Seeking the exclusive binding region of phenylalkylamine derivatives on human T-type calcium channels via homology modeling and molecular dynamics simulation approach

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Abstract
Pharmaceutical features of phenylalkylamine derivatives (PAAs) binding to calcium channels have been studied extensively in the past decades. Only a few PAAs have the binding specificity on calcium channels, for example, NNC 55-0396. Here, we created the homology models of human Ca\textsubscript{3.2}, Ca\textsubscript{3.3} and use them as a receptor on the rigid docking tests. The nonspecific calcium channel blocker mibebradil showed inconsistent docking preference across four domains; however, NNC 55-0396 had a unique binding pattern on domain II specifically. The subsequent molecular dynamics (MD) simulations identified that Ca\textsubscript{3.1}, Ca\textsubscript{3.2}, and Ca\textsubscript{3.3} share domain II when Ca\textsuperscript{2+} appearing in the neighbor region of selective filters (SFs). Moreover, free-energy perturbation analysis suggests single mutation of lysine at P-loop domain III, or threonine at the P-loop domain II largely reduced the total amount of hydration-free energy in the system. All these findings suggest that P-loop and segment six domain II in the T-type calcium channels (TCCs) are crucial for attracting the PAAs with specificity as the antagonist.

**KEYWORDS**
homology modeling, phenylalkylamine, selective binding, T-type calcium channels, virtual drug screening
1 | INTRODUCTION

T-type calcium channels (TCCs) belong to the one sort of voltage-dependent calcium channel family. They are known to be activated by membrane depolarization, conducting inward currents with a small single-channel conductance. Roles of TCCs in controlling the hormone and neurotransmitter release under various conditions have been extensively studied in the past decades.\textsuperscript{1-5} The distribution of TCCs can be found in pancreatic β-cells,\textsuperscript{6} heart,\textsuperscript{7} and neuron cells.\textsuperscript{8} In β-cells, overexpressed Cav3.2 followed by generating frequent random calcium spikes.\textsuperscript{9} Humans with gradually raised calcium concentration will more likely develop type 2 diabetes in later life.\textsuperscript{10} In cardiomyocytes, the TCC currents significantly affect the later stage of the action potential.\textsuperscript{11} In neuron cells, the development of chronic neuropathic pain due to spinal cord injury is contributed by the increased activity of TCCs.\textsuperscript{12}

When the inhibitor binds to the calcium channel, it cuts down the Ca\textsuperscript{2+} pathway by allosterically changing the pore conformation or physical blocks in the pore as a plug.\textsuperscript{13} As the earliest launched TCC inhibitor, mibefradil was initially developed for blocking L-type calcium channel (LCC) and showed the promising effect of blocking TCCs in vitro.\textsuperscript{14} Such phenylalkylamine derivatives (PAAs) are more likely to behave as a physical plug when interacting with TCCs. Unfortunately, mibefradil was withdrawn from the market due to the interaction with other drugs due to its effect on P450.\textsuperscript{15} Because phenylalkylamine-based TCC antagonists are derived from LCC blockers, only some of them have specificity to block the TCC in vivo.\textsuperscript{16} Recently discovered drug Z944\textsuperscript{16} showed an excellent specificity and potent to block the TCC for treating epilepsy\textsuperscript{17} and neuronal pain.\textsuperscript{18} In contrast to phenylalkylamines that physically block the pore, Z944 was reported to change the conformation of TCC by shifting the α to π helix at domain II and further shut down the calcium currents.\textsuperscript{19}

In this study, we created two homology models for human Ca\textsubscript{3.2} (UniProt id: O95180) and Ca\textsubscript{3.3} (UniProt id: Q9P0X4) in terms of the structure of human Ca\textsubscript{3.1} at apostate. A total of three TCC structures were used in subsequent molecular dynamics (MD) simulations. Mibefradil, NNC-55-0396 were selected as the representative ligands in the MD to mimic the real cases when TCCs interact with different sorts of an antagonist. Also, the external electric force field MD and steering MD simulation was utilized to find the pathway of Ca\textsuperscript{2+} permeation when it penetrates the channel pore with and without introducing the blockers.

2 | MATERIALS AND METHODS

2.1 | Protein comparative modeling

Two human TCC 3D structures (α\textsubscript{1},H and α\textsubscript{1},l) were created by using the comparative modeling package from Rosetta.\textsuperscript{20} The Cryo-EM structure of human Ca\textsubscript{3.1}(PDB id: 6KZO) at the apostate was selected as input for Rosetta to create the homology model of α\textsubscript{1},H and α\textsubscript{1},l. The energy-based clustering method\textsuperscript{21} was applied to categorize the predicted models before the quality evaluation process. To filter out the low quality of structure, PROCHECK\textsuperscript{22} and WHATCHECK\textsuperscript{23} were employed on clustered data. Finally, the structures that hold the lowest Rosetta energy score will be selected as targets for the next tests, that is, rigid docking and MD simulations.

2.2 | Rigid docking with selected PAAs

The 3D structure of mibefradil and NNC-55-0396 was downloaded from the PubChem online database. We used Open Babel\textsuperscript{24} to convert the compound format and Frog\textsuperscript{25} to find the coordinate of compound conformers in 3D space. To find out the possible binding sites between existing antagonists and TCCs, we used AutoDock Vina\textsuperscript{26} to simulate the rigid docking process. The searching box was set at the center of the SF with the grid spacing 0.375 Å and 40 grid points along X, Y, and Z directions. The number of predict binding modes is set as 10, and the random seed number was set as −1460306363.

2.3 | MD simulations

2.3.1 | Ion channel membrane assembly

The CHARMM-GUI Membrane Builder\textsuperscript{27} was used to build the membrane system. The missing residues were modeled by GalxyFill. The heterogeneous lipid bilayer was created by choosing phosphatidylcholine lipid and phosphatidylethanolamine lipid ratio as 2 over 1 on both inner and outer leaflets of the membrane. The water thickness was set as 22.5 Å. The system size along the X and Y dimensions is set as 120 Å. About 150 mM calcium chloride solution was added into the 3D rectangle computational domain. To generate the parameter and force field file for selected TCC blockers, the Ligand Reader and Modeler\textsuperscript{28} was used. Overall, the system was formed as an isothermal-isobaric ensemble at temperature 310 K.

2.3.2 | Classic MD simulation

The force field to support all MD simulations was set as CHARMM36m. All MD productions were conducted on the precompiled NAMD-2.14b GPU-acceleration Linux version. Since MD cases prepared by CHARMM-GUI Membrane Builder need to be equilibrated before long-term standard running. The energy minimization process was performed under gradually reduced restrain forces over six-constitute steps. To keep the system stable, the integration time step was set as 2 femtoseconds (fs) for the first and second steps, then adjust to 1 fs for the others. As long-term standard MD production, the integration time step was set as 2 fs and the position and velocity of all atoms in the system will be recorded every 10 picoseconds (fs). The maximum periodic electrostatics calculation was based on the 1 Å grid size with periodic boundary conditions. To
calculate the nonbound interactions, a cutoff of 12 Å is used with
switch distance 10 Å as input conditions for solving Langevin equa-
tions at temperature 310 K and 1 atm. For long-term production,
the trajectory information was collected over 100 nanoseconds (ns)
timescale with zero restrictions added to it.

2.3.3 | Simulation under the external electric force field

The Cryo-EM structure of human α,βG at the apo state shows the
opening channel pore without ligand binding. To simulate the open
channel process, an external voltage was required to apply to the
lipid membrane. The external electric potential $E_z$ along $L_z$ can be
defined by the equation: $E_z = \frac{V}{L_z \times 4(\pi)}$, where $V = -60 \text{ mV}$ and
$L_z$ is the average length of TCCs in Z direction at the last 10 ns from
standard MD simulation. The numerical value of 43.17 was the force
conversion coefficient used by NAMD-2.14b. Total MD simulation
time under $-60 \text{ mV}$ for all TCCs was set as 100 ns.

2.3.4 | Structure identification through RMSD

For all the MD simulation cases, the structure clustering analysis was
conducted when the overall system reaches equilibrium, typically, in
the last 50 ns. The optimized number of clusters for a given length of
MD data was determined by a comparison of the clustering results
from various methods.31

2.3.5 | Steering MD

To simulate the Ca$^{2+}$ influx through the channel pore, the steering
molecular dynamics (SMD) method was employed. The clustering
analysis of external electrical force field simulation provided the
three most typical conformations for running SMD. We picked up
one Ca$^{2+}$ which most close to the SF as a pulling target. Then
we applied a constant pulling velocity: $V = -4e-5 \text{ Å/fs}$ with spring
constant $K = 4 \text{ kcal/mol Å}$ along $z$ direction for 3 ns simulation. To
minimize the artificial impact from fixed atoms in SMD simulation,
we fixed the C-alpha atoms at four geometrical symmetrical amino
acids that away from channel pore, in which Pro$^{116}$, Pro$^{849}$, Gln$^{1213}$,
and Val$^{1704}$ for α,βG; Pro$^{116}$, Pro$^{801}$, Pro$^{1316}$, and Thr$^{1635}$ for α,H;
Pro$^{116}$, Pro$^{649}$, Pro$^{213}$, and Thr$^{1528}$ for α,l, respectively. With Tcl
scripts to define all necessary parameters, the repeating rate of each
structure was set as 20.

2.3.6 | Finding calcium ion interaction pathway

The trajectory files generated by NAMD over the entire time-span
were analyzed in PyContact.32 To find the interactions between Ca$^{2+}$
and select molecules, the maximal interatomic distance for contact
and select molecules, the maximal interatomic distance for contact

| TABLE 1 | Protein basic local alignment search tool (BLAST) results on the identity score of a given length of amino acids between template proteins and targets. |
|-----------------|-----------------|-----------------|
|                | H-Ca $^{3.1}$ (full length) | R-Ca $^{1.1}$ (full length) | Bac-Ca $^{Ab}$ (full length) |
| H-Ca $^{3.2}$  | 62.58%            | 31.47%           | 23.93%          |
| H-Ca $^{3.3}$  | 57.02%            | 31.09%           | 24.46%          |

Abbreviations: Bac-Ca$^{Ab}$ (SKMH), bacterial Ca$^{Ab}$; H-Ca$^{3.1}$, human Ca$^{3.1}$; R-Ca$^{1.1}$, rabbit Ca$^{1.1}$ (3JBR).

scoring was set as 5 Å, the cutoff angle for hydrogen bonds was set
as 120°, and maximal distance between the hydrogen bond receptor
and the H-atom was set as 2.5 Å. Accumulate score was used to eval-
uate the interaction strength between Ca$^{2+}$ and selected molecular.

2.3.7 | Free energy perturbation analysis

The alchemical free-energy perturbation (FEP) method33 was used to
estimate the influence of Gibbs-free energy by mutation of pre-
identified binding associated amino acids.19 $K_0^{1446}$ and $T_0^{921}$ at α,βG,
$K_0^{1405}$ and $T_0^{2784}$ at α,l, and $K_0^{1302}$ and $T_0^{742}$ at α,l are mutated
into alanine. Two FEP simulations were conducted for molecular in
vacuum or immersed in bulk water. In every simulation, 20 equally
stratified external parameters $\lambda$ in the range [0, 1] were used to
sample the energy variation in 50 ps with coupled intramolecular
interactions. The net solvation free energy change was computed
as follows: $\Delta \Delta G_{\text{mutation}} = \Delta G_{\text{hydr}}^1 - \Delta G_{\text{hydr}}^2$, where $\Delta G_{\text{hydr}}^1$ is hydration
energy in the solvation state and $\Delta G_{\text{hydr}}^2$ is the hydration energy at
the isolate state. The results of FEP were processed by ParseFEP
in VMD 1.9.3 at temperature 310 K, Gram–Charlier order at 3, with
Gaussian approximation and BAR estimator.35

2.4 | Nomenclature of targets and ligands

Key protein targets and ligands in this article are hyperlinked
to corresponding entries in http://www.guidetopharmacology.org,
the common portal for data from the IUPHAR/BPS Guide to
PHARMACOLOGY,36 and are permanently archived in the Concise
Guide to PHARMACOLOGY 2019/20.37

3 | RESULTS

3.1 | Homology modeling

Compare to other calcium channel templates, 6KZO returns the
standard protein BLAST identity value as 62.58%, 57.02% for the
full length of amino acid (AA) sequences (Table 1). We only consid-
ered amino acids aligned with 139 to 1906 to generate the homology
model for α,l and α,l to reduce the computational cost. The detailed
alignment of amino acids for three TCC can be found in the supporting
information (Figure S1). Based on the target protein AA sequences, the energy-based clustering method has identified 10 and 19 featured structures for $\alpha_1$H and $\alpha_1$I, respectively. The Rosetta Comparative Modeling method uses the continuous alignment results of $\alpha_1$H and $\alpha_1$I to $\alpha_1$G for generating the homology models. The AAs that are not included in 6KZO will affect the quality of predicted protein structures. We found that after trimmed these regions out, the highest Ramachandran Z-score can go to 99.5% (Figure 1A1), and side-chain planarity can reach 0.42 for selected $\alpha_1$H homology modeling result (Figure 1A2). Meanwhile, the highest Z-score and side-chain planarity is 99.7% (Figure 1B1) and 0.425 for trimmed $\alpha_1$I homology modeling results. The full structure comparison before and after trimmed for the unnecessary region of $\alpha_1$H and $\alpha_1$I is summarized in Table S1.

### 3.2 | Rigid docking with selected PAAs

Vina virtual screening process finds a similar binding pattern between receptor and ligands. The hydroxy-group of S4p1776 domain IV at $\alpha_1$G forms hydrogen bonds with a hydrogen atom at ammonia.
(N$_3$) on the imidazole ring in the benzimidazole moiety of mibebradil. One hydrogen atom from the central ammonium of NNC 55-0396 could also form a hydrogen bond with carbonyl-oxygen in the amide of N$^{2p952}$ domain II at $\alpha$$_1$G (Figure 2A); For $\alpha$$_1$H, one oxygen from carboxyl groups on the side-chain of mibebradil forms a hydrogen bond within 3.5 Å at S$^{4p1776}$ from domain IV. Ammonia (N$_3$) on the imidazole ring in the benzimidazole moiety of NNC 55-0396 generates a connection with hydrogen atom from K$^{3p1405}$ from domain III (Figure 2B); For $\alpha$$_1$I, one hydrogen atom from ammonia (N$_3$) on the imidazole ring in the benzimidazole moiety of NNC 55-0396 could bind to the lipophilic side-chain of L$^{2p776}$ at domain II. In comparison, the mibebradil is less likely to place at the center of the channel pore after binding to the amino acid (Figure 2C). The predicted binding affinities between testing drugs and TCCs from 2.67 to 25.99 mM (Figure 2D) are matched to other published studies.$^{14,38–40}$

3.3 | MD simulations and analysis

The time-dependent root-mean-square-displacement (RMSD) plot for C-alpha and whole protein indicates that all three TCC membrane protein complex reaches the equilibrium at 20 ns (Figure 3A–$\alpha$$_1$G, Figure 3A–$\alpha$$_1$H, and Figure 3A–$\alpha$$_1$I). Employing the first-five ranking eigenvalue,$^{41}$ the structural variation and conformation

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**FIGURE 2** Comparison of amino acid binding sites between mibebradil and NNC 55-0396 across three T-type calcium channel (TCC) structures. Cyan: domain I; green: domain II; yellow: domain III; and red: domain IV. Sidechains of four key amino acids are given with partially displayed domains for better visualization. (A) mibebradil (magenta) binds to S$^{4p1776}$ and NNC 55-0396 (wheat) binds to N$^{2p952}$ at $\alpha$$_1$G; (B) mibebradil (marine) binds to S$^{4p1776}$ and NNC 55-0396 (yellow) binds to K$^{3p1405}$ at $\alpha$$_1$H; (C) mibebradil (green) binds to N$^{1s314}$ and NNC 55-0396 (violet) binds to L$^{2p776}$ at $\alpha$$_1$I. (D) predict the binding affinity of mibebradil and NNC 55-0396 varied across $\alpha$$_1$G (red), $\alpha$$_1$H (blue), and $\alpha$$_1$I (black) (unit: kcal/mol).
changes among $\alpha_1G$, $\alpha_1H$, and $\alpha_1I$ are 55.6% (Figure 4B), 61.6% (Figure S2B), and 56.5% (Figure S3B), respectively, while the proportion of variance decreasing monotonically with increased eigenvectors. For the first-three ranking eigenvalue, 50.9% $\alpha_1G$ structure was covered. And this number increased to 58.2% on $\alpha_1H$ and 51.4% on $\alpha_1I$. Conducting principal component analysis (PCA) with three eigenvectors, we did not observe significant conformation changes in the transmembrane part at the last 50 ns simulation for all three TCC structures. The local structural differences among each cluster can be quickly distinguished by checking the shape of the transmembrane region between segment 5 and segment 6 (Figure 3B, clusters 1–3; Figure S2B, clusters 1–3; Figure S3B, clusters 1–3). For $\alpha_1G$, expect for P-loop domain I which dynamic behavior is represented by high RMSD value, the rest of the domains have almost identical structure variation patterns in both sorted and unsorted trajectory data within 50 ns (Figure S4A); There is no clear structure variation of P-loop domains I and IV for $\alpha_1H$ till the last 5 ns (Figure S4A); For $\alpha_1I$, P-loop domain IV has more unstable local fluctuations than other domains (Figure S4C).

The classic MD simulation (0-100 ns) with the following EEF MD (100–200 ns) simulation did not detect the penetration of
calcium ion through the membrane in α₁G, α₁H, and α₁I. For α₁G, Ca³⁷ binds to E¹p³⁵⁴ domain I at \( t = 30 \) ns (Figure 4A), and Ca¹⁵ binds to E²p⁹⁲³ domain II at \( t = 8 \) ns (Figure 4B). Before simulation reaches steady-state, Ca³⁷ is initially shared by two SFs at domains II and III for 25 ns, then it jumps to E¹p³⁵⁴ domain I (Figure S5A and B). SF at domain IV has less attraction to calcium ions. For α₁H, Ca⁵⁸ binds to E²p⁸⁸⁰ domain I at \( t = 0 \) (Figure 4C) and Ca³⁸ binds to E²p⁸⁷⁶ domain II after 25 ns (Figure 4D). SF D³⁵p¹⁴⁰⁶ at domain III and D⁴p¹⁷¹⁰ at domain IV show limited binding affinity to Ca²⁺ (Figure S5C and D). For α₁I, Ca³¹ binds to E²p⁷⁴⁴ at domain II (Figure 4E) with unstable binding behavior and Ca⁵⁶ firmly binds to E⁴p¹⁶⁰¹ domain IV (Figure 4F) after 25 ns. The SF at domains I and III shows fewer interests in binding Ca²⁺ (Figure S5E and F). At the end of EEF MD simulation, α₁G and α₁H have two calcium ions bind to the domain I and II (Figure S5A and B); however, two calcium ions go to domains II and IV at α₁I (Figure S5C).

Charged amino acids or amino acids with polarity potential are more likely to interact with the Ca²⁺ when the interatomic distance...
is smaller than 10 Å. Our SMD tests find $T_{2921}$, $Q_{2922}$, and $D_{2924}$ at domain II from $\alpha_1G$; $T_{2674}$, $Q_{2672}$, and $Q_{2675}$ at domain II from $\alpha_1H$; $T_{2742}$, $Q_{2743}$ at domain II; and $K_{3p1302}$ at domain III from $\alpha_1I$ can interact with pulling $Ca^{2+}$ due to the short atom–atom distance (Figure 5D–F). During the 3 ns SMD simulation, a maximum pulling force is less than −15 pN along the z-axis for all three TCCs structures.

The reaction coordinate in the tests is fixed in the negative direction along the z-axis; three clusters have different initial z-coordinate of $Ca^{2+}$, which yield the varied initial force profile. For all three TCC structures, throughout the $Ca^{2+}$ penetration, the absolute value of interatomic force first decreases in the SF region, the increase when $Ca^{2+}$ wants to escape from the SFs. Eventually, it reaches the
maximum before leaving the intercellular gate. Expect for cluster 1s at $\alpha_1H$ and $\alpha_1I$, an arch-shape force was generated when Ca$^{2+}$ moving towards the negative direction of Z (Figure 6).

Replace K$^{3p1462}$ in $\alpha_1G$ with Phe or Gly will shift the activation and decrease the drug sensitivity.$^{19}$ FEP analysis finds in water a mutation caused total Gibbs-free energy to increase by 43.83, 40.52, and 36.9 kcal/mol on $\alpha_1G$, $\alpha_1H$, and $\alpha_1I$, respectively. In vacuum, these numbers become 73.96, 43.58, and 62.27 kcal/mol on $\alpha_1G$, $\alpha_1H$, and $\alpha_1I$, respectively. Thus, the contribution of mutation lysine (K$^{3p1462}$, K$^{3p1405}$, and K$^{3p1302}$) at P-loop domain III on whole protein hydration-free energy $\Delta\Delta G_{(K-A)}$ yields: $-30.13$, $-3.06$, and $-25.37$ kcal/mol. FEP analysis on $\tau^{2p921}$, $\tau^{2p874}$, and $\tau^{2p742}$ also finds the polarity of threonine in response to the mutation test. In water, $\Delta G$ variation is $13.08$, $-25.47$, and $14.65$ kcal/mol for $\alpha_1G$, $\alpha_1H$, and $\alpha_1I$, respectively. These numbers become $13.8$, $11.46$, and $14.54$ kcal/mol (Table 2). Details of how Gibbs-free energy in response to selection of $\lambda$ during the alchemical reaction in water or vacuum are given in Figure S6. The quality control of FEP analysis is given in Figure S7.

**Figure 6** Nonlinear increased pulling force (blue) in response to moving one Ca$^{2+}$ across three channels, (A) $\alpha_1G$, (B) $\alpha_1H$, and (C) $\alpha_1I$. The error bar is colored by gray for $\alpha_1G$ and green for selected cluster 1 structure, yellow for selected cluster 2 structure, and magenta for selected cluster 3 structure from last 50 ns of external electric force field (EEF) molecular dynamics (MD) simulation trajectory of $\alpha_1H$, and $\alpha_1I$. 

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**TABLE 2** 

| Channel  | Gibbs-Free Energy (kcal/mol) |
|----------|------------------------------|
| $\alpha_1G$ | $-30.13$ |
| $\alpha_1H$ | $-3.06$ |
| $\alpha_1I$ | $-25.37$ |

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**TABLE 2** 

| Channel  | Gibbs-Free Energy (kcal/mol) |
|----------|------------------------------|
| $\alpha_1G$ | $13.08$ |
| $\alpha_1H$ | $-25.47$ |
| $\alpha_1I$ | $14.65$ |

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**TABLE 2** 

| Channel  | Gibbs-Free Energy (kcal/mol) |
|----------|------------------------------|
| $\alpha_1G$ | $13.8$ |
| $\alpha_1H$ | $11.46$ |
| $\alpha_1I$ | $14.54$ |
TABLE 2 Free Gibbs energy (ΔG) and free hydration energy (ΔΔG) variation introduced by mutation lysine/threonine into alanine (unit: kcal/mol).

|          | K1462/K1405/ K1302 | T921/T874/T742 |
|----------|---------------------|-----------------|
|          | ΔG(Water) | ΔG(Vacuum) | ΔΔG | ΔG(Water) | ΔG(Vacuum) | ΔΔG |
| αi,G     | 43.83      | 73.96       | -30.13 | 13.08      | 13.8        | -0.72 |
| αi,H     | 40.52      | 43.58       | -3.06  | -25.47     | 11.46       | -36.93 |
| αi,I     | 36.9       | 62.27       | -25.37 | 14.65      | 14.54       | 0.11  |

4 | DISCUSSION

Emerging evidence points out the pathological role of TCCs that is associated with the progression of different diseases. Previous works suggest that nitrile and isopropyl groups in phenylalkylamine serve a role to guide the drug to the position of Ca2+: this function remains positive if the nitrile is replaced with other high electronegative potential elements such as oxygen or sulfur. The recently deposited crystallization structure of human Ca3.1 offers a great opportunity to explore the Ca2+ binding mechanism on all TCCs.19

It is suggested that at least 20,000 decoys should be generated if we go through the de novo homology modeling process.43 Using the human Ca3.1 structure as a template, we set the sampling number at 15,000 for each comparative modeling case. Although the homology modeling template is constructed based on the splice variant containing a deletion of amino acids within the I–II linker, there are not AAs breaks from SF to the intercellular gates. Therefore, delete the regions that are not aligned with αi,G from selected clustering results should not affect the docking preference of Ca2+ to αi,H and αi,I. Vina does predict binding free energy between ligand and receptor less accurate than MM-PBSA.44 However, it gives a direct visualization for filtering out the incorrect binding poses between ligand and receptor. Using the predefined homology models, we confirmed that the mibebradil and NNC 55-0396 have different binding regions on the same structure but consistent binding regions from αi,H to αi,I. This finding partially explained why NNC 55-0396 is an exclusive TCCs blocker.14 The existence of positively charged lysine at P-loop domain III will largely reduce the probability of Ca2+ binds to SF at domain III. Our MD simulations further confirmed that SF at P-loop domain II is the common region to dock Ca2+ across all TCCs structure. Two AAs (K1302/Thr1462) that affect the binding affinity of z944 on αi,G19 also show a clear variation of free hydration energy if mutate to alanine in water. Additional tests results for free hydration energy variation due to the mutation of calcium-binding associated amino acids are available upon request. We used water as solvation in FEP analysis because only K1462 has a direct impact on αi,G current and conductance. More sophisticated conditions should add into solvate for calculating the influence of polarized AA if it has the potential to be the target site to bind.

DISCLOSURE

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

YL, GYL, and ZC: Design, programming, and writing; KZ, A.E., ML, NZ: Supervision.

ETHICS

N/A.

OPEN RESEARCH BADGES

This article has earned Open Data, Open Materials and Preregistered Research Design badges. Data, materials and the preregistered design and analysis plan are available in the article.

DATA AVAILABILITY STATEMENT

Human Ca3.1(6KZO), rabbit Ca1.1 (3JBR), and bacterial CaAb (5KMH) Cryo-EM structures were downloaded from RCSB PDB. Amino acid sequences of Ca3.2 (O95180) and Ca3.3 (Q9POX4) were downloaded from UniProt. The 2D structures of mibebradil and NNC 55-0396 were downloaded from PubChem. The homology modeling and MD simulation data are available on request from the authors.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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