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ORIGINAL RESEARCH

In silico predictive model to determine vector-mediated transport properties for the blood-brain barrier choline transporter

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delivery vector to the central nervous system (CNS). We therefore initiated molecular docking studies with the AutoDock and AutoDock Vina (ADVina) algorithms to develop predictive models for compound screening and to identify structural features important for binding to this transporter. The binding energy predictions were highly correlated with $r^2=0.88$, F=692.4, standard error of estimate =0.775, and P-value<0.0001 for selected BBB-ChT-active/inactive compounds (n=93). Both programs were able to cluster active (Gibbs free energy of binding <-6.0 kcal*mol⁻¹) and inactive (Gibbs free energy of binding >-6.0 kcal*mol⁻¹) molecules and dock them significantly better than at random with an area under the curve value of 0.86 and 0.84, respectively. In ranking smaller molecules with few torsional bonds, a size-related bias in scoring producing false-negative outcomes was detected. Finally, important blood-brain barrier parameters, such as the logBB_{massive} and logBB_{active} values, were assessed to predict compound transport to the CNS accurately. Knowledge gained from this study is useful to better understand the binding requirements in BBB-ChT, and until such time as its crystal structure becomes available, it may have significant utility in developing a highly predictive model for the rational design of drug-like compounds targeted to the brain.

Abstract: The blood-brain barrier choline transporter (BBB-ChT) may have utility as a drug

Keywords: blood-brain barrier choline transporter, central nervous system, drug delivery vector, molecular docking, virtual screening, Gibbs free energy of binding, diffusion

Introduction

In the drug design and discovery process, the drug permeation across the blood-brain barrier (BBB) is a pivotal task for neuropharmaceuticals to reach their site of action within the central nervous system (CNS). The BBB consists of the brain capillary endothelial cells connected by tight junction proteins, such as occludin and claudins^{1,2} that circumferentially surround the cell margin restricting passage especially for hydrophilic and positively charged drugs into the CNS.³⁻⁵ While careful chemical modifications are helpful to increase the octanol-water partitioning coefficient (logP) for such drugs to improve their brain accumulation, there would be a total decreased exposure of them to the CNS due to the excessive partitioning of these compounds to other tissues.⁶

An alternative way for a charged molecule to access the brain could be achieved via the BBB native nutrient transporters, such as the blood-brain barrier choline transporter (BBB-ChT). This transporter is responsible for delivery of a positively charged choline molecule into the CNS, where it acts as a structural component of cell membranes and precursor for the neurotransmitter acetylcholine.⁷

Pharmacological applications of using BBB-ChT for drug delivery to the brain may encompass treatment strategies for traumatic brain injury, hypoxia, or ischemia after

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Advances and Applications in Bioinformatics and Chemistry 2014:7 23-36

http://dx.doi.org/10.2147/AABC.S63749

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a stroke, and other neurodegenerative disorders, including Alzheimer's and Huntington's diseases.^{8,9} Recent research has determined that this transporter system might be a suitable drug delivery vector, because it is only occupied at 25 percent of Michaelis–Menten constant (K_m) with choline plasma concentration.^{7,10} However, the membrane-associated and polytopic state of this choline transporter affects its successful cloning¹⁰ and crystallization,¹¹ providing no detailed information on its three-dimensional (3D) structure. For that reason, the neuronal high affinity choline transporter 1 (ChT1) was already employed by other researchers in many molecular docking and 3D structure-activity relationship (3D-QSAR) studies as a substitute for the BBB-ChT to determine structural requirements for various drug-like substances to permeate across the BBB.^{11–13}

Although these studies were shown to be useful in determining lead molecules only for small datasets including novel semi-rigid cyclic and acyclic bis- and mono-quaternary compounds¹³, they are all lacking a systematic approach in terms of the exhaustive molecular docking using different algorithms and important BBB permeability parameters, such as the brain-to-plasma concentration ratio. Therefore, to find possible candidates for the BBB-ChT-mediated transport, the researchers for the current study performed structure-based virtual screening of the BBB-ChT-active/ inactive molecules using the appropriate scoring functions to calculate binding affinities and to correlate them to the molecular physicochemical properties for more accurate BBB permeability prediction. The applicability of the current model will allow identifying prospective drug-like molecules that have desirable BBB-ChT binding properties prior to their chemical synthesis, eliminating the urgency for conventional time and resource-consuming quantitative structure-activity relationship (QSAR) techniques.

Computational methods

The ChT1 homology model to mimic the BBB-ChT protein was constructed using the A-chain of the sodium/galactose symporter (Protein Data Bank ID: 3DH4) as a template¹⁴ by the Iterative Threading Assembly Refinement (I-TASSER) server.¹⁵ The Volume, Area, and Dihedral Angle Reporter server,¹⁶ which is an improved version of the PROCHECK software (European Bioinformatics Institute, Cambridge, UK),¹⁷ was implemented for the stereochemical validation of the ChT1 molecule to investigate the ϕ – ψ dihedral angles in a Ramachandran plot.

Altogether, observed statistics showed that 87% (508 residues) and 8% (44 residues) of all observed residues were in core and allowed regions. The expected values for the

comparison were 90% (522 residues) and 7% (41 residues), respectively, for the same regions obtained elsewhere, from the literature.^{18,19} The BBB-ChT-active/inactive chemical compound database, where most of them are positively charged, included 93 molecules compiled from different literature sources^{12,20–22} in the PubChem BioAssay server. Among them, 44 molecules (47.3%) were the BBB-ChT binders (active substances); 49 molecules (52.7%) were the BBB-ChT non-binders (inactive substances). Prior to the virtual screening procedure, the PyRx software (Scripps Research Institute, San Diego, CA, USA) was used to optimize the dataset.²³

After the conversion, all molecules were inspected manually to detect atoms with improper valence due to mixed aromatic-Kekulé representation. Gasteiger charges were added, and polar hydrogen atoms were assigned. The rotatable bonds were set up, and structure data files were converted into the Protein Data Bank partial charge and atom type format. Rigid-flexible molecular docking was applied to the center of the ChT1 transport channel using Cartesian coordinates: x=0.32 Å; y=0.94 Å; and z=-0.44 Å. The AutoDock and AutoDock Vina (ADVina) docking engines (Scripps Research Institute) were implemented via the Raccoon v1.0 (Scripps Research Institute) and iDOCK (Department of Computer Science and Engineering, Chinese University of Hong Kong, Hong Kong) modifications optimized to perform virtual screening.

AutoDock v.4.2.5.1 was used in the study since its previous version incorrectly calculates part of the intermolecular desolvation energy term. The docking grid with a dimension size of $60 \times 60 \times 60$ Å for AutoDock and $22.5 \times 22.5 \times 22.5$ Å for ADVina was used in the study. The docking output results were represented by the docking scores as Gibbs free energy of binding (ΔG_{bind}), and they were further converted to the predicted inhibition constants (Ki_{pred}). The Ki_{pred} parameters for all the docked poses were calculated from the ΔG_{bind} values as follows:

$$Ki_{pred} = exp([\Delta G_{bind}^*1,000]/[R*T])$$
 (1)

where R (gas constant) is 1.98 cal(mol*K)⁻¹, and T (room temperature) is 298.15 Kelvin.

The python summarize_results4.py script available from MGLTools (Scripps Research Institute) was used to analyze, summarize, and cluster the AutoDock results. Virtual screen performances were evaluated using areas under the receiver operating characteristic (ROC) curve (AUC) and the Boltzmann-enhanced discrimination of ROC (BEDROC) metrics.²⁴ The AUC was calculated by summation while BEDROC 20 (α value is 20.0) values were determined using the R scripts kindly provided by Hiroaki

Yabuuchi from Kyoto University (Department of Genomic Drug Discovery Science, Graduate School of Pharmaceutical Sciences, Kyoto University, Kyoto, Japan). The electrostatic potential maps were calculated with Delphi v5.1 (Computational Biophysics and Bioinformatics, Clemson University, Clemson, SC, USA) using a finite difference solution to the Poisson–Boltzmann equation.²⁵ The calculated octanol-water partitioning coefficient (ClogP) and polar surface area (PSA) for analyzed compounds were determined from molecular interaction fields with the VolSurf+ program (Molecular Discovery, Perugia, Italy). The decimal logarithm value of brain-to-plasma concentration ratio (logBB_{passive}), based mainly on the passive transport (diffusion) for the dataset of BBB-ChTactive/inactive compounds, was calculated from the ClogP and PSA parameters using empirical Clark's equation,²⁶

$$\log BB_{\text{nassive}} = 0.152 \text{Clog P} - 0.0148 \text{PSA} + 0.139 \quad (2)$$

The MATLAB R2012a software (MathWorks, Natick, MA, USA) was used to calculate linear relationships (Table S1) devised from the logarithmic value of the brain-to-plasma concentration ratios ($logBB_{passive}$) for analyzed compounds concerning the BBB-ChT transport by using the following equation,

$$\log BB_{active} = \log BB_{passive} pKi_{exp}$$
(3)

where pKi_{exp} is a negative decimal logarithm of the experimentally determined inhibition constant.

Molecular graphics and visualization were performed with the UCSF Chimera v.1.7 software (Resource for Biocomputing, Visualization, and Informatics, University of California, San Francisco, CA, USA). Statistical analyses were performed using a linear regression analysis, followed by graphic representations using GraphPad Prism v.4 (Graph-Pad Software, Inc., La Jolla, CA, USA). The differences were considered statistically significant at *P*-value <0.0001.

Results and discussion

Our search for a CNS-active hit/lead molecule focuses on a substance that allows for better BBB transfer after binding to the BBB-ChT. Therefore, we wanted to improve a substrate selection and filtering by molecular docking prescreening. For this, we compiled a dataset of the BBB-ChT-active/ inactive substances, including 93 molecules in total. The first ten BBB-ChT-active bis-pyridinium cyclophanes were determined by their binding affinity to the BBB-ChT in a rat brain and were assessed according to their inhibition of [³H] choline uptake.²¹ The other 29 various molecules, comprising

four active and 25 inactive compounds, were analyzed by the same uptake assay, involving in situ brain perfusion studies of male rats.²⁰ Also, 17 bis-azaaromatic quaternary ammonium salts, among them 15 active and two inactive molecules, were synthesized as ligands for the BBB-ChT protein.²²

Finally, 37 chemical compounds including conformationally flexible, semi-rigid, and cyclic quaternary ammonium analogs, among them 15 actives and 22 inactives, were taken from the BBB-ChT 3D-QSAR studies¹² to enrich the entire dataset. A standard rigid-flexible docking technique produced two main outcomes: a particular conformational sampling as a docking pose of the chemical compound with the ChT1 transport channel; and a scoring function (ΔG_{hind}) depicting the protein-ligand interaction strength. Since the true positives and the true negatives are known in this study, the AUC and BEDROC 20 values were quantified from random rankings; statistical significance was estimated by a bootstrap method described in the literature by Efron.²⁷ The results showed a better performance for AutoDock with an AUC value of 0.82 and a standard error of 0.045 than for ADVina, which has an AUC value of 0.81 and a standard error of 0.046. From Figure 1A, it is clear that both programs perform well with all the points on a curve above diagonal, a random AUC selection performance presented with only a 0.5 value.

However, the AutoDock performance indicators show that this method slightly outperforms ADVina, although the advantage is insignificant. In terms of early detection, as determined using the BEDROC 20 measure, both programs performed significantly better than random with BEDROC 20 values of 0.97 for AutoDock and 0.92 for ADVina. This new metric also takes into account the shape of the ROC curves,²⁴ resulting in higher values due to the curves' steep elevation – meaning that known actives are identified at the top of the dataset. The AutoDock and ADVina scoring functions are both weighted functions containing hydrogen bonding and torsional penalty values. While these latter parameters usually differ,^{28,29} it is important to estimate the overall scoring deviations.

A comparison of the predicted docking energies from both programs is shown in Figure 1B, demonstrating a strong correlation between the docking results. As evident by a high Pearson's chi-squared test (r^2) of 0.88, an *F* of 692.4, and a standard error of estimation of 0.775, there is a clear association between the predictions from both algorithms. Based on this correlation in terms of the ΔG_{bind} value, it was expected that the compound conformations would also tend to be similar. All docking poses were ranked according to a score

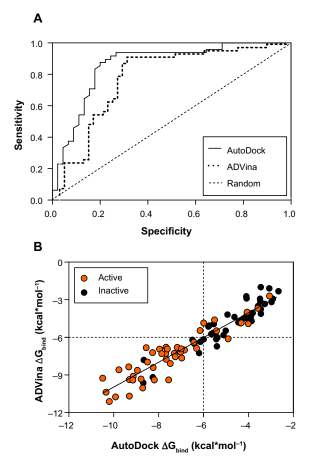


Figure I ROC curves (A) and predicted binding energies (B) from the AutoDock (Scripps Research Institute, San Diego, CA, USA) and ADVina (Scripps Research Institute) runs for 93 analyzed compounds.

Notes: AUC value for each docking run and random selection are shown in the legend. The thresholds are depicted as dashed lines.

Abbreviations: ROC, receiver operating characteristic; AUC, areas under the ROC curve; ΔG_{bind} . Gibbs free energy of binding.

that the docking program assigns to each pose, estimating the ΔG_{bind} values in the range from -10.49 to -2.66 kcal*mol⁻¹ for AutoDock and -11.117 to -2.0 kcal*mol⁻¹ for ADVina (Table S2). Three false-positive AutoDock (ADVina) docking

results with minimal ΔG_{bind} values ranging from -8.68 (-9.62) to -6.49 (-6.21) kcal*mol⁻¹ were detected for chemical compounds 31, 46, and 47 from the entire dataset. On the contrary, the false-negative AutoDock (ADVina) docking outcomes with significantly higher ΔG_{bind} values ranging from -4.32 (-4.92) to -3.06 (-2.7) kcal*mol⁻¹ were determined for five chemical structures, including compounds 10 (choline), 20, 36, 79, and 85, respectively. All these eight molecules were further extracted as outliers from the BBB-ChT compound database to determine the active and inactive compound clustering for AutoDock and ADVina docking runs based on their ΔG_{bind} values. Notably, the substances with ΔG_{bind} deviating slightly from the threshold value (ΔG_{bind} =-6.0 kcal*mol⁻¹) were not excluded as outliers from the analysis.

It can be seen from Figure 2A and B that the lowestbinding energies were generated by the active compounds. Interestingly, most of the active molecules were found to be largely below the $-6.0 \text{ kcal*mol}^{-1}$ threshold with an average ΔG_{bind} value of $-7.92\pm-1.394 \text{ kcal*mol}^{-1}$ for AutoDock and $-7.88\pm1.606 \text{ kcal*mol}^{-1}$ for ADVina. In contrast, the majority of inactive molecules was observed to be above the threshold level with an average ΔG_{bind} value of $-4.55\pm1.057 \text{ kcal*mol}^{-1}$ for AutoDock and $-4.61\pm1.32 \text{ kcal*mol}^{-1}$ for ADVina.

These results indicate again that both docking algorithms perform at the same level with very similar ΔG_{bind} values emphasizing the importance of molecular properties of the analyzed compounds to further explain the clustering process. It was also observed that docking accuracy depends on the size of the compound.³⁰ Therefore, a comparison of the number of heavy atoms presented in each compound, plotted against predicted energetics, revealed a strong correlation coefficient of 0.86, an *F* of 557.7, and a standard error of estimation of 0.796 for AutoDock; ADVina represented an r^2 of 0.87, an *F* of 595, and a standard error

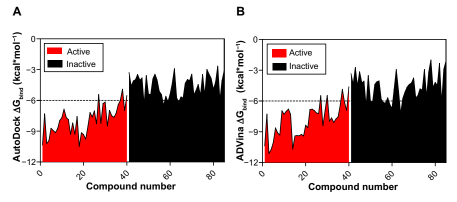


Figure 2 Clustering of BBB-ChT-active/inactive chemical compounds either for AutoDock (Scripps Research Institute, San Diego, CA, USA) (A) or ADVina (Scripps Research Institute) (B) runs, based on their minimal and maximal ΔG_{bind} values. Note: The threshold is depicted as a dashed line.

Abbreviations: BBB-ChT, blood-brain barrier choline transporter; ΔG_{bind} , Gibbs free energy of binding.

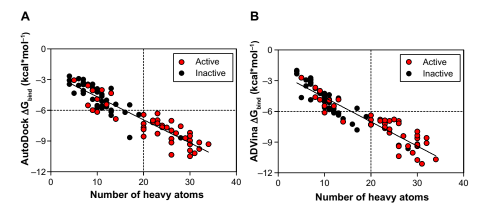


Figure 3 Predicted binding energies for the dataset of BBB-ChT-active/inactive substances as a function of heavy atoms in the compound are plotted for AutoDock (Scripps Research Institute, San Diego, CA, USA) (A) and ADVina (Scripps Research Institute) (B).

Note: The thresholds are depicted as dashed lines.

Abbreviations: BBB-ChT, blood-brain barrier choline transporter; ΔG_{hind} , Gibbs free energy of binding.

of estimation of 0.814. Figure 3A and B shows that all compounds were divided into two major clusters. Namely, the nine active as false-negatives with 44 inactive molecules landed in the cluster with a small number of heavy atoms (<20). Likewise, the five inactive as false-positives with 35 active molecules landed in the cluster with a greater number of heavy atoms (≥ 20). Since the number of atoms and torsions (rotatable bonds) is primarily associated with a larger search space, the clustering of the chemical compounds shows that the false-negative docking results occur in a molecular size-dependent manner and that high levels of ΔG_{hind} correlate with low numbers of heavy atoms in the molecule. Moreover, this size-related bias in scoring was previously detected for the AutoDock and ADVina algorithms through a virtual screening of the Diversity Set II (DSII) and the Database of Useful Decoys (DUD) compound libraries.³¹ Further analysis based on heavy atom count came up with an AUC value of 0.85, a standard error of 0.043 (Figure 4A), and a BEDROC 20 of 0.97, respectively. The AUC for heavy atom count ranking was very similar to the AUC from the AutoDock and ADVina runs,

showing that molecular docking contributes no net signal over heavy atom count.

To accurately assess the correlations between the compound ranking and the molecular properties of chemical compounds, we estimated the root-mean-square deviation ($RMSD_{LC}$) difference between the lowest energy conformation and the reference ligand conformation in the largest cluster. The $RMSD_{LC}$ parameters represented a uniform distribution, which is not dependent on the number of atoms and torsions in the molecule (Figure 4B). On the other hand, the latter two values significantly correlated in a direct manner with the compound ranking; the ranking increased together with a increasing number of atoms and torsions in the docked molecule.

The next step was to validate the data through the correlation between the docking results and experimental affinities (Ki_{exp}) for the BBB-ChT-active ligands (n=29) except 15 cyclic quaternary ammonium analogs. The selection was based on the Ki values for choline (Ki_{choline}), which were in the range of 41–42 μ M for the active bis-pyridinium cyclophanes, bis-azaaromatic quaternary ammonium salts, and

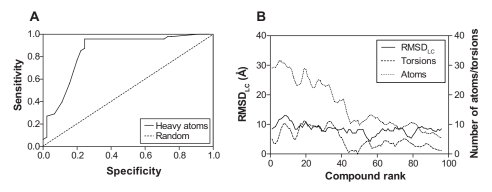


Figure 4 ROC curve of heavy atom count (A) and schematic data smoothing (B) of 93 analyzed compounds for the comparison of the RMSD_{LC} values, number of atoms, and torsions in the molecule.

Abbreviations: ROC, receiver operating characteristic; RMSD_{LC}, the RMSD difference between the lowest energy conformation in the largest cluster and the reference ligand conformation; RMSD, root-mean-square deviation.

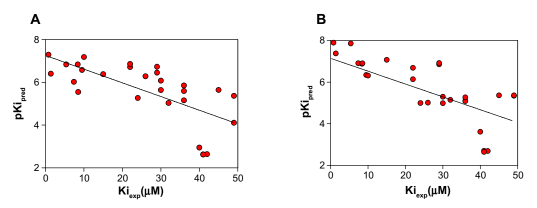


Figure 5 Negative decimal logarithm of predicted inhibition constants (pKi_{pred}), calculated from the ΔG_{bind} values of AutoDock (Scripps Research Institute, San Diego, CA, USA) (A) and ADVina (Scripps Research Institute) (B) runs, between 29 BBB-ChT-active compounds and target protein and plotted against experimentally determined inhibition constants (Ki_{exp}).

Abbreviations: BBB-ChT, blood-brain barrier choline transporter; ΔG_{bind} Gibbs free energy of binding.

various molecules from the 3D-QSAR dataset,²⁰ in contrast to the Ki_{choline} of 0.68 μ M for ammonium analogs. Regardless of the moderate correlation coefficient (r^2 =0.47) between the pKi_{pred} values, calculated from the appropriate ΔG_{bind} parameters and Ki_{exp} for both algorithms (Figure 5A and B), it was previously reported that scoring tests on 90 proteinligand complexes from the Protein Data Bank demonstrated a statistically significant correlation even with r^2 in the range from 0.45 to 0.55, respectively.³²

To gain some insight into the binding characteristics of BBB-ChT-active compounds, we docked them into its binding cavity. During the docking process, the protein was considered to be rigid while the ligands were flexible. As it has been already suggested that the BBB-ChT pore is extremely important for a translocation of positively charged drugs across the BBB,²² the analyzed active substances occupied negatively charged portions of the pore within the same binding cavity close to the center of the transporting channel (Figure 6). In accordance with our results, the hypothetical model for the choline transporter binding site²⁰ explains this phenomenon due to strong ionic interaction between the trimethylammonium moiety of ligand and corresponding amino acid residues of protein. In addition, bulk cavity methyl acceptors might play some role to establish a conformational congruence with ligand methyl groups and potentiate the BBB-ChT affinity.

The ClogP, PSA, and logBB values for BBB-ChT-active/ inactive compounds were obtained to evaluate their ability to interact at the CNS level and possess optimal BBB permeation properties. All compounds include ClogP values in the range of -5.87 to 5.204 and PSA values in the range of 3.24-64.67 Å². For a CNS-active molecule to permeate the BBB, an area <60 Å² is usually needed, and molecules with a PSA of >120-140 Å² tend to be poor in permeating cell membranes.³³ The ClogP values specify the lipophilic character of the examined compounds, which should be high enough for the molecule (ClogP >0) to cross the BBB. The results indicate that most of the BBB-ChT-active compounds have suitable PSA (PSA <60 Å²), but most of them are not lipophilic enough (ClogP <0) to permeate the BBB successfully (Figure 7A). Judging by low ClogP, these drugs will probably retain at the hydrophilic compartment, such as blood serum. However, the ClogP and PSA alone have proven insufficient for the accurate evaluation of the BBB permeation, since they correlate poorly with the logBB values in our previous studies.^{5,34}

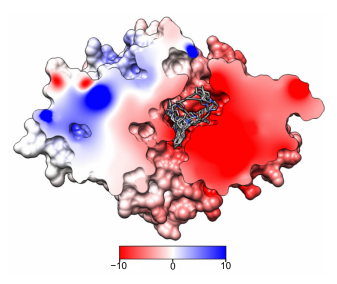
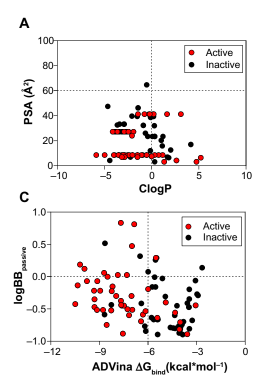
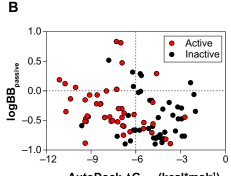


Figure 6 AutoDock (Scripps Research Institute, San Diego, CA, USA) rigid-flexible molecular docking of BBB-ChT-active compounds (44 molecules) into the binding site of the ChTI homology model.

Notes: The molecular surface is divided by the frontal plane to visualize the protein pore. Hydrogen bonds are omitted for clarity. Red and blue colors show negative and positive potentials, while the zero potential is in white. Hydrogen atoms are omitted for clarity. **Abbreviations:** BBB-ChT, blood-brain barrier choline transporter; ChTI, choline transporter 1.





AutoDock ΔG_{bind} (kcal*mol⁻¹)

Figure 7 Relationship between molecular properties, such as ClogP, PSA (**A**), and Gibbs free energy of binding (ΔG_{bind}), and decimal logarithm of brain-to-plasma concentration ratio (logBB_{passive}) based on passive transport, such as diffusion (**B** and **C**).

Notes: Thresholds are shown as dashed lines. AutoDock, Scripps Research Institute (San Diego, CA, USA). ADVina, Scripps Research Institute. Abbreviations: ClogP, calculated octanol-water partitioning coefficient; PSA, polar surface area.

According to the CNS ± activity classification for different chemical compounds, molecules with logBB >0 can cross the BBB readily, while drugs with logBB <0 cannot.³⁵ Therefore, we calculated logBB_{passive} value for the BBB-ChT-active/ inactive compounds using Equation 2 (Table S3), which was empirically devised from a smaller dataset (n=55; r^2 =0.79; standard error of estimate (SEE) =0.35).²⁶ It can be seen from Figure 7B and C that most of the substances have negative log-BB_{passive} parameters except for the active compounds 2–5, 17, 24, 41, and 84. Based upon these results, the bis-pyridinium cyclophane compounds 3–5 (5b–5c) with the best Ki_{exp} values (Table 1) were subjected to further analysis to define their blood–brain concentrations according to Equation 3. Centrally acting cholinesterase inhibitor donepezil hydrochloride (Aricept) with experimental logBB of 0.89 was used as a reference substance. As demonstrated in Figure 8A and B, the compound distribution into the brain, governed by linear relationship, was significantly improved due to enhanced permeation rate across the BBB after considering the passive (logBB_{passive}) and BBB-ChT-active transport (logBB_{active}).

Conclusion

In this current study, we report on the development of an in silico structure-based predictive model to determine

Compound	Κi_{exp} (μΜ)	рКі _{ехр}	$\Delta G_{bind}^{AutoDock}$ (kcal*mol ⁻¹)	$\Delta G_{\text{bind}}^{\text{ADVina}}$ (kcal*mol ⁻¹)	ClogP	PSA (Ų)	logBB _{passive}
3 (5d)#	33.8	4.47	-10.2	-11.11	1.16	8.66	0.187
4 (5c)	0.8	6.09	-9.92	-10.74	0.74	8.66	0.123
5 (5b)	1.4	5.85	-8.71	-10.04	0.313	8.66	0.058
Donepezil	274*	3.65	-	-	3.08-4.11**	38.8***	0.89****

Table I Experimental and predicted parameters to assess BBB permeation for CNS- and BBB-ChT-active compounds

Notes: "alternative drug name as denoted in Zhang et al,²¹ *Kang et al,³⁶ **Thevis et al,³⁷ Xia et al,³⁸ Choi et al,³⁹ ***Goh et al,⁴⁰ ****Muehlbacher et al.⁴¹ AutoDock, Scripps Research Institute (San Diego, CA, USA). ADVina, Scripps Research Institute.

Abbreviations: BBB, blood–brain barrier; CNS, central nervous system; BBB-ChT, blood–brain barrier choline transporter; $K_{i_{exp}}$, experimentally determined inhibition constant; ClogP, calculated octanol-water partitioning coefficient; PSA, polar surface area; $logBB_{passive}$, decimal logarithm of brain-to-plasma concentration ratio, based on passive transport; $\Delta G_{bind}^{AutoDock}$, Gibbs free energy of binding calculated by AutoDock; ΔG_{bind}^{ADVina} , Gibbs free energy of binding calculated by ADVina.

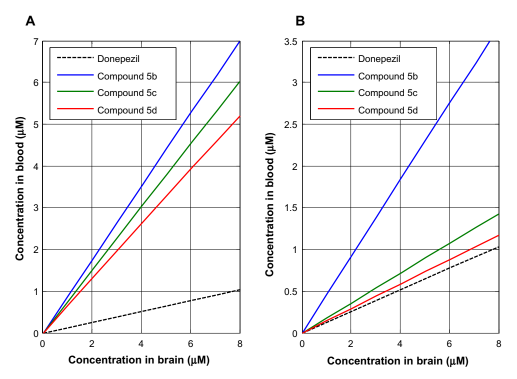


Figure 8 Brain and blood concentration relationship for the bis-pyridinium cyclophane compounds devised from the decimal logarithm of brain-to-plasma concentration ratio based on diffusion alone (A) and together with BBB-ChT active transport (B).

Notes: Donepezil is used as a reference substance (dashed line). The logBB parameter is described as linear cumulative distribution function. **Abbreviation:** BBB-ChT, blood-brain barrier choline transporter.

vector-mediated transport properties for drug-like chemical compounds using the BBB-ChT system. Surprisingly, this transporter has not been cloned, expressed, or crystalized. However, the homology model of ChT1 was implemented in virtual screening as a substitute for the BBB-ChT protein due to the absence of its 3D crystal structure. The molecular docking studies were initiated with the AutoDock and ADVina search algorithms and provided highly correlated ΔG_{bind} values with r^2 =0.88, F=692.4, SEE=0.775, and P-value<0.0001 for the compiled BBB-ChT database. Both programs were able to cluster active ($\Delta G_{bind} < -6.0 \text{ kcal*mol}^{-1}$) and inactive ($\Delta G_{bind} > -6.0 \text{ kcal*mol}^{-1}$) molecules and dock them significantly better than at random with an AUC of 0.82 for AutoDock and 0.81 for ADVina.

In the molecular docking of smaller compounds with few torsional bonds, a size-related bias in scoring was detected, which affects the ligands comprising <20 heavy atoms. This bias was responsible for a failure to preferentially rank only active compounds at the top, producing false-negative outcomes. Finally, important BBB parameters, such as the logBB_{passive} and logBB_{active} values, were assessed to evaluate the role of active transport for compounds to cross the BBB. Information obtained from this study is useful to determine the binding requirements in BBB-ChT, and until such time as

its crystal structure becomes available, it may have significant utility in elaborating a highly predictive model for the rational design of drug-like compounds targeted to the brain.

Acknowledgments

Special thanks are extended to Dr Ellaine Salvador from Würzburg University and Anna Poon from the City College of New York for their assistance in the writing of this paper. The authors are grateful to the Bundesministerium für Bildung und Forschung (BMBF) for its support of this work by providing the BMBF01 grant to Carola Förster.

Disclosure

The authors report no conflicts of interest in this work.

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Supplementary materials

Table SI MATLAB script to generate Figure 8A and B

clear clear all subplot(1,2,1) x=linspace(0,100); yl=x/7.7625; y2=x/1.1429; y3=x/1.3274; y4=x/1.5382; plot(x,y1,'black--',x,y2,x,y3,x,y4,'LineWidth', 2); ylabel('Concentration in blood (\muM)'); xlabel('Concentration in brain (\muM)'); legend('Donepezil','Compound 5b','Compound 5c','Compound 5d', 'Location', 'NorthWest') ylim([0 7]) xlim([0 8]) grid on subplot(1,2,2) x=linspace(0,100); y5=x/7.7625; y6=x/2.1842; y7=x/5.6114; y8=x/6.8531; plot(x,y5,'black--',x,y6,x,y7,x,y8,'LineWidth', 2); ylabel('Concentration in blood (\muM)'); xlabel('Concentration in brain (\muM)'); legend('Donepezil','Compound 5b','Compound 5c','Compound 5d', 'Location', 'NorthWest') ylim([0 3.5]) xlim([0 8]) grid on

Table S2 Docking poses	ranked according to a sco	re that the docking program	assigns to each pose

#COMP	#RUNS	#LC	RMSD_LC	#ATS	#TORS	ΔG_{bind} (AutoDock)	ΔG_{bind} (ADVina)	BBB-ChT Activity
I	10	10	7.1249	26	0	-10.34	-10.378	I
2	10	3	4.6183	20	8	-7.3	-7.272	I
3	10	10	10.7815	31	0	-10.2	-11.117	1
4	10	10	9.5712	30	0	-9.92	-10.741	1
5	10	7	8.1711	29	0	-8.71	-10.037	1
6	10	2	11.7575	30	11	-8.95	-8.628	1
7	10	2	16.1779	32	13	-9.13	-9.098	I
8	10	3	10.3328	26	7	-8.78	-9.323	I
9	10	I	14.653	26	13	-7.96	-6.939	I
10	10	4	8.4713	8	3	-3.58	-3.676	I
11	10	4	10.4312	7	3	-3.27	-3.335	0
12	10	2	10.0631	11	6	-4.62	-4.379	0
13	10	10	8.7533	7	3	-4.09	-2.73	0
14	10	10	5.8992	10	5	-3.99	-4.566	0
15	10	9	8.4537	7	3	-3.47	-3.147	0
16	10	3	9.5477	9	4	-4.03	-4.23 I	0
17	10	2	5.7884	20	8	-7.68	-7.295	I
18	10	4	5.4073	9	3	-3.81	-4.05	0
19	10	6	11.8551	19	6	-6.41	-6.506	0
20	10	7	5.2819	11	6	-4.01	-4.915	I
21	10	10	9.2404	6	I	-3.57	-3.508	0
22	10	6	4.2376	12	2	-5.38	-6.014	0
23	10	9	4.2149	12	2	-5.42	-6.077	0
24	10	3	6.0864	14	7	-6.85	-7.003	I
25	10	5	5.9122	10	5	-3.9	-4.371	0
26	10	3	4.4788	9	3	-3.44	-4.12	0
27	10	5	10.6334	9	4	-4.01	-4.045	0
28	10	7	4.3613	11	2	-5.11	-5.816	0
29	10	6	4.2999	12	2	-5.43	-6.148	0
30	10	9	5.847	13	I	-6.49	-6.208	0
31	10	2	10.2889	17	4	-8.66	-7.807	0
32	10	5	6.5735	11	2	-5.59	-5.68	0
33	10	3	8.0609	13	4	-6.14	-6.355	0
34	10	5	6.706	14	4	-5.4	-6.71	0
35	10	7	10.3509	7	0	-4.35	-4.863	0
36	10	3	6.2092	13	8	-4.32	-4.841	I
37	10	3	9.576	6	3	-2.91	-2.928	0
38	10	6	10.0262	10	I	-5.74	-5.06	0
39	10	6	5.5151	16	2	-6.2	-7.241	0
40	10	I	9.3143	25	12	-7.67	-6.785	I
41	10	Ι	7.1695	20	8	-7.86	-7.294	I
42	10	2	14.6284	34	8	-9.3	-10.685	I
43	10	2	8.4975	30	8	-8.19	-9.398	I
44	10	4	11.394	30	8	-9.3	-9.363	I
45	10	2	8.7913	28	8	-7.55	-9.389	I
46	10	2	9.1544	28	8	-8.68	-9.623	0
47	10	2	11.9626	30	10	-8.26	-9.183	0
48	10	-	8.74	30	10	-10.49	-9.253	I
49	10	3	10.1642	26	7	-9.15	-9.409	1
50	10	2	11.2523	30	,	-9.33	-8.35	1
51	10	1	12.2826	32	13	-9.77	-8.587	
52	10		6.8438	20	7	-8.55	-6.822	
53	10		8.9795	20	9	-0.55 -7.17	-6.795	
		1	13.5064	24	11	-7.61	-6.922	•

(Continued)

#COMP	#RUNS	#LC	RMSD_LC	#ATS	#TORS	$\Delta \mathbf{G}_{bind}(\mathbf{AutoDock})$	$\Delta \mathbf{G}_{bind}(\mathbf{ADVina})$	BBB-ChT Activity
55	10	2	10.8102	26	13	-7.02	-7.164	1
56	10	I.	10.7718	23	10	-8.27	-7.211	I
57	10	7	4.9855	12	3	-5.42	-5.478	I
58	10	3	7.3153	27	11	-8.24	-8.406	I
59	10	2	8.1588	23	10	-6.31	-6.876	I
60	10	8	3.6022	12	I	-5.65	-5.737	0
61	10	6	8.7255	11	I	-5.78	-5.135	0
62	10	5	8.9773	10	0	-6.15	-5.457	I
63	10	I	8.894	30	14	-8.5	-7.968	I
64	10	6	7.925	23	8	-6.97	-7.648	I
65	10	2	7.6806	24	8	-7.48	-8.086	I
66	10	2	8.7331	24	7	-7.66	-7.744	I
67	10	4	8.1205	23	7	-7.26	-7.543	I
68	10	I.	8.0574	20	7	-6.43	-6.385	I
69	10	2	6.632	11	5	-4.32	-4.648	0
70	10	6	6.3117	10	5	-3.96	-4.371	0
71	10	4	5.6526	10	4	-4.08	-4.242	0
72	10	6	8.9682	9	0	-5.99	-4.854	I
73	10	4	10.9724	8	4	-3.45	-3.755	0
74	10	2	5.7092	17	12	-5.47	-5.7	0
75	10	4	6.2461	12	L	-5.01	-5.464	0
76	10	6	5.5277	10	L	-4.9	-6.115	0
77	10	2	9.5978	22	15	-6.99	-6.914	I
78	10	3	6.7502	11	5	-4.45	-4.802	0
79	10	6	10.7114	9	4	-4.06	-3.983	I
80	10	6	10.1741	10	L	-5.54	-4.931	0
81	10	10	8.6855	7	3	-3.48	-2.925	0
82	10	3	8.3284	10	0	-5.54	-5.621	0
83	10	4	4.8109	7	3	-2.97	-3.182	0
84	10	4	10.3719	8	0	-5.51	-4.602	I
85	10	7	9.2977	5	0	-3.06	-2.701	I
86	10	10	7.9856	4	0	-3.45	-2	0
87	10	7	3.9228	13	2	-3.86	-4.699	0
88	10	3	9.4446	9	4	-4.93	-4.186	0
89	10	4	10.2794	10	I	-4.21	-4.49	0
90	10	6	8.8939	4	0	-2.66	-2.323	0
91	10	5	5.1308	13	I	-5.99	-6.16	0
92	10	10	8.7583	7	3	-4.08	-2.89	0
93	10	8	7.4217	4	0	-3.07	-2.095	0

Notes: AutoDock, Scripps Research Institute (San Diego, CA, USA). ADVina, Scripps Research Institute.

Abbreviations: #COMP, number of compound; #RUNS, total number of runs found in all the dlg files in the specified directory; #LC, number of largest cluster; RMSD_LC, the RMSD difference between the lowest energy conformation in the largest cluster and the reference ligand conformation; #ATS, number of atoms; #TORS, number of torsions; ΔG_{bind} , Gibbs free energy of binding in kcal*mol⁻¹; 1, BBB-ChT active; 0, BBB-ChT inactive; BBB-ChT, blood–brain barrier choline transporter; dlg, docking log; RMSD, root-mean-square deviation.

Table S3 Calculated logBB
 passive
 value for BBB-ChT-active/inactive

 compounds using Equation 2
 2

Fable S3 (Continued)	Fable	S 3	(Continued)	
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compounds us	sing Equation 2			#COMP ClogP PSA logBB				
#COMP	ClogP	PSA	logBB _{passive}	54	-3.022	8.66	-0.448512	
1	-0.956	8.66	-0.13448	55	-2.04	8.66	-0.299248	
2	-0.006	7.57	0.02605199	56	-3.513	8.66	-0.523144	
3	1.159	8.66	0.187	57	-2.464	27.2	-0.638088	
4	0.736	8.66	0.122704	58	1.625	41.26	-0.224648	
5	0.313	8.66	0.05840815	59	-0.764	41.26	-0.587776	
6	-1.506	8.66	-0.21808	60	-2.638	27.2	-0.664536	
7	-0.524	8.66	-0.068816	61	-2.438	27.2	-0.634136	
8	-3.47	8.66	-0.516608	62	-2.086	24.04	-0.533864	
9	-2.04	8.66	-0.299248	63	3.098	41.26	-0.000752	
10	-4.123	27.2	-0.890256	64	-0.271	41.26	-0.51284	
11	-1.527	39.44	-0.676816	65	0.152	41.26	-0.448544	
12	-3.442	27.2	-0.786744	66	0.214	41.26	-0.43912	
13	-1.434	46.25	-0.763468	67	-0.209	41.26	-0.503416	
14	-3.405	27.2	-0.78112	68	-1.445	41.26	-0.691288	
15	-0.882	32.26	-0.472512	69	-2.942	33.27	-0.80058	
16	-3.896	27.2	-0.855752	70	-3.141	27.2	-0.740992	
17	5.204	6.48	0.834104	71	-3.41	33.27	-0.871716	
18	-2.857	27.2	-0.697824	72	-1.503	6.97	-0.192612	
19	0.196	39.34	-0.41344	72	0.076	23.47		
20	-2.914	27.2	-0.706488	73	1.674	6.97	-0.196652 0.290292	
21	-2.935	6.97	-0.410276	75	-4.655	47.42		
22	-2.63	27.2	-0.66332	76			-1.270376	
23	-2.63	27.2	-0.66332		-2.864	6.97	-0.399484	
24	4.746	3.24	0.81244	77 78	2.628	4.33	0.474372	
25	-3.141	27.2	-0.740992	78 79	-2.908	33.27	-0.795412	
26	-3.574	27.2	-0.806808	80	-3.632	27.2	-0.815624	
27	-3.632	27.2			0.323	23.47	-0.15926	
28	-3.832 1.191	23.47	-0.815624	81	-0.882	32.26	-0.472512	
28		23.47	-0.027324	82	2.015	12.63	0.258356	
30	-2.63		-0.66332	83	-0.414	23.47	-0.271284	
31	1.801 4.148	6.48 17.07	0.316848 0.51686	84	1.324	3.24	0.292296	
32	1.013	20.31		85	-3.162	6.97	-0.44478	
33		20.31	-0.007612	86	-0.661	26.02	-0.346568	
34	-3.825	32.6	-0.84496	87	-0.525	64.67	-0.897916	
	-3.635		-0.896	88	-2.254	6.97	-0.306764	
35	-4.436	4.33	-0.599356	89	-0.116	38.65	-0.450652	
36	-1.932	27.2	-0.557224	90	0.317	3.24	0.139232	
37	0.286	32	-0.291128	91	1.13	15.87	0.075884	
38	0.323	23.47	-0.15926	92	-1.434	46.25	-0.763468	
39	-2.096	39.73	-0.767596	93	-0.143	12.03	-0.06078	
40	-2.531	8.66	-0.37388		#COMP, number of co			
41	-0.006	7.57	0.02605199		cient; PSA, polar surfa			
42	-2.384	8.66	-0.351536	brain to piasma co brain barrier choli	oncentration ratio base ne transporter.	on passive transpo	, DDD-CITT, DIOOC	
43	-1.07	8.66	-0.151808					
44	-3.164	8.66	-0.470096					
45	-5.874	8.66	-0.882016					
46	-3.932	8.66	-0.586832					
47	-2.936	8.66	-0.43544					
48	-2.86	8.66	-0.423888					
49	-3.47	8.66	-0.516608					
50	-1.506	8.66	-0.21808					
51	-0.524	8.66	-0.068816					
52	-4.986	8.66	-0.74704					
53	-4.004	8.66	-0.597776					
			(Continued)					

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