Discovery of CNS-Like D3R- Selective Antagonists Using 3D Pharmacophore Guided Virtual Screening

June Hyeong Lee,1 Sung Jin Cho,2,3 Mi-hyun Kim1*

1Gachon Institute of Pharmaceutical Science & Department of Pharmacy, College of Pharmacy, Gachon University, 191 Hambakmoero, Yeonsu-gu, Incheon, Republic of Korea, 2CimplSoft, Thousand Oaks, CA 91320, 3CimplRx, Euni-ro, Seoul, Republic of Korea

* Author for correspondence
E-mail: Mi-hyun Kim: kmh0515@gachon.ac.kr; Sung Jin Cho: sjcho@cimplsoft.com

Abstract
The dopamine D3 receptor is an important CNS target for the treatment of a variety of neurological diseases. Selective dopamine D3 receptor antagonists modulate the improvement of psychostimulant addiction and relapse. In this study, five and six featured pharmacophore models of D3R antagonists were generated and evaluated with the post-hoc score combining two survival scores of active and inactive. Among Top 10 models, APRRR215 and AHPRRR104 were chosen based on the coefficient of determination (APRRR215: R²\text{training} = 0.80; AHPRRR104: R²\text{training} = 0.82) and predictability (APRRR215: Q²\text{test} = 0.73, R²\text{predictive} = 0.82; AHPRRR104: Q²\text{test} = 0.86, R²\text{predictive} = 0.74) of their 3D-quantitative structure–activity relationship models. Pharmacophore-based virtual screening of a large compound library from eMolecules (> 3 million compounds) using two optimal models expedited the search process 100-fold speed increase compared to the docking-based screening (HTVS scoring function in Glide) and identified a series of hit compounds having promising novel scaffolds. After the screening, docking scores, as an adjuvant predictor, were added to two fitness scores (from the pharmacophore models) and predicted Ki (from PLSs of the QSAR models) to improve accuracy. Final selection of the most promising hit compounds were also evaluated for CNS-like properties as well as expected D3R antagonism.

Keywords: Pharmacophore, 3D-QSAR, virtual screening, D3R selective antagonist, molecular docking, CNS-like

Introduction
Dopamine receptors are a class of G protein-coupled receptors in the central nervous system (CNS). There are at least five subtypes of dopamine receptors: D1R, D2R, D3R, D4R, and D5R. The D1R and D5R are members of the D1-like family of dopamine receptors, whereas the D2R, D3R, and D4R are members of the D2-like family [1]. The D3R is found within a key neuronal network involved in motivation and cognition. In contrast to the D2R, the D3R does not appear to play a role in the regulation of movements [2, 3]. Furthermore, the D3R subtype is considered to represent an important target for the treatment of a variety of neurological diseases, such as schizophrenia and Parkinson’s disease [4]. D3R-selective antagonists can be potential therapeutic agents for the treatment of psychostimulant addiction and relapse [5-9]. However, many clinically approved drugs targeting the D3R do not show significant selectivity over the D2R and other receptors [9]. Thus, it is necessary to develop selective D3R antagonists.

In this study, we have conducted computer-aided drug identification to investigate D3R-selective antagonists for use in CNS diseases. After examining structural differences between the D3R (PDB name: 3PBL) and the D2R proteins [10], binding modes of three ligands (dopamine, eticlopride, and R-22) with D3R were comparatively analysed using docking: (1) dopamine (as the endogenous ligand), (2) eticlopride (as a non-selective D2R/D3R antagonist [11]), and (3) R-22 (as a selective D3R antagonist). According to the literature [12],...
our docking approach identified a common binding site, a deep pocket common to all three compounds, and a shallow pocket occupied by some antagonists (Figure 1). R-22, a selective D3R antagonist, occupied the deep pocket and was found bound to the outer binding pocket to a greater extent [12]. The residue information derived from the docking study was used to develop novel molecules [13], and models including pharmacophore features were built from the dataset containing selected D3R antagonists with high efficacy, selectivity, and metabolic stability.

Figure 1: Superimposed 3PBL protein-ligand docking of dopamine (green; endogenous D3R agonist), eticlopride (yellow; non-selective D3R antagonist), and R-22 (sky blue; selective D3R antagonist).

Rational drug design is based on the notion that molecules are active at a particular receptor because they have numerous chemical features and complementary geometries that facilitate their interaction with the binding site [14]. These pharmacophore features reflect the atoms present, their locations, and six key representation of non-covalent interactions: hydrogen bond acceptor, hydrogen bond donor, hydrophobicity, positive & negative ionizable moieties, and aromatic ring. A pharmacophore hypothesis is a collection of 3D vectors which represent these features. To discover pharmacophores in the absence of reliable co-crystallographic data, the active analog approach is typically used to facilitate the identification of common protein-ligand interactions. [15]. The user must set a minimum number of features to be used to build a pharmacophore model without overfitting. The common pharmacophore generation step is designed to identify top-ranked active compounds. Scoring is based on the vector score, site score, volume score, selectivity score, and clustering. The scoring of active compounds and overall ranking of plausible models may not be enough to select more plausible models in describing the binding mode of active compounds. By including the scoring of inactive compounds, pharmacophore models were compared. From chosen pharmacophore models, 3D-QSAR models could be developed using the conformers of compounds in the training set, which match the pharmacophore on sites. The employed fitness indicates an alignment tendency of each molecule. After processing, generating, and validating models, two
optimal D3R-antagonist models were used to screen compounds obtained from commercial databases. This process is shown as the workflow in Figure 2.

![Figure 2: Workflows of this study.](image)

**Materials and methods**

**Data collection and molecular docking.**

A set of 25 novel D3R-selective antagonists (Table 1) [16] and their inhibition constant (Ki) data (nM) were utilized for the development of a pharmacophore model and building pharmacophore-based three-dimensional quantitative structure–activity relationship (3D-QSAR) model [16]. The negative logarithms of the measured Ki values (pKi for human dopamine D3R binding data in human embryonic kidney cells) were converted to μM, which was used in this study. Before building a pharmacophore model, compounds were processed by Ligprep to attach hydrogens, convert 2D structures to 3D, and generate stereoisomers [17]. In addition, we applied the target pH of 7 ± 2 using Epik, and specified chirality using Force field OPLS_2005 in Maestro 10.4, Schrödinger [18]. After ligand preparation, each ligand had five conformers and every conformer was used for docking to D3R. For the docking, various docking models between R-22 and the apo protein of 3PBL were generated using (1) residue specific centroid selection, (2) adjustment of grid size & shape, (3) constraint on the residues having expectable non-covalent interaction, and (4) three scoring functions. The optimal model was selected based on the docking score and similarity of the pose compared to one found in the literature [12]. The best docking condition of R-22 was used to dock 50 conformers to generate the best pose of each conformer.
Table 1. Twenty-five selective D3R-antagonists used to build the pharmacophore model [14]

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Identification of pharmacophore hypotheses.

Pharmacophore models were built using Pharmacophore Alignment and Scoring Engine (PHASE) running on Maestro 10.4 (Schrödinger). The best docking pose was considered to be a bioactive conformer of each conformer and used for the development of the pharmacophore model. The activities of compounds were scaled from a minimum value of 1.01 to a maximum value of 3.93 (to be consistent with ones in the table), with an activity threshold of 1.35; this meant that compounds with a Ki of < 44 nM were considered to be antagonists (actives). The pharmacophore features used for hypothesis generation were hydrogen bond acceptor (A), hydrogen bond donor (D), hydrophobic group (H), positively ionisable (P), negatively ionisable (N), and aromatic rings (R) defined by a set of chemical structure patterns. For the current dataset of 50 conformers, five or six features were chosen for model construction. The pharmacophore feature of active ligands that contain identical sets of features with very similar spatial arrangements were grouped together to give rise to a common pharmacophore hypothesis. In the present study, pharmacophore-based QSAR modeling was conducted by dividing the dataset into a 35-member training set (70%) and a 15-member test set (30%) in a random manner. The chemical features associated with ligand-protein binding and the training set data are used to create a model, and the test set is used to evaluate this model [19].

Internal and external QSAR model validation.

The success of a virtual screening campaign using a QSAR model depends heavily on the quality of the model. Several statistical parameters in PHASE, such as R2 for the training set and Q2 for the test set, the standard deviation, root mean square error, and variance ratio (F), can be used to evaluate the robustness of a QSAR model [20-23]. To test the reliability of QSAR models, compounds that were not used for model development must be used: (1) test set (15 compounds), (2) 3rd set (out of initial dataset). The external validation of the 3rd set was conducted using D3R antagonists extracted from the ChEMBL database. ChEMBL compounds with D3R Ki data were imported and filtered using a cut-off of pKi < 0 uM. The selected compounds were then docked to 3PBL. This external validation was followed by advanced screening using ligands that could generate conformers, and that matched a minimum of 4 out of 5, or 5 out of 6 site points. The predictive values derived from this screening process were then compared with the experimental values.

The pharmacophore models were further validated by a receiver operator characteristic curve (ROC) analysis. Therefore, it was necessary to prepare a test set that included active and inactive compounds. These were imported into ChEMBL, and processed on the basis that the pKi values of active compounds (n = 436) were > 1.5, the pKi values of inactive compounds (n = 450) were < 1.0, the molecular weight was between 400 and 500, the ALogP was between
-2 and 6, the number of hydrogen bond donors is < 4, and the number of hydrogen bond acceptors is < 8. The ROC curve analysis describes the sensitivity (true positive rate, Se) for any possible change in the number of selected compounds as 1 – Sp (specificity, which is defined as the true negative rate) [24].

\[
\begin{align*}
\text{Se} &= \frac{\text{number of selected actives}}{\text{total number of actives}} = \frac{TP}{TP + FN} \\
\text{Sp} &= \frac{\text{number of discarded inactives}}{\text{total number of inactives}} = \frac{TN}{TN + FP}
\end{align*}
\]

In the ROC analysis, true positive (TP) means that the active compound is measured as active in the test. Likewise, true negative (TN) means to infer that inactive compound is measured as inactive. In contrast, false positive (FP) means that inactive is measured to active and false negative (FN) means that active is measured to inactive. A model with fewer FP and FN errors is superior model, and selectivity and specificity are the ratios of these. R software (version 3.3.2 probably need a reference) was used to plot ROC curves, calculate the area under the curve (AUC), and generate box plots, to compare the range of experimental and predictive values.

**Pharmacophore-based virtual screening**

The validated pharmacophore models were chosen for queries in our pharmacophore-based virtual screening [15]. Screening was conducted to identify hit compounds with chemical features corresponding to those of the template [25]. If a molecule can be fitted inside pharmacophore features, it could be considered a hit molecule based on the fitness score [26]. Existing conformers of all ligands in the database were included and screened for matches on at least 4 out of 5 or 5 out of 6 site points using the advanced pharmacophore method in PHASE (Schrödinger) running on a Linux-x86_64. The output of this QSAR model represented a maximum of 100,000 hits (≤ 1 hit per molecule) and considered atom types when computing volume scores. The commercially available compound library from eMolecules was used as the source of screening compounds to find novel hit compounds.

**Hit selection**

The screened in silico hits were docked to the D3R using the best docking condition identified when R22 was docked to the apo protein (3PBL). The processed compounds were then subjected to filtering, and the selected hit molecules were grouped by K-means clustering using binary fingerprints generated with MACCs in Canvas. They were divided into 10 clusters based on their molecular similarity. Representative compounds from these clusters were then selected after considering their estimated efficacy, docking scores, and CNS drug-like properties.
Figure 3: Docking pose showing the interaction between R-22 and 3PBL residues. In the left-hand picture, the blue line shows Pi-Pi stacking, the purple line shows hydrogen bond interaction, and the yellow-green line shows hydrophobic interaction.

A summary of the hit selection workflow is summarized in Figure 3. Hit compounds identified by screening using two models were sorted by fitness score in descending order; then, they were filtered using the following drug-like criteria based on Lipinski’s rule: AlogP < 5; molecular refractivity of 40 ~ 130; nitrogen-containing; Molecular weight (MW) of 180 ~ 500 and preferably around 400; Polar surface area (PSA) < 90; Hydrogen bonding donor (HBD) < 5; and Hydrogen bonding acceptor (HBA) < 10 \[\text{27}\]. Filtered hits were processed through Ligprep for conversion to 3D, docked to 3PBL to investigate their interaction with the target protein, and filtered using the secondary criteria of a docking score < -7, a ligand efficiency < -0.3 \[\text{28}\], and a charge range from -1 to 1. Qikprop in Maestro is used to apply CNS-like property filtering criteria of QPlogBB > -0.523, BB > 0.3 \[\text{29-31}\], and CNS ≥ 1; the score value of -2 is CNS inactive, and the score value of 2 is CNS active \[\text{32}\].

Results and Discussion
Docking analysis of D3R selective antagonists
The high degree of sequence identity within the transmembrane helixes of the D2R and D3R (78%) implies that these receptors have similar binding sites \[\text{33}\]. Due to this reason, it has been difficult to develop D3R-selective antagonists with drug-like physicochemical properties \[\text{34, 35}\]. To clarify the ligand-protein binding features of the D3R, we performed ligand docking to 3PBL using dopamine, eticlopride (D2R/D3R dual antagonist) \[\text{11}\], and the D3R-selective antagonist, R-22 under the various condition including (1) the centre, size, and shape of grid box, (2) constraints on non-covalent interactions with reported key residues (eg., Glu90, Asp110, Ser192, Ser193, Phe197, Tyr373, and His349), and (3) three scoring functions. As the result, the proposed binding mode of R-22 in the literature \[\text{12}\] was reproducible under the optimal docking condition showing the non-covalent interactions with important residues included Asp110, Tyr365, Tyr373, and Phe346 (Figure 3). From the docking condition, docked dataset compounds also showed the occupation of the outer shallow pocket as well as the deep pocket where the endogenous agonist binds to. Based on the binding mode, it is reasonable to develop selective D3R antagonists that target the shallow pocket. Conversely, the deep pocket is a common binding region where agonists and antagonists interact. It is a common expectation that docking within the deep pocket is necessary for targeting this protein, and the pharmacophore should therefore interact with both the shallow and deep pockets. In addition, ligands including the hydroxy group in the linker showed a reduced potency.
Pharmacophore modeling and 3D-QSAR models

The knowledge from docking analysis made us consider models containing five or six features to describe the interactions with two pockets. After building models to explore every plausible hypotheses from training set, the next scoring technique was applied to evaluate the models according to site, vector, volume, selectivity, conformational energy and activity.

Survival score = $W_{\text{site}} S_{\text{site}} + W_{\text{vec}} S_{\text{vec}} + W_{\text{vol}} S_{\text{vol}} + W_{\text{sel}} S_{\text{sel}} + W_{\text{m rew}} - W_{E \Delta E} + W_{\text{act}} A$

The selection of superior hypothesis was committed based on outstanding post-hoc survival score combining two survival scores of active and inactive. Top 10 pharmacophore models showing superior post-hoc scores were used to generate 3D-QSAR models. Internal validation of the chosen PLS models produced reliable coefficient of determination (R squared of each hypothesis > 0.7, Max. 0.88) and their high calibrated values (Q squared of each hypothesis > 0.6, Max. 0.73) in Table2. In addition, Pearson correlation constants also showed more than 0.8 in all of our chosen models. Finally, two models (APRRR215 and AHPRRR104) were selected for virtual screening based on these statistical analyses and the appropriate location of features (Figure 4). In the simple regression analysis of the experimental activities with predicted activities, the best model, APRRR215, presented $\text{Act}_{\text{pred.}} = 0.80 \text{ Act}_{\text{exp.}} + 0.49$ (R predictive squared = 0.80) in the training set and $\text{Act}_{\text{pred.}} = 0.62 \text{ Act}_{\text{exp.}} + 1.10$ (R predictive squared = 0.78) in the test set (Figure 5). Another chosen model, AHPRRR104, indicated that $\text{Act}_{\text{pred.}} = 0.82 \text{ Act}_{\text{exp.}} + 0.45$ (R predictive squared = 0.82) in the training set and $\text{Act}_{\text{pred.}} = 0.66 \text{ Act}_{\text{exp.}} + 0.98$ (R predictive squared = 0.74) in the test set (Figure 6).

Table 2. Statistic values of QSAR model
Figure 4: Representative 3D-QSAR models: APRRR215 (upper) and AHPRRR104 (lower).

Figure 5: APRRR215 regression lines for the training (blue) and test (red) sets.

Figure 6: AHPRRR104 regression lines for the training (blue) and test (red) sets.

The external validation of the best model, APRRR215 was further conducted using 3rd data set (known D3 antagonists with structural diversity) from the ChEMBL database.
Visualization of the validation was presented in ROC analysis to show how effectively the pharmacophore models distinguished between active and inactive compounds. Sensitivity (in other words, true positive rate, recall, hit rate) and specificity (in other words, true negative rate) are general indices to show the predictive power of a validated model. The ROC curve is a graphical representation of the false positive rate, \((1 – \text{specificity})\) of x-axis and the true positive rate, sensitivity of y-axis for each of the possible cut-offs \([36]\). The accuracy of this test is measured by the AUC of the ROC curve. According to the general judgement on an AUC value of ROC curve (AUC of poor model = 0.5, AUC of a moderate accurate model = 0.5 to 0.7, and very accurate if the AUC > 0.9) \([37]\), the AUC of APRRR215 was 0.696 showing moderate usefulness in global scaffolds (Figure 7). In addition, the distribution of the experimental and predictive values was visually compared through box-and-whisker plot in Figure 8. Even though the model presented the limit to residual between experimental value and predicted value of active compounds, median values of two distributions could be well-matched and two distributions were identical after normalization.

![Figure 7: APRRR215 ROC curve and AUC.](image)

![Figure 8: APRRR 215 skeletal box-and-whisker plot.](image)
Identification of CNS-like D3R antagonist through virtual screening.

The two reasonable pharmacophore hypotheses (APRRR215 and AHPRRR104) showing best statistics in their 3D-QSAR were used as templates for virtual screening [19]. The screening identified in silico hit compounds with chemical features corresponding to those of the template [19, 23, and 26]. Some of these hits might be similar to known active compounds, while others might have more novel scaffolds [23, 38]. The identification of hit compounds having different scaffolds was the primary aim of our research, development of novel D3R-selective antagonists. Our pharmacophore-based virtual screening played the role of high-throughput filter against compounds that lack the features essential for binding. The validated two best pharmacophore models allowed a researcher to efficiently screen a large commercial compound library from eMolecule (more than 3 millions), and the results were filtered to identify compounds with CNS drug-like properties (Figure 9).

Figure 9: Hit selection workflow
APRRR215 and AHPRRR104 resulted in 1,000,000 and 15,205 hits, respectively, and these were filtered by drug-like criteria, MW, AlogP, HBD, HBA, molecular refractivity, nitrogen-containing and PSA. In particular, PSA is commonly used to indicate whether a CNS drug can penetrate the cell membrane through passive diffusion. The PSA of a CNS drug is required to be < 90 angstroms to penetrate the blood brain barrier [39]. The numbers of filtered hit molecules were 48,379 for APRRR215 and 6,823 for AHPRRR104, with 5,770 compounds in common. Even though drug property filtering had already been performed in the pharmacophore hit selection steps, the CNS drug-like properties were reconsidered for the 3D conformation and filtered. In addition, binding poses and docking scores of 5,770 compounds with D3R (apo protein of PDB: 3PBL) were considered to present the most selective D3R antagonists interacting with both the deep and shallow pockets. Consequently, 374 compounds were extracted as filtered hits. The primary aim of this study was to find compounds with different scaffolds from the known D3R antagonists (a template), sharing the pharmacophore features. We identified 10 (a representative member of each cluster) novel hits predicted to act as dopamine D3R antagonists with appropriate estimated efficacies and pharmacokinetic properties (Table 3). The final selection also was experimentally validated by efficiency of the in silico D3R antagonists, of which beta arrestin assay (PathHunter®) was measured in D3R overexpressed cell line.
Table 3. Structure, estimated efficacy and ADME properties of selected hit compounds

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<th>Ro3</th>
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The current limitation of this study & future possibility

The external validation result of APRRR215 indicated that the QSAR model could show moderate predictive power with discriminating ability in binominal decision but could not predict pKi value on D3R with high accuracy. Plausible reasons for the large deviation between experiment and prediction in 3rd set are the following: (1) current models built from the long-shape drug scaffold can permit narrow deviation of a drug shape for the prediction, (2) D2R/D3R- selectivity ratio was used for choosing 3rd set (D2R/D3R ratio > 10 in active) but two variables, K_i value and the selectivity, can’t be independent and the selectivity as a confounding factor perfectly can’t be controlled in our current 3D-model, (3) flexibility of 3D-coformation, in particular, the flexibility of D3R antagonists in shallow pocket can make quantitative prediction difficult. In the future, plausible trials and advances from the trials are here: (1) more than two dependant variables (activity, D2/D3-selectivity, and so on), (2)
extension of dataset, and (3) well-designed conformational sampling of data set for building an advanced model [40].

Despite the limitation of current models, we can consider the advantages and contribution of the models. Firstly, our pharmacophore model can effectively contribute to finding novel compounds with high 3D similarity but low 2D similarity. In general, when designing a new scaffold from a known template molecule, medicinal chemists tend to consider 2D similarity than 3D. However, if a drug is 2D-dissimilar to existing drugs for the cognate target but has high 3D similarity, there is greater likelihood to obtain a novel pharmacological effect [38, 41, and 42]. In addition, the measured distances between the pharmacophores are consistent with proposed binding conformer of R-22 and each pharmacophore feature could describe non-covalent bonding interaction between R-22 and corresponding residues. Secondly, in the view of screening efficiency, the pharmacophore based screening was 100-fold faster than high-throughput docking without any pre-processing in one sub job. Under the condition of Intel Xeon E5-2650v2 (20M Cache, 2.60 GHz, 8.00 GT/s, 8x4 cores), HTVS docking of 100,000 chemicals from eMolecules (> 3 million compounds) consumed total 57959 seconds (16 hours) CPU time in 16 parallel sub jobs. Under the same resource, the pharmacophore based screening of whole eMolecule chemicals could completed within 72 hours in one job. Because parallel (thread) processing of both screenings does not have dependency between sub jobs, the pharmacophore based screening was also superior to docking in the speed of total job. Finally, double filtering through two screening and data fusion of scored values (e.g., the fitness score of pharmacophore model, predicted pKi of QSAR, and docking score) can improve predictive power [43, 44].

**Conclusion**

In this study, we have developed pharmacophore models of D3R selective antagonists to efficiently identify a series of compounds with novel scaffolds, having both predicted D3R antagonism and CNS drug-like properties. The models were built from docking conformers, and evaluated by combined survival scores of active and inactive of the models. And then two best models (APRRR215 and AHPRRR104) were chosen by predictive power among their 3D-QSAR models generated from Top10 among five/six-featured models. In the view of screening speed and performance, the pharmacophore models were efficient at finding 2D-novel and 3D-similar scaffolds with known D3R antagonists. Our docking model, as a second predictor, could compensate the limited accuracy of fitness score or predicted pKi through the data combination of the scored values. Moreover, CNS-like property was used as an additional criterion with the two models for choosing promising hit compounds. In conclusion, our current work shows that (1) efficient screening of a large compound database (more than 300 millions) is possible, and (2) the interpretation of models is well matched in pharmacophore features and docking poses.

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**Competing interests**

The authors declare no competing financial interests.

**Author contributions**

M.-h. K. and S. J. C. conceived and designed the study. Under the designed study, J. L. built his models, validated them and acquired in-silico hits through his models. S. J. C. validated J. L.’s models and implemented current optimal models in 3rd party program. M.-h. K. and J. L.
wrote the manuscript. S. J. C. revised the manuscript. All the authors read and approved the final manuscript.

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