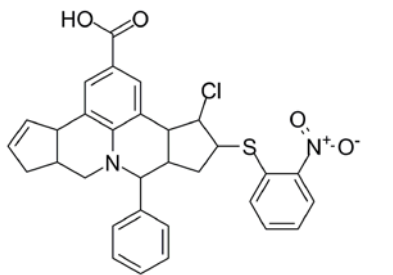
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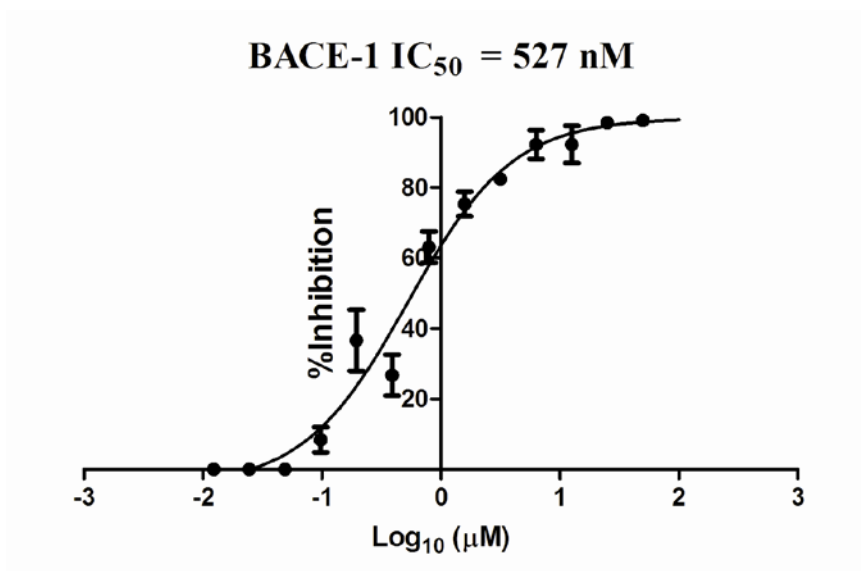
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Scheme 1. Flowchart of virtual screening and bioassays for discovering BACE-1 inhibitors.

Table 1. BACE-1 inhibition results of compound 2 and 15 .			
Compd	Structure	% inhibition at 100 μ M ^[a]	IC ₅₀ ^[b]
2		100	0.53
15		80	9.3

[a] Initial screen were carried out in duplicate and results were presented as the average value. [b] IC₅₀ was done in duplicate. Values were expressed as the average of two assays.

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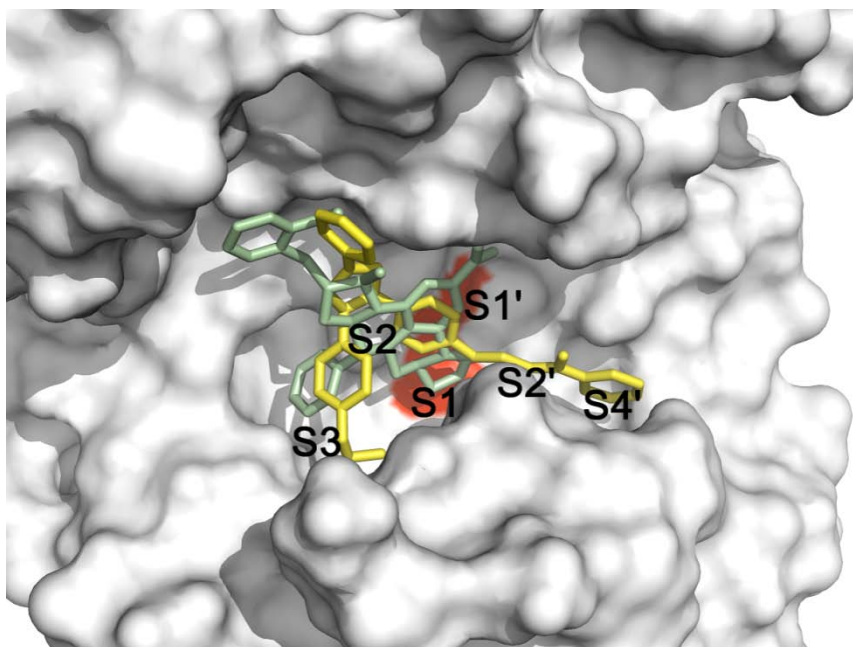
1 a.

2 **Figure 1.** IC₅₀ plot of compound **2**. Measurement was done in duplicate and error bar was shown.

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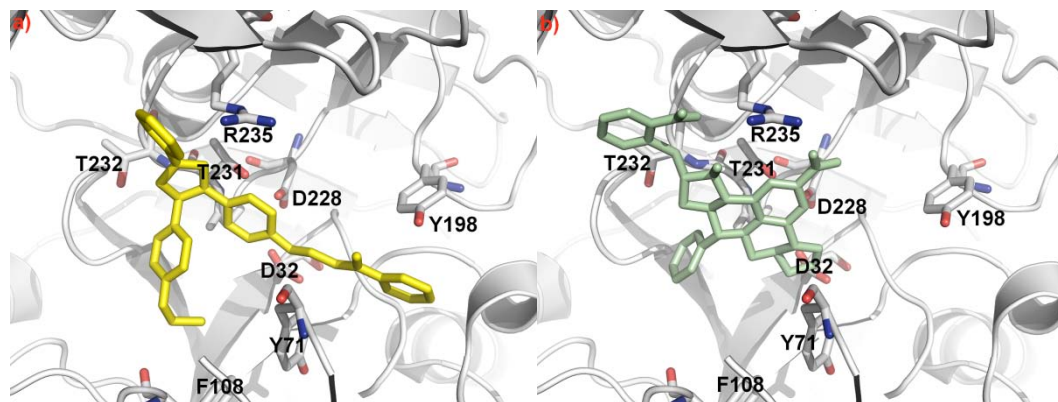
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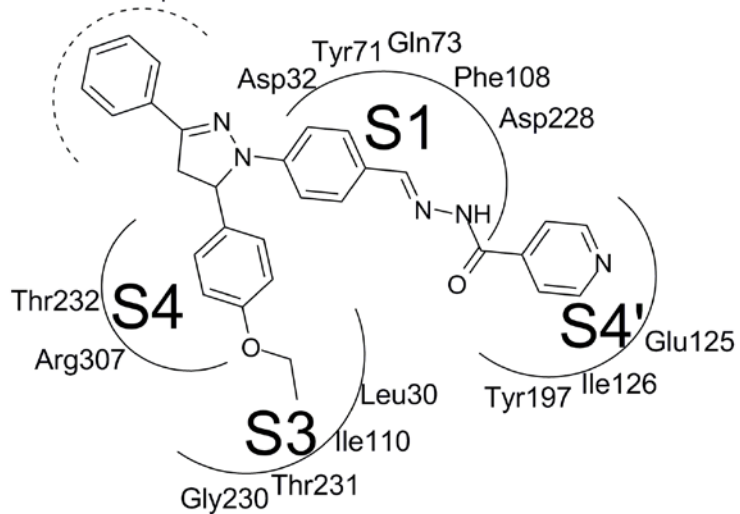
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7 **Figure 2.** Superimposed binding of **2** and **15** derived from docking by GOLD. Compound **2** is colored by yellow
 8 and Compound **15** is indicated using green. Protein surface is colored white. Surfaces of catalytic aspartic acids 32
 9 and 228 are colored in red. Two residues, Thr 72 and Gln 73, were deleted for a whole view of active site. Sub-sites
 10 of BACE-1 are labeled in black. The picture was generated by Pymol. (DeLano, W.L. The PyMOL Molecular
 11 Graphics System (2002) DeLano Scientific, San Carlos, CA, USA.).



1
2 **Figure 3.** Cartoon representation of BACE-1 active site. Key residues are shown in stick and colored by atom type.
3 a) Molecular docking pose of compound **2** b) Molecular docking pose of compound **15**. The binding mode was
4 derived from GOLD and the picture was generated by Pymol software.

Solvent-Exposure



5
6 **Figure 4.** Docked binding mechanism of **2** in BACE-1 active site.

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1 Reference and note

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