Exploration of binding and inhibition mechanism of a small molecule inhibitor of influenza virus H1N1 hemagglutinin by molecular dynamics simulation

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Influenza viruses are a major public health threat worldwide. The influenza hemagglutinin (HA) plays an essential role in the virus life cycle. Due to the high conservation of the HA stem region, it has become an especially attractive target for inhibitors for therapeutics. In this study, molecular simulation was applied to study the mechanism of a small molecule inhibitor (MBX2329) of influenza HA. Behaviors of the small molecule under neutral and acidic conditions were investigated, and an interesting dynamic binding mechanism was found. The results suggested that the binding of the inhibitor with HA under neutral conditions facilitates only its intake, while it interacts with HA under acidic conditions using a different mechanism at a new binding site. After a series of experiments, we believe that binding of the inhibitor can prevent the release of HA1 from HA2, further maintaining the rigidity of the HA2 loop and stabilizing the distance between the long helix and short helices. The investigated residues in the new binding site show high conservation, implying that the new binding pocket has the potential to be an effective drug target. The results of this study will provide a theoretical basis for the mechanism of new influenza virus inhibitors.

Influenza virus is the causative agent of influenza, an infectious disease which usually leads to symptoms such as high fever, cough, headache, muscle and joint pain, sore throat, nasal discharge, and even a fatal illness similar to pneumonia1–4. Influenza viruses are divided into three types, type A, type B and type C, with influenza A virus presenting serious threats to public health worldwide due to its high mutation rate5–7. At present, two classes of drugs are approved by the Food and Drug Administration for treatment or chemoprophylaxis of influenza: matrix protein 2 (M2) inhibitors amantadine and rimantadine and the neuraminidase (NA) inhibitors (NAIs) oseltamivir and zanamivir8, 9. However, with the wide use of these drugs, drug-resistant strains have appeared in succession10. Therefore, new antiviral targets with novel inhibition mechanism need to be developed.

Hemagglutinin (HA), a viral receptor-binding and membrane-fusion glycoprotein involved in the invasion of influenza into host cells, plays an essential role in the life cycle of influenza A virus3, 11. HA is a trimer of identical subunits, each of which consists of a variable membrane-distal receptor-binding globular domain (HA1) and a more conserved membrane-proximal helix-rich stem structure (HA2)12. Under acidic conditions, the residues on the outer surface of HA1 can be protonated easily, which leads to the large gathering of positive charges on the surface of HA13, 14. With gradually increased charges, disaggregation of HA first occurs as the positive charges repel each other, followed by the entry of water into the interior of the protein causing further structural changes of HA213. In the HA2 subunit, one short and one long α-helical segment are connected by a loop (helix-loop-helix...
ions was added to randomly replace the water molecules (18 sodium ions for INT-HAWashington complex, 15 sodium ions for INT-HAFlorida complex and 33 chloride ions for INT-HA1ruz complex). First, energies of the complex were minimized. The grid Vina, a grid of 24 × 24 × 24 points in the x, y, and z-axis directions was built with a grid spacing of 1 Å, the center of which was treated as the mass center of one monomer of HA, and the exhaustiveness was set to 2031. The grid detail used in the docking simulations is shown in Fig. S3.

**Molecular dynamics simulation.** The three molecular systems are listed in Table 1. All of the complex systems were subjected to molecular dynamics simulation in a periodic boundary condition using the GROMACS 4.5.2 software package with SPC (simple point charge) water model. The Gromos 53 A6 force field was used to describe both the receptors and ligand. The parameterization of the INT was produced by Automated Topology Builder (ATB) server. To keep each system at an electrically neutral state, an appropriate amount of ions was added to randomly replace the water molecules (18 sodium ions for INT-HAWashington complex, 15 sodium ions for INT-HAFlorida complex and 33 chloride ions for INT-HA1ruz complex). First, energies of the complex systems were minimized. Then, the systems were subjected to molecular dynamics simulation in a periodic boundary condition using the GROMACS 4.5.2 software package with SPC (simple point charge) water model. The Gromos 53 A6 force field was used to describe both the receptors and ligand. The parameterization of the INT was produced by Automated Topology Builder (ATB) server. To keep each system at an electrically neutral state, an appropriate amount of ions was added to randomly replace the water molecules (18 sodium ions for INT-HAWashington complex, 15 sodium ions for INT-HAFlorida complex and 33 chloride ions for INT-HA1ruz complex).
on HA and INT to allow for relaxation of the solvent molecules in two phases. The solvent molecules were equili-
ibrated with fixed protein at 300 K, taking the initial velocities from a Maxwellian distribution. Subsequently, the pro-
tein and inhibitor were relaxed step by step and heated up to 300 K. The long-range electrostatics were de-
scribed with the particle mesh Ewald algorithm with interpolation order of 4, a grid spacing of 0.16 nm and the
Coulomb cutoff distance of 1.0 nm. The LINCS (Linear Constraint Solver) algorithm was used to constrain all
bonds. Temperature and pressure coupling types were set with V-rescale and Parrinello-Rahman, respectively.
In the NVT ensemble, the temperature of the systems reached a plateau at the desired value (reference tempera-
ture = 300 K; time constant = 0.1 ps). In addition, the equilibration of pressure (reference pressure = 1.0 bar; time
constant = 2.0 ps) was performed under the NPT ensemble. Finally, molecular dynamics simulations for collect-
ing data with a time step of 2 fs and coordinates every 2 ps were initiated.

**Calculation of binding free energy and ligand-residue interaction decomposition.** In our study, the binding free energies were calculated using the MM-PBSA method supplied with the Amber 10 package. A total number of 100 snapshots were chosen evenly from the molecular dynamics trajectory. The MM-PBSA method can be conceptually summarized as:

\[ \Delta G_{\text{bind}} = \Delta G_{\text{complex}} - [\Delta G_{\text{protein}} + \Delta G_{\text{lig}}] \]  

(1)

\[ \Delta G_{\text{bind}} = \Delta H - T \Delta S \]  

(2)

where \( \Delta H \) of the system consists of the enthalpy changes in the gas phase upon complex formation (\( \Delta E_{\text{MM}} \)) and the solvated free energy contribution (\( \Delta G_{\text{sol}} \)), while \(-T \Delta S\) refers to the entropy contribution to the binding. Equation (2) can then be approximated as shown in Eq. (3):

\[ \Delta G_{\text{bind}} = \Delta E_{\text{MM}} + \Delta G_{\text{sol}} - T \Delta S \]  

(3)

In addition, the change in flexibility of HA and inhibitor upon binding is similar in all complexes. Based on previous studies, the entropy differences should be very small. Thus, the calculation of the solute entropy term was omitted in the present study.

\[ \Delta E_{\text{MM}} \] is the summation of the van der Waals (\( \Delta E_{\text{vdw}} \)) and the electrostatic (\( \Delta E_{\text{ele}} \)) interaction energies.

\[ \Delta E_{\text{MM}} = \Delta E_{\text{vdw}} + \Delta E_{\text{ele}} \]  

(4)

In addition, \( \Delta G_{\text{sol}} \), which denotes the solvation free energy, can be computed as the summation of an electrostatic component (\( \Delta G_{\text{el, electro}} \)) and a nonpolar component (\( \Delta G_{\text{nonpolar, sol}} \)), as shown in Eq. (5).

\[ \Delta G_{\text{sol}} = \Delta G_{\text{el, electro}} + \Delta G_{\text{nonpolar, sol}} \]  

(5)

The interactions between the inhibitor and each residue in the binding site were analyzed using the MM-PBSA decomposition process applied in the MM-PBSA module in Amber 10. The binding interaction of each ligand-residue pair includes three terms: the van der Waals contribution (\( \Delta E_{\text{vdw}} \)), the electrostatic contribution (\( \Delta E_{\text{ele}} \)) and the solvation contribution (\( \Delta E_{\text{sol}} \)). All energy components were calculated using the same snapshots as the free energy calculation.

**Results and Discussion**

Influenza virions that are attached to the host cell membrane via HA are phagocytosed and gain entry into the
cell through endocytosis. During this endocytosis process, small molecules generally can enter the endosome
at the same time. Thus, the effective combination of Inhibitors of HA under neutral conditions outside the cell
is crucial for the intake of INT. After entering into the endosome, HA will undergo large conformational changes
under acidic conditions, which is aimed at promoting the membrane fusion between the virus and host cell.
An efficient HA entry inhibitor must be able to stop the progression of changes in this molecule. In order to investi-
gate if INT could stably bind with HA under neutral conditions and be efficiently carried into the endosome to
exert its inhibition at low-pH conditions, the optimal binding mode, binding force and binding behavior of INT
with HA were determined by a series of molecular dynamics simulations and MM-PBSA calculation.

**Analysis of reliability of investigated complex systems.** The validation was carried out using
Ramachandran plot calculations computed with the Procheck program by examining the detailed
residue-by-residue stereochemical quality of HA structures, and the results are shown in Fig. 1. Altogether, 100%
of the investigated residues were in allowed regions, which confirmed the availability of the modeled HA protein
systems. AutoDock Vina was used for obtaining the initial structure of INT-HA complexes. The docking results
were in agreement with those from the mutation experiments and molecular docking reported by Arnab.

For the accuracy of dynamics sampling, the complex structure of INT-HA Florida was regarded as a replicate
system of INT-HA Washington under the neutral condition molecular simulations. In order to monitor if two sys-
tems possess similar dynamics behaviors, the cross-correlations of their Cα atomic displacements, a parameter
which can reflect the detailed atomic dynamic state, were analyzed during the simulations.

The relevant results are illustrated in Fig. 2. Highly positive regions (red) indicate a strong correlation in the movement of residues,
whereas negative regions (blue) are associated with strong anti-correlated motion of the residues. The color depth of diagonal could represent the movement degree of atoms. As shown in Fig. 2a and b, the similarity could be observed in the behavior of Cα atoms from HA_Washington HA1 and HA_Florida HA1. Also the similar behavior could be discovered from Cα atoms from HA_Washington HA2 and HA_Florida HA2 as shown in Fig. 2c and d. As expected, no obvious differences in the movement of atoms occurred between the two complexes during the timescale of simulations. Moreover, according to the structures of both simulations, the conformations of two repeated simulations were found to be identical. Two independent dynamics samplings yielded similar results. Thus, in the following sections, results from only INT-HA_Washington are discussed.

**Binding mode of complex system under neutral conditions.** For convenience, INT-HA_Washington is simplified as INT-HA in the following section. After 200 ns simulations, the root-mean-square deviation (RMSD) of backbone Cα atoms and all atoms of HA were first investigated to evaluate if the complex system could reach an equilibrium during the simulation. As shown in Fig. 3, the RMSD curve of the INT-HA complex was found to reach the equilibrium at about 40 ns, suggesting that the structures could be applied to analyze the optimal binding mode between INT and HA.

In order to obtain the most stable complex structure, cluster analysis of INT-HA was investigated to determine the optimal binding mode. In addition, the comparison of free energy surfaces between the first 100 ns and further one with extended 100 ns were applied to confirm the convergence of complexes (Fig. S5). Cluster analysis is a method that can categorize similar low-energy conformations into the same designated area by analyzing the relative free energy surfaces along the first two principle components (PC-1, PC-2) of the complex during simulation. Two principal amounts (PC-1 and PC-2) were obtained as the reaction coordinate that renders the potential of the mean force ($\Delta F = -RT \times \ln P$, where $P$ is the relative probability in a region) free-energy plane. The first few PCs describe the slow-motion modes which are related to the functional motions of a biomolecular system. In the cluster analysis plot (Fig. 4a), the conformations found in the blue area indicated more stable and lower energy states than those found in the red area. And these lower energy conformations generally could be chosen as the best analysis subjects for the binding modes. As shown in Fig. 4a, more than 80% of the conformations converged at the blue area, and the structure from this area could be applied to the binding mode analysis. The detailed binding mode of INT-HA based on the above analysis is shown in Fig. 4b. The results revealed that INT is bound in the pocket located on the interface between HA1 and HA2. This pocket is surrounded by the side chains of HA1-Val31, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51. The side chain of HA1-Val31 extends towards C9 of INT, and the side chain of HA1-Leu290 is close to C7 of INT. HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51 together approach the seven-member heterocycle of INT.

In order to confirm if INT could stably bind with HA in this pocket during the simulation, the distances between the above residues and INT were calculated by using the distance analysis program of GROMACS. According to the corresponding results shown in Fig. S3, all of the distances monitored were around 0.3–0.4 nm during the 200 ns simulation, which indicated the accuracy of the binding pose. INT is a molecule with an aminoalkyl phenol ether scaffold and has no hydrogen donor or receptor. Thus, the stabilization of INT in the binding pocket most likely contributes to nonpolar interactions. It is well known that the van der Waals radius of a carbon atom is 0.2 nm, and distances of 0.3–0.4 nm favor the formation of van der Waals interactions between two carbon atoms. According to the distances between INT and residues around the pocket shown in Fig. 5, we speculate that van der Waals interactions provide the major contribution to the bonding of INT and HA. This speculation was verified as described in the next section.
Verification of binding mode of INT-HA under neutral conditions. In order to further validate the binding modes and obtain detailed information on the key residues of the binding pocket in the complex, the electrostatic, van der Waals, solvation and total contribution of the residues to the binding free energy of the INT-HA complex were calculated with the MM-PBSA method. To achieve more accurate results, MM-PBSA calculations were carried out twice. The snaps used in calculations were respectively extracted from equilibrium stages within first and second 100 ns. By comparing the energy of the two results, similarity could be observed (Fig. S6), thus, only one group of data from the equilibrium stages within the first 100 ns was shown in this section. The contributions from the residues around INT are shown in Fig. 6 and Table 2.

The contribution of selected residues surrounding the INT-HA complex binding pocket can be clearly seen from Fig. 6 and Table 2. As shown, HA1-Val31, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51 have favorable total energy contributions, suggesting that these residues participate in the complex binding. More importantly, for aforementioned residues van der Waals interactions play a major role in the total energy contribution as evidenced by lower values of $\Delta E_{\text{vdW}}$ ($-1.72$, $-0.65$, $-0.52$, $-0.54$, $-0.73$ and $-1.08$ kcal·mol$^{-1}$, respectively). The free energy calculation results are consistent with the speculation above that van der Waals interactions are a major contributor to the bonding of INT and HA.
The total binding energy of the INT-HA complex systems, an important standard measure of binding affinity between INT and HA, were calculated as well. Total binding energies of INT-HA\textsubscript{Washington} and INT-HA\textsubscript{Florida} were calculated by MM-PBSA at the same time. As listed in Table 3, the results show that the values of $\Delta G_{\text{total}}$ for INT-HA\textsubscript{Washington} and INT-HA\textsubscript{Florida} are $-16.15$ and $-16.02$ kcal·mol$^{-1}$, respectively, indicating that INT has similar binding interactions with HA\textsubscript{Washington} and HA\textsubscript{Florida}. The calculated result is qualitatively consistent with the experimental results showing that INT could inhibit HA of A/Washington/10/2008(H1N1) and A/Florida/21/2008(H1N1) at similar levels (IC$_{50}$ of 0.29 µM and 0.3 µM) \textit{in vitro}\textsuperscript{16}.

Contributions from total free energy were analyzed in detail as well. The $\Delta G_{\text{total}}$ can be divided into polar and nonpolar energies. The free energies of INT binding to two HAs were found to be primarily derived from the nonpolar energies (equal to $\Delta G_{\text{nonp}} + \Delta E_{\text{vdw}}$), which were $-21.95$ kcal·mol$^{-1}$ in the INT-HA\textsubscript{Washington} complex and $-22.75$ kcal·mol$^{-1}$ in the INT-HA\textsubscript{Florida} complex. However, for both complexes, the polar energy (equal to $\Delta G_{\text{pol}} + \Delta E_{\text{ele}}$) showed negative contributions at $-5.80$ and $-6.73$ kcal·mol$^{-1}$ for INT-HA\textsubscript{Washington} and INT-HA\textsubscript{Florida}, respectively. In addition, the values of $\Delta E_{\text{vdw}}$ for INT-HA\textsubscript{Washington} and INT-HA\textsubscript{Florida} were calculated at $-18.29$ and $-19.42$ kcal·mol$^{-1}$, respectively, indicating that van der Waals interactions are dominant in the total energy.

Figure 3. RMSD plot during molecular dynamics simulations.

Figure 4. (a) Relative free energy surfaces along the first two principle components (PC-1, PC-2) of INT-HA under neutral conditions. (b) Predicted binding modes of INT-HA under neutral conditions.
**Figure 5.** Distances between INT and residues (HA1-Val31, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51) around the binding site under neutral conditions.

**Figure 6.** Decomposition of the binding energy on per residue basis on INT-HA under neutral conditions.
The binding and dynamics behavior of INT at low pH conditions. In the above discussion, effective binding of INT with HA under neutral conditions was demonstrated. Nevertheless, the ability of INT to maintain the interaction with HA at a low pH is crucial to its inhibition and determines whether it has the potential to be a HA entry inhibitor. Prior experimental data indicated that INT indeed could inhibit different influenza virus H1N1 strains in vitro, but its mechanism of inhibition at low pH at the atom level has not been explored. In this section, the binding and dynamics behavior of INT at low-pH conditions will be discussed.

As described in method, HA$_{ruz}$ was chosen as the low-pH model in this section based on the calculated pK$_a$ of the titratable residues by Zhou et al. In order to determine whether the INT binding would affect the protonation state of the residues, the titratable residues’ locations of HA$_{ruz}$ were investigated. Most titratable residues were located far away from the INT binding site except HA1-Lys278, HA1-Glu302 and HA2-Glu56. Subsequently, the pK$_a$ of the above three residues were calculated via PRoPKA with or without INT (Table S2). It’s further illustrated that the bound of INT could not affect the protonation state of the residues. Thus, protonation state of HA$_{ruz}$ could be assigned according to the results of Zhou et al. directly (Table S1).

In order to facilitate the comparison, for the investigated complex in this study only one monomer of the HA trimer was bound with INT, and the other two unbound monomers still could disaggregate at low pH as usual. In order to monitor the location of INT at low-pH conditions, distances between INT and its binding residues (HA1-Val31, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51) in the neutral state were calculated again. As shown in Fig. 7, almost all the distances between INT and its binding residues (in neutral state) were disrupted at different degrees at 0–45 ns and increased sharply at about 46 ns, except for HA1-Leu290. The disrupted distances kept at about 65–200 ns with small fluctuation. These observations suggest that INT had relocated out of the pocket formed by the side chain of HA1-Val31, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51 and moved to a new low potential energy point at 65–200 ns. The distance between INT and Leu290-HA1 was always maintained at about 0.3–0.4 nm, which indicated that the new binding location was still nearby. In order to confirm this phenomenon, an independent repeated trial was conducted with a different computing resource with the same parameters, and the same results were obtained (Fig. S7). Therefore, we suspected that the binding of INT with HA under neutral conditions only facilitates its intake, and another mechanism exists for the INT binding and inhibition of HA at low-pH conditions.

In order to explore the binding and dynamic behavior of INT at low pH, a cluster analysis of INT-HA$_{ruz}$ was performed. As shown in Fig. 8a, unlike with INT-HA under neutral conditions (Fig. 4a), mainly five blue areas could be found in the free energy surface plot of the complex at low pH. Most conformations of 1–8 ns, 9–28 ns, 29–64 ns, 65–145 ns and 146–200 ns were clustered in regions 1, 2, 3, 4 and 5, respectively. Five representative conformations extracted from different areas are shown in Fig. 8b. After analysis, INT could bind with HA in at least four dominant positions at low-pH conditions. In the initial stage, INT bound between HA1 and the short helix.

### Table 2. Decomposition of binding energy on per residue basis on INT-HA under neutral conditions.

| Residue      | $\Delta E_{ele}$ (kcal mol$^{-1}$) | $\Delta E_{vdw}$ (kcal mol$^{-1}$) | $\Delta G_{np}$ (kcal mol$^{-1}$) | $\Delta G_{PB}$ (kcal mol$^{-1}$) | $\Delta G_{total}$ (kcal mol$^{-1}$) |
|--------------|----------------------------------|----------------------------------|---------------------------------|----------------------------------|-----------------------------------|
| Val31(HA1)   | −1.72                            | −0.36                            | 0.02                            | 2.06                             | 1.27                              |
| Leu290(HA1)  | −0.65                            | −0.02                            | −0.08                           | −0.75                            | −1.04                             |
| Thr316(HA1)  | −0.52                            | 0.08                             | 0.06                            | −0.38                            | −0.09                             |
| Ile47(HA2)   | −0.54                            | 0.03                             | −0.09                           | −0.60                            | −0.07                             |
| Thr48(HA2)   | −0.73                            | 0.05                             | −0.09                           | −0.78                            | −0.07                             |
| Val51(HA2)   | −1.08                            | −0.05                            | −0.07                           | −1.2                             | −0.07                             |

### Table 3. Calculation of binding free energy using MM/PBSA under neutral conditions.

| Energy components (kcal mol$^{-1}$) | INT-HA$_{Washington}$ | INT-HA$_{Florida}$ |
|-------------------------------------|-----------------------|--------------------|
| $\Delta E_{ele}$                    | −1.04                 | 0.19               |
| $\Delta E_{vdw}$                    | −18.29                | −19.42             |
| $\Delta G_{np}$                     | 6.84                  | 6.54               |
| $\Delta G_{PB}$                     | −3.66                 | −3.33              |
| Nonpolar                            | −21.95                | −22.75             |
| Polar                               | 5.80                  | 6.73               |
| $\Delta G_{total}$                  | −16.15                | −16.02             |
of HA2, but INT moved to the HA head region gradually over time. According to the cluster analysis, INT kept the fourth binding state (Fig. 8b (4 and 5)) with HA after 65 ns, which is in agreement with the results from Fig. 7.

Due to the bulky head region of HA1 being rich in protonated residues at low pH, a large repulsion promoted the structural disaggregation at the initial stage of the simulation. As the initial binding site of INT was far away from the positively charged head, INT could not prevent the disaggregation at the initial stage. We propose that following the disaggregation of HA1 and the shift of charged state of HA2, a new binding site might be revealed and thus enabled the purposeful relocation of INT.

The new detailed binding mode of the INT-HA1ruz complex at low pH was investigated and verified by simulation and MM-PBSA calculation twice as described above (Fig. S8). The relevant results are provided in Fig. 9 and Table 4. As shown, a new pocket bound by INT at low pH was found to be surrounded by HA1-Lys278, HA1-Leu290, HA1-Thr316, HA2-Ile47, HA2-Thr48 and HA2-Val51. In detail, the seven-member heterocycle of INT could locate between side chains of HA1-Lys278 and HA1-Ser289. Side chain of HA1-Pro304 is close to C9 of INT and the side chain of HA2-Thr60 is adjacent to C8 of INT. HA1-Leu290, HA1-Pro291, HA2-Ile55 and HA2-Glu56 together are approaching the six-member ring of INT. The exposed new binding site is still located on the interface between HA1 and HA2. According to the data calculated by MM-PBSA, van der Waals interactions were confirmed to play a major role in the total energy contribution for the residues as well as evidenced by lower values of $\Delta E_{\text{vdw}}$ (−1.07, −1.58, −1.72, −2.03, −1.43, −0.58, −1.03 and −0.98 kcal·mol$^{-1}$, respectively) as shown in Table 4. Although there are four polar residues, HA1-Lys278, HA1-Ser289, HA2-Glu56 and HA2-Thr60, locating around the new binding site, the polarity of these residues still could not contribute to INT binding due to the molecular property of INT. Furthermore, all the polar part of the side chains from the above residues were redirected to the outside of the binding pockets. The sequence conservation of eight residues around the new binding site was evaluated by sequence alignment of the different H1N1 influenza virus strains in the recent 30-year period of time. The results (Table 5) indicated that all the investigated residues in the new binding site are highly conserved, implying that the new binding pocket has the potential to be an effective drug target.
Mechanism of INT Inhibiting HA disaggregation at low pH conditions. Based on analysis of its amino acid sequence, the HA2 polypeptide was found to contain the well-known heptad-repeat module of hydrophobic/hydrophilic amino-acids, which is crucial for the formation of coiled-coil structures. In order to drive the loop-to-helix transition and the formation of a coiled-coil structure, it is necessary for water molecules to directly interact with the hydrophilic residues\(^1\). Therefore, the ability of INT to stabilize the connection between HA1 and HA2 and to effectively prevent water from entering into the interior of HA2 is crucial for the inhibitory activity of INT. Because INT is located on the interface between HA1 and HA2, it is considered to not only be able to stabilize HA1 and HA2 but also prevent the entry of water into the interior of HA and further influence the rearrangement of HA2.

In order to investigate the different degrees of water accessibility, the hydrophobic areas of HA2 subunits with or without INT were monitored during 200 ns dynamics simulation at low pH. As shown in Fig. 10, the hydrophobic area of HA2 subunits bound with INT always stabilized at about 70 nm\(^2\), but those of the other two free monomers (HA monomers bound without INT) clearly decreased at a value below 70 nm\(^2\) as the simulation. This reduced hydrophobic area of the free HA2 subunit indicated that the solution gained access to the inside of the protein due to the lack of protection by INT. By contrast, the HA2 subunit from monomers bound with INT consistently retained the larger hydrophobic area throughout the simulation.

A helix-loop-helix structure provides stability for HA2, and the entry of water must lead to a structural change of this molecule to some extent. We investigated the different H1N1 influenza virus strains in recent 30-year period of time. *Low entropy data means high conservation.

| Residue    | \(\Delta E_{\text{int}}\) (kcal mol\(^{-1}\)) | \(\Delta E_{\text{vdw}}\) (kcal mol\(^{-1}\)) | \(\Delta E_{\text{ele}}\) (kcal mol\(^{-1}\)) | \(\Delta E_{\text{sol}}\) (kcal mol\(^{-1}\)) | \(\Delta E_{\text{total}}\) (kcal mol\(^{-1}\)) |
|------------|--------------------------------------|------------------|------------------|------------------|------------------|
| Lys278(HA1) | -1.07                                | -0.63            | 0.72             | -0.98            |
| Ser289(HA1) | -1.58                                | 0.06             | 0.91             | -0.62            |
| Leu290(HA1) | -1.72                                | -0.78            | 1.05             | -1.45            |
| Pro291(HA1) | -2.03                                | -0.11            | 0.34             | -1.80            |
| Pro304(HA1) | -1.43                                | 0.14             | -0.10            | -1.39            |
| Ile55(HA2)  | -0.58                                | -0.28            | 0.47             | -0.39            |
| Glu56(HA2)  | -1.03                                | 0.35             | -0.11            | -0.79            |
| Thr60(HA2)  | -0.98                                | -0.07            | 0                | -1.06            |

| Residue    | Entropy data* | Residue    | Entropy data* |
|------------|---------------|------------|---------------|
| Lys278(HA1)| 0.42889       | Pro304(HA1)| 0.06495       |
| Ser289(HA1)| 0.04048       | Ile55(HA2)| 0.05048       |
| Leu290(HA1)| 0.03910       | Glu56(HA2)| 0.05353       |
| Pro291(HA1)| 0.03809       | Thr60(HA2)| 0.05977       |

*Low entropy data means high conservation.
shown in Fig. 11a(1), due to the bound INT, HA preferably maintained the initial helix-loop-helix structure, while the short helixes of the two free HA clearly changed. The secondary structures of the three short helixes were monitored during the 200 ns simulation. The command “do_dssp” of Gromacs was used for secondary structure calculation. As shown in Fig. 11b, the great structural fluctuations in the short helixes of free HAs could be observed at about 30 ns, and the distorted short helixes could endure till 200 ns. These distortions were mainly composed of the formations of the coil, bend and turn during the conformation simulation. By contrast, it's found that the corresponding secondary structure of HA could display much more stability in the presence of INT. Except for the influence of INT on the short helix, we suspected that the binding of INT also could maintain the stability of the loops connecting the short and long helices. The stability of loops from each HA2 subunit was measured by the RMSD of loops. As shown in Fig. 12a, the RMSD of the loop from the monomer bound with INT was very stable and held the value at about 0.2 nm, indicating that the binding of INT could maintain the rigidity of the loop. RMSD of loops from the two free monomers fluctuated more obviously, which indicated a disruption in the rigidity of the loops. Differences in the loops can be observed in Fig. 11a as well.

In addition, the side chain dynamic behavior of a group of reported residues (Glu101 and Lys50) was also monitored. The salt bridge between Glu101 and Lys50 is crucial for the stabilization of short and long helices of HA2, and the break of this salt bridge would promote the loop-helix transformation. Therefore, stabilization in the distance between these two residues is an important measure of protective efficacy of INT. From the results shown in Fig. 12b, the distances between Glu101 and Lys50 in the three monomers were found to be different. In the two free monomers, this distance was clearly increased and not favorable for keeping the salt bridge. By contrast, the corresponding distance in the monomer bound with INT was always stable at a lower value, which indicated that the salt bridge of this monomer was protected well and the helix-loop-helix was more stable.
According to the above analysis, we may conclude that INT has the potential to effectively prevent water from entering into the interior of HA, maintain the rigidity of the HA2 loop and stabilize the distance between long helix and short helix, and further obstruct the loop-to-helix transition.

**Conclusion**

In the current study, the mechanism of binding and inhibition of INT, a potential influenza virus HA inhibitor, was explored by molecular dynamics simulation and MM-PBSA calculation. The behaviors of INT under neutral and acidic conditions were investigated at the same time, and an interesting dynamic binding mechanism of INT was found. We propose that the binding of INT with HA under neutral conditions allows for its intake, while the interaction of INT with HA at low-pH conditions at a new binding site employs a different mechanism to carry out its inhibitory effect. After a series of confirmation, we believe that the binding of INT can prevent HA1 from disassociating from HA2, further maintain the rigidity of the HA2 loop and stabilize the distance between the long helix and short helix.

The investigated residues in the new binding site are highly conserved, implying that the new binding pocket has the potential to be an effective drug target. The results in this study will be useful for the development of efficient drug targets and provide a theoretical basis for the mechanism of new influenza virus inhibitors.

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Author Contributions
S.S.G. and Y.M.S. designed the experiments. S.S.G., T.A.W., X.P.T., X.Q.Z. and M.D.Q. carried out the experiments. S.S.G., Y.M.S., Z.Y.K., Y.J.Y., H.Z., S.W., H.L., and W.K. analyzed and discussed the results. S.S.G. wrote the manuscript. All authors have reviewed the manuscript.

Additional Information
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