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Identification of the Inhibitory Mechanism of FDA Approved Selective Serotonin Reuptake Inhibitors: Insight from a Molecular Dynamics Simulation Study

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ABSTRACT

Antidepressants selectively inhibiting serotonin reuptake (SSRIs) represent a highly-effective drug class, and novel therapeutic strategies were proposed to improve SSRIs' drug efficacy. The knowledge of the inhibitory mechanism of FDA approved SSRIs could provide great insights and act as important starting points to discover privileged drug scaffold with improved efficacy. However, the structure of human serotonin transporter (hSERT) has not yet been determined and the inhibitory mechanism underlying SSRIs is still need to be further explored. In this study, the inhibitory mechanism of 4 approved SSRIs treating major depression (fluoxetine, sertraline, paroxetine and escitalopram) was identified by integrating multiple computational methods. Firstly, a recently published template with high sequence identity was adopted for the first time to generate hSERT's homology model. Then, docking poses of 4 SSRIs were used as initial conformation for molecular dynamics (MD) simulation followed by MM/GBSA binding free energy calculation and per-residue free energies decomposition. Finally, a binding mode shared by 4 studied SSRIs was identified by hierarchically clustering per-residue free energies. The identified binding mode was composed of collective interactions between 3 chemical groups in SSRIs and 11 hot spot residues in hSERT. 6 out of these 11 were validated by previous mutagenesis studies or pharmacophore models, and the remaining 5 (Ala169, Ala173, Thr439, Gly442 and Leu443) found in this work have not yet been identified as common determinants of all 4 studied SSRIs in binding hSERT. Moreover, changes in SSRIs' binding induced by mutation on hot spot residues were further explored, and 3 mechanisms underlining their drug sensitivity were summarized. In summary, the identified binding mode provided important insights into the inhibitory mechanism of approved SSRIs treating major depression, which could be further utilized as framework for assessing and discovering novel lead scaffold.

KEYWORDS:

Major depression; Binding mode; Selective serotonin reuptake inhibitor; Molecular dynamics; Binding free energy

INTRODUCTION

Antidepressants inhibiting serotonin reuptake represent a highly-effective class of therapeutics treating major depression and generalized anxiety disorder¹, and are the first-line and the most prescribed class of antidepressant in the United States². Typical antidepressants of this class include selective serotonin reuptake inhibitors (SSRIs), serotonin-norepinephrine reuptake inhibitors (SNRIs)³ and others⁴⁻⁷. So far, this class has resulted in >10 marketed antidepressants approved by U. S. Food and Drug Administration (*Figure 1*) and several candidates in clinical trial. As the primary target of SSRIs^{8, 9}, human serotonin transporter (hSERT) plays a key role in regulating the duration and intensity of serotonin signal in the synaptic cleft¹⁰, and is thus closely related to the pathology of mood and anxiety disorders¹¹.

In order to deal with the delayed onset of action¹² and the partial- or non-response¹³ of SSRI treating depression, novel therapeutic strategies were frequently proposed as "improvement" to the drug efficacy^{14, 15}. These strategies include designing dual- and triple-acting antidepressants¹⁵, multi-targeting hSERT and acid sphingomyelinase¹⁶, enhancing SSRIs' efficacy by co-targeting low-affinity high-capacity hSERT¹⁷, and others^{18, 19}. The binding mode of certain SSRI to hSERT was reported to be crucial for revealing its specific inhibitory mechanism and drug efficacy²⁰⁻²². A detailed understanding of the binding mode shared by SSRIs of clinical importance, especially FDA approved 4 SSRIs (*Figure 1*), could provide more insights into the identification of privileged drug-like scaffold with improved drug efficacy^{21, 23}.

The determination of the crystal structure of hSERT's bacterial homolog LeuT^{24, 25} provided a template to elucidate binding mode of SSRIs-hSERT via computational methods²⁶ in the past decade. The most frequently used methods include induced fit docking (IFD)^{20-22, 27, 28}, pharmacophore model²⁹⁻³¹ and molecular dynamics (MD)³². IFD focuses on considering the flexibility of hSERT's amino acid side chains lining the binding cavity²⁷, while pharmacophore model is primarily designed for screening new SSRIs³¹. For a better mobile recognition mechanism of SSRIs-hSERT, MD simulation was used to reveal the binding of substrates and drugs to hSERT's S1 site^{26, 33-35}, a reported primary binding site of SSRIs' competitive inhibition³⁶⁻³⁸. However, escitalopram is so far the only approved SSRI, the binding mode of which to S1 site was analyzed by MD simulation^{32, 39}. Currently, no comprehensive study on binding modes of SSRIs-hSERT has been carried out. Thus, a

detailed understanding of the inhibitory mechanism underlining SSRIs' pharmacological activity and target recognition is in urgent need^{20, 36}.

In this study, we carried out an inhibitory mechanism identification of 4 FDA approved SSRIs treating major depression (fluoxetine, sertraline, paroxetine and escitalopram) by integrating multiple computational methods. Firstly, a newly reported template of *drosophila* dopamine transporter (dDAT)³⁶ with much higher sequence identity (53%) than LeuT (23%) was used for the first time to generate the homology model of hSERT. Then, SSRIs were docked into hSERT and used as initial conformation for MD simulation. Finally, a binding mode (at atomic level) shared by 4 studied SSRIs was identified by clustering per-residue binding free energies. As a validation, drug sensitive mutations inducing >10-fold shifts on inhibitory potency (Ki)⁴⁰ were further explored. The binding mode identified in this study provided important insights into the inhibitory mechanism of approved SSRIs, which could be further utilized as a useful framework for assessing and discovering novel lead scaffold²³.

METHODS

Homology Modeling

As illustrated in *SI, Figure S1*, a recently determined 3.0Å X-ray crystal structure (PDB code 4M48³⁶) of dDAT (from Glu26 to Asp599) showed a much higher sequence identity (53%) to hSERT than LeuT²⁴ (23%). By using it as template, the automated mode in SWISS-MODEL⁴¹ was applied to construct homology model of hSERT. The constructed model was from Glu78 to Pro617, which covers all 12 transmembrane regions and corresponding intervening loops. Then, Ramachandran plot in PROCHECK⁴² was applied to validate the constructed model. Finally, two functional Na⁺ in LeuBAT³⁷ (a LeuT variant engineered to harbor hSERT-like pharmacology by mutating key residues around the primary binding pocket) were manually fitted into their corresponding binding sites in hSERT via PyMOL's⁴³ structure superimposition.

Molecular Docking

To get their initial poses, 4 SSRIs were docked into the modeled hSERT using Glide⁴⁴ with standard precision. Residues identified as key determinants of SSRIs-hSERT binding^{21, 40} were used to define the docking grid box. Docking poses of fluoxetine, sertraline and paroxetine with the most similar conformations as those in LeuBAT³⁷, and docking pose of escitalopram with the most similar orientation with those of fluoxetine, sertraline and

paroxetine were selected as their initial binding poses for MD simulation. Moreover, to assess the validity of the docking approach in the studied model, a cross-docking strategy was adopted by docking SSRI into the structure of LeuBAT obtained with other SSRI bound (e.g. docking fluoxetine into the structure of LeuBAT obtained with sertraline bound). The co-crystallized poses of fluoxetine, sertraline and paroxetine (PDB codes 4MM8, 4MM5 and 4MM4) in LeuBAT³⁷ were superimposed to their corresponding cross-docking poses. Although, no co-crystallized structure of escitalopram in LeuBAT has been reported, its resulting docking pose is oriented in a very similar way as fluoxetine, sertraline and paroxetine in LeuBAT, which forms an ionic interaction between the ammonium group and residue Asp98. The detailed information can be found in *SI*, *Methods*.

Protein-Ligand/Membrane System Setup

SSRIs-hSERT complexes obtained by docking were embedded into the explicit POPC lipid bilayer using the Membrane Builder in CHARMM-GUI⁴⁵⁻⁴⁷. TIP3P water⁴⁸ of 20Å thickness was then placed above and below the constructed bilayer. Environmental salt concentration was kept at 0.15M by adding Na⁺ and Cl⁻. The overall system contained a total of ~96,000 atoms per periodic cell, and the box size was set as $83\text{\AA} \times 83\text{\AA} \times 127\text{\AA}$. The detailed information can be found in *SI, Methods*.

MD Simulation

MD simulation was performed within AMBER14⁴⁹ using GPU-accelerated PMEMD. AMBER force field *ff14SB*⁵⁰ and *Lipid14*⁵¹ were used for protein and lipids, respectively. The ions parameters for TIP3P water were collected from Joung & Cheatham⁵². The force field parameters for fluoxetine, sertraline, escitalopram, paroxetine and cholesterol were described by the General AMBER Force Field⁵³ and the charges were assigned using Restrained Electrostatic Potential partial charges⁵⁴ with Antechamber⁵⁵. Geometry optimization and the electrostatic potential calculations were performed with Gaussian09 at the HF/6-31G* level⁵⁶. For each simulation, a sophisticated protocol including minimization, heating and equilibration was employed. Then, 150ns MD simulation was conducted in NPT ensemble under a temperature of 310K and a pressure of 1 atm using a periodic boundary condition. Meanwhile, the direct space interaction was calculated by considering the long range electrostatic interaction (cutoff = 10.0Å) using particle-mesh Ewald method⁵⁷. All bonds involving hydrogen were constrained by SHAKE algorithm⁵⁸ allowing an integration time step of 2fs. The detailed information can be found in *SI, Methods*.

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Binding Free Energy Calculation

Binding free energy ($\Delta G_{MM/GBSA}$) of SSRIs to hSERT excluding entropic contribution was calculated using the single-trajectory based MM/GBSA method^{59, 60}. A total of 500 snapshots were taken from the last 50 ns MD simulation. For each snapshot, the SSRIs-hSERT binding free energy was calculated by:

$$\Delta G_{MM/GBSA} = \Delta E_{vdW} + \Delta E_{ele} + \Delta G_{pol} + \Delta G_{nonpol}$$
(1)

In Eq. (1), ΔE_{vdW} and ΔE_{ele} are the van der Waals and the electrostatic components in the gas phase, and ΔG_{pol} and ΔG_{nonpol} are the polar and the non-polar solvent interaction energies. ΔG_{nonpol} was calculated by $\Delta G_{nonpol} = 0.0072 \times \Delta SASA$ by the linear combination of pairwise overlaps method (LCPO)⁶¹, where SASA is the solvent accessible area. The detailed information can be found in *SI*, *Methods*.

Per-residue Free Energy Decomposition Analysis

To quantitatively evaluate the contribution to SSRIs binding, the total binding free energy was decomposed on a per-residue basis. The corresponding binding free energy $(\Delta G_{MM/GBSA}^{per-residue})$ excluding entropic contribution was given by:

$$\Delta G_{MM/GBSA}^{\text{per-residue}} = \Delta E_{vdW}^{\text{per-residue}} + \Delta E_{ele}^{\text{per-residue}} + \Delta G_{pol}^{\text{per-residue}} + \Delta G_{nonpol}^{\text{per-residue}}$$
(2)

The definition of quantities in Eq. (2) is similar to that in Eq. (1), except for the non-polar solvent interaction energy ($\Delta G_{nonpol}^{per-residue}$) which is calculated by recursive approximation of a sphere around an atom, starting from an icosahedron (ICOSA)⁴⁹. The detailed information can be found in *SI*, *Methods*.

Hierarchical Clustering with Ward Algorithm

Energy contributions of certain residues to 4 studied SSRIs calculated in the previous section were used to generate a 4 dimensional vector. Then, the per-residue energy contribution vector-based hierarchical clustering tree of 245 residues with contributions to at least one studied SSRI in hSERT's binding (contribution \neq 0 kcal/mol) was generated using the R statistic analysis software⁶² with the similarity levels among vectors measured by the Manhattan distance:

$$Distance(a, b) = \sum_{i} |a_{i} - b_{i}|$$
(3)

where i denotes each dimension of per-residue energies a and b. Cluster algorithm used here is the Ward's minimum variance method⁶³, which is designed to minimize the total

within-cluster variance. In this study, a Ward's minimum variance module in R package was used⁶². The hierarchical tree graph was generated using EMBL's online tree generator (iTOL⁶⁴ version 3.0). Per-residue binding free energy contributions favoring SSRI's binding were displayed in red, with the highest contribution set as exact red and lower contributions gradually fading towards white (no contribution). Meanwhile, per-residue energy contributions hampering SSRI's binding were shown in blue, with the highest one set as exact blue and lower ones gradually fading towards white.

RESULTS AND DISCUSSION

Homology Modeling and SSRIs-hSERT Complexes Construction

As shown in *SI*, *Figure S2*, superimposition of the modeled hSERT and the X-ray crystal structure of $dDAT^{36}$ showed high degree of homology. The structure of modeled hSERT consists of 12 transmembrane α -helices, and the corresponding residues within these helices agree well with the reported experimental results⁶⁵. The stereo chemical quality and accuracy of the predicted model was evaluated by Ramachandran plot in PROCHECK⁴², and 99.5% of the modeled residues were located in the "allowed region" (*SI, Figure S3*), indicating a reliable homology model.

Based on the modeled hSERT, molecular docking was carried out to determine initial poses of SSRIs in hSERT. Results (*SI, Figure S4*) show that fluoxetine, sertraline, escitalopram and paroxetine fit in the binding site primarily surrounded by TM1, TM3, TM6 and TM8. To assess the validity of the docking approach in the studied model, a cross-docking strategy was adopted by docking SSRI into the structure of LeuBAT obtained with other SSRI bound (for example, docking fluoxetine into the structure of LeuBAT obtained with sertraline bound). The co-crystallized poses of fluoxetine, sertraline and paroxetine (PDB codes 4MM8, 4MM5 and 4MM4) in LeuBAT³⁷ were superimposed to their corresponding cross-docking poses. Both poses kept consistence with each other (*SI, Figure S5*), which guaranteed correct initial poses generated in this work. So far, no co-crystallized structure of escitalopram in LeuBAT has been reported, but its resulting docking pose is oriented in a very similar way as fluoxetine, sertraline and paroxetine in LeuBAT, which forms an ionic interaction between the ammonium group and residue Asp98.

Assessing the Binding Mode of SSRIs-hSERT

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<u>Simulation Stability Analysis</u>. Initial SSRIs-hSERT complex generated by docking was assessed by 150ns MD simulation to reach equilibration state, which could be monitored by the structural root mean square deviation (RMSD) of both protein backbone atoms and ligand heavy atoms as a function of time. According to RMSD analysis (*Figure 2*), each simulation of hSERT in complex with 4 studied SSRIs reached equilibration around 100ns, and a time scale of 150ns was therefore set in this work to guarantee a stable state for the SSRIs-hSERT complex.

Binding Free Energy Analysis. Binding free energies of hSERT in complex with fluoxetine, sertraline, escitalopram and paroxetine were calculated, and the results ($\Delta G_{MM/GBSA}$) were -41.52, -44.32, -49.21 and -51.14 kcal/mol respectively. Meanwhile, experimental binding free energies (ΔG_{exp}) were also estimated based on reported Ki values⁴⁰ using $\Delta G_{exp} = -RTln(K_i)$. As shown in *Table 1*, $\Delta G_{MM/GBSA}$ in this work were overestimated comparing to those of experiment⁴⁰. For ligands with similar structures and binding modes, entropy contribution could be omitted if one is only interested in the relative order of binding affinities⁶⁶, so differences of energies calculated in this study ($\Delta\Delta G_{MM/GBSA}$) and estimated based on experiments $(\Delta\Delta G_{exp})$ among SSRIs (*Table 1*) could help to understand whether the overestimation came from the exclusion of entropy. As illustrated in SI, Figure S6, $\Delta\Delta G_{MM/GBSA}$ correlates very well with $\Delta\Delta G_{exp}$ (R² = 0.93). The ascending trend of energy difference $(\Delta\Delta G_{exp})$ from experiment⁴⁰ was reproduced very well by $\Delta\Delta G_{MM/GBSA}$ in this work, though their values were still estimated higher than experiment⁴⁰. The higher energy calculated in this work was in accordance with the reported over estimation of binding affinity by the MM/GBSA methods⁶⁷⁻⁶⁹. Calculated contributions of each energy term in Eq. (1) were listed in SI, Table S1. As shown by numbers in bold, the binding of SSRI to hSERT was primarily driven by hydrophobic (ΔE_{vdW}) and coulomb interaction energies (ΔE_{ele}), but hampered by the polar solvent energy (ΔG_{pol}).

Analyzing the Binding Mode of SSRIs-hSERT

In this study, the binding mode of each SSRIs-hSERT was obtained by molecular docking and further assessed by MD simulation. As illustrated in *Figure S7*, slight conformation shifts between 4 SSRIs' docking poses and their representative snapshots in the equilibrated MD trajectories were observed, but the key interactions such as the salt bridge and the hydrogen bond between the ligands and Asp98 of hSERT were preserved.

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Representative snapshots of SSRIs-hSERT complexes from equilibrated MD trajectories are also shown in *SI*, *Figure S8*, illustrating the orientations of each SSRI and its interacting residues. However, to fully understand the binding mode of SSRIs-hSERT, it is essential to quantitatively distinguish contributions of each amino acid to the drug binding. Thus, plots showing binding free energy contributions of each amino acid were generated by a per-residue free energy decomposition analysis (*Figure 3*). To the best of our knowledge, *Figure 3* is the first reported SSRIs' binding free energy contributions at a per-residue basis.

As illustrated in *Figure 3*, there were 11, 12, 14 and 14 residues identified as high contribution ones (with the absolute energy contribution ≥ 0.5 kcal/mol) for binding fluoxetine, sertraline, escitalopram and paroxetine respectively. On one hand, energy contributions of different residues to the same SSRI vary greatly (for example, from -0.57 kcal/mol for Ala173 to -3.68 kcal/mol for Tyr95 in fluoxetine's binding), and energy contributions of the same residue to different SSRIs also differ significantly (the polar residue Asp98's contributions ranged from -1.83 kcal/mol for escitalopram to -2.75 kcal/mol for fluoxetine, as an example). On the other hand, *Figure 3* also infers a certain level of similarity among 4 studied SSRIs, which inspired us to conduct a deeper exploration of binding mode shared by 4 SSRIs.

Identifying the Binding Mode Shared by Approved SSRIs

Binding mode shared by approved SSRIs to hSERT could provide a useful framework from which novel lead scaffolds can be assessed and discovered²³. To identify the binding mode, the hierarchical clustering with ward algorithm⁶³ was applied to find hot spot residues from those per-residue binding free energies generated in the previous section. In particular, there were 541 residues with available binding free energy contribution to 4 studied SSRIs, 296 of which have no contribution (= 0 kcal/mol) to the binding of any studied SSRIs. Then, energy contributions of the remaining 245 residues were used for clustering analysis. As illustrated by the hierarchical tree in *Figure 4*, 4 distinct residue groups (A, B, C and D) were identified. Binding free energy contributions favoring SSRI's binding were displayed in red, with the highest contribution (-3.68 kcal/mol) set as exact red and lower contributions gradually fading towards white (contribution = 0 kcal/mol). Meanwhile, binding free energy contributions hampering SSRI's binding were shown in blue, with the highest one (0.22 kcal/mol) set as exact blue and lower ones gradually fading towards white. It is necessary to clarify that the absolute value of the highest contribution favoring SSRI's binding (exact red) is 16 times stronger than that hampering the binding (exact blue).

It is clear to see that binding free energy contributions of residues in group A (Tyr95, Asp98, Ala169, Ile172, Ala173, Tyr176, Phe341, Ser438, Thr439, Gly442 and Leu443) are consistently higher across 4 studied SSRIs in favoring the binding than those in group B, C and D. For each SSRI, the sum of energy contributions of residues in group A constitutes the primary portion of the total energy (77.04% for flouxetine, 74.01% for sertraline, 60.19% for escitalopram and 71.73% for paroxetine). Therefore, these 11 residues in hSERT were identified as hot spots for the binding of 4 SSRIs. Moreover, energy contributions of residues in subgroup A1 (Tyr95, Asp98, Ile172 and Tyr176) are more significant than those in subgroup A2 (Ala169, Ala173, Phe341, Ser438, Thr439, Gly442 and Leu443), and the sum of energy contributions of residues in subgroup A1 must be subgroup A1 consists of 48.55%, 37.68%, 35.34% and 40.87% of total energies for flouxetine, sertraline, escitalopram and paroxetine, respectively. Thus, residues in subgroup A1 and A2 could be classified as hot spots with "strong" and "relatively strong" contributions.

Moreover, the root mean square fluctuation (RMSF) versus the protein residue numbers of hSERT was calculated and illustrated in *SI*, *Figure S9*. It is observed that the hot spot residues of hSERT (Tyr95, Asp98, Ala169, Ile172, Ala173, Tyr176, Phe341, Ser438, Thr439, Gly442 and Leu443) show relatively rigid behaviors for the studied 4 complexes. The relatively low RMSF values of those residues suggest that SSRIs binding could stabilize the key residues located at the drug binding site. This is in accordance with the observed stabilization effect in previously study⁷⁰.

Conformational features of how SSRIs accommodate into 11 hot spot residues are shown in *Figure 5*. A pocket was defined by hot spot residues with a slight conformational shift, and all SSRIs fitted the pocket in a resembled orientation. Therefore, the binding mode of SSRIs-hSERT recognition was generalized and schematically represented in *Figure 6*. As shown, the binding mode was defined by collective hydrophilic and hydrophobic interactions between 3 chemical groups (R1, R2 and R3) and 11 hot spot residues (Tyr95, Asp98, Ala169, Ile172, Ala173, Tyr176, Phe341, Ser438, Thr439, Gly442 and Leu443). As shown in *Figure 5* and *Figure 6*, hot spot residues with strong and relatively strong contributions were illustrated in black and gray color, respectively. Chemical groups in *Figure 6* here were highlighted in blue and green. In particular, R1 formed salt-bridge and hydrogen bond interactions with Asp98; R2 interacted with Ala169, Ile172, Tyr176 and Phe341 via

hydrophobic contacts; R3 formed hydrophobic interactions with Tyr95, Ala173, Ser438, Thr439, Gly442 and Leu443, and also contacted with Ile172, Tyr176 and Phe341.

Among those 11 identified hot spot residues, 6 (Tyr95, Asp98, Ile172, Tyr176, Phe341 and Ser438) were collectively validated by previous mutagenesis studies^{20-22, 40, 71, 72} as sensitive to 4 approved SSRIs. Co-crystallized structures of SSRIs in hSERT's homolog^{37,73} could shade great light on their binding mechanism, but so far no structure of escitalopram in hSERT's homolog has been reported. Based on co-crystallized structures of 3 SSRIs (fluoxetine, sertraline and paroxetine), 7 residues (Tyr95, Asp98, Ile172, Ala173, Tyr176, Phe341 and Gly442) were suggested as primary binding by visualizing interaction distances between SSRIs and LeuBAT³⁷. Besides of mutagenesis studies and crystal structures, computational methods such as induced fit docking^{20-22, 27, 28}, pharmacophore model²⁹⁻³¹ and molecular dynamics³² collectively identified 4 residues (Tyr95, Asp98, Ile172 and Tyr176) as features for 4 SSRIs' binding^{21, 22, 31, 71, 72}. All primary binding residues found in co-crystallized structures and features identified by computational methods are covered by those 11 hot spot residues of this study. Besides those 6 residues (Tyr95, Asp98, Ile172, Tyr176, Phe341 and Ser438) identified by mutagenesis studies and computational methods, 5 hot spot residues found in this work have not yet been identified by previous studies as common determinants of 4 approved SSRIs in binding hSERT. However, among these 5 hot spots, 2 (Ala173 and Gly442) were suggested to interact with fluoxetine, sertraline and paroxetine³⁷, 1 (Thr439) was found as sensitive to fluoxetine²² and escitalopram⁷², and 2 (Ala169 and Leu443) were only reported to be in close proximity to the cyano group of escitalopram^{21,72}. Thus, Ala169, Ala173, Thr439, Gly442 and Leu443 were identified for the first time as common determinants of 4 FDA approved SSRIs treating major depression in binding hSERT.

Besides group A in *Figure 4*, group B (especially subgroup B1) represents residues with significant differences in binding free energy contribution. The sum of energy contributions of residues in subgroup B1 (Ala96, Leu99, Gly100, Trp103, Tyr175, Phe335, Ser336, Gly338 and Asp437) consists of 8.09%, 8.60%, 25.41%, and 10.33% of total energies for fluoxetine, sertraline, escitalopram and paroxetine, respectively. Particularly, Phe335 and Leu99 were main energy contributors to escitalopram's binding. Take the residue Phe335 as an example, its contribution to escitalopram's binding was relatively strong (-1.99 kcal/mol), while its contributions to other 3 SSRIs' binding were much weak with the highest contribution to paroxetine (-0.42 kcal/mol). As illustrated in *Figure S8C*, the strong contribution of Phe335

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may come from its interaction with escitalopram's cyano group which is in the close vicinity of Phe335. Meanwhile, Gly442 and Leu443 showed stronger interaction with fluoxetine, sertraline and paroxetine than escitalopram, and Ala96 showed stronger interaction with fluoxetine than other SSRIs. Understanding energy differences at per-residue basis could help to interpret drug selectivity, which may eventually aid the development of future selective antidepressants.

Group C represents residues with contributions hampering the binding of 4 SSRIs to hSERT. It is necessary to clarify that the absolute value of the highest contribution favoring SSRI's binding (exact red) is 16 times stronger than that hampering the binding (exact blue), which makes the contribution favoring the binding overwhelming. Moreover, Arg104 and Asn368 were identified in *Figure 4* as residue favoring (fluoxetine, sertraline and paroxetine) or hampering (escitalopram) the binding of SSRIs to hSERT.

In-silico Analysis of Drug Sensitivity Profile on Hot Spot Residues

Analyzing Drug Sensitive Mutation via Binding Free Energy. Analysis of the sensitivity profiles could provide insight into SSRIs' binding mode²¹. Sensitivity of certain residues to SSRI's binding can be reflected by comparing the difference in energy contributions before and after *in-silico* mutation on the corresponding residue. It is easy to understand that mutations on hot spot residues are critical to drug sensitivity. Therefore, this study investigated the influence of hot spot residues-mutation on SSRIs' binding. In particular, 12 single-point mutant complexes were analyzed individually by adding 20ns simulation using the assessed wild type model as the starting point (*SI, Figure S8*). Binding free energies were also calculated (*SI, Table S1*). As a comparison, impacts of hSERT mutations on binding affinities of SSRIs from previous experimental study⁴⁰ were listed in *SI, Table S1*. A significant decrease in Ki from wild type to mutant in experiment⁴⁰ was reproduced very well by $\Delta G_{MM/GBSA}$, whereas two exceptions (Phe/Tyr341 and Ser/Thr438) from paroxetine's binding were also observed.

Exploring the Mechanism of Hot Spot Residues Mutation Induced Drug Sensitivity.

Drug sensitive mutation on hot spot residues led to conformational changes in hSERT's binding pocket, and orientation shifts of SSRIs to accommodate into the pocket were observed (*Figure 7 (A1-C1)* and *SI, Figure S9*). This structural rearrangement further led to changes in binding free energy contribution of each amino acid (*Figure 7 (A2-C2)* and *SI, Figure S10*). By correlating mutations on hot spot residues with changes in binding free

energy contribution, 3 mechanisms underlining mutation-induced drug sensitivity were summarized.

In the first mechanism, mutation-induced drug sensitivity comes directly from a significant binding free energy contribution loss in mutational residues. Taking the Tyr/Ala95 of fluoxetine's binding as an example, in spite of an increase (-0.69 kcal/mol) in Ser438, binding free energy contribution was dramatically decreased (3.73 kcal/mol) in the mutational residue Tyr/Ala95, which resulted in unfavorable binding (*Figure 7 A2*). The detailed structural rearrangements (*Figure 7 A1*) further clarified that reduction in the side chain size from Try to Ala was the main reason of mutation-induced drug sensitivity.

In the second mechanism, mutation-induced drug sensitivity originates from binding free energy contribution loss in residues other than the mutational one. Taking the Ile/Met172 of fluoxetine's binding as an example, in contrast to unnoticeable energy change in the mutational residue Ile/Met172, binding free energy contribution was decreased by 0.87 kcal/mol in Tyr176 leading to unfavorable binding (*Figure 7 B2*). As shown in *Figure 7 B1*, a clear increase in the distance between fluoxetine and its interacting residue Tyr176 was induced by Ile/Met172 mutation, which resulted in the decrease in Tyr176's binding free energy contribution.

In the third mechanism, mutation-induced drug sensitivity comes from binding free energy contribution loss from both mutational and other residues. Taking the Phe/Tyr341 of escitalopram's binding as an example, besides a great decrease by 1.14 kcal/mol in the mutational residue Phe/Tyr341, binding free energy contribution was also decreased by 0.82 and 1.05 kcal/mol in Leu99 and Ser438 respectively (*Figure 7 C2*). Phe/Tyr341 mutation induced an clear structural rotation of escitalopram in the binding pocket, and leaded to unfavorable interactions between drug and two other residues (Leu99 and Ser438 in *Figure 7 C1*), which contributed to the mutation-induced drug sensitivity.

As shown in *Figure 7* and *SI*, *Figure S9* and *S10*, all residues with significant change in binding free energy contribution between the mutant and wild type hSERT were from those 11 hot spot residues or residues in their close vicinity, which further validates the binding mode of SSRIs' binding to hSERT identified in this study.

CONCLUSION

In this study, identification of the inhibitory mechanism from 4 approved SSRIs treating major depression was carried out by integrating multiple computational methods. A recently reported template dDAT was successfully adopted to generate the homology model of hSERT, and the binding mode shared by 4 studied SSRIs was identified by hierarchically clustering per-residue binding free energies of 245 residues with contributions to at least one studied SSRI in hSERT's binding. The identified binding mode was defined by interactions between SSRIs and 11 hot spot residues (Tyr95, Asp98, Ala169, Ile172, Ala173, Tyr176, Phe341, Ser438, Thr439, Gly442 and Leu443) in hSERT. 6 out of these 11 hot spot residues were validated by previous mutagenesis studies or pharmacophore models. 5 hot spot residues (Ala169, Ala173, Thr439, Gly442 and Leu443) found in this work have not yet been identified as common determinants of all 4 studied SSRIs in binding hSERT. In the last section of this study, changes in SSRIs' binding induced by mutation on hot spot residues were further explored, and 3 mechanisms underlining their drug sensitivity were summarized. The binding mode identified in this study provided significant insights into the inhibitory mechanism of approved SSRIs, which could be utilized as a useful framework for assessing and discovering novel lead scaffold.

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TABLES

Table 1. Comparison between the calculated and experimental binding free energies among 4 studied SSRIs to the wild type hSERT (ΔG is in kcal/mol and K_i value is in nM)

Systems	K _i ^a	$\Delta G_{ m exp}{}^b$	$\Delta\Delta G_{ m exp}$	$\Delta G_{\text{MM/GBSA}}{}^{c}$	$\Delta\Delta G_{ m mm/gbsa}$
Fluoxetine	255 ± 61	-9.00	0	-41. 52 ± 0. 10	0
Sertraline	242 ± 33	-9.03	-0.03	-44. 32 ± 0. 12	-2.80
Escitalopram	32 ± 1	-10. 23	-1.23	-49. 21 ± 0. 12	-7.69
Paroxetine	24 ± 6	-10. 40	-1.40	-51. 14 ± 0. 12	-9.62

^{*a*} Experimental value from Sørensen's work⁴⁰.

 $^{\textit{b}}$ Estimated binding free energy based on K_i values using $\Delta G_{exp} = -RTln(K_i).$

^c Calculated binding free energy in this work.

FIGURES



Figure 1. Structures of the SSRI and SNRI antidepressants approved by FDA.



Figure 2. Root mean square deviations of protein backbone atoms (A) and ligand heavy atoms (B) as a function of time in MD simulations.



Figure 3. Per-residue decomposition of binding free energy contributions of 4 SSRI-hSERT complexes.



Figure 4. Hierarchical clustering tree of 245 residues with contributions to at least one studied SSRI in hSERT's binding by their per-residue energy contributions. Per-residue binding free energy contributions favoring SSRI's binding were displayed in red, with the highest contribution set as exact red and lower contributions gradually fading towards white (no contribution). Per-residue energy contributions hampering SSRI's binding were shown in blue, with the highest one set as exact blue and lower ones gradually fading towards white.



Figure 5. The binding mode of SSRIs (A) fluoxetine, (B) sertraline, (C) escitalopram and (D) paroxetine to 11 hot spot residues in hSERT. Residues and SSRIs were shown in stick representation, and salt bridges and hydrogen bonds were depicted as blue and red dotted lines, respectively.



Figure 6. Schematic representation of the binding mode of 4 studied SSRIs to hSERT identified in this work.



Figure 7. Three mechanisms underlining drug sensitivity induced by mutations on hot spot residues. Structural superimposition of (A1) fluoxetine binding to wild type and Tyr/Ala95 mutant hSERT, (B1) fluoxetine binding to wild type and Ile/Met172 mutant hSERT and (C1) escitalopram binding to wild type and Phe/Tyr341 mutant hSERT. Contribution changes in SSRI's binding (A2-C2) by mutation on hot spot residues measured by the per-residue binding free energy corresponding to A1-C1. Residues affected were shown as a stick representation in wild type (light brown) and mutant (lightblue) models. The mutation residues were highlighted in red dash line circle and font.



Graphical Table of Contents

A binding mode shared by 4 FDA approved SSRIs treating major depression was identified by integrating multiple computational methods.